The Crystal Structure of an Integral Membrane Fatty Acid α-Hydroxylase*

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Neuronal electrical impulse propagation is facilitated by the myelin sheath, a compact membrane surrounding the axon. The myelin sheath is highly enriched in galactosylceramide (GalCer) and its sulfated derivative sulfatide. Over 50% of GalCer and sulfatide in myelin is hydroxylated by the integral membrane enzyme fatty acid 2-hydroxylase (FA2H). GalCer hydroxylation contributes to the compact nature of the myelin membrane, and mutations in FA2H result in debilitating leukodystrophies and spastic paraparesis. We report here the 2.6 Å crystal structure of sphingolipid α-hydroxylase (Scs7p), a yeast homolog of FA2H. The Scs7p core is composed of a helical catalytic cap domain that sits atop four transmembrane helices that anchor the enzyme in the endoplasmic reticulum. The structure contains two zinc atoms coordinated by the side chains of 10 highly conserved histidines within a dimetal center located near the plane of the cytosolic membrane. We used a yeast genetic approach to confirm the important role of the dimetal-binding histidines in catalysis and identified Tyr-322 and Asp-323 as critical determinants involved in the hydroxylase reaction. Examination of the Scs7p structure, coupled with molecular dynamics simulations, allowed for the generation of a model of ceramide binding to Scs7p. Comparison of the Scs7p structure and substrate-binding model to the structure of steroyl-CoA desaturase revealed significant differences in the architecture of the catalytic cap domain and location of the dimetal centers with respect to the membrane. These observations provide insight into the different mechanisms of substrate binding and recognition of substrates by the hydroxylase and desaturase enzymes.

Electrical impulse propagation in the central and peripheral nervous system is facilitated by the myelin sheath, a highly organized multilamellar membrane that wraps around the neuronal axon (1). The myelin sheath is highly enriched in glycosphingolipids, with galactosylceramide (GalCer) and its sulfated derivative sulfatide representing the major glycolipid components. GalCer is generated from a ceramide-based precursor composed of a long chain base (LCB) moiety that is derived from palmitoyl-CoA and a very long chain fatty acid (VLCFA) moiety that is typically 26 carbons long (2). Over 50% of GalCer and sulfatide within the myelin sheath are hydroxylated at C-2 of the VLCFA moiety by the enzyme FA2H (3, 4). No other mammalian tissues contain such high levels of hydroxylated lipid. The hydroxylation of the VLCFA moiety, together with the amide and hydroxyl groups associated with the GalCer and sulfatide headgroups, leads to the formation of extensive hydrogen bonding networks that significantly contribute to the tightly organized and compact nature of the myelin membrane (1, 5). Mutations in FA2H cause hereditary leukodystrophy and spastic paraparesis in humans underscoring this enzyme's importance in the nervous system (6–9).

FA2H belongs to a superfamily of integral membrane di-iron-containing enzymes that hydroxylate or desaturate lipid-based substrates via a reaction mechanism that is dependent on NADH and oxygen (10). Members of this superfamily participate in an electron transport chain that moves electrons from NADH through cytochrome b₅ reductase and cytochrome b₅, with termination at the di-iron center of the enzyme for utilization in subsequent catalysis. Although enzymes within the membrane-bound arm of the superfamily do not share extensive sequence similarity, particular features of their sequences are highly conserved. For example, distinct sequence motifs, termed “histidine boxes,” contain histidine residues that are thought to coordinate two iron atoms at the active center of the enzyme (11, 12). Site-directed mutagenesis of these histidines has shown that they are essential for catalysis (12). In addition, hydropathy plots suggest that the membrane-bound family members contain four transmembrane helices that orient the enzymes such that their N and C termini are located on the cytosolic side of the membrane (13). Many of the integral membrane enzymes of the superfamily, including FA2H, have the intermediate cytochrome b₅ electron donor directly fused to the catalytic domain at the N terminus (14).

The Saccharomyces cerevisiae endoplasmic reticulum (ER) contains five members of the membrane-bound hydroxylase/desaturase family, including sphingolipid α-hydroxylase (Scs7p), the yeast homolog of FA2H (11, 15). In S. cerevisiae, the catalytic domains of these enzymes are distributed throughout the membrane (16). The Scs7p hydroxylase catalyzes the hydroxylation of VLCFA at C-2, and the desaturase catalyzes the desaturation of VLCFA at C-14 (17). The crystal structures of three family members, Scs7p, Hsc8p, and Scs16p, have been determined previously (18–20). While these structures provided insights into the catalytic mechanisms of this superfamily, the structures lack substrate binding specificity. The crystal structure of Scs7p reveals two equivalents of hydroxylation, with the catalytic dimetal center containing five histidines and two zinc ions.

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The atomic coordinates and structure factors (codes 4ZR0 and 4ZR1) have been deposited in the Protein Data Bank (http://wwpdb.org/).

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2 The abbreviations used are: GalCer, galactosylceramide; bO, n-octyl-β-D-glucopyranoside; ER, endoplasmic reticulum; FA2H, fatty acid 2-hydroxylase; LCB, long chain base; m, mouse; PEG, polyethylene glycol; sc, Saccharomyces cerevisiae; VLCFA, very long chain fatty acid; scScs7p, S. cerevisiae Scs7p.
Scs7p is responsible for adding a hydroxyl group to the \( \alpha \)-carbon of the VLCFA moiety of the ceramide substrate to generate inositol phosphoceramide, one of three major sphingolipids in yeast (15). Scs7p and human FA2H share 36% sequence identity. The other four members of the enzyme family in yeast are C-5 sterol desaturase (Erg3p) (16), methyl sterol oxidase (Erg25p) (17), \( \Delta 9 \) desaturase (Ole1p) (18), and sphenolipid C-4 hydroxylase (Sur2p) (15). When compared with other eukaryotic members of the hydroxylase/desaturase family, all five enzymes exhibit sequence conservation of the “histidine box” motifs and similar hydropathy plots indicative of the presence of four putative transmembrane helices. Scs7p and Ole1p are the only family members in yeast that contain a cytochrome \( b_5 \) domain fused to the catalytic domain. The cytochrome \( b_5 \) domain of Scs7p is fused at the N terminus of the enzyme, whereas the \( b_6 \) domain of Ole1p is fused at the C terminus.

Only one integral membrane representative of the hydroxylase/desaturase superfamily, steroyl-CoA desaturase (SCD1) (19, 20), has been successfully crystallized due in large part to the difficulty of producing suitable quantities of stable protein for structural studies. We report here the 2.6-A crystal structure of scScs7p, which is the first hydroxylase enzyme within the hydroxylase/desaturase superfamily to be characterized at the molecular level. The core structure is composed of a helical catalytic cap domain that sits atop four transmembrane helices. The dimetal-binding site is located within the catalytic cap domain and is occupied by zinc atoms coordinated by the highly conserved histidine side chains from four histidine box motifs. A phenotypic assay was used in conjunction with site-directed mutations to confirm the role of the dimetal-binding histidine residues. This approach revealed that Tyr-322 and Asp-323 are critical determinants involved in the hydroxylase reaction. Although global folds of the scScs7p and SCD1 enzymes are similar, the two structures exhibit significant differences in the architecture of the cytosolic cap domain, the location of the dimetal center, and the modes of substrate access and binding. Comparison of the scScs7p and SCD1 structures, coupled with molecular dynamics simulations, allows modeling of a ceramide substrate below the dimetal center of scScs7p.

**Experimental Procedures**

**Construct Design and Mutagenesis**—For protein expression, the SCS7 gene, encoding sphingolipid \( \alpha \)-hydroxylase (Scs7p), was PCR-amplified from *S. cerevisiae* genomic DNA and subsequently inserted into vector pSGP46, containing both C-terminal ZZ and His\(_{10}\) affinity tags along with a 3C protease site for tag removal, via ligation-independent cloning as described previously (21). A deletion construct, with N-terminal residues 2–95 removed (\( \Delta 95 \)scScs7p), was also engineered using the QuikChange mutagenesis kit (Stratagene, La Jolla, CA) with the wild-type scScs7p in vector pSGP46 as the template. All constructs were confirmed by DNA sequencing.

**Protein Expression and Purification**—The overproduction of scScs7p and \( \Delta 95 \)scScs7p for structural studies followed the general procedures outlined previously (21), with some modifications. Both scScs7p and \( \Delta 95 \)scScs7p in pSGP46 were transformed into *S. cerevisiae* strain BJ5460 (ATCC 208285; MAT\( a \) ura3-52 trp1 lys2-801 leu2 \( \Delta 1 \) his3-\( \Delta 200 \) pep4::HIS3 prb1 \( \Delta 1 \Delta 6 R \) can1 GAL\(^+\)). A 1-ml culture of cells expressing either scScs7p or \( \Delta 95 \)scScs7p in BJ5460 was utilized to inoculate 100 ml of SD-Ura media. After overnight growth, the 100-ml culture was added to 8 liters of SD-Ura media in a BioFlo 3000 fermentor (New Brunswick Scientific) and grown at 30 °C with agitation at 500 rpm to an \( A_{600} \) of 1.0. At this time, an additional 2 liters of 5× YP media (1% yeast extract; 2% peptone) was added to the fermentor. Expression of scScs7p and \( \Delta 95 \)scScs7p from pSGP46 was controlled by the ADH2 promoter, which is induced when glucose is completely depleted from the growth medium. Thus, the glucose concentration of the medium was monitored utilizing a glucose monitor. Upon depletion of glucose, the cells were grown for 16–18 h and harvested via centrifugation.

For purification of scScs7p and \( \Delta 95 \)scScs7p, the cell pellets from a 10-liter fermentor growth was resuspended in 50 mM Tris, pH 8.0, 150 mM NaCl at a ratio of 1 ml of resuspension buffer per 0.35 g wet cell pellet weight. The cells were lysed by four passes at ~24,000 p.s.i. through a Microfluidics M110 Microfluidizer. PMSF was added to the cell lysate at a ratio of 1.7 mg of PMSF/liter of cell pellet utilized. The lysed cells were centrifuged at 4000 × g for 10 min at 4 °C to remove cell debris. The supernatant was subsequently ultracentrifuged at 125,000 × g for 120 min at 4 °C to isolate the membrane fraction. The membrane fraction was resuspended in 50 mM Tris, pH 8.0, 150 mM NaCl at a ratio of 10 ml of buffer/g of membrane, along with the addition of 2 tablets of Complete Protease Inhibitor mixture (Roche Applied Science), using a homogenizer. For solubilization, \( \beta \)-octyl glucoside (\( \beta \)OG) was added to a final concentration of 2% (w/v). The mixture was stirred at room temperature for 90 min, followed by ultracentrifugation at 125,000 × g for 40 min at 4 °C to remove insoluble debris. Affinity purification was carried out in batch utilizing the C-terminal ZZ tag in conjunction with IgG-Sepharose 6 Fast Flow resin (GE Healthcare) pre-equilibrated in 50 mM Tris, pH 8.0, 150 mM NaCl, 1% \( \beta \)OG (w/v). A total of 10 ml of pre-equilibrated IgG-Sepharose 6 Fast Flow resin was added to the solubilized membrane fraction, followed by incubation with rocking for 2.5 h at 4 °C. The resin was then poured into a 13 × 2.5-cm column and washed with 100 ml of 50 mM Tris, pH 8.0, 150 mM NaCl, 1% \( \beta \)OG (w/v). The protein was subsequently cleaved from the IgG-Sepharose resin by adding 1 mg of rhinovirus 3C protease, engineered with N-terminal His\(_{6}\) and maltose-binding protein affinity tags (22), in 10 ml of 50 mM Tris, pH 8.0, 150 mM NaCl, 1 mM EDTA, 1% \( \beta \)OG (w/v) and incubating for 16 h at 4 °C. The cleaved protein was eluted from the IgG-Sepharose resin by washing with 100 ml of 50 mM Tris, pH 8.0, 150 mM NaCl, 1% \( \beta \)OG (w/v) to the combined elution and wash fractions and incubating for 1 h with rocking at 4 °C. The slurry was poured into a 13 × 1-cm column and washed with 10 column volumes of 50 mM Tris, pH 8.0, 150 mM NaCl, 1% \( \beta \)OG (w/v). The wash fractions were pooled, concentrated to ~2 ml using an Amicon Ultra-15 concentrator with a 50-kDa molecular mass cutoff (30 kDa for \( \Delta 95 \)scScs7p; Millipore), and subjected to size-exclusion chromatography using a
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Superdex 200 HiLoad 16/60 column (GE Healthcare) equilibrated in 20 mM Hepes, pH 7.0, 150 mM NaCl, 1% βOG (w/v). Peak fractions were pooled and concentrated to ~3 mg/ml for crystallization trials. 10-Liter fermenter growths consistently yielded ~2.5 mg of pure scScs7p and ∆95scScs7p.

Crystallization—Initial crystallization screening was conducted using the 1536 condition tailored membrane protein screen (23) in the High-Throughput Crystallization Laboratory at the Hauptman-Woodward Institute (24). Crystals of scScs7p were grown at 14 °C using the microbatch-under-oil method by combining 2 μl of protein solution at 3.3 mg/ml with 2 μl of 20% polyethylene glycol (PEG) 3350 (w/v), 100 mM Hepes, pH 7.0, 300 mM ammonium sulfate and then covering the drop with 50 μl of mineral oil. Crystals appeared within 2 days and grew to ~100 μm in a week. Crystals of ∆95scScs7p were also grown at 14 °C using the microbatch-under-oil method by combining 2 μl of protein solution at 2.5 mg/ml with 2 μl of 10% PEG 2000, 100 mM sodium citrate, pH 5.6, 50 mM ammonium sulfate and then covering the drop with 50 μl of mineral oil. Crystals appeared within 2 days and grew to ~300 μm within a week. We noted that scScs7p and ∆95scScs7p crystals deteriorated over time in the crystallization trays. Thus, crystals were harvested after 1 week of growth and frozen in liquid nitrogen for diffraction analysis. For cryopreservation, crystals were harvested into the crystallization mixtures described above that also included 20% ethylene glycol. To generate the heavy atom derivative of scScs7p, crystals were harvested from drops and soaked in a solution of 11% PEG 3350, 50 mM Hepes, pH 7.0, 150 mM ammonium sulfate, 75 mM NaCl, and 1% βOG (w/v), which was saturated with K₂HgI₄ for 25 min. The crystals were then back-soaked into 11% PEG 3350, 50 mM Hepes, pH 7.0, 150 mM ammonium sulfate, 75 mM NaCl, 1% βOG (w/v), and 20% ethylene glycol for 2 min, followed by freezing in liquid nitrogen.

Structure Solution and Refinement—X-ray diffraction data were collected on beamline 17-ID at the Advanced Photon Source (Argonne National Laboratory) using a Pilatus 6 M detector and processed with HKL2000 (25). Data collection statistics are reported in Table 1. Single isomorphous replacement with anomalous scattering (SIRAS) phasing was utilized to solve the structure of scScs7p. PHENIX.autosol (26) located the positions of eight mercury atoms within the asymmetric unit using data to 6 Å resolution. These initial phases were then used in conjunction with the native scScs7p dataset to further improve the phases and extend the data to 3.8 Å in PHENIX. At this stage, the experimental map exhibited clear molecular boundaries for the two molecules of scScs7p in the asymmetric unit. Manual model building was carried out using COOT (27). The mercury atoms were found to interact with the cysteine residues in the protein, which aided in building a model with proper amino acid register. An x-ray fluorescence scan revealed a 2:1 ratio of zinc to iron in the crystals (data not shown). We therefore modeled two zinc atoms in the structure instead of the iron atoms that were expected to be present based on previous biochemical studies on enzymes of the hydroxylase/desaturase family. No electron density was observed for the N-terminal 116 residues corresponding to the cytochrome b₅ domain and linker. The cytochrome b₅ domain is present in the crystals based both on their brown color and on a characteristic signal present at the iron-edge in an x-ray fluorescence scan. We subsequently carried out molecular replacement with PHASER (28) utilizing a cytochrome b₅ search model, but we were unsuccessful in locating this domain. Thus, we conclude that the N-terminal cytochrome b₅ domain is disordered within scScs7p crystals. The final model was refined in PHENIX.refine to R and Rfree values of 24.6 and 27.1%, respectively, utilizing NCS restraints, individual B-factor, and TLS refinement. The structure of ∆95scScs7p was solved to 2.6 Å resolution using single-wavelength anomalous dispersion methods at the zinc edge to generate initial phases. Two molecules were found in the asymmetric unit, corresponding to scScs7p monomers composed of residues 117–384 and residues SGLEV from the C-terminal affinity tag. The final model was refined to R and Rfree values of 19.7 and 23.9%, respectively utilizing individual B-factor and TLS refine-ment (Table 1). The two monomers were virtually identical, with a root mean square deviation between monomers of 0.33 Å (268 Ca atoms).

Yeast Viability Assay—S. cerevisiae exhibits sensitivity toward the antifungal cyclic lipodepsinonapeptide syringomycin E, produced by Pseudomonas syringae pv. syringae (29). We replaced SC57 with the hygromycin B resistance marker to generate an scs7Δ strain in BY4742 (MATa, his3Δ1, leu2Δ0, ura3Δ2, trp1Δ1, can1Δ1, ade2Δ1, gal1Δ3Δ4, lys2Δ800, 300 mM sodium citrate, pH 5.6, 50 mM ammonium sulfate, followed by covering the drop with 50 μl of mineral oil. Crystals appeared within 2 days and grew to ~100 μm in a week. Crystals of ∆95scScs7p were also grown at 14 °C using the microbatch-under-oil method by combining 2 μl of protein solution at 2.5 mg/ml with 2 μl of 10% PEG 2000, 100 mM sodium citrate, pH 5.6, 50 mM ammonium sulfate, followed by covering the drop with 50 μl of mineral oil. ∆95scScs7p crystals grew to a size of ~300 μm within a week. We noted that scScs7p and ∆95scScs7p crystals deteriorated over time in the crystallization trays. Thus, crystals were harvested after 1 week of growth and frozen in liquid nitrogen for diffraction analysis. For cryopreservation, crystals were harvested into the crystallization mixtures described above that also included 20% ethylene glycol. To generate the heavy atom derivative of scScs7p, crystals were harvested from drops and soaked in a solution of 11% PEG 3350, 50 mM Hepes, pH 7.0, 150 mM ammonium sulfate, 75 mM NaCl, and 1% βOG (w/v), which was saturated with K₂HgI₄ for 25 min. The crystals were then back-soaked into 11% PEG 3350, 50 mM Hepes, pH 7.0, 150 mM ammonium sulfate, 75 mM NaCl, 1% βOG (w/v), and 20% ethylene glycol for 2 min, followed by freezing in liquid nitrogen.

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lys2Δ0, ura3Δ0). The scs7Δ strain was used to assay the function of constructs used for crystallization and to analyze the effect of point mutations on scs7p activity. Genes encoding both the scs7p-ZZ-His10 and Δ95scs7p-ZZ-His10 crystallization constructs were cloned into the BamHI and EcoRI sites of vector pRS316-ADH1 (30). The variants H173A, H244A, H249A, H264A, H268A, H271A, H272A, Y319A, Y322A, D323A, H326A, H330A, H331A, H345A, H348A, and H349A were created using the QuickChange mutagenesis kit with wild-type scs7p in pRS316-ADH1 as the template. All constructs were confirmed by DNA sequencing. Both wild-type and mutant scs7p constructs were transformed into the scs7Δ strain. To test for growth on syringomycin E, yeast cells were grown in SD-Ura media overnight at 30 °C to an A\text{600} of ~1.0. The cells were subsequently diluted to 0.3 A\text{600} units, and 2 μl of each dilution was spotted onto SD-Ura plates and SD-Ura plates containing 0.5 μg/ml syringomycin E. The plates were incubated at 30 °C for 2 days and analyzed for growth. For each mutant, at least two colonies from independent transformations were chosen to ensure that they behaved similarly on plates containing syringomycin E. The expression of all mutant constructs was confirmed via Western blot analysis using the protocol outlined previously (31) (data not shown). A total of 15 μl of each mutant construct was run out on a 4–20% SDS-polyacrylamide gel, followed by transfer to a PVDF membrane and probing with anti-mouse IgG. Western blot analysis was performed three independent times to confirm expression of all the mutant constructs.

**Modeling of the Enzyme-Substrate Complex**—To gain insight into how substrate binds to scs7p, molecular dynamics was performed with GROMACS (32) utilizing the CHARMM36 force field (33) and SPCE water model. Topology and parameter files for the phytoceramide substrate were generated with SwissParam (34). The model of Δ95scs7p was inserted into a pre-equilibrated lipid bilayer comprised of 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine using the CHARMM-GUI (35) and solvated in an 11 × 11 × 10-nm³ rectangular box. Charge was neutralized by replacing seven water molecules with seven chloride ions, followed by energy minimization utilizing a steepest decent protocol. The energy-minimized system was then subjected to a 100-ns NVT equilibration using a modified Berendsen thermostat at a temperature of 310 K, followed by a 1-ns NPT equilibration using the Parrinello-Rahman barostat. Molecular dynamics simulations for Δ95scs7p within the 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine bilayer were performed until a stable structure with the proper metal coordination was identified. The stabilized Δ95scs7p structure was then utilized to generate the enzyme-substrate complex. Phytoceramide, containing a 26-carbon VLCFA moiety, was used as the model substrate based on the findings of Beeler and co-workers (11). The substrate was modeled in two different orientations, with the VLCFA moiety placed in the upper channel and the LCB moiety in the lower channel and vice versa. Both ligand-bound models were subsequently equilibrated at 310 K prior to running a final 100-ns MD simulation. All simulations were carried out on the cluster at the Center for Computational Resources at the University at Buffalo.

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**Results**—We cloned Scs7p orthologs from nine different yeast species and subsequently expressed these constructs in *S. cerevisiae* to identify protein constructs that exhibited enhanced stability and suitability for crystallization (21). Purified protein from the *S. cerevisiae* Scs7p (scs7p) ortholog was readily crystallized. The sequence of scs7p shares 36% sequence identity with *Homo sapiens* FA2H (Fig. 1). Crystallization utilizing the microbatch-under-oil method generated brown-colored crystals of scs7p that diffracted to 3.8 Å resolution. Although residues 117–384, corresponding to the hydroxylase domain, were successfully built into the electron density map, there was no interpretable electron density for the N-terminal 116 residues. This region, corresponding to the heme-containing cytochrome *b*₅ domain and linker peptide, is disordered within the crystals. To improve resolution, we engineered a hydroxylase domain construct with the N-terminal cytochrome *b*₅ domain removed (Δ95scs7p). Crystals of the Δ95scs7p construct were colorless, confirming the absence of the heme-containing cytochrome *b*₅ domain, and diffracted to 2.6 Å resolution. The resulting electron density map was significantly higher in quality than that of the full-length construct, allowing for a more accurate model of the hydroxylase domain to be built.

The structure of the hydroxylase domain of scs7p is composed of 13 helical secondary structural elements resembling a mushroom in overall shape (Fig. 2). The four predicted transmembrane helices (TM1–TM4) form the stalk, and the other nine helices (H1–H9) form the catalytic cap of the mushroom. TM2 and TM4 contain 27 and 26 residues, respectively, whereas TM1 and TM3 are shorter and comprise 21 and 18 residues. Short 3-residue loops connect TM1/TM2 and TM3/TM4 on the luminal side of the ER membrane. Consistent with previous structural predictions for the membrane-bound members of the hydroxylase/desaturase family, the N and C termini are located on the cytosolic side of the ER membrane (13).

The catalytic cap domain is defined by three stretches of residues according to their location with respect to the transmembrane helices. The N-terminal stretch comprises 91 residues (Phe-117 through Ala-197) prior to TM1. There is a short stretch of 38 residues (Phe-248 through Pro-284) between TM2 and TM3, followed by a 54-residue stretch (His-331 through Glu-384) beyond TM4 at the C terminus. The helices within the catalytic cap domain can be divided into two layers. Helices H1, H2, and H4 are located the apex of the catalytic cap domain, whereas helices H3, H5, H6, H7, H8, and H9 lie closer to cytosolic face of the membrane. Moreover, helices H3, H5, H7, and
H9 are amphipathic in nature, and their observed orientation suggests that they provide additional stabilization of the catalytic cap domain on the surface of the membrane. Indeed, residues within helices H5, H7, and H9 form stabilizing interactions with detergent molecules utilized to stabilize the protein during purification and crystallization (Fig. 2C).

**Dimetal Center—** Despite the fact that Scs7p and FA2H are classified as di-iron enzymes within the hydroxylase/desaturase superfamily, their structures have not been extensively studied. The secondary structure for the hydroxylase domain is displayed above the sequence alignment. Helices from the catalytic cap domain are colored red, and transmembrane helices are colored purple. Helix segments within helices are depicted in light blue. Completely conserved residues within the alignment are shaded green, and highly conserved residues are shaded in lime. Histidine box motifs are outlined in blue with the conserved histidines shaded in red. Residues lining the substrate binding channels are denoted with a blue diamond.
Figure 2. Structure of scScs7p. Schematic representations of the secondary structural elements of the scScs7p hydroxylase domain placed within a model lipid bilayer (gray lines) are depicted. Transmembrane helices TM1–TM4 in the stalk domain are colored in gray, and the helices at the apex of the catalytic cap domain (H1, H2, and H4) are colored green. The remaining helices (H3, H5, H6, H7, H8, and H9) are colored blue. The two zinc atoms within the dimetal center are colored magenta. 2Foc – F, electron density (green) for unidentifed lipid (or βOG), carried along from the expression and purification process, is shown bound in the lower and upper channels of the enzyme for monomer A (A) and monomer B (B). C, view of A and B rotated 180° depicting βOG detergent molecules bound near the amphipathic helices at the proposed membrane interface. 2Foc – F, electron density contoured at 1σ for βOG is shown in green. Unidentified lipid and βOG molecules are depicted as sticks, with carbon and oxygen atoms colored yellow and red, respectively.

Superfamily (13), we routinely observed the presence of zinc atoms in purified enzyme preparations and within crystals of scScs7p. X-ray fluorescence measurements, coupled with diffraction data collection at the zinc absorption edge yielded anomalous difference peaks that unambiguously identified the metal atoms in the crystal as zinc. Two zinc atoms were modeled within the core of the catalytic cap domain (Fig. 3A). These reside 6.25 Å apart and are coordinated by the side chains of 10 highly conserved histidine residues (Fig. 1). Based on its depth and spatial location with respect to the resolved detergent molecules in the structure, the dimetal-binding site is proposed to lie at the surface of the cytosolic side of the ER membrane. The zinc-coordinating histidine residues are located in four motifs, defined as Ia, Ib, Ia, and Iib (11). Motifs Ia and Ia are located at the C-terminal ends of TM2 and TM4, respectively, whereas motifs Ib and Iib are located in H7 and H8 within the catalytic cap domain (Fig. 1). Each zinc atom is coordinated to the Ne2 atoms of five histidine side chains, with coordination exhibiting square pyramidal geometry as follows: zinc1 with His-244 (TM2), His-249 (TM2), His-268 (H7), His-319 (H8), and His-348 (H8); and zinc2 with His-271 (H7), His-326 (TM4), His-330 (TM4), His-345 (H8), and His-349 (H8). The average Zn-Ne2 distance is 2.20 Å (Fig. 3B).

In vitro assay of the hydroxylase activity of scScs7p was not pursued because of the difficulties associated with reconstituting the necessary electron transport chain and the extremely hydrophobic nature of the ceramide substrate. We therefore turned to a phenotypic assay of mutant SCS7 alleles in whole yeast cells as a means to evaluate the roles of various residues in scScs7p function. S. cerevisiae exhibits sensitivity toward the antifungal cyclic lipopeptidopeptide syringomycin E, produced by P. syringae pv. syringae (29). We generated an scs7Δ strain and evaluated point mutations by transforming the deletion strain with a multicopy plasmid containing a normal or mutated copy of SCS7 expressed under control of the ADH1 promoter. Cells expressing Scs7p with normal activity from this plasmid do not grow on plates containing syringomycin E, i.e. syringomycin E sensitivity is observed. Mutant constructs that affect hydroxylase activity lose their sensitivity to syringomycin E and thus grow on plates containing the lipopeptide. Mutation of each of the 10 zinc-coordinating histidines to alanine resulted in the reduction or complete loss of this sensitivity on plates containing syringomycin E (Fig. 3, C and D), reinforcing the important role that these histidine residues play with respect to the catalytic activity of the enzymes of the hydroxylase/desaturase superfamily (12).

Putative Substrate Channels—Electron density corresponding to the hydrocarbon tail of an unidentified lipid molecule or a βOG detergent molecule is observed within two orthogonal channels that intersect just below the dimetal center (Fig. 2). The lower channel runs perpendicular to the lipid bilayer and is formed by transmembrane helices TM2, TM3, and TM4. The upper channel runs parallel to the lipid bilayer and is formed by amphipathic helices H5 and H7, along with the N-terminal portion of TM3. The lower channel is lined with hydrophobic residues, including Trp-236 (TM2), Phe-248 (TM2), Phe-288 (TM3), Leu-291 (TM3), Phe-295 (TM3), Val-299 (TM3), and Phe-312 (TM4). Surprisingly, there are a significant number of residues that are hydrophilic or have side chains with the potential to form ionic or hydrogen bonds within the lower channel. These include His-244 (TM2), Tyr-219 (TM4), Tyr-322 (TM4), and Asp-323 (TM4), which are all highly conserved in membrane-bound hydroxylase enzymes across species (Fig. 1). His-244 and Phe-248 are part of histidine motif Ia. Conversely, the upper channel is lined solely by hydrophobic residues Leu-191 (H5), Leu-266 (H7), Leu-267 (H7), Met-283 (TM3), Pro-284 (TM3), and Leu-287 (TM3).

A pocket is formed at the intersection of the two channels, directly below the dimetal center. The unique secondary structure observed for helix H7, which harbors histidine motif Ib, plays a vital role in the formation of this pocket, which could accommodate the hydrophilic headgroup of a ceramide sub-
strate. Helix H7 is kinked by 70° at Leu-267, due to the insertion of one turn of the H9266-helix. As a result, the histidine side chains in motif Ib are properly aligned near the dimetal center in an appropriate configuration for coordination. This 70° kink presumably provides access for the substrate headgroup to the dimetal center for initiation of the hydroxylase reaction. Interestingly, there is no significant sequence identity in the region of H7 outside of the Scs7p/FA2H family of enzymes, suggesting that the observed secondary structure for H7 is unique to both the substrate specificity and function of hydroxylation (Fig. 1).

N-terminal Cytochrome b₅ Domain Interaction—The hydroxylase reaction proceeds upon the transfer of an electron to the dimetal center via the fused N-terminal cytochrome b₅ domain. Electron transfer is thought to occur through an interaction that involves an anionic surface on cytochrome b₅ composed of the carboxylate side chains of conserved glutamic acid residues and the heme moiety (40). The interaction surface for the cytochrome b₅ domain on the hydroxylase domain of scScs7p is unknown. Although the N-terminal cytochrome b₅ domain is present in crystals of full-length scScs7p, this domain is disordered in the structure, suggesting that the interaction between the two domains is transient in nature. We constructed point mutants containing the substitutions H173A, H264A, and H331A at sites that are located outside of the histidine box motifs to gain insight into which surface of the hydroxylase domain of Scs7p might interact with cytochrome b₅. The side chains of all three histidines are surface accessible, with His-173 and His-264 located on one side of the dimetal center and His-331 on the opposite side (Fig. 3D). Only the H173A mutation resulted in the loss of syringomycin E sensi-
tivity, suggesting its involvement in electron transfer from cytochrome b5 (Fig. 3C). His-173 is highly conserved among the hydroxylase enzymes within this family (Fig. 1).

Comparison of Hydroxylase and Desaturase Scaffolds—There are only 21 residues within the 268-residue hydroxylase domain of scScs7p that are identical compared with SCD1, a desaturase representative within the hydroxylase/desaturase superfamily (19, 20). Despite the less than 8% sequence identity, scScs7p and SCD1 exhibit a similar tertiary scaffold, predominantly consisting of helical secondary structure. Both enzymes are composed of a cytosolic catalytic cap domain that sits atop a stalk of four transmembrane helices, with the N- and C-terminal ends of the enzymes located on the cytosolic side of the ER membrane (Fig. 4). The residues involved in metal coordination at the dimetal centers of scScs7p and SCD1 share the highest sequence and structural conservation between enzymes (Fig. 5). Both structures have zinc atoms bound within their dimetal centers, despite the utilization of different expression systems in the generation of the proteins used for structural studies (19, 20). The zinc atoms are similarly spaced ~6.5 Å apart from each other. Nine of the 21 residues that are conserved between the enzymes are zinc-coordinating histidine residues located in the histidine box motifs. Zinc is coordinated in virtually the same manner in both structures, exhibiting square pyramidal geometry.

Although the global architecture is similar between enzymes, the root mean square deviation between scScs7p and SCD1 is strikingly poor (3.9 Å for 138 equivalent Ca atoms). In the SCD1 structure, the transmembrane helices are splayed apart at the cytosolic face of the ER membrane, with the catalytic cap domain of the enzyme nestled within. In contrast, the transmembrane helices of scScs7p exhibit a closed conformation, attributable primarily to differences in helical geometry observed in TM2 and TM3 between the structures. There is a single turn of 3-helix in the last turn of TM2 in scScs7p that is lacking in the SCD1 structure, resulting in a kink that positions TM2 in closer proximity to the N-terminal end of TM3. TM3 in SCD1 is kinked due to the presence of Pro-226, whereas there is no kink in TM3 in the scScs7p structure (Fig. 4). A hydrogen bond formed between the carbonyl oxygen of Pro-226 and the side chain of Arg-249 in TM4 is hypothesized to stabilize the kink within the membrane interior of SCD1 (19). Neither Pro-
226 nor Arg-249 are conserved in the membrane-bound hydroxylase enzymes (Fig. 1).

There are also significant differences between scScs7p and SCD1 in the catalytic cap domain. The catalytic cap domain of scScs7p is composed of nine helices within three stretches of 91, 38, and 54 residues (Fig. 4). The N-terminal stretch contains five helices, whereas the central and C-terminal stretches each contain two helices. Four of the nine helices in the catalytic cap domain, H3, H5, H7, and H9 are amphipathic in nature. In contrast, only the 93-residue central and 90-residue C-terminal stretches of SCD1 contain helices, of which three, H1, H7, and H9, are amphipathic (Fig. 4). The different locations of the helices within the catalytic caps result in vastly different domain architectures for each enzyme. For SCD1, the two domains, C1 and C2, lie on opposite sides of the dimetal center. It is hypothesized that the cytochrome b5 domain accesses the dimetal center for subsequent electron transfer via a groove between C1 and C2 (19). In contrast, the helices in the N- and C-terminal stretches of scScs7p serve to bury helices H6 and H7 in the central stretch, suggesting an alternative binding mode for cytochrome b5 and mechanism for electron transfer (Fig. 4).

The major difference between the dimetal centers of scScs7p and SCD1 is the location of these groups with respect to the cytosolic membrane. In SCD1, TM2 and TM4 extend three helical turns out of the membrane and form interactions with residues within the catalytic cap domain (19, 20). Zinc-coordinating histidine residues His-120, His-125, and His-269 in SCD1 are contained within the portions of TM2 and TM4 that extend out of the membrane. As a result, the dimetal center is located ~8.5 Å above the plane of the catalytic cap. In stark contrast, the dimetal center of scScs7p resides only ~2.8 Å above the plane of the cytosolic membrane (Fig. 4). The two factors responsible for the difference in location of the dimetal center in scScs7p are the presence of the π-helix turn in TM2 that bends the end of the helix closer to the membrane and the increased number of amphipathic helices in the catalytic cap that stabilize this domain closer to the membrane interface. The observed distance differences in the location of the dimetal center with respect to the membrane may play a role in the different catalytic activities carried out by each enzyme as well as their substrate specificities.

The serendipitous capture of the steroyl-CoA substrate in SCD1 crystal structures provides insight into the similarities and differences in substrate binding between the hydroxylase and desaturase enzymes. In SCD1, the CoA moiety binds to the catalytic cap domain, between helices H2 and H4 and near the end of TM4 (19, 20). The adenosine, diphosphate, and pantetheine groups of the CoA moiety form electrostatic and hydrogen bonds with polar residues that include Asn-75, Asn-148, Arg-155, Asp-156, Arg-188, Lys-189, and Lys-194. Collectively, this highly conserved cluster of residues forms an electrostatic surface within the catalytic cap domain of SCD1 that aids in substrate recognition and binding (19, 20). There is no equivalent electrostatic patch on the surface of the catalytic cap in scScs7p, as the corresponding residues in scScs7p for Asn-148, Arg-155, and Asp-156 are hydrophobic residues Ile-259, Leu-266, and Leu-267 (Fig. 4). The lack of conservation of these residues, coupled with the significantly different architectures observed for the catalytic cap domains of SCD1 and scScs7p, suggests that fatty acyl-CoA substrates are not hydroxylated by scScs7p.

The fatty acyl tail of steroyl-CoA binds within a kinked hydrophobic tunnel in SCD1 that is composed of residues from TM2 and TM4. The kink in the tunnel is generated by the positioning of the side chains of Trp-149 and Thr-257, two highly conserved residues in the desaturase branch of the hydroxylase/desaturase family. In addition to forming a kink, these residues cap the top of the tunnel. As a consequence, the shape of the tunnel forces the acyl chain to bind in a conformation poised for desaturation at the carbon 9–10 bond. Tyr-104, also highly conserved among desaturase enzymes, is located at the base of the tunnel, where it serves as a plug that restricts fatty acyl chains with more than 18 carbons from binding within the tunnel (19, 20). Although we do not observe bound substrate in scScs7p, there is an equivalent channel lined by residues located within TM2, TM3, and TM4 that is occupied by a lipid mole-
cule. The channel in scScs7p is more open below the dimetal center as there are no residues equivalent to Trp-149 and Thr-257. Moreover, the equivalent residue to Tyr-104 in scScs7p is Gly-232, which would not be expected to impose any restriction on the length of fatty acyl chain that can bind within the channel. The implications that these differences between SCD1 and scScs7p have for substrate specificity, recognition, and binding are discussed below.

**Modeling Ceramide Binding to scScs7p and Implications for Catalysis**—We carried out molecular dynamics simulations to gain further insight into how a ceramide substrate could bind to scScs7p. Prior to the simulations, models were created that placed the VLCFA moiety in the lower channel bound by TM2, TM3, and TM4 and LCB moiety of the ceramide in the upper channel bound by H5, H7, and TM3 and vice versa. Three different starting positions, each of which differed by a shift of the substrate headgroup by a few angstroms with respect to the dimetal center in a direction perpendicular to the membrane, were utilized for both binding modes. The molecular dynamics simulations resulted in ending positions for the ceramide substrate that were very similar to the starting positions. In each case, the VLCFA and LCB moieties remained in their respective channels, and the headgroup was located below the dimetal center. There were no significant changes observed in the scScs7p structure over the course of the simulations. Models that placed the VLCFA moiety in the upper channel resulted in the positioning of C-2 of the VLCFA far away from the dimetal center, with distances from C-2 to the zinc atoms of 9.0 and 9.2 Å. These long distances do not favor hydroxylation and thus the binding pose that places the VLCFA moiety in the upper channel is unlikely to be the catalytically competent conformation for the substrate. When the VLCFA moiety was placed in the lower channel and the LCB moiety in the upper channel, the simulations placed C-2 much closer to the dimetal center, with distances of 5.2 and 5.9 Å (Fig. 6). In this orientation, the pro-(R) hydrogen of C-2 lies 4.9 and 5.3 Å away from the zinc atoms, consistent with the C-9/C-10 distances to the dimetal center for steryl-CoA bound to SCD1 (4.7 and 5.4 Å) (19, 20) and with the production of (R)-enantiomers by FA2H (41).

The highly conserved residues Tyr-319, Tyr-322, and Asp-323 are located at the C-terminal end of TM4 (Fig. 1), where they introduce functional groups that could participate in the stabilization of the ceramide headgroup through interaction with the hydroxyl groups and amine group. In our model, the majority of the interactions made between the ceramide headgroup and scScs7p are hydrogen bonds to protein main chain nitrogen and carbonyl atoms. Interestingly, the phenolic oxygen of Tyr-322 is positioned 4 Å away from the nitrogen atom in the ceramide headgroup. Mutation of Tyr-322 to alanine results in the loss of syringomycin E sensitivity (Fig. 3). Because the side chain of Asp-323 is not predicted to interact with the ceramide substrate, it may be that this residue is involved in water egress during the hydroxylase reaction.

**Discussion**

We report here the x-ray crystal structure of the integral membrane enzyme sphingolipid α-hydroxylase Scs7p, a representative of the hydroxylase branch of the hydroxylase/desaturase superfamily. Scs7p is the yeast ortholog of human FA2H, which is responsible for hydroxylating GalCer and sulfatide within the myelin sheath. The enzyme adopts a mushroom-like shape similar to the overall fold of the related steryl-CoA desaturase, but with significant differences from these structures relevant to the differing substrate specificities and catalytic action of the two enzymes.

We routinely observed the presence of zinc atoms in purified enzyme preparations and within crystals of scScs7p, despite the fact that scScs7p is classified as a di-iron enzyme. We made significant attempts to increase the iron content within the enzyme, including reduction of the zinc content in the expression media, stripping out zinc with subsequent addition of iron during purification, and manipulation of the yeast expression conditions to favor iron incorporation over zinc (42). Although these attempts led to the production of highly purified enzyme with iron incorporated at the dimetal center up to ~50%, we were unsuccessful at achieving 100% replacement with iron. Moreover, the incorporation of zinc into scScs7p is not a result of the use of S. cerevisiae as an overexpression system, as zinc atoms were also observed bound to scScs7p when we utilized an insect cell expression system. Our results are consistent with the observation of zinc bound within the dimetal centers of the structures of human and mouse SCD1, both of which were determined utilizing protein derived from an insect cell expression system (19, 20). It is quite possible that zinc simply serves
as a suitable surrogate for iron during overexpression, given their similar size and charge properties (19). In addition to the structures of SCD1 and that of scScs7p presented here, there are other examples of zinc replacement at iron-binding sites, including the structure of quinol-dependent nitric-oxide reductase (43).

A surprising aspect of the studies reported here is that the Δ95scScs7p crystallization construct exhibits syringomycin E
sensitivity, indicative of hydroxylase function. This indicates that the electron transfer to the truncation mutant must be provided by some endogenous source of cytochrome b₅ (Fig. 3) (14). Indeed, the Saccharomyces castellii and Candida glabrata Scs7p orthologs do not contain the His-72 residue in their cytochrome b₅ domains, which is required for heme binding (Fig. 1) (44). In contrast to these observations for scScs7p, the fused cytochrome b₅ domain is required for activity of other family members such as yeast Δ9 desaturase (Ole1p), which when similarly truncated is not functional in cells, implying that it cannot be rescued by an alternative cytochrome b₅ (45). Similarly, in humans, FA2H containing point mutations in the cytochrome b₅ domain causes mild forms of hereditary leukodystrophies (45). It is not clear what evolutionary advantage is attained through the fusing of a cytochrome b₅ domain to a hydroxylase or desaturase domain. One hypothesis is that the fusion provides a kinetic advantage for enzyme function (44).

Although it is clear that FA2H and Scs7p hydroxylate C-2 of a VLCFA moiety, the breadth of the substrate specificity of these enzymes has not been definitively established. Candidate substrates include free fatty acids, fatty acyl-CoA derived substrates, and ceramide. Establishing substrate specificities is complicated by the difficulties associated with reconstituting the electron transport chain for in vitro assays and by the hydrophobic nature of the substrates. Lipid analysis of gene deletions in S. cerevisiae suggests that phytoceramide is the preferred substrate for Scs7p (15), and these findings are further supported by experiments carried out with rat brain microsomes, which indicate that free fatty acid substrates and fatty acyl-CoA-derived substrates are not effectively modified (46). The structural architecture observed in scScs7p provides strong evidence that ceramide is an optimal substrate for the enzyme. The elongated shape and open pocket below the dimetal center provide suitable space for the binding of a substrate that contains both an LCB and VLCFA moiety. The lack of a polar electrostatic patch in scScs7p and the different structural architectures observed between the catalytic cap domains of scScs7p and SCD1 would seem to rule out the binding of a fatty acyl-CoA derived substrate.

Molecular dynamics simulations suggest that ceramide binds optimally to scScs7p with the VLCFA moiety located in the lower channel and the LC moiety in the upper channel. Detergent or lipid molecules are bound in both of these channels in the scScs7p crystal structure, further indicating that these channels bind the VLCFA and LC moieties of ceramide. Additional support for the location of VLCFA in the lower channel comes from inspection of the equivalent hydrophobic channel in SCD1, responsible for binding the fatty acyl tail of steryl-CoA. The SCD1 hydrophobic channel is plugged at its base by Tyr-104. Interestingly, Tyr-104 is known to be responsible for limiting the substrate specificity for SCD1 to substrates containing fatty acyl chain lengths between 14 and 19 carbons (19). Although Tyr-104 is highly conserved in the desaturase branch of the hydroxylase/desaturase family, the marine copepod desaturase from Calanus hyperboreus has a threonine at this position, which allows it to desaturate substrates containing fatty acyl chains with lengths between 22 and 26 carbons (47). No such plug is present in the lower channel in the scScs7p structure, where the equivalent residue to Tyr-104 is Gly-232, suggesting that a VLCFA moiety can be accommodated. Moreover, Gly-232 is highly conserved in the hydroxylase branch of the hydroxylase/desaturase superfamily (Fig. 1).

Occupation of the lower and upper channels by the VLCFA and LCB moieties, respectively, places the C-2 of the VLCFA below the dimetal center in an orientation that is poised for hydroxylation, with distances to the zinc atoms of 5.2 and 5.9 Å. These distances are consistent with the spacing observed between the metal atoms and C-9 and C-10 for steryl-CoA bound to SCD1 (19, 20). Tyr-319, Tyr-322, and Asp-323, highly conserved residues within the hydroxylase branch of the hydroxylase/desaturase family, are located near the substrate headgroup. Both Y322A and D323A mutant constructs resulted in the loss of syringomycin E sensitivity in the cellular activity assay, suggesting a role for each in catalysis. Tyr-322 is positioned 4 Å away from the nitrogen atom in the ceramide headgroup. The equivalent residue in SCD1 is Trp-262, which forms a hydrogen bond with the carbonyl oxygen of the fatty acyl chain within the steryl-CoA substrate. Given both the observed loss of syringomycin E sensitivity and its location with respect to the substrate, we hypothesize that Tyr-322 forms a similar interaction with the ceramide headgroup. Tyr-319 and Asp-323 interact as part of the helical geometry of TM4. The observed interaction between Tyr-319 and Asp-323 could play a role in stabilizing the turn of helix in TM4 and subsequently the position of Tyr-322 for optimal interaction with the substrate headgroup. We hypothesize that the side chain of Asp-323 plays a role in the egress of water, generated during the hydroxylase reaction, to the aqueous cytosol.

The differences in substrate specificities between SCD1 and scScs7p provide insight into the entry and egress of ceramide into the enzyme for subsequent catalysis. In SCD1, binding of the CoA moiety of the substrate to the polar surface within the catalytic cap domain drives insertion of the fatty acyl tail into the kinked hydrophobic pocket that is capped by residues Trp-149 and Thr-257 (19, 20). The side chains of Trp-149 and Thr-257 are stabilized by hydrogen bonds with Gln-143. Bai et al. (19) propose that breaking the hydrogen bond between Gln-143 and Thr-257 provides an opening between TM4 and the loop between H1 and H2 that would allow the desaturated product to be released from the pocket. The area below the
dimensional center in the scScc7p is not capped as it is in the SCD1 structure and thus is more open. We propose that the predominantly hydrophobic ceramide substrate would access and depart the active site directly from within the confines of the lipid bilayer through an opening between transmembrane helices TM2 and TM3. Direct entry from the lipid bilayer would orient the LCB and VLCPA moieties in the appropriate channels and the headgroup below the dimetal center, analogous to the positioning of the ligand sphingonosine 1-phosphate upon binding to its cognate G protein-coupled receptor (48). The observed difference in proximity of the dimetal centers of scScc7p and SCD1 to the plane of the cytosolic membrane (2.8 Å versus 8. Å) provides additional support for different modes of substrate access and egress for the enzymes.

In summary, the crystal structure of scScc7p presented here provides the first structural snapshot of a hydroxylase enzyme representative of the membrane-bound hydroxylase/desaturase superfamily. The location of the metal-binding histidine boxes with respect to the positions of the four transmembrane helices and the placement of the metal center at the surface of the membrane facing the cytosol confirms previous structural predictions made for scScc7p. Examination of the structure, along with molecular dynamics simulations, suggests that ceramide binds below the dimetal center, with the VLCPA placed in the lower channel and the LCB moiety placed in the upper channel. Furthermore, results of cell-based activity assays suggest roles for highly conserved residues Tyr-322 and Asp-323 of channel. Furthermore, results of cell-based activity assays suggest roles for highly conserved residues Tyr-322 and Asp-323 of channel. Furthermore, results of cell-based activity assays suggest roles for highly conserved residues Tyr-322 and Asp-323 of channel. Furthermore, results of cell-based activity assays suggest roles for highly conserved residues Tyr-322 and Asp-323 of channel.

Author Contributions—M. G. M. conceived and coordinated the study and wrote the paper. M. E. D., S. M. C., and G. Z. were responsible for expression, solubilization, purification, and functional analysis. All authors reviewed the results and approved the final version of the manuscript.

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