Characterization of the Human Sigma-1 Receptor Chaperone Domain Structure and Binding Immunoglobulin Protein (BiP) Interactions

Jose Luis Ortega-Roldan, Felipe Ossa, and Jason R. Schnell

From the Department of Biochemistry, University of Oxford, Oxford OX1 3QU, United Kingdom

Background: Sigma-1 receptor is a ligand-regulated membrane protein chaperone involved in BiP regulation and the ER stress response.

Results: The chaperone domain of human sigma-1 receptor is mostly helical with short extended regions.

Conclusion: Regions of the sigma-1 receptor chaperone domain implicated in ligand and cholesterol binding can be mapped to separate helices.

Significance: A structural framework for delineating sigma-1 receptor BiP and ligand interactions is presented.

The sigma-1 receptor (S1R) is a ligand-regulated membrane protein chaperone involved in the ER stress response. S1R activity is implicated in diseases of the central nervous system including amnesia, schizophrenia, depression, Alzheimer disease, and addiction. S1R has been shown previously to regulate the Hsp70 binding immunoglobulin protein (BiP) and the inositol triphosphate receptor calcium channel through a C-terminal domain. We have developed methods for bacterial expression and reconstitution of the chaperone domain of human S1R into detergent micelles that enable its study by solution NMR spectroscopy. The chaperone domain is found to contain a helix at the N terminus followed by a largely dynamic region and a structured, helical C-terminal region that encompasses a membrane associated domain containing four helices. The helical region at residues 198–206 is strongly amphipathic and proposed to anchor the chaperone domain to micelles and membranes. Three of the helices in the C-terminal region closely correspond to previously identified cholesterol and drug recognition sites. In addition, it is shown that the chaperone domain interacts with full-length BiP or the isolated nucleotide binding domain of BiP, but not the substrate binding domain, suggesting that the nucleotide binding domain is sufficient for S1R interactions.

The sigma-1 receptor (S1R) is a ligand-regulated membrane protein chaperone involved in the ER stress response and interorganelle communication (1–3). S1R is localized to mitochondria-associated ER membranes (4, 5), which are sites for regulation of mitochondrial bioenergetics via ER calcium release (6). S1R is expressed primarily in cerebral cortex, hippocampus, and cerebellum Purkinje cells (7, 8), and has been proposed as a target for treatment of central nervous system diseases, including amnesia, pain, schizophrenia, clinical depression, Alzheimer disease, stroke, and addiction (9, 10).

S1R activity is modulated by N,N-dimethyltryptamine (11), progesterone (12), and sphingosine (13). In addition, S1R is regulated by a large number of exogenous small molecules, including opiates, antipsychotics, antidepressants, antihistamines, phenycyclidine-like compounds, β-adrenergic receptor ligands, and cocaine (reviewed in Ref. 10).

Activated S1R dissociates ankyrin from the inositol triphosphate receptor (3, 14), which results in calcium release at the mitochondria-associated ER membrane that is efficiently taken up by mitochondria to increase energy production. S1R appears also to have other roles in the stress response. ER calcium depletion or agonist binding dissociates S1R from the Hsp70 protein BiP, resulting in activation of protein chaperoning activity in both BiP and S1R (1). Chaperone activity and BiP binding (1) and the interaction of S1R with inositol triphosphate receptor (14) have been localized to the C-terminal domain of S1R, and truncation of the C terminus leads to dysfunction in mitochondrial stress response (15).

The chaperone domain is C-terminal to two putative transmembrane domains (residues 11–29 and 80–100) but contains a predicted membrane associated region (residues 176–204) containing two cholesterol recognition motifs (CRM) (Fig. 1) (16, 17). We have studied the S1R chaperone domain (residues 112–223; S1R(cd)) reconstituted into detergent micelles by solution NMR. S1R(cd) solubilized in dodecylphosphocholine (DPC) adopts a conformation competent to bind BiP in a calcium-dependent manner. Analysis of the structure and dynamics indicates that S1R(cd) contains five helices and at least two short extended regions. Three of the helices in the C-terminal membrane-associated domain correspond closely to regions previously implicated in cholesterol and drug interactions.
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EXPERIMENTAL PROCEDURES

Protein Sample Production—An ACA-free gene construct (GeneArt) containing a (His)₆-tag, a factor Xa cleavage site, and residues 112–223 of human S1R was subcloned into the pCOLD-I vector (Takara) and confirmed by sequencing. The N-terminal sequence preceding residues 112–223 of S1R was MNHKVHHHHHHIEGRHM. The S1R(cd) plasmid and a pMazF plasmid containing the gene for the RNA interferase MazF (Takara) were transformed into C43(DE3) cells. Transformed cells were grown to an OD of 0.8–0.9, cold shocked on ice, and incubated for 45–60 min at 15 °C before induction with 2 mM isopropyl β-D-thiogalactopyranoside (IPTG). The expression proceeded for 16 h at 15 °C. Both membranes and inclusion bodies were collected with a 40,000 rpm spin and incubated overnight in a solution containing 6 mM guanidine, 200 mM NaCl, 1% Triton X-100, and 20 mM Tris, pH 8.0. S1R(cd) was separated by nickel affinity chromatography and dialyzed against water to remove guanidine. The precipitated protein was resolubilized in hexafluoro-2-propanol and dialyzed against NMR buffer containing 20 mM HEPES, pH 6.5. Based on integration of peaks in 1H one-dimensional spectra calibrated against samples with known detergent concentrations, the final amount of DPC in S1R(cd) NMR samples was ~30 mM. Where indicated, certain samples of S1R(cd) were run over gel filtration rather than dialyzed to reduce the DPC concentration to 5 mM. Final protein concentrations for all NMR samples were ~250 μM.

NMR Spectroscopy—All NMR experiments were collected at 37 °C at field strengths of 500, 600, 750, or 950 MHz (1H). The 750 and 950 MHz spectrometers were equipped with room temperature probes (home-built), and the 500 and 600 MHz spectrometers were equipped with cryogen-cooled probes (Bruker). NMR spectra were processed using NMRPipe (20) and analyzed using NMRView or CARA. Backbone 1H, 15N, and 13Cα, and side chain 13Cβ assignments were obtained from triple resonance HNCA, CBCA(CO)NH, HNCCAB, and HNCO spectra on 15N, 13C, and 2H-labeled S1R(cd). Assignments were confirmed and extended to sidechain protons with 15N-edited NOEY (90- or 180-ms mixing times at 950 or 600 MHz, respectively) and 15N-edited TOCSY spectra (40-ms mixing time; 600 MHz). 15N R1, R2, and 1H, 15N heteronuclear NOE experiments were collected on a 0.27 mM 15N-labeled S1R(cd) sample at 600 MHz using the following relaxation delays: 4, 350, and 480 ms (T1) and 0, 40, and 60 ms (T2).

Water-soluble Paramagnetic Studies—Mn2⁺-EDDA2⁻ was prepared as described (21). 1H, 15N HSQCs were recorded before and after addition of 1 mM Mn2⁺-EDDA2⁻. The accessibility to water was taken to be inversely proportional to the ratio of cross-peak intensity in the absence (I) and presence (I₀) of Mn2⁺-EDDA2⁻.

BiP Interactions—BiP titrations were carried out by preparing concentrated stocks of BiP constructs in 100 mM NaCl, 30 mM DPC, 5 mM 2-mercaptoethanol, and 20 mM HEPES, pH 6.5. Isotope-labeled samples of S1R(cd) in identical conditions were titrated with BiP, and 1H, 15N SOFAST-HMQC spectra (22) were collected to monitor changes in cross-peak intensities and chemical shifts.

Circular Dichroism—CD spectra were collected on a Jasco J-815 circular dichroism spectropolarimeter by collecting spectra from 190–250 nm with 10 accumulations (far UV) or 250–350 nm with 10 accumulations. Samples for CD contained 17.5 μM (far UV) and 240 μM (near UV) S1R(cd) in 5 mM DPC and 20 mM potassium phosphate at pH 6.5.

RESULTS

S1R(cd) Sample Production and Structural Overview—An S1R construct (S1R(cd)) containing the residues following the second putative transmembrane domain of S1R was produced with an N-terminal (His)₆-tag using the single protein production approach (23) and purified to homogeneity (Fig. 2). Attempts to reconstitute S1R(cd) into aqueous solution in the absence of lipids or detergent resulted in protein aggregation. Therefore, S1R(cd) was reconstituted in the presence of DPC films of protein and DPC that were solubilized in hexafluoro-2-propanol and dried down under a nitrogen stream (19). The film was resolubilized in urea and reconstituted by dialysis against NMR buffer containing 20 mM HEPES, pH 6.5. Based on integration of peaks in 1H one-dimensional spectra calibrated against samples with known detergent concentrations, the final amount of DPC in S1R(cd) NMR samples was ~30 mM. Where indicated, certain samples of S1R(cd) were run over gel filtration rather than dialyzed to reduce the DPC concentration to 5 mM. Final protein concentrations for all NMR samples were ~250 μM.

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Regions of extended structure could be determined with less confidence than helical regions, although an extended conformation is likely for residues ~145–147 and ~153–155. Chemical shift indices analysis and intense $daN(i + l)$ NOEs suggested the possibility of a third $\beta$-strand in residues ~160–162, but the RCI indicated a high degree of flexibility in these residues.

Backbone Dynamics of S1R(cd)—Backbone amide dynamics of S1R(cd) were probed by measuring $^{15}$N transverse ($R_2$) and longitudinal ($R_1$) relaxation rates and $^1$H–$^{15}$N heteronuclear NOEs (Fig. 6). Small $^{15}$N $R_2$ values and negative or small heteronuclear NOEs were found throughout the region of residues 142–163, consistent with the low amount of structure in this region, although increased heteronuclear NOEs in residues 145–147 correlated with the chemical shift-based prediction that these residues are in a stable extended conformation. In addition, the relaxation properties of some of the interhelical regions at the C terminus exhibited decreased heteronuclear NOEs and $^{15}$N $R_2$ relaxation rates, indicating increased flexibility, particularly in Ile-178 between helices 2 and 3, Val-190 between helices 3 and 4, and Arg-211 between helices 4 and 5.

$R_2/R_1$ ratios correlate with the effective rotational correlation time (24) and are plotted in Fig. 6D for S1R(cd). The average $R_2/R_1$ value for helical residues was 14.0 ± 2.2, which corresponds to a rotational correlation time ($\tau_c$) of ~11.6 ns. This $\tau_c$ is nearly twice that expected for a 14.7-kDa protein at 37 °C (~6.1 ns), suggesting that the protein is tightly associated with a detergent micelle. Although residues in helices 1, 2, 3, and 5 exhibited similar mean $\tau_c$ values (10.9–11.5 ns), residues in helix 4 had a mean $\tau_c$ value of 12.4 ns. Although effects from rotational anisotropy or contributions to the $R_2$ from conformational exchange could not be ruled out, the higher $\tau_c$ value for helix 4 was consistent with this helixanchoring S1R(cd) to the detergent micelle (see below).

Water and Micelle Interactions of S1R(cd)—Based on observation of cross-peaks at the water proton resonance frequency in the NOESY, a large number of backbone amides within the region of residues 140–160 had apparent amide proton chemical exchange with water. Therefore, the rates of backbone amide exchange with water were measured using CLEANEX experiments (Fig. 7A) (25). In the region between helices 1 and 2, residues 137, 139–144, 149–151, and ~155–161 had the largest exchange rates. Exchange rates in the predicted extended regions (residues 145–147 and 153–155) were low, suggesting the presence of $\beta$-sheet structure. Increases in exchange were also observed in the interhelical regions in the C-terminal half of S1R(cd) and helped to assign breaks in the helices.

Low amide proton exchange rates may be due to hydrogen bonding and/or inaccessibility to bulk water. Therefore, we measured line-broadening effects from the water-soluble paramagnetic agent Mn$^{2+}$EDDA$^{-}$−. Broadening of many resonances corresponding to the N-terminal half of S1R(cd) was observed, indicating few continuous regions of protection from water. In contrast, the two segments of residues ~174–184 and ~197–206 showed relatively high levels of protection (i$H_i$/i$H_{N}$ ~0.5), indicating interactions with the detergent micelle and/or other regions of the protein.
FIGURE 3. $^1$H, $^{15}$N HSQC spectrum (600 MHz, $^1$H) of S1R(cd) in 5 mM DPC at 37 °C. Backbone amide resonance assignments are indicated. The positions of the cross-peaks corresponding to residues Arg-114 and Gly-118, which are weak, are indicated with dashed line circles. The five cross-peak pairs expected from asparagine and glutamine H$_2$N(e2) groups of the S1R(cd) construct are indicated with horizontal lines. Inset, the four tryptophan HN(e1) cross-peaks.

FIGURE 4. Circular dichroism spectra of S1R(cd) in 5 mM DPC and 20 mM HEPES, pH 6.5, plotted as the mean residue molar ellipticity ([θ] MR). A, far UV spectrum of S1R(cd) (17.5 μM) at 37 °C. B, near UV spectra of S1R(cd) (240 μM) at 37 °C (solid line) and 95 °C (dashed line).
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Regions of interactions of S1R(cd) with the detergent micelle were assessed from an 15N-labeled NOESY (90 ms) recorded on an 15N-labeled and partially deuterated sample at high field (950 MHz) to resolve cross-peak overlaps and reduce spin diffusion (Fig. 5A). The largest clusters of residues with NOEs to detergent occurred in residues 183–189 in helix 3 and residues 197–204 in helix 4. Helix 4 was also strongly protected from the water-soluble paramagnetic agent. The region ~198–206 forms an amphipathic helix, with residues Leu-199, Phe-200, Leu-203, and Tyr-206 forming the hydrophobic face (26).

S1R(cd) Interactions with BiP—Previous work by Hayashi et al. (1) showed that S1R residues 116–223 are sufficient for binding to the Hsp70 protein BiP. Therefore, the NMR spectrum of 15N-labeled S1R(cd) was monitored as a function of unlabeled BiP concentration (Fig. 8). The S1R(cd) cross-peak intensities decreased in a BiP concentration-dependent manner, suggesting that a BiP-S1R(cd) complex was formed. After several hours, a white precipitate formed, indicating that the complex was unstable in solution. By contrast, BiP alone in the titration buffer (containing 30 mM DPC) was more stable. It was speculated that binding of the S1R(cd) micelle complex to BiP may increase the local concentration of DPC to destabilize BiP.

The changes in 1H-15N cross-peak intensities upon addition of full-length BiP were plotted as a function of residue number at molar ratios of 0.5 and 1 ([BiP]/[S1R(cd)]) (Fig. 8A). Decreases in cross-peak intensities were higher for residues in the structured helical regions and smaller in the disordered regions at the N and C termini and residues ~140–160, suggesting that these disordered residues retained some flexibility in the complex.

BiP chaperone (27) and ATPase (1) activity has been shown to be inhibited by calcium, and the interaction between BiP and S1R is proposed also to be modulated by calcium under physiological conditions, with the presence of calcium leading to increased association and inactivation of S1R and BiP (1). Therefore, a second titration of full-length BiP against S1R(cd) was carried out in the presence of 2.5 mM calcium chloride. A comparison of the cross-peak intensities for helical residues as a function of BiP indicates an increased association of S1R(cd) for full-length BiP in the presence of calcium (Fig. 8B). No chemical shift changes were observed upon addition of calcium to S1R(cd) alone, consistent with a calcium binding site on BiP (18).

To test whether the NBD of BiP was sufficient for S1R binding, the isolated domain was titrated against S1R(cd). Addition of the NBD resulted in similar changes in intensities as full-length BiP (Fig. 8B), indicating that the BiP NBD was sufficient for S1R interactions. The NBD domain was generally less stable than full-length BiP and more unstable in the presence of calcium, preventing evaluation of the effects of calcium on the NBD interaction with S1R(cd). In contrast, no changes in intensities or chemical shifts were observed after addition of the SBD (Fig. 8D).

DISCUSSION

Most membrane-associated proteins remain difficult to express at sufficient levels to enable inexpensive isotope labeling. We have found that the chaperone domain of human S1R could be efficiently expressed in Escherichia coli using the single protein production approach (23). In single protein production, the cells remain metabolically active and express the target protein, but cell growth is halted, which appears to greatly diminish expression toxicity. The approach permitted 10-fold condensation of the cultures when transferring the cells into

FIGURE 5. A, summary of NOE cross-peaks to backbone amide protons. Amide proton to amide proton NOEs (dNN(i+1), dNN(i+2), and dNN(i+3)), as well as cross-peaks at the position of the water proton (dN H2O), were measured in a three-dimensional 15N-edited NOESY with 180-ms mixing time at 600 MHz (1H) on an 15N-labeled S1R(cd) in 5 mM DPC. Amide proton to DPC methyl proton NOEs (dN DPC (CH3)), as well as cross-peaks at the position of the water proton (dN H2O), were measured in a three-dimensional 15N-edited NOESY with 90 ms of mixing time at 950 MHz (1H) on an 15N- and 2H-labeled (~70%) S1R(cd) in 30 mM DPC. The x axis (arbitrary units) is the cross-peak intensity normalized to that of the diagonal. No attempt was made to deconvolute overlapped cross-peaks. B, the chemical shift indices (CSI) of 1H, 15N, 13C, and 13Cβ are shown in the top four panels. In panels five and six are shown the secondary structure and random coil index derived from chemical shift analysis using TALOS+ (49) and the method of Berjanskii et al. (50), respectively. For the TALOS+ plot, values above or below the line indicated that the calculated ω/ψ values correspond to helical (H) or extended conformation (E), respectively. Shown schematically at the top are the S1R(cd) residues determined to be helical (cylinders) or extended (arrow) (see main text for full description of secondary structure assignment).
labeled medium and resulted in yields of 8–10 mg of pure S1R(cd) per liter of labeled medium.

The S1R chaperone domain is predicted to contain a C-terminal membrane-associated region (residues 180–203). Consistent with this prediction, solubilization of S1R(cd) required the presence of lipid or detergent, suggesting that membrane interactions may be required also for proper folding of S1R(cd) in vivo.

For the NMR studies reported here, S1R(cd) was reconstituted from 8 M urea into DPC, which contains a phosphocholine headgroup and has been used extensively for studying folded integral membrane and membrane-associated proteins (28–31). The S1R(cd) samples permitted measurement of high quality NMR spectra, and near UV CD indicated the presence of temperature-sensitive tertiary structure. The finding that the S1R(cd) construct purified from E. coli and reconstituted with DPC micelles interacted with BiP in a calcium-dependent manner indicated that S1R(cd) adopted a native-like conformation in the NMR conditions, at least in respect to those elements required for the BiP interaction.

The N-terminal half of S1R(cd) contains a ~17-residue helix, and at least two short β-strands. The region containing the

![Graphs and data](images)
extended regions, residues ~140–160, is highly dynamic, based on secondary chemical shifts, relaxation measurements, and amide proton exchange rates.

The putative membrane associated C-terminal half of S1R(cd) is largely helical, with four helices separated by short, flexible, water-exposed linkers. By combining information from NOEs to detergent and resonance broadening from a water-soluble paramagnetic agent Mn²⁺ EDDA²⁻. A decreased ratio of the peak intensity after (I) and before (I₀) addition of Mn²⁺ EDDA²⁻ indicates increased water accessibility. Measurement error has been estimated from the spectral noise. Data for resonances that exhibit high water exchange rates as indicated by CLEANEX ratios of I/I₀ > 0.4 (dashed line in A) were excluded from the analysis due to potential artifacts in cross-peak intensities due to changes in relaxation properties.

The chaperone domain contains several regions implicated in cholesterol and drug interactions (Fig. 9). Based on a cholesterol recognition consensus sequence (32, 33) (L/V)X₁₋₅YX₁₋₅(R/K) two cholesterol recognition motifs have been identified within the structured region of S1R(cd) (17, 34, 35). The first motif (CRM1; residues 171–175; amino acid sequence, VEYGR) corresponds approximately to helix 2. The second motif (CRM2; residues 199–208; amino acid sequence, LFYTLRSYAR) contains two overlapping cholesterol recognition motifs (Leu-199/Tyr-201/Arg-204 and Leu-203/Tyr-206/Arg-208), and corresponds to the central region of helix 4. Residues in the vicinity of both
FIGURE 8. S1R(cd) interactions with BiP. A, backbone amide cross-peak intensities are plotted as a function of residue number after addition of 0.5 (black bars) and 1.0 (red bars) molar equivalents of full-length BiP (no calcium). The intensities are normalized to the cross-peak intensities in the absence of BiP. B, the normalized average backbone amide cross-peak intensity of S1R(cd) as a function of the molar ratio of BiP NBD (no calcium) or full-length BiP to S1R(cd) in the absence or presence of 2.5 mM calcium chloride. The intensity average is for only the helical regions. C, spectral overlays of 15N-labeled S1R(cd) with (red) or without (black) full-length BiP (1:1) in the presence of 2.5 mM calcium. D, spectral overlays of 15N-labeled S1R(cd) with (red) and without (black) addition of the BiP SBD (1:1) in the presence of 2.5 mM calcium.

FIGURE 9. The helical (cylinders) and extended (arrow) residues of S1R(cd) determined from a combination of chemical shifts, amide-water proton exchange, 15N relaxation rates, and NOEs. Residues and regions previously implicated in cholesterol binding, 15N relaxation rates, and NOEs. Residues and regions previously implicated in cholesterol binding, 15N relaxation rates, and NOEs. CRM1 and CRM2, cocaine binding (SBDLII; residues Asp-188 and Val-190 are indicated by ↓), and haloperidol binding (residues Asp-126 and Glu-172 are indicated by an asterisk). Residues 198–206, which are proposed to have the strongest interactions with the ER membrane, are indicated. The amino acid sequence and corresponding residue numbers are shown at the bottom.
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CRM1 and CRM2 exhibited heightened protection from the water-soluble paramagnetic agent, although a larger number of NOEs to DPC were detected in the CRM2, suggesting a more intimate interaction with the detergent micelle at this site.

Although the chaperone domain does not bind drugs in the absence of the N-terminal transmembrane domain (1, 35), the secondary structure determined here may reflect the structural propensities of the intact receptor. Studies using a chemically reactive affinity probe have provided information on the binding site of cocaine in guinea pig S1R, which is 98% similar to human S1R (35–37). Those studies defined a steroid binding-like domain (SBDLII) at residues 176–194 and located residues Asp-188 and Val-190 close to the cocaine interaction site. SBDLII together with a steroid binding-like domain (denoted SBDLI; residues 91–109) in the second putative transmembrane helix stabilizes cocaine binding (35). SBDLII corresponds closely to helix 3 (residues 180–189) and the adjacent residues connecting helix 3 to helices 2 and 4 (Fig. 9). Asp-188 and Val-190, which are proposed to be close to the cocaine binding site, are at the C-terminal end of helix 3. Both Asp-188 and Val-190 are solvent exposed in S1R(cd), and Val-190 is one of the most flexible residues within the C-terminal helical region of S1R(cd). It is unknown whether the flexibility observed here is preserved in full-length S1R, but such flexibility may facilitate binding-induced conformational changes necessary to mediate S1R downstream interactions.

Mutational studies of S1R have also implicated Asp-126 and Glu-172 in haloperidol binding (8). Asp-126 and Glu-172 are found in helices 1 and 2 of S1R(cd), respectively, suggesting that these helices may interact in the helical boundary conformation of the receptor.

BiP is an ER resident chaperone that regulates the unfolded protein response in addition to assisting protein folding (reviewed in Ref. 38). S1R is proposed to sequester BiP in the absence of ER stress. Upon depletion of ER calcium or S1R ligand binding the S1R-BiP complex dissociates leading to chaperone activity and downstream signaling of S1R, including inositol triphosphate receptor-mediated calcium release (1, 3). We have shown here that S1R(cd) under NMR conditions interacts with BiP in a calcium-dependent manner and that the NBD of BiP is sufficient for these interactions. The finding that S1R(cd) interacts with the NBD of BiP is consistent with the known regulatory interactions with other Hsp proteins. For example, the co-chaperone BAG1 binds to the NBD of Hsc70 (42, 43), and bacterial GroPE binds to the NBD of DnaK (44, 45).

The role of the region 140–160 remains unknown. This region did not appear to tightly associate with BiP in our studies, and substitution of acidic residues within this region has been shown to have no impact on haloperidol binding (8). Based on sequence analysis (39), several short β-strands are predicted for this region in residues 143–145, 151–153, and 159–164, which corresponds approximately to those regions observed to be extended here (residues 145–147, 153–155; secondary chemical shifts also indicate transient extended conformation in residues 160–162). However, sequence analysis also predicts a high degree of disorder in this region (40), which is confirmed by the NMR data. By contrast, residues 124–137 and 168–173 are predicted from sequence to be extended but found here to be helical. DPC has previously been shown to disrupt β-sheets in a concentration-dependent manner (41). However, the concentrations used here are as much as 10-fold lower than in that study, and the spectra of S1R(cd) in 5 and 30 mM DPC are essentially identical.

In summary, we have studied the chaperone domain of S1R by solution NMR and have characterized its secondary structure and dynamics. S1R(cd) is composed of five helices and at least two short extended regions in a dynamic region between helices 1 and 2. Three of the helices in the C-terminal membrane-associated region map to residues previously identified as important in cholesterol and cocaine binding. A fourth helix (helix 4) is implicated in membrane association. In addition, we have shown that the NBD domain of BiP is sufficient for interaction with the S1R chaperone domain. These results advance our understanding of S1R and are likely to be useful in refining models of the S1R drug binding sites (46). Future studies are needed to identify tertiary interactions in the S1R chaperone domain and to further delineate S1R/BiP and S1R ligand and cholesterol interactions.

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