



Review

G protein-coupled receptors: the evolution of structural insight

Samantha B. Gacasan, Daniel L. Baker, and Abby L. Parrill *

Department of Chemistry, University of Memphis, 3744 Walker Ave, Memphis, TN 38152, USA

* **Correspondence:** Email: aparrill@memphis.edu; Tel: +901-678-3371.

Abstract: G protein-coupled receptors (GPCR) comprise a diverse superfamily of over 800 proteins that have gained relevance as biological targets for pharmaceutical drug design. Although these receptors have been investigated for decades, three-dimensional structures of GPCR have only recently become available. In this review, we focus on the technological advancements that have facilitated efforts to gain insights into GPCR structure. Progress in these efforts began with the initial crystal structure determination of rhodopsin (PDB: 1F88) in 2000 and has continued to the most recently published structure of the A_{1A}R (PDB: 5UEN) in 2017. Numerous experimental developments over the past two decades have opened the door for widespread GPCR structural characterization. These efforts have resulted in the determination of three-dimensional structures for over 40 individual GPCR family members. Herein we present a comprehensive list and comparative analysis of over 180 individual GPCR structures. This includes a summary of different GPCR functional states crystallized with agonists, dual agonists, partial agonists, inverse agonists, antagonists, and allosteric modulators.

Keywords: G protein-coupled receptors; fusion proteins; x-ray crystallography; ligand binding; agonists; antagonists; inverse agonists; allosteric modulators; extracellular loops; intracellular loops; 7TM domain

Abbreviations: A_{1A}R: Adenosine A_{1A} receptor; A_{2A}R: adenosine A_{2A} receptor; ADRB₁: adrenergic β₁ receptor; ADRB₂: adrenergic β₂ receptor; AT₁R: angiotensin II receptor type 1; AT₂R: angiotensin II receptor type 2; CB₁: cannabinoid receptor 1; CCR2: chemokine receptor 2; CXCR4: chemokine receptor 4; CCR5: chemokine receptor 5; CCR9: chemokine receptor 9; CRF₁R: corticotropin-releasing factor 1; δ-OR: δ-opioid receptor; D₃R: dopamine D₃ Receptor; ET-B:

endothelin receptor type-B; CXCR1: chemokine receptor 1; EL: extracellular loop; FFAR₁: free fatty acid receptor 1; GPCR: G protein-coupled receptors; GCGR: glucagon receptor; mGlu₁: glutamate receptor 1; mGlu₅: glutamate receptor 5; H₁R: histamine H1 receptor; κ-OR: κ-opioid receptor; LPA₁: lysophosphatidic acid receptor 1; μ-OR: μ-opioid receptor; M₁R: muscarinic receptor 1; M₂R: muscarinic receptor 2; M₃R: muscarinic receptor 3; M₄R: muscarinic receptor 4; NTSR₁: neurotensin receptor 1; NOP: nociceptin receptor; OX₁: orexin receptor 1; OX₂: orexin receptor 2; PAR1: protease-activated receptor 1; P2Y₁: purinergic P2 receptor 1; P2Y₁₂: purinergic P2 receptor 12; Rho: rhodopsin; 5-HT_{1B}: serotonin 5HT1B; 5-HT_{2B}: serotonin 5HT2B; SMO: smoothed receptor; S1P₁: sphingosine 1-phosphate receptor 1; TACR₁: tachykinin receptor 1; TM: transmembrane; IL: intracellular loop; HHV-5 US28: viral GPCR US28

1. Introduction

G protein-coupled receptors (GPCR) comprise a superfamily of over 800 proteins that are the largest family of cell surface receptors in the human genome [1,2,3]. These proteins share a characteristic seven transmembrane spanning, alpha-helical structure [4]. The GPCR superfamily is commonly subdivided, based on sequence comparisons, into five distinct families: *Rhodopsin* (class A), *Adhesion* (class B), *Secretin* (class B), *Glutamate* (class C), and *Frizzled/Taste2* (class F) [3,5]. More details on the sequence-based analyses that led to these phylogenetic divisions are further discussed in the section titled “Phylogenetic classification/structure” in this review. GPCR have been implicated in numerous biological processes such as cognitive responses [6], cardiovascular functions [7], and cancer growth and development [8]. GPCR implication in human disease is reflected by the estimated 50% of pharmaceutical drugs that interact with these receptors [9].

GPCR mediate signal transduction cascades initiated by numerous extracellular molecules through which they produce downstream physiological responses [4,8,10]. However, receptor activity is not solely stimulated by binding of extracellular ligands. Constitutively active basal signaling [11] has been demonstrated in over 60 wild-type GPCR, including ADRB₂; A_{2A}R; and CB₁ [12]. Some GPCR, such as taste receptors [13,14], adopt active state conformations upon interaction with other receptors [15]. Further details regarding GPCR activation can be found in the “GPCR function” section.

Although these receptors have diverse roles in cellular signaling and in physiology and pathophysiology, they share similar structural topology. This topology includes common structural features shown in Figure 1 that include extracellular loops (EL1–3) and intracellular loops (IL1–3) that alternately connect a characteristic seven transmembrane (7TM) α-helical bundle (Figure 1B) [3,4]. Crystallographic approaches described in more detail in the section titled “Structural characterization” have revealed structural similarities shared by many GPCR found mainly within the 7TM domain. This structural homology, described in more detail in later sections titled by class, suggests similar means for cell signaling. Although GPCR have a shared topology and 7TM structure, there is diversity within the superfamily in regards to sequence composition and length, producing structural variations that are associated with functional specificity in regards to ligand binding or G protein

coupling for different types of GPCR. Select structural differences are also described in more detail in later sections titled by class.

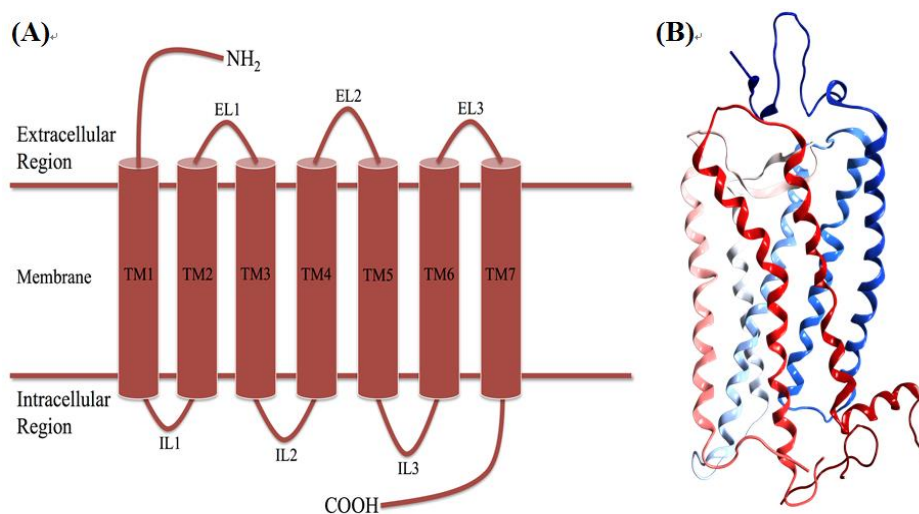


Figure 1. (A) Topological cartoon representing the N-terminal domain, EL 1–3, 7TM domain in cylinders, IL1–3, and C-terminal domain. (B) Ribbon structure of rhodopsin (PDB: 1F88) demonstrating the arrangement of the 7TM α -helical domains within the characteristic membrane-spanning bundle. Here the helices are colored blue starting at the N-terminus fading to red at the C-terminus.

2. GPCR Function

Although this review focuses on GPCR structure and structural characterization, GPCR play a significant role in signal transduction cascades that warrants a generalized overview of their functional and regulatory mechanisms. GPCR are a critical mediator in overall cell signaling, both through ligand-dependent and ligand-independent mechanisms. The majority of these receptors relay signals initiated by binding of extracellular ligands. These ligands are either naturally produced (endogenous) or externally administered (exogenous). Ligands are classified as agonists, inverse agonists, or antagonists. Agonists bind to the receptors and induce a conformational change to an active state, which increases signaling effects. Inverse agonists shift the receptor conformational equilibrium toward inactive conformations, thereby inhibiting basal activity. Antagonists, on the other hand, prevent binding of agonists or inverse agonists without affecting the dynamic conformational equilibrium, which prevents agonist-dependent receptor activation [16]. Changes in conformation within the TM domain, in turn, initiate a conformational change in the intracellular region. IL2 and IL3 have been found to contain critical interaction sites for G proteins or other cytoplasmic effectors [17]. This is illustrated in the crystal structure of ADRB₂-G α_s complex (PDB: 3SN6). Residues located in IL2, TM5, and TM6 of ADRB₂ demonstrate an interaction interface with the α 4 helix, α 5-helix, α N- β 1 junction, and β 3 strand of G α_s [18]. In the classical understanding of GPCR signaling, agonist binding promotes the formation of the activated receptor-G protein complex that modulates functional effects. A detailed analysis of G protein

structure, regulation, and their involvement in signaling has been effectively summarized by Gilman [19]. The heterotrimeric G protein complex is made up of α , β , and γ subunits where the α subunit falls into four subfamilies: $G\alpha_s$, which stimulates adenylyl cyclase activity; $G\alpha_i$, which inhibits adenylyl cyclase activity; $G\alpha_q$, which activates phospholipase C β ; and $G\alpha_{12/13}$, which promotes Rho activation. An activated receptor can interact with heterotrimeric G protein complexes. Receptors function as guanine nucleotide exchange factors (GEF) to promote dissociation of GDP from the $G\alpha$ subunit in exchange for GTP (Figure 2A) [4]. In the activated heterotrimeric G protein complex, the $G\alpha$ -GTP bound subunit dissociates from the $\beta\gamma$ subunit resulting in propagation of signaling cascades. However, subsequent studies have shown some receptors are able to reach an activated state through the formation of dimers and oligomers without agonist binding [4,16]. Alternatively, GPCR activation can occur through interactions with adaptor and scaffolding proteins. Through specific binding sites, adaptor/scaffolding proteins, such as A-kinase anchoring proteins (AKAP) and β -arrestins (Figure 2B), mediate interactions between the receptor and downstream second messengers by scaffolding a network of proteins which then operate as a large molecular complex [4]. Furthermore, receptors including the cannabinoid receptor 1 (CB₁) exhibit high basal signaling, independent of agonist activation [20]. High basal signaling might be indicative of conformational flexibility [21]; however, a study with a constitutively active ADRB₂ mutant has linked conformational flexibility with structural instability [11].

Receptor signaling is modulated by various mechanisms to control cellular functions. Three such processes are highlighted here—deactivation, desensitization, and receptor internalization. At the receptor level, members of the RGS family of proteins serve as GTPase activating proteins (GAPs) and accelerate deactivation by enhancing the rate of $G\alpha$ -catalyzed hydrolysis of GTP to GDP by factors up to 2000-fold (Figure 3A) [22]. In other cases of continuous agonist stimulation, the receptor can be phosphorylated by downstream second-messenger protein kinases or a family of G protein-coupled receptor kinases (GRK). Following phosphorylation, β -arrestin is recruited to the receptor leading to the inhibition of receptor-G protein interaction via steric hindrance. The receptor- β -arrestin complex is targeted and removed from the cell surface through endocytosis. The receptor is then either degraded or recycled back to the cell surface in the inactive conformation (Figure 3B). Receptor internalization is mediated by β -arrestin, clathrin-coated pits, and caveolae through diverse mechanisms [23]. Further in-depth evaluations of GPCR function can be found in excellent reviews by Pierce, et al. [4] and Syrovatkina, et al. [16].

3. Structural Characterization

Due to its high endogenous concentration, the GPCR rhodopsin was purified from bovine rod outer segment (ROS) membranes [24] leading to its crystallization in 2000 (Figure 4; PDB: 1F88) [25]. To date, rhodopsin is the only GPCR that has been purified from its natural source for structural studies. The next GPCR successfully crystallized was ADRB₂ (PDB: 2RH1 [26], 2R4R/2R4S [27]) seven years later. This delay was due to a need for numerous technological advances required to express, purify and crystallize these lipid-soluble targets. The most critical of these advances were improved lipid phases [28,29,30], lipidic cubic phase (LCP) methods for crystallization [31], and the incorporation of soluble fusion partners [31,32].

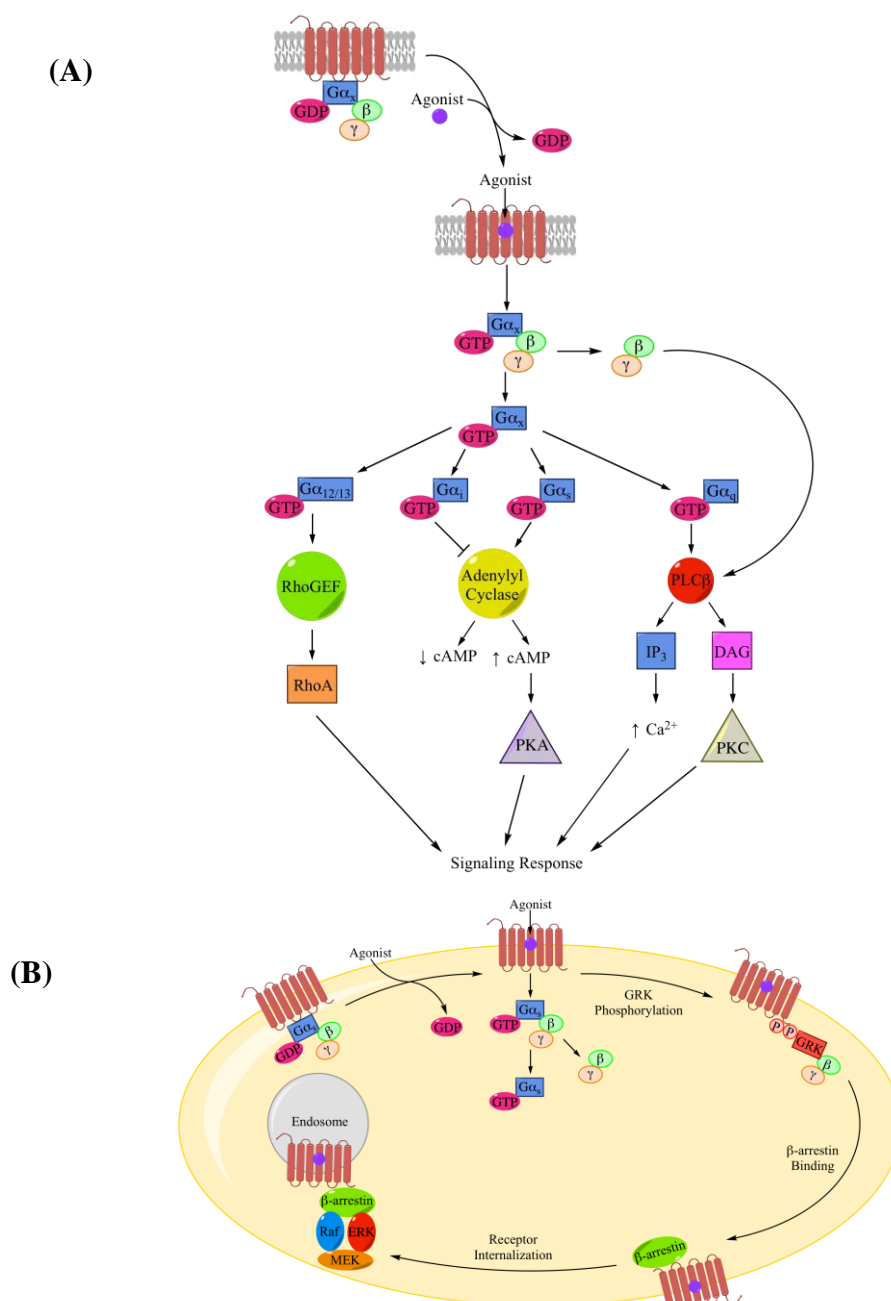


Figure 2. (A) A classical example of GPCR signaling pathway. Receptors, such as $ADRB_2$, are in an inactive state until an agonist binds to activate the receptor. The activated receptor-G protein complex is formed resulting in GDP to GTP exchange by the G protein complex followed by the dissociation of the $G\alpha$ subunit from the $G\beta\gamma$ dimer. Both $G\alpha$ and $G\beta\gamma$ subunits, in turn, activate downstream effectors. $G\alpha_x$ represents the general $G\alpha$ subunit followed by diagrams for $G\alpha_{12/13}$, $G\alpha_i$, $G\alpha_s$, and $G\alpha_q$ pathways. (B) β -arrestins can function as adaptor/scaffolding molecules activating the ERK (MAPK) cascade. GRK phosphorylates the activated receptor recruiting β -arrestin to the phosphorylation sites along with Raf, ERK (MAPK), and MEK. This association triggers activation of the ERK (MAPK) cascade leading to cytosolic signaling pathways. Figures 2A and 2B were adapted from Pierce, et al. [4].

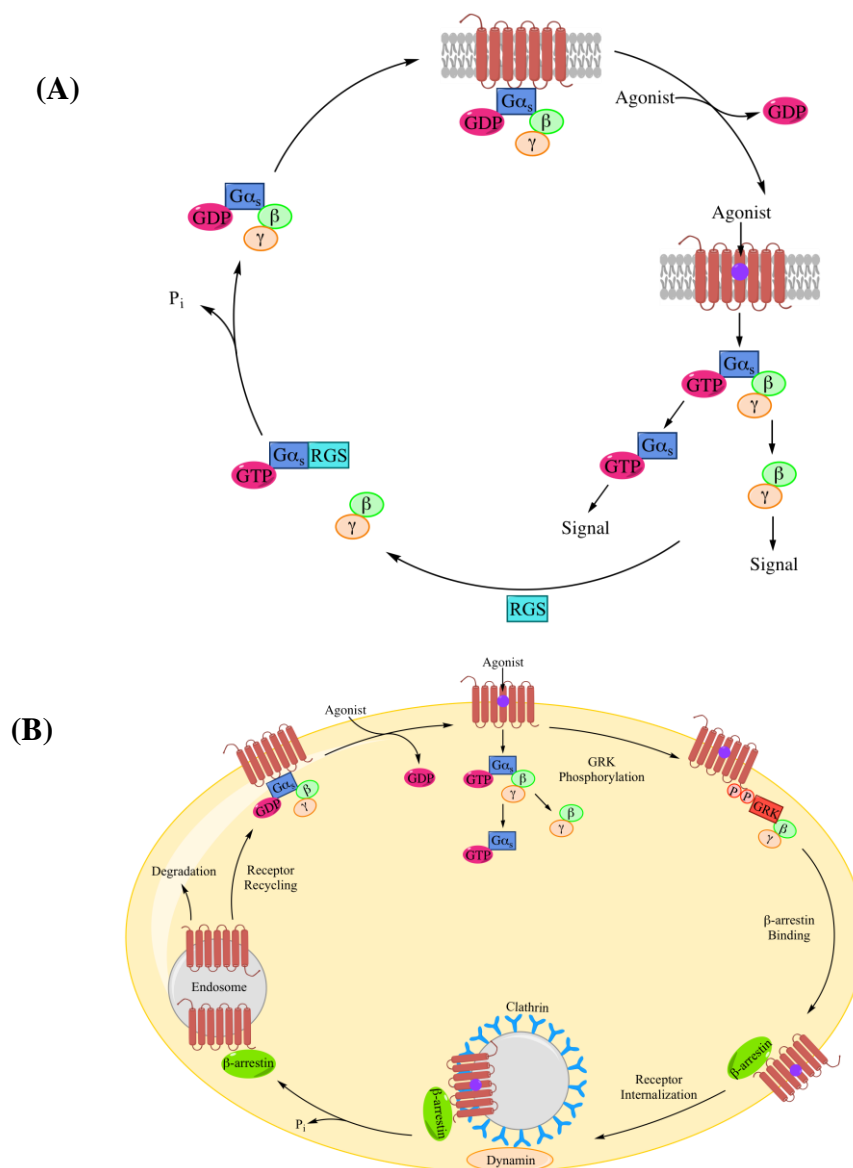


Figure 3. (A) Receptor deactivation is mediated by the RGS family of proteins. RGS alters the conformation of $G\alpha$ -GTP complex making it a better hydrolase, which accelerates the hydrolysis of GTP to GDP. (B) GPCR desensitization through internalization occurs when GRK phosphorylates the activated receptor promoting β -arrestin binding, which sterically hinders receptor-G protein interaction. The receptor is either degraded or recycled back to the cell surface. Figure 3A was adapted from Neubig, et al. [33] and 3B was adapted from Pierce, et al. [4].

Purifying hydrophobic membrane proteins requires additional steps compared to water-soluble proteins. Prior to purification, membrane proteins must be removed from their native lipid environment. Although solubilizing detergents are required to enable efficient extraction of GPCR from the membranes, it is essential to select a mild detergent that discourages protein denaturation. It is important to note that detergent protocols used for one receptor may not be appropriate to other receptors, thus making detergent screening a critical step in the obtaining viable protein.

Rhodopsin (PDB: 1F88) was effectively solubilized in a mixed micellar solution containing heptane-1,2,3-triol (HPTO), nonyl glucoside, and zwitterionic lauryldimethylamine-N-oxide (LDAO) [24]. Both ADRB₂ structures from 2007 (PDB: 2RH1 [26], 2R4R/2R4S [27]) were solubilized in dodecylmaltoside. Often, detergents used for extraction are not sufficient to stabilize protein and a detergent exchange is required [31]. This method was observed in the crystallization trials of the ADRB₂-Gs complex (PDB: 3SN6) where the complex was formed in dodecylmaltoside solution and exchanged into maltose neopentylglycerol detergent (MNG-3) [18].

Crystallization of membrane proteins has been accomplished through the use of different lipid phases including micelles, bicelles, and nanodiscs. These different lipid phases allow for protein stabilization and promote crystal lattice formation [28,29,30]. Detergent micelles surround the hydrophobic GPCR structural regions and allow crystal lattice contacts to form between exposed protein loop structures [28]. Lipid-based bicelles and nanodiscs, on the other hand, act as lipidic mimics of a native bilayer environment. Bicelles are formed via the combination of a detergent or short-chain lipid with a long chain lipid, such as dimyristoyl phosphatidylcholine (DMPC) [29]. A nanodisc is a non-covalently assembled phospholipid bilayer encircled by membrane scaffold proteins, such as plasma lipoproteins [34]. Rhodopsin (PDB: 1F88 [24]) and ADRB₂ (PDB: 2R4R/2R4S [27]) were crystallized in micelles and DMPC/CHAPSO bicelles, respectively. Nanodiscs were used in the crystallization of the nanobody-stabilized ADRB₂ (PDB: 3P0G).

Following many detergent optimization efforts, rhodopsin was crystallized by vapor diffusion [24]. Vapor diffusion forces protein solutions to reach supersaturation prior to crystallization. In this method, protein solutions are mixed with a crystallization solution and positioned in a sitting-drop or hanging-drop orientation in an airtight chamber that also contains the crystallization solution. The chemical equilibrium between the drop and the chamber results in the evaporation of volatile species, which diffuse from the drop to the well. This diffusion leads to a slow saturation of protein within the drop allowing the protein-detergent complexes to form crystal lattice contacts with neighboring molecules [35]. In lipidic cubic phase (LCP) methods, proteins are placed in a membrane-like environment where they can diffuse and interact with each other to form crystal lattice contacts on both complementary hydrophobic and hydrophilic regions [31]. Crystals of rhodopsin (PDB: 1F88 [25]) and ADRB₂ (PDB: 2R4R/2R4S [27]) were grown by hanging drop diffusion, whereas ADRB₂ (PDB: 2RH1 [26]) and ADRB₂-Gs complex (PDB: 3SN6 [18]) crystallization experiments implemented LCP methods.

In determining crystallographic structures, molecular replacement has been successfully utilized to help in determining the phase of an unknown target by applying the phase of a closely related and previously characterized target. This approach was implemented in the analysis of the ADRB₂ structure [26]. Multi-wavelength anomalous scattering (MAD) and single-wavelength anomalous scattering (SAD) methods have impacted atomic-level structure determinations of GPCR when a heavy atom anomalous scatterer is incorporated into the protein structure. Structural determination using MAD phasing was implemented in the first crystal structure of rhodopsin (PDB: 1F88) [25].

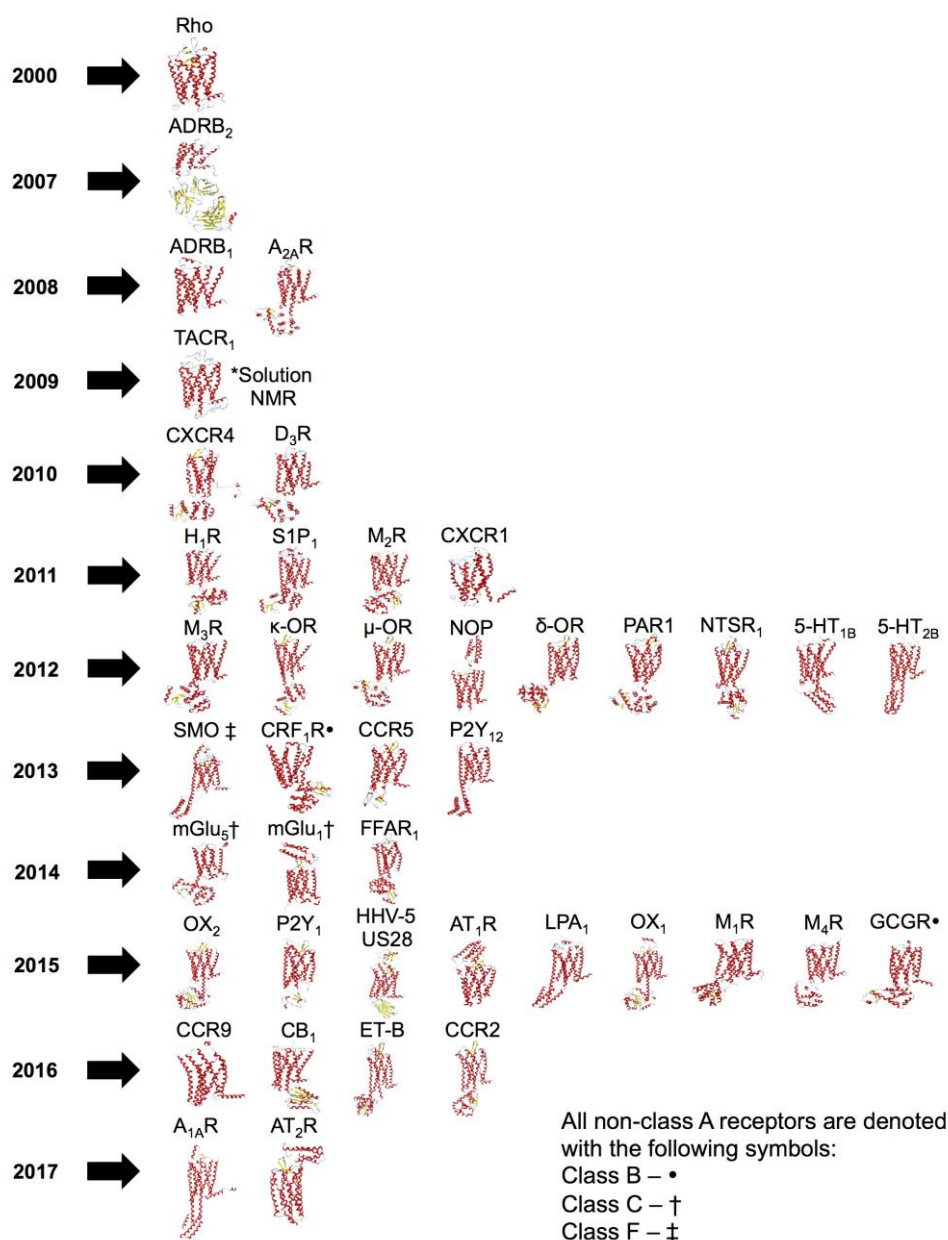


Figure 4. Timeline of availability for individual representative GPCR crystal structure structures including protein fusion partners. Rho (1F88), ADRB₂ (2R4R), ADRB₁ (2VT4), A_{2A}R (3EML), TACR₁ (2KS9), CXCR4 (3ODU), D₃R (3PBL), H₁R (3RZE), S1P₁ (3V2W), M₂R (3UON), CXCR1 (2LNL), M₃R (4DAJ), κ-OR (4DJH), μ-OR (4DKL), NOP (4EA3), δ-OR (4EJ4), PAR1 (3VW7), NSTR₁ (3ZEV), 5-HT_{1B} (4IAQ), 5-HT_{2B} (4IB4), SMO (4JKV), CRF₁R (4K5Y), CCR5 (4MBS) P2Y₁₂ (4NTJ), mGlu₅ (4OO9), mGlu₁ (4OR2), FFAR₁ (4PHU), OX₂ (4S0V), P2Y₁ (4XNW), HHV-5 US28 (4XT1), AT₁R (4YAY), LPA₁ (4Z34), OX₁ (4ZJ8), M₁R (5CXV), M₄R (5DSG), GCGR (5EE7), CCR9 (5LWE), ET-B (5GLH), CCR2 (5T1A), CB₁ (5TGZ), A_{1A}R (5UEN), AT₂R (5UNF).

Developments in protein engineering and heterologous protein expression have accelerated structural determinations of GPCR. Site-directed mutagenesis has been beneficial for the stabilization of receptor structure, as well as determination of the functional activity of many GPCR [31]. Mutagenesis has also enabled the introduction of fusion partners into protein sequences. Fusion partners are designed to be highly stable, compact, and easily crystallized. These partners maintain essential surface contacts and increase protein solubility, making them suitable replacements for flexible domains [31,32]. The position of the fusion partner and the number of residues conjoining the fusion partner to the protein may alter expression and/or stability; nevertheless, optimizing the number of linker residues can minimize adverse effects. A Fab5 epitope was engineered into IL3 of ADRB₂ (PDB: 2R4R/2R4S) by mutagenesis, which allowed a Fab5 monoclonal antibody to bind to the epitope [27]. Fab5 stabilized the receptor conformation and increased the polar surface area necessary for crystal lattice contacts. Since the initial use of the Fab5 antibody, many GPCR have been successfully engineered with fusion partners accelerating the number solved of crystallographic structures in the last two decades as illustrated in the timeline in Figure 4. T4 lysozyme has been the most widely used fusion partner in multiple receptors such as adenosine A_{2A} receptor (A_{2A}R; PDB: 3EML) [36] and metabotropic glutamate receptor 5 (mGlu₅; PDB: 4OO9) [37]. Thermostabilized apocytochrome b₅₆₂RIL (BRIL) is a highly versatile fusion partner that has been appended to the N-terminal domain of nociception receptor 1 (NOP; PDB: 4EA3) [38] and C-terminal domain of smoothed receptor (SMO; PDB: 4JKV) [39]. *Pyrococcus abyssi* glycogen synthase (PGS) and rubredoxin emerged as fusion partners later in the timeline of GPCR characterization. These tools have been successfully used in analysis of the orexin 1 (OX₁; PDB: 4ZJ8) [40] and purinergic receptor 1 (P2Y₁; PDB: 4XNW) receptors [41]. Although the addition of fusion protein sequences have been effective in the analysis of many GPCR, it is important to note that engineering designs are often specific for one protein and require optimization for each application to new receptor sequences.

GPCR loops and termini, which are characteristically flexible and difficult to crystallize, can result in additional challenges. Proteins truncated or modified at these areas can reduce flexibility. However, simple truncations can also reduce the polar surface area of the protein, which is a characteristic crucial for forming the crystal lattice contacts required for crystal formation. Rhodopsin has a relatively short IL3 with ~3 amino acids and C-terminal domain with ~25 amino acids, in contrast, ADRB₂ has a lengthy IL3 with ~28 amino acids and a C-terminal domain ~70 amino acids [42]. Two early structures of ADRB₂ were engineered differently in these regions to achieve crystal lattice formation during crystallization. In the ADRB₁-Fab5 complex (PDB: 2R4R/2R4S), truncating the final 48 amino acids in the unstructured C-terminal domain optimized crystal size and uniformity [27]. In another structure of ADRB₂ (PDB: 2RH1), IL3 was completely removed and replaced with the soluble fusion partner, T4 lysozyme, which promoted the growth of diffraction quality crystals [26]. Amino acid truncations have also been found to have a variable effect on protein expression. Truncations at the N-terminus reduced expression. However, replacing the N-terminus with BRIL maintained similar expression as constructs with the complete N-terminus [38]. This was the case with NOP (PDB: 4EA3) where an N-terminal replacement with BRIL and a 31 amino acid deletion at the C-terminus significantly increased protein expression [31,38].

Table 1. A comprehensive list of GPCR crystal structures according to date deposited into the Protein Data Bank as of April 19, 2017.

GPCR Crystal Structures							
Individual Receptors	Family	Class /Subclass	Species	Receptor	PDB [43] ID	Resolution (Å)	Date Deposited
1	Rhodopsin	Class A α	Bovine	Rho	1F88 [25]	2.80	6/29/00
2	Rhodopsin	Class A α	Bovine	Rho	1HZX [44]	2.80	1/26/01
3	Rhodopsin	Class A α	Bovine	Rho	1JFP [45]	Solution NMR	6/21/01
4	Rhodopsin	Class A α	Bovine	Rho	1L9H [46]	2.60	3/23/02
5	Rhodopsin	Class A α	Bovine	Rho	1LN6 [47]	Solution NMR	5/3/02
6	Rhodopsin	Class A α	Bovine	Rho	1GZM [48]	2.65	5/24/02
7	Rhodopsin	Class A α	Bovine	Rho	1U19 [49]	2.20	7/15/04
8	Rhodopsin	Class A α	Bovine	Rho	2G87 [50]	2.60	3/2/06
9	Rhodopsin	Class A α	Bovine	Rho	2HPY [51]	2.80	7/18/06
10	Rhodopsin	Class A α	Bovine	Rho	2I35 [52]	3.80	8/17/06
11	Rhodopsin	Class A α	Bovine	Rho	2I36 [52]	4.10	8/17/06
12	Rhodopsin	Class A α	Bovine	Rho	2I37 [52]	4.15	8/17/06
13	Rhodopsin	Class A α	Bovine	Rho	2J4Y [53]	3.40	9/7/06
14	Rhodopsin	Class A α	Bovine	Rho	2PED [54]	2.95	4/2/07
15	Rhodopsin	Class A α	Bovine	Rho	3C9L [55]	2.65	2/16/08
16	Rhodopsin	Class A α	Bovine	Rho	3C9M [55]	3.40	2/16/08
17	Rhodopsin	Class A α	Bovine	Rho	3CAP [56]	2.90	2/20/08
18	Rhodopsin	Class A α	Bovine	Rho	3DQB [57]	3.20	7/9/08
19	Rhodopsin	Class A α	Bovine	Rho	2X72 [58]	3.00	2/22/10
20	Rhodopsin	Class A α	Bovine	Rho	3OAX[59]	2.60	8/5/10
21	Rhodopsin	Class A α	Bovine	Rho	3PQR [60]	2.85	11/26/10
22	Rhodopsin	Class A α	Bovine	Rho	3PXO [60]	3.00	12/10/10
23	Rhodopsin	Class A α	Bovine	Rho	4A4M [61]	3.30	10/17/11
24	Rhodopsin	Class A α	Bovine	Rho	4J4Q [62]	2.65	2/7/13
25	Rhodopsin	Class A α	Bovine	Rho	4BEY [63]	2.90	3/12/13
26	Rhodopsin	Class A α	Bovine	Rho	4BEZ [63]	3.30	3/12/13
27	Rhodopsin	Class A α	Bovine	Rho	4PXF [64]	2.75	3/23/14
28	Rhodopsin	Class A α	Bovine	Rho	4X1H [65]	2.29	11/24/14
29	Rhodopsin	Class A α	Human	Rho	4ZWJ [66]	3.30	5/19/15
30	Rhodopsin	Class A α	Bovine	Rho	5DYS [67]	2.30	9/25/15
31	Rhodopsin	Class A α	Bovine	Rho	5EN0 [67]	2.81	11/8/15
32	Rhodopsin	Class A α	Bovine	Rho	5TE3 [68]	2.70	9/20/16
33	Rhodopsin	Class A α	Bovine	Rho	5TE5 [68]	4.01	9/21/16
34	Rhodopsin	Class A α	Turkey	ADRB ₁	2VT4 [69]	2.70	5/9/08
35	Rhodopsin	Class A α	Turkey	ADRB ₁	2Y00 [70]	2.50	11/30/10
36	Rhodopsin	Class A α	Turkey	ADRB ₁	2Y01[70]	2.60	11/30/10
37	Rhodopsin	Class A α	Turkey	ADRB ₁	2Y02 [70]	2.60	11/30/10
38	Rhodopsin	Class A α	Turkey	ADRB ₁	2Y03 [70]	2.85	11/30/10
39	Rhodopsin	Class A α	Turkey	ADRB ₁	2Y04 [70]	3.05	11/30/10
40	Rhodopsin	Class A α	Turkey	ADRB ₁	2YCW [71]	3.00	3/17/11

41	Rhodopsin	Class A α	Turkey	ADRB ₁	2YCX [71]	3.25	3/17/11
42	Rhodopsin	Class A α	Turkey	ADRB ₁	2YCY [71]	3.15	3/17/11
43	Rhodopsin	Class A α	Turkey	ADRB ₁	2YCZ [71]	3.65	3/17/11
44	Rhodopsin	Class A α	Turkey	ADRB ₁	3ZPQ [72]	2.80	3/1/13
45	Rhodopsin	Class A α	Turkey	ADRB ₁	3ZPR [72]	2.70	3/1/13
46	Rhodopsin	Class A α	Turkey	ADRB ₁	4AMI [73]	3.20	3/11/12
47	Rhodopsin	Class A α	Turkey	ADRB ₁	4AMJ [73]	2.30	3/12/12
48	Rhodopsin	Class A α	Turkey	ADRB ₁	4BVN [74]	2.10	6/26/13
49	Rhodopsin	Class A α	Turkey	ADRB ₁	4GPO [75]	3.50	8/21/12
50	Rhodopsin	Class A α	Turkey	ADRB ₁	5A8E [76]	2.40	7/15/15
51	Rhodopsin	Class A α	Turkey	ADRB ₁	5F8U [77]	3.35	12/9/15
52	Rhodopsin	Class A α	Human	ADRB ₂	2R4R [27]	3.40	8/31/07
53	Rhodopsin	Class A α	Human	ADRB ₂	2R4S [27]	3.40	8/31/07
54	Rhodopsin	Class A α	Human	ADRB ₂	2RH1 [26]	2.40	10/5/07
55	Rhodopsin	Class A α	Human	ADRB ₂	3D4S [78]	2.80	5/14/08
56	Rhodopsin	Class A α	Human	ADRB ₂	3KJ6 [79]	3.40	11/2/09
57	Rhodopsin	Class A α	Human	ADRB ₂	3NY8 [80]	2.84	7/14/10
58	Rhodopsin	Class A α	Human	ADRB ₂	3NY9 [80]	2.84	7/14/10
59	Rhodopsin	Class A α	Human	ADRB ₂	3NYA [80]	3.16	7/14/10
60	Rhodopsin	Class A α	Human	ADRB ₂	3P0G [81]	3.50	9/28/10
61	Rhodopsin	Class A α	Human	ADRB ₂	3PDS [82]	3.50	10/24/10
62	Rhodopsin	Class A α	Human	ADRB ₂	3SN6 [18]	3.20	6/28/11
63	Rhodopsin	Class A α	Human	ADRB ₂	4GBR [83]	3.99	7/27/12
64	Rhodopsin	Class A α	Human	ADRB ₂	4LDE [84]	2.79	6/24/13
65	Rhodopsin	Class A α	Human	ADRB ₂	4LDL [84]	3.10	6/24/13
66	Rhodopsin	Class A α	Human	ADRB ₂	4LDO [84]	3.20	6/24/13
67	Rhodopsin	Class A α	Human	ADRB ₂	4QKX [85]	3.30	6/10/14
68	Rhodopsin	Class A α	Human	ADRB ₂	5D5A [86]	2.48	8/10/15
69	Rhodopsin	Class A α	Human	ADRB ₂	5D5B [86]	3.80	8/10/15
70	Rhodopsin	Class A α	Human	ADRB ₂	5D6L [Ma, P; Caffrey, M]	3.20	8/12/15
71	Rhodopsin	Class A α	Human	ADRB ₂	5JQH [87]	3.20	5/5/16
72	Rhodopsin	Class A α	Human	H ₁ R	3RZE [88]	3.10	5/11/11
73	Rhodopsin	Class A α	Human	D ₃ R	3PBL [6]	2.89	10/20/10
74	Rhodopsin	Class A α	Human	5-HT _{1B}	4IAQ [89]	2.80	12/7/12
75	Rhodopsin	Class A α	Human	5-HT _{1B}	4IAR [89]	2.70	12/7/12
76	Rhodopsin	Class A α	Human	5-HT _{2B}	4IB4 [90]	2.70	12/7/12
77	Rhodopsin	Class A α	Human	5-HT _{2B}	4NC3 [91]	2.80	10/23/13
78	Rhodopsin	Class A α	Human	5-HT _{2B}	5TVN [92]	2.90	11/9/16
79	Rhodopsin	Class A α	Human	M ₁ R	5CXV [93]	2.70	7/29/15
80	Rhodopsin	Class A α	Human	M ₂ R	3UON [94]	3.00	11/16/11
81	Rhodopsin	Class A α	Human	M ₂ R	4MQS [95]	3.50	9/16/13
82	Rhodopsin	Class A α	Human	M ₂ R	4MQT [95]	3.70	9/16/13
83	Rhodopsin	Class A α	Rat	M ₃ R	4DAJ [96]	3.40	1/12/12
84	Rhodopsin	Class A α	Rat	M ₃ R	4U14 [97]	3.57	7/15/14
85	Rhodopsin	Class A α	Rat	M ₃ R	4U15 [97]	2.80	7/15/14

86	Rhodopsin	Class A α	Rat	M ₃ R	4U16 [97]	3.70	7/15/14
87	Rhodopsin	Class A α	Human	M ₄ R	5DSG [93]	2.60	9/17/15
88	Rhodopsin	Class A α	Human	A _{1A} R	5UEN [98]	3.20	1/3/17
89	Rhodopsin	Class A α	Human	A _{2A} R	3EML [36]	2.60	9/24/08
90	Rhodopsin	Class A α	Human	A _{2A} R	3PWH [99]	3.30	12/8/10
91	Rhodopsin	Class A α	Human	A _{2A} R	3QAK [100]	2.71	1/11/11
92	Rhodopsin	Class A α	Human	A _{2A} R	2YDO [101]	3.00	3/23/11
93	Rhodopsin	Class A α	Human	A _{2A} R	2YDV [101]	2.60	3/24/11
94	Rhodopsin	Class A α	Human	A _{2A} R	3REY [99]	3.31	4/5/11
95	Rhodopsin	Class A α	Human	A _{2A} R	3RFM [99]	3.60	4/6/11
96	Rhodopsin	Class A α	Human	A _{2A} R	3VG9 [102]	2.70	8/4/11
97	Rhodopsin	Class A α	Human	A _{2A} R	3VGA [102]	3.10	8/4/11
98	Rhodopsin	Class A α	Human	A _{2A} R	3UZA [103]	3.27	12/7/11
99	Rhodopsin	Class A α	Human	A _{2A} R	3UZC [103]	3.34	12/7/11
100	Rhodopsin	Class A α	Human	A _{2A} R	4E1Y [104]	1.80	4/6/12
101	Rhodopsin	Class A α	Human	A _{2A} R	4UG2 [105]	2.60	3/21/15
102	Rhodopsin	Class A α	Human	A _{2A} R	4UHR [105]	2.60	3/25/15
103	Rhodopsin	Class A α	Human	A _{2A} R	5IU4 [106]	1.72	3/17/16
104	Rhodopsin	Class A α	Human	A _{2A} R	5IU7 [106]	1.90	3/17/16
105	Rhodopsin	Class A α	Human	A _{2A} R	5IU8 [106]	2.00	3/17/16
106	Rhodopsin	Class A α	Human	A _{2A} R	5IUA [106]	2.20	3/17/16
107	Rhodopsin	Class A α	Human	A _{2A} R	5IUB [106]	2.10	3/17/16
108	Rhodopsin	Class A α	Human	A _{2A} R	5K2A [107]	2.50	5/18/16
109	Rhodopsin	Class A α	Human	A _{2A} R	5K2B [107]	2.50	5/18/16
110	Rhodopsin	Class A α	Human	A _{2A} R	5K2C [107]	1.90	5/18/16
111	Rhodopsin	Class A α	Human	A _{2A} R	5K2D [107]	1.90	5/18/16
112	Rhodopsin	Class A α	Human	A _{2A} R	5G53 [108]	3.40	5/19/16
113	Rhodopsin	Class A α	Human	A _{2A} R	5UIG [109]	3.50	1/13/17
114	Rhodopsin	Class A α	Human	S1P ₁	3V2W [110]	3.35	12/12/11
115	Rhodopsin	Class A α	Human	S1P ₁	3V2Y [110]	2.80	12/12/11
116	Rhodopsin	Class A α	Human	LPA ₁	4Z34 [111]	3.00	3/30/15
117	Rhodopsin	Class A α	Human	LPA ₁	4Z35 [111]	2.90	3/30/15
118	Rhodopsin	Class A α	Human	LPA ₁	4Z36 [111]	2.90	3/30/15
119	Rhodopsin	Class A α	Human	CB ₁	5TGZ [112]	2.80	9/28/16
120	Rhodopsin	Class A α	Human	CB1	5U09 [113]	2.60	11/23/16
121	Rhodopsin	Class A β	Rat	NTSR ₁	4GRV [114]	2.80	8/27/12
122	Rhodopsin	Class A β	Rat	NTSR ₁	3ZEV [115]	3.00	12/7/12
123	Rhodopsin	Class A β	Rat	NTSR ₁	4BUO [115]	2.75	6/21/13
124	Rhodopsin	Class A β	Rat	NTSR ₁	4BV0 [115]	3.10	6/24/13
125	Rhodopsin	Class A β	Rat	NTSR ₁	4BWB [115]	3.57	7/1/13
126	Rhodopsin	Class A β	Rat	NTSR ₁	4XEE [116]	2.90	12/23/14
127	Rhodopsin	Class A β	Rat	NTSR ₁	4XES [116]	2.60	12/24/14
128	Rhodopsin	Class A β	Rat	NTSR ₁	5T04 [117]	3.30	8/16/16
129	Rhodopsin	Class A β	Human	OX ₁	4ZJ8 [40]	2.75	4/29/15
130	Rhodopsin	Class A β	Human	OX ₁	4ZJC [40]	2.83	4/29/15
131	Rhodopsin	Class A β	Human	OX ₂	4S0V [118]	2.50	1/6/15

132	Rhodopsin	Class A β	Human	TACR ₁	2KS9 [119]	Solution NMR	12/31/09
133	Rhodopsin	Class A β	Human	TACR ₁	2KSA [119]	Solution NMR	12/31/09
134	Rhodopsin	Class A β	Human	TACR ₁	2KSB [119]	Solution NMR	12/31/09
135	Rhodopsin	Class A β	Human	ET-B	5GLI [120]	2.5	7/11/16
136	Rhodopsin	Class A β	Human	ET-B	5GLH [120]	2.8	7/11/16
137	Rhodopsin	Class A γ	Human	CXCR1	2LNL [121]	Solid-State NMR	12/31/11
138	Rhodopsin	Class A γ	Human	CCR2	5T1A [122]	2.81	8/18/16
139	Rhodopsin	Class A γ	Human	CXCR4	3ODU [123]	2.50	8/11/10
140	Rhodopsin	Class A γ	Human	CXCR4	3OE0 [123]	2.90	8/12/10
141	Rhodopsin	Class A γ	Human	CXCR4	3OE6 [123]	3.20	8/12/10
142	Rhodopsin	Class A γ	Human	CXCR4	3OE8 [123]	3.10	8/12/10
143	Rhodopsin	Class A γ	Human	CXCR4	3OE9 [123]	3.10	8/12/10
144	Rhodopsin	Class A γ	Human	CXCR4	4RWS [124]	3.10	12/5/14
145	Rhodopsin	Class A γ	Human	CCR5	4MBS [125]	2.71	8/19/13
146	Rhodopsin	Class A γ	Human	CCR9	5LWE [126]	2.80	9/16/16
147	Rhodopsin	Class A γ	Human	NOP	4EA3 [38]	3.01	3/22/12
148	Rhodopsin	Class A γ	Human	NOP	5DHG [127]	3.00	8/30/15
149	Rhodopsin	Class A γ	Human	NOP	5DHH [127]	3.00	8/31/15
150	Rhodopsin	Class A γ	Human	κ -OR	4DJH [128]	2.90	2/1/12
151	Rhodopsin	Class A γ	Mouse	μ -OR	4DKL [129]	2.80	2/3/12
152	Rhodopsin	Class A γ	Mouse	μ -OR	5C1M [130]	2.10	6/15/15
153	Rhodopsin	Class A γ	Mouse	δ -OR	4EJ4 [131]	3.40	4/6/12
154	Rhodopsin	Class A γ	Human	δ -OR	4N6H [132]	1.80	10/12/13
155	Rhodopsin	Class A γ	Human	δ -OR	4RWA [133]	3.28	12/1/14
156	Rhodopsin	Class A γ	Human	δ -OR	4RWD [133]	2.70	12/2/14
157	Rhodopsin	Class A γ	Human	AT ₁ R	4YAY [134]	2.90	2/18/15
158	Rhodopsin	Class A γ	Human	AT ₁ R	4ZUD [135]	2.80	5/15/15
159	Rhodopsin	Class A γ	Human	AT ₂ R	5UNF [136]	2.80	1/30/17
160	Rhodopsin	Class A γ	Human	AT ₂ R	5UNG [136]	2.80	1/30/17
161	Rhodopsin	Class A γ	Human	AT ₂ R	5UNH [136]	2.90	1/30/17
162	Rhodopsin	Class A γ	HHV-5	HHV-5 US28	4XT1 [137]	2.89	1/22/15
163	Rhodopsin	Class A γ	HHV-5	HHV-5 US28	4XT3 [137]	3.80	1/22/15
164	Rhodopsin	Class A δ	Human	P2Y ₁	4XNW [138]	2.70	1/16/15
165	Rhodopsin	Class A δ	Human	P2Y ₁	4XNV [138]	2.20	1/16/15
166	Rhodopsin	Class A δ	Human	P2Y ₁₂	4NTJ [41]	2.62	12/2/13
167	Rhodopsin	Class A δ	Human	P2Y ₁₂	4PXZ [139]	2.50	3/25/14
168	Rhodopsin	Class A δ	Human	P2Y ₁₂	4PY0 [139]	3.10	3/25/14
169	Rhodopsin	Class A δ	Human	PAR1	3VW7 [140]	2.20	8/7/12
170	Rhodopsin	Class A δ	Human	FFAR ₁	4PHU [141]	2.33	5/7/14
171	Secretin	Class B	Human	GCGR	4L6R [142]	3.30	6/12/13
172	Secretin	Class B	Human	GCGR	5EE7 [143]	2.50	10/22/15
173	Secretin	Class B	Human	CRF ₁ R	4K5Y [144]	2.98	4/15/13
174	Secretin	Class B	Human	CRF ₁ R	4Z9G [145]	3.18	4/10/15
175	Glutamate	Class C	Human	mGlu ₁	4OR2 [146]	2.80	2/10/14

176	Glutamate	Class C	Human	mGlu ₅	4OO9 [37]	2.60	1/31/14
177	Glutamate	Class C	Human	mGlu ₅	5CGC [147]	3.10	7/9/15
178	Glutamate	Class C	Human	mGlu ₅	5CGD [147]	2.60	7/9/15
179	Frizzled/ Taste2	Class F	Human	SMO	4JKV [39]	2.45	3/11/13
180	Frizzled/ Taste2	Class F	Human	SMO	4N4W [148]	2.80	10/8/13
181	Frizzled/ Taste2	Class F	Human	SMO	4O9R [85]	3.20	1/2/14
182	Frizzled/ Taste2	Class F	Human	SMO	4QIM [148]	2.61	5/31/14
183	Frizzled/ Taste2	Class F	Human	SMO	4QIN [148]	2.60	5/31/14
184	Frizzled/ Taste2	Class F	Human	SMO	5L7D [149]	3.20	6/3/16
185	Frizzled/ Taste2	Class F	Human	SMO	5L7I [149]	3.30	6/3/16

There have been challenges in GPCR structure determination due to their hydrophobic nature and instability outside of the hydrophobic membrane environment [150,151]. Since the first complete x-ray crystallographic structure of a GPCR (bovine rhodopsin; PDB: 1F88) was solved in 2000 [25], there are now over 180 comprehensive structures of GPCR in the Protein Data Bank (www.rcsb.org [43]) listed in Table 1. More than 150 of these structures co-crystallized with ligands are described in more detail in Table 2. However, less than 45 distinct family members are represented out of a superfamily of over 800 proteins. This limitation in representation suggests that significantly more work is yet to be done.

4. Phylogenetic Classification/Structure

Prior to the availability of three-dimensional structural information, GPCR classification methods initially utilized primary amino acid sequences to phylogenetically categorize receptors. This approach ultimately laid the foundation for the most-commonly used GPCR classification system. The sequences of seven hydrophobic regions (represented as cylinders in Figure 1A) were used to design a fingerprinting method to identify sequences not previously categorized as rhodopsin-like receptors [152]. In the method, an individual conserved hydrophobic region served as a single “feature”. Multiple features grouped together comprised a signature “fingerprint” within a sequence. A feature, when used by a scanning algorithm to search a database, is then referred to as a “discriminator” and fingerprints are referred to as “composite discriminators. The search results using the discriminators generate an output hit list for each hydrophobic region. A hit list correlation was used to differentiate between true members, where all features of the fingerprints matched, and noise, where zero, one or two random matches occur. A second database was built from the sequences resulting from the previous search and scans were repeated with the new discriminators. This process was repeated until the composite discriminator continuously distinguished between true

members and noise. As a result, 240 sequences were identified as members of the superfamily. Sequences for pheromone receptors did not match any of the discriminators and cAMP receptors exhibited only two matches, thus falling within the noise [152]. These sequences were previously identified as GPCR [153] suggesting they were members of distantly related families [152]. The hit-list fingerprint correlation was later expanded to distinguish partial matches. Upon this expansion, sequences belonging to the GPCR superfamily increased from 240 to 393. In addition, the pheromone, cAMP, and secretin-like families were established as rhodopsin-like receptors [154].

The rhodopsin-like family rapidly became overly complex as more diverse receptors were discovered, leading to the establishment of the GPCR superfamily and formation of the “class” system. This A-F system includes GPCR found in both vertebrates and invertebrates [3]. In 2001, the first draft of the human genome became available [1,2] and allowed further GPCR to be classified using the most prevalent classification system with most of the human GPCR categorized into five families shown in italics: *Glutamate* (class C), *Rhodopsin* (class A), *Adhesion* (class B), *Frizzled/Taste2* (class F), and *Secretin* (class B); also recognized as GRAFS [3].

5. Rhodopsin (Class A)

The *Rhodopsin* (class A) family, the largest of the GPCR classes [3], is divided into α , β , γ , and δ subgroups [5]. The α -subgroup is the largest group in class A and is comprised of the prostaglandin, amine, opsin, melatonin, and MECA receptor clusters [3]. Generally, ligands bind to these receptors within a pocket inside the transmembrane cavity involving residues contained in TM3, TM5, and TM6 [155–158]. An example of this target binding domain can be found in the ADRB₂ structure in complex with carazolol (PDB: 2RH1) [26]. Biogenic monoamine receptors, such as the adrenergic; cannabinoid; muscarinic; serotonin; dopamine; and histamine receptors, are important drug targets for cardiovascular drugs, antipsychotics, and anti-histamines [159,160]. Novel drugs that lack receptor specificity for particular amine receptors pose a risk for cardiovascular side effects due to off-target interactions with adrenergic receptors that show expression in many tissues [161]. Since the α -subgroup receptors outside the amine receptor subset are divergent from the amine receptors, side effects of novel aminergic drugs are most likely to occur only through the amine subset [5].

The β -subgroup of the *Rhodopsin* (class A) family has no branching subgroups and includes the hypocretin receptor, neuropeptide FF receptor, tachykinin receptor, cholecystokinin receptor, neuropeptide Y receptor, endothelin-related peptide receptor, gastrin-releasing peptide receptor, neuromedin receptor, thyrotropin releasing hormone receptor, ghrelin receptor, arginine vasopressin/oxytocin receptor, gonadotropin-releasing hormone receptor, and orphan receptors. Orphan receptors have been established as GPCR based on DNA sequence but have unidentified endogenous ligands [162]. Members of the β -subgroup mainly bind peptides with a high specificity-binding profile and are pursued as drug targets for treatments including pulmonary arterial hypertension [163] and hormone-related cancer [164]. Agonist drug design for β -subgroup members has been a challenge since it is difficult to design novel ligands that are flexible enough to mimic the magnitude of interaction sites engaged by endogenous peptide agonists [3].

Table 2. A comprehensive list of GPCR crystallized with ligands deposited into the Protein Data Bank as of April 19, 2017.

	PDB [43] ID	Class/ Subclass	GPCR	Ligand Name	Ligand Type
1	1F88 [25]	Class A α	Rho	11- <i>cis</i> -Retinal	
2	1GZM [48]	Class A α	Rho	11- <i>cis</i> -Retinal	
3	1HZX [44]	Class A α	Rho	11- <i>cis</i> -Retinal	
4	1JFP [45]	Class A α	Rho	all- <i>trans</i> -Retinal	
5	1L9H [46]	Class A α	Rho	11- <i>cis</i> -Retinal	
6	1LN6 [47]	Class A α	Rho	all- <i>trans</i> -Retinal	
7	1U19 [49]	Class A α	Rho	11- <i>cis</i> -Retinal	
8	2G87 [50]	Class A α	Rho	all- <i>trans</i> -Retinal	
9	2HPY [51]	Class A α	Rho	all- <i>trans</i> -Retinal	
10	2I35 [52]	Class A α	Rho	11- <i>cis</i> -Retinal	
11	2J4Y [53]	Class A α	Rho	11- <i>cis</i> -Retinal	
12	2PED [54]	Class A α	Rho	9- <i>cis</i> -Retinal	
13	2X72 [58]	Class A α	Rho	11- <i>cis</i> -Retinal	
14	3C9L [55]	Class A α	Rho	11- <i>cis</i> -Retinal	
15	3C9M [55]	Class A α	Rho	11- <i>cis</i> -Retinal	
16	3OAX [59]	Class A α	Rho	11- <i>cis</i> -Retinal	
17	3PQR [60]	Class A α	Rho	all- <i>trans</i> -Retinal	
18	3PXO [60]	Class A α	Rho	all- <i>trans</i> -Retinal	
19	4A4M [61]	Class A α	Rho	all- <i>trans</i> -Retinal	
20	5DYS [67]	Class A α	Rho	all- <i>trans</i> -Retinal	
21	5EN0 [67]	Class A α	Rho	all- <i>trans</i> -Retinal	
22	5TE5 [68]	Class A α	Rho	(2E)-{(4E)-4-[(3E)-4-(2,6,6-trimethylcyclohex-1-en-1-yl)but-3-en-2-ylidene] cyclohex-2-en-1-ylidene}acetaldehyde	
23	4J4Q [62]	Class A α	Rho	B-Octylglucoside	Agonist-stabilized active receptor conformation
24	2VT4 [69]	Class A α	ADRB ₁	Cyanopindolol	Antagonist
25	2Y00 [70]	Class A α	ADRB ₁	Dobutamine	Partial Agonist
26	2Y01 [70]	Class A α	ADRB ₁	Dobutamine	Partial Agonist
27	2Y02 [70]	Class A α	ADRB ₁	Carmoterol	Agonist
28	2Y03 [70]	Class A α	ADRB ₁	Isoprenaline	Agonist
29	2Y04 [70]	Class A α	ADRB ₁	Salbutamol	Partial Agonist
30	2YCW [71]	Class A α	ADRB ₁	Carazolol	Antagonist
31	2YCX [71]	Class A α	ADRB ₁	Cyanopindolol	Antagonist
32	2YCY [71]	Class A α	ADRB ₁	Cyanopindolol	Antagonist
33	2YCZ [71]	Class A α	ADRB ₁	Iodocyanopindolol	Antagonist
34	3ZPQ [72]	Class A α	ADRB ₁	4-(Piperazin-1-yl)-1H-indole	Antagonist
35	3ZPR [72]	Class A α	ADRB ₁	4-Methyl-2-(piperazin-1-yl)quinoline	Antagonist

36	4AMI [73]	Class A α	ADRB ₁	Bucindolol	Agonist
37	4AMJ [73]	Class A α	ADRB ₁	Carvedilol	Agonist
38	4BVN [74]	Class A α	ADRB ₁	Cyanopindolol	Antagonist
39	5A8E [76]	Class A α	ADRB ₁	7-methylcyanopindolol	Inverse Agonist
40	5F8U [77]	Class A α	ADRB ₁	4-[[{(2S)-3-(tert-butylamino)-2-hydroxypropyl]oxy}-3H-indole-2-carbonitrile	Antagonist
41	2RH1 [26]	Class A α	ADRB ₂	Carazolol	Antagonist
42	3D4S [78]	Class A α	ADRB ₂	Timolol maleate	Antagonist
43	3NY8 [80]	Class A α	ADRB ₂	ICI 118,551	Antagonist
44	3NY9 [80]	Class A α	ADRB ₂	Ethyl 4-((2S)-2-hydroxy-3-[(1-methylethylamino)propyl]oxy)-3-methyl-1-benzofuran-2-carboxylate	Antagonist
45	3NYA [80]	Class A α	ADRB ₂	Alprenolol	Antagonist
46	3POG [81]	Class A α	ADRB ₂	BI 167107	Agonist
47	3PDS [82]	Class A α	ADRB ₂	FAUC50	Agonist
48	3SN6 [18]	Class A α	ADRB ₂	BI 167107	Agonist
49	4GBR [83]	Class A α	ADRB ₂	Carazolol	Antagonist
50	4LDE [84]	Class A α	ADRB ₂	BI 167107	Agonist
51	4LDL [84]	Class A α	ADRB ₂	Hydroxybenzyl isoproterenol	Agonist
52	4LDO [84]	Class A α	ADRB ₂	Epinephrine	Agonist
53	4QKX [85]	Class A α	ADRB ₂	FAUC37	Agonist
54	5D5A [86]	Class A α	ADRB ₂	Carazolol	Antagonist
55	5D5B [86]	Class A α	ADRB ₂	Carazolol	Antagonist
56	5D6L [Ma, P; Caffrey, M]	Class A α	ADRB ₂	Carazolol	Antagonist
57	5JQH [87]	Class A α	ADRB ₂	Carazolol	Antagonist
58	3RZE [88]	Class A α	H ₁ R	Doxepin (E/Z)	Antagonist
59	3PBL [6]	Class A α	D ₃ R	Eticlopride	Antagonist
60	4IAQ [89]	Class A α	5-HT _{1B}	Dihydroergotamine	Agonist
61	4IAR [89]	Class A α	5-HT _{1B}	Ergotamine	Agonist
62	4IB4 [90]	Class A α	5-HT _{2B}	Ergotamine	Agonist
63	4NC3 [91]	Class A α	5-HT _{2B}	Ergotamine	Agonist
64	5TVN [92]	Class A α	5-HT _{2B}	(8 α)-N,N-diethyl-6-methyl-9,10-didehydroergoline- 8-carboxamide	Agonist
65	5CXV [93]	Class A α	M ₁ R	Tiotropium	Antagonist
66	3UON [94]	Class A α	M ₂ R	3-quinuclidinyl-benzilate	Antagonist
67	4MQS [95]	Class A α	M ₂ R	Iperoxo	Agonist
68	4MQT [95]	Class A α	M ₂ R	Iperoxo/LY2119620	Agonist/ Allosteric modulator
69	4DAJ [96]	Class A α	M ₃ R	Tiotropium	Antagonist
70	4U14 [97]	Class A α	M ₃ R	Tiotropium	Antagonist
71	4U15 [97]	Class A α	M ₃ R	Tiotropium	Antagonist
72	4U16 [97]	Class A α	M ₃ R	N-methyl scopolamine	Antagonist
73	5DSG [93]	Class A α	M ₄ R	Tiotropium	Antagonist

74	5UEN [98]	Class A α	A _{1A} R	DU172	Antagonist
75	2YDO [101]	Class A α	A _{2A} R	Adensoine	Agonist
76	2YDV [101]	Class A α	A _{2A} R	NECA	Agonist
77	3EML [36]	Class A α	A _{2A} R	ZM241385	Antagonist
78	3PWH [99]	Class A α	A _{2A} R	ZM241385	Antagonist
79	3QAK [100]	Class A α	A _{2A} R	UK-432097	Agonist
80	3REY [99]	Class A α	A _{2A} R	XAC	Antagonist
81	3RFM [99]	Class A α	A _{2A} R	Caffeine	Antagonist
82	3UZA [103]	Class A α	A _{2A} R	6-(2,6-Dimethylpyridin-4-yl)-5-phenyl-1,2,4-triazin-3-amine	Antagonist
83	3UZC [103]	Class A α	A _{2A} R	4-(3-amino-5-phenyl-1,2,4-triazin-6-yl)-2-chlorophenol	Antagonist
84	3VG9 [102]	Class A α	A _{2A} R	ZM241385	Antagonist
85	3VGA [102]	Class A α	A _{2A} R	ZM241385	Antagonist
86	4E1Y [104]	Class A α	A _{2A} R	ZM241385	Antagonist
87	4UG2 [105]	Class A α	A _{2A} R	CGS21680	Agonist
88	4UHR [105]	Class A α	A _{2A} R	CGS21680	Agonist
89	5G53 [108]	Class A α	A _{2A} R	DB03719	Agonist
90	5IU4 [106]	Class A α	A _{2A} R	ZM241385	Antagonist
91	5IU7 [106]	Class A α	A _{2A} R	Compound 12c	Antagonist
92	5IU8 [106]	Class A α	A _{2A} R	Compound 12f	Antagonist
93	5IUA [106]	Class A α	A _{2A} R	Compound 12b	Antagonist
94	5IUB [106]	Class A α	A _{2A} R	Compound 12x	Antagonist
95	5K2A [107]	Class A α	A _{2A} R	ZM241385	Antagonist
96	5K2B [107]	Class A α	A _{2A} R	ZM241385	Antagonist
97	5K2C [107]	Class A α	A _{2A} R	ZM241385	Antagonist
98	5K2D [107]	Class A α	A _{2A} R	ZM241385	Antagonist
99	5UIG [109]	Class A α	A _{2A} R	5-amino-N-[(2-methoxyphenyl)methyl]-2-(3-methylphenyl)-2H-1,2,3-triazole-4-carboximidamide	Antagonist
100	3V2W [110]	Class A α	S1P ₁	ML056	Antagonist
101	3V2Y [110]	Class A α	S1P ₁	ML056	Antagonist
102	4Z34 [111]	Class A α	LPA ₁	ONO9780307	Antagonist
103	4Z35 [111]	Class A α	LPA ₁	ONO-9910539	Antagonist
104	4Z36 [111]	Class A α	LPA ₁	ONO-3080573	Antagonist
105	5TGZ [112]	Class A α	CB ₁	AM6538	Antagonist
106	5U09 [113]	Class A α	CB ₁	Taranabant	Antagonist
107	3ZEV [115]	Class A β	NTSR ₁	Neurotensin	Agonist
108	4BUO [115]	Class A β	NTSR ₁	Neurotensin	Agonist
109	4BV0 [115]	Class A β	NTSR ₁	Neurotensin	Agonist
110	4BWB [115]	Class A β	NTSR ₁	Neurotensin	Agonist
111	4GRV [114]	Class A β	NTSR ₁	Neurotensin	Agonist
112	4XEE [116]	Class A β	NTSR ₁	Neurotensin	Agonist
113	4XES [116]	Class A β	NTSR ₁	Neurotensin	Agonist
114	5T04 [117]	Class A β	NTSR ₁	Neurotensin	Agonist
115	4ZJ8 [40]	Class A β	OX ₁	Suvorexant	Dual Antagonist

116	4ZJC [40]	Class A β	OX ₁	SB-674042	Antagonist
117	4S0V [118]	Class A β	OX ₂	Suvorexant	Dual Antagonist
118	5GLH [120]	Class A β	ET-B	Endothelin-1	Agonist
119	5T1A [122]	Class A γ	CCR2	BMS-681/CCR2-RA-[R]	Orthosteric/allos- teric antagonist
120	3ODU [123]	Class A γ	CXCR4	IT1t	Antagonist
121	3OE0 [123]	Class A γ	CXCR4	CVX15	Antagonist
122	3OE6 [123]	Class A γ	CXCR4	IT1t	Antagonist
123	3OE8 [123]	Class A γ	CXCR4	IT1t	Antagonist
124	3OE9 [123]	Class A γ	CXCR4	IT1t	Antagonist
125	4MBS [125]	Class A γ	CCR5	Maraviroc	Antagonist
126	5LWE [126]	Class A γ	CCR9	Vercirnon	Antagonist
127	4EA3 [38]	Class A γ	NOP	Banyu Compound-24 (C-24)	Antagonist
128	5DHG [127]	Class A γ	NOP	Compound-35 (C-35)	Antagonist
129	5DHH [127]	Class A γ	NOP	SB-612111	Antagonist
130	4DJH [128]	Class A γ	κ -OR	JDTic	Antagonist
131	4DKL [129]	Class A γ	μ -OR	Beta-FNA	Antagonist
132	5C1M [130]	Class A γ	μ -OR	BU72	Agonist
133	4EJ4 [131]	Class A γ	δ -OR	Naltrindole	Antagonist
134	4N6H [132]	Class A γ	δ -OR	Naltrindole	Antagonist
135	4RWA [133]	Class A γ	δ -OR	Bifunctional Peptide H-Dmt(1)-Tic (2)- Phe(3)-Phe(4)-NH ₂ (DIPP-NH ₂)	Antagonist
136	4RWD [133]	Class A γ	δ -OR	Bifunctional Peptide H-Dmt(1)-Tic (2)- Phe(3)-Phe(4)-NH ₂ (DIPP-NH ₂)	Antagonist
137	4YAY [134]	Class A γ	AT ₁ R	ZD7155	Antagonist
138	4ZUD [135]	Class A γ	AT ₁ R	Olmesartan	Inverse Agonist
139	5UNF [136]	Class A γ	AT ₂ R	N-benzyl-N-(2-ethyl-4-oxo-3-{{[2'- (2H-tetrazol-5-yl)[1,1'-biphenyl]-4- yl]methyl}-3,4-dihydroquinazolin- 6-yl}thiophene-2-carboxamide	
140	5UNG [136]	Class A γ	AT ₂ R	N-benzyl-N-(2-ethyl-4-oxo-3-{{[2'- (2H-tetrazol-5-yl)[1,1'-biphenyl]-4- yl]methyl}-3,4-dihydroquinazolin-6- yl}thiophene-2-carboxamide	
141	5UNH [136]	Class A γ	AT ₂ R	N-[(furan-2-yl)methyl]-N-(4-oxo- 2-propyl-3-{{[2'-(2H-tetrazol-5-yl) [1,1'-biphenyl]-4-yl]methyl}-3,4- dihydroquinazolin-6-yl}benzamide	
142	4XNW [138]	Class A δ	P2Y ₁	MRS2500	Antagonist
143	4XNV [138]	Class A δ	P2Y ₁	BPTU	Antagonist
144	4NTJ [41]	Class A δ	P2Y ₁₂	AZD1283	Antagonist
145	4PXZ [139]	Class A δ	P2Y ₁₂	2MeSADP	Agonist
146	4PY0 [139]	Class A δ	P2Y ₁₂	2MeSADP	Agonist
147	3VW7 [140]	Class A δ	PAR1	Vorapaxar	Antagonist
148	5EE7 [143]	Class B	GCGR	MK-0893	Antagonist
149	4K5Y [144]	Class B	CRF ₁ R	CP-376395	Antagonist

150	4Z9G [145]	Class B	CRF ₁ R	CP-376395	Antagonist
151	4OR2 [146]	Class C	mGlu ₁	FITM	Negative Allosteric Modulator
152	4O09 [37]	Class C	mGlu ₅	Mavoglurant	Negative Allosteric Modulator
153	5CGC [147]	Class C	mGlu ₅	HTL14242	Antagonist
154	5CGD [147]	Class C	mGlu ₅	HTL14242	Antagonist
155	4JKV [39]	Class F	SMO	LY2940680	Antagonist
156	4N4W [148]	Class F	SMO	SANT-1	Antagonist
157	4O9R [85]	Class F	SMO	Cyclopamine	Antagonist
158	4QIM [148]	Class F	SMO	Anta XV	Antagonist
159	4QIN [148]	Class F	SMO	SAG1.5	Agonist
160	5L7D [149]	Class F	SMO	Cholesterol	
161	5L7I [149]	Class F	SMO	Vismodegib	Antagonist

The γ -subgroup of *Rhodopsin* (class A) contains the SOG, MCH, and chemokine receptor clusters, which bind both peptide and small ligand-like compounds. The SOG receptor cluster members include GPR54, the somatostatin receptors (SSTRs), and the opioid receptors [3]. The opioid receptors are important drug targets for the treatment of pain, cough, and alcoholism [159]. The MCH cluster includes receptors that branch off the SOG cluster. These receptors bind melanin-concentrating hormone (MCH), whereas the chemokine receptor cluster contains the chemokine receptors, the angiotensin/bradykinin-related receptors, and additional orphan receptors. Most ligands that interact with these receptors are peptides such as the chemokines and angiotensin. The chemokine receptors are drug targets due to their roles in acute and chronic inflammation [3] and as co-receptors engaged by some HIV strains [165]. While there are chemokine receptor-targeting drugs in clinical stages, maraviroc, an antagonist for CCR5, is the only currently FDA-approved drug on the market [166] targeting this class.

The δ -subgroup of *Rhodopsin* (class A) includes four main groups, the MAS-related receptor, glycoprotein receptor, purine receptor, and the olfactory receptor clusters. The MAS-receptor cluster includes the MAS1 oncogene receptor and the MAS-related receptors. The glycoprotein receptor cluster contains the classic glycoprotein hormone receptors and the leucine-rich-repeat-containing G protein-coupled receptors, whereas the purine receptor cluster is the largest in the δ -subgroup made up of the formyl peptide receptors, the nucleotide receptors, and a number of orphan receptors [3].

5.1. Extracellular region

The extracellular region of GPCR contains the N-terminus and three loops (ECL1–3) that shape the opening to the ligand-binding pocket. The ligand-binding region is found within the extracellular region of the 7TM bundle. EL2 (Figure 5B) is known to vary in length between GPCR classes resulting in distinct conformations, while EL1 (Figure 5A) and EL3 (Figure 5C) are short and often have disordered structures [167]. A highly conserved disulfide bond between EL2 and C3.25 (see

description of the Ballesteros-Weinstein numbering system in the 7TM region section) has been observed in a majority of crystallized GPCR structures. This covalent modification limits movement and stabilizes the conformation of EL2 [167,168]. Compared to other classes, class A receptors have a shorter N-terminus [168].

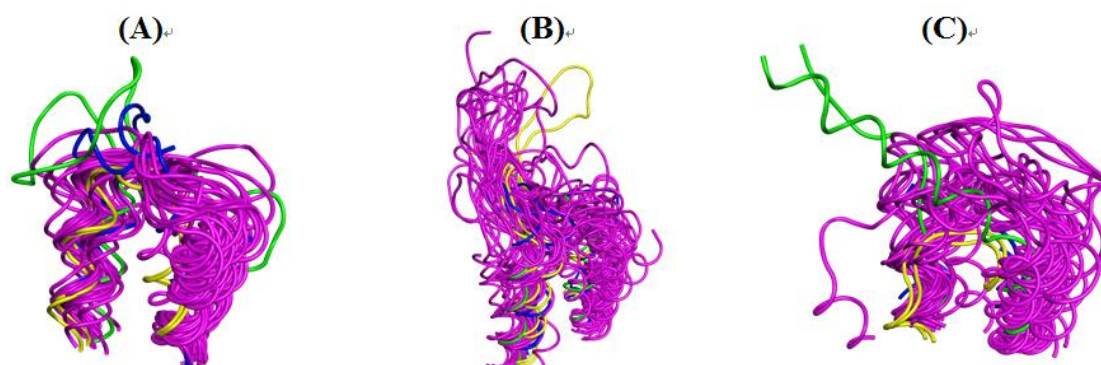


Figure 5. Superpositions of 31 class A (fuchsia), 2 class B (blue), 2 class C (yellow), and 1 class F (green) extracellular loops that are connected to the first two membrane-embedded helical turns are shown in ribbon structures where structures chosen did not have an N-terminal fusion partner. (A) EL1 and (C) EL3 are short in classes A through C and form mainly disordered structures that are mostly conserved among all classes. (B) EL2 is the longest of the EL, and varies in length and conformation in different GPCR. Class A GPCR: Rho (1F88), ADRB₁ (2VT4), ADRB₂ (2R4R), H₁R (3RZE), D₃R (3PBL), 5-HT_{1B} (4IAQ), 5-HT_{2B} (4IB4), M₁R (5CXV), M₂R (3UON), M₃R (4DAJ), M₄R (5DSG), A_{2A}R (3EML), S1P₁ (3V2W), LPA₁ (4Z32), CB₁ (5TGZ), NTSR₁ (3ZEV), OX₁ (4ZJ8), OX₂ (4S0V), TACR₁ (2KSP), CXCR1 (2LNL), CXCR4 (3ODU), CCR5 (4MBS), CCR9 (5LWE), κ-OR (4DJH), μ-OR (4DKL), δ-OR (4EJ4), HHV-5 US28 (4XT1), P2Y₁ (4XNW), P2Y₁₂ (4NTJ), PAR1 (3VW7), FFAR₁ (4PHU). Class B GPCR: GCGR (5EE7), CRF₁R (4K5Y). Class C GPCR: mGlu₁ (4OR2), mGlu₅ (4OO9). Class F GPCR: SMO (4QIN).

5.2. 7TM region

Despite differences in primary structure, a majority of the proteins in class A/*Rhodopsin* share conserved residues found within the 7TM domain [5]. The Ballesteros-Weinstein numbering system is often used to assign numbers to common residues that are conserved in both sequence and structure-based alignments of different receptors [169]. This indexing system consists of two numbers separated by a period. The first value represents the TM helix in which the residue is found and has values ranging from 1 to 7. The second number denotes the residue position relative to the most conserved residue within that TM segment, which is defined as position 50. The value decreases

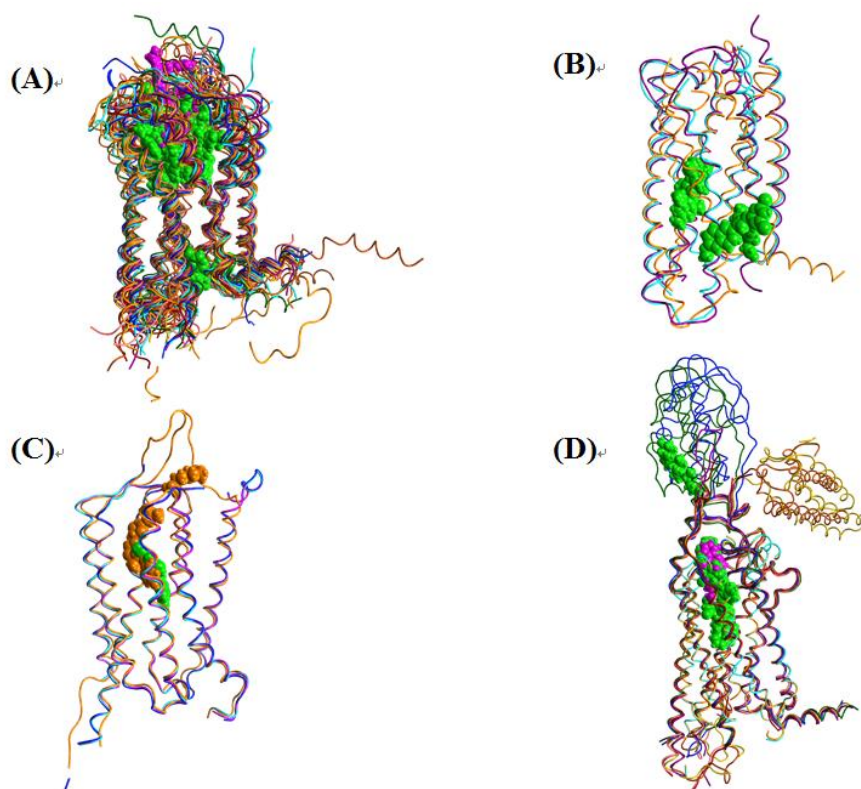


Figure 6. Superpositions of GPCR represented as thin ribbon structures with ligands as space-filling atoms colored accordingly by antagonists as green, agonists as fuchsia, and negative allosteric modulators as orange. (A) Structures of 29 individual class A GPCR. (B) Structures of 3 class B GPCR. (C) Structures of 4 class C GPCR. (D) Structures of 7 class F GPCR. Class A GPCR: Rho (1F88), ADRB₁ (2VT4), ADRB₂ (2RH1), H₁R (3RZE), D₃R (3PBL), 5-HT_{1B} (4IAQ), 5-HT_{2B} (4IB4), M₁R (5CXV), M₂R (3UON), M₃R (4DAJ), M₄R (5DSG), A_{2A}R (2YDO), S1P₁ (3V2W), LPA₁ (4Z34), CB₁ (5TGZ), NTSR₁ (3ZEV), OX₁ (4ZJ8), OX₂ (4S0V), CCR4 (3ODU), CCR5 (4MBS), CCR9 (5LWE), NOP (4EA3), κ-OR (4DJH), μ-OR (4DKL), δ-OR (4EJ4), AT₁R (4YAY), P2Y₁ (4XNW), P2Y₁₂ (4NTJ), PAR1 (3VW7). Class B GPCR: GCGR (5EE7), CRF₁R (4K5Y, 4Z9G). Class C GPCR: mGlu₁ (4OR2, 5CGC, 5CGD), mGlu₅ (4OO9) Class F GPCR: SMO (4JKV, 4N4W, 4OR9, 4QIM, 4QIN, 5L7D, 5L7I).

as you move through the amino acid sequence toward the N-terminus and increases as you move through the amino acid sequence toward the C-terminus. Class A/*Rhodopsin* receptors have highly conserved residues in each TM segment: N1.50, D2.50, R3.50, W4.50, P5.50, P6.50, and P7.50 [169]. In addition to these residues, class A/*Rhodopsin* contains two fingerprint regions: the D/ERY motif at positions 3.49–3.51 and the NPXXY motif at positions 7.49–7.53 [3]. In 2000, x-ray crystallography studies on rhodopsin (PDB: 1F88) confirmed that conserved residues formed interhelical networks important for stabilization and activation [25] as had previously been suggested [170]. Seven years later, two structures of ADRB₂ (PDB: 2RH1, 2R4R/2R4S) were determined and revealed TM structural similarity to rhodopsin [26,27]. The basic canonical architecture of this region has now been

observed in numerous examples of crystallized GPCR as is shown in Figure 4. It is interesting to note that an allosteric sodium ion binding pocket involving two conserved residues D2.50 and S3.39 was identified in the 1.8Å crystal structure of A_{2A} (PDB: 4EIY) [104,171]. This central Na⁺ pocket was formed by D2.50, S3.39, and three water molecules. Liu, et al., compared their inactive A_{2A} structure to an active A_{2A} (PDB 3QAK [100]) structure and found that the size of the central pocket in the active form could only support three water molecules without sufficient room for Na⁺ coordination. This suggests that Na⁺ may stabilize the inactive conformation of A_{2A} [104]. After the discovery of the Na⁺ pocket in A_{2A}, the same characteristic was seen in other inactive GPCR crystal structures. These other structures include representatives of the α , γ , and δ subgroups of class A [ADRB1 (PDB: 4BVN [74]), delta-opioid (δ -OR; PDB: 4N6H [132]), and protease-activated receptor 1 (PAR1; PDB: 3VW7 [140])], suggesting the sodium ion binding pocket is a common feature in class A GPCR.

Conformational changes in the 7TM bundle are required for signal transduction across the cell membrane following ligand binding. Experimentally determined crystal structures of ADRB₂ reflect a variety of activation states. In these structures, TM5, TM6, and TM7 have a critical function in GPCR activation due to clear differences in helical arrangement. The active state of ADRB₂ (PDB: 3P0G) exhibits altered conformations of TM5 and TM7 and a prominent outward shift of TM6 [18] in contrast to the inactive state (PDB: 2RH1) [26].

The 7TM region forms distinctive ligand-binding pockets for different receptors where varying size, shape, and electrostatics provide receptor-ligand selectivity. Aminergic and nucleotide receptors have small binding pockets buried inside the 7TM bundle while peptide receptors have large, more accessible binding pockets near the extracellular surface [167]. These diverse ligand binding pocket profiles are demonstrated with 29 class A GPCR in Figure 6A.

5.3. Intracellular region

The intracellular region of GPCR includes the C-terminus and three loops (ICL1–3) that interact with G proteins, β -arrestins, and other downstream effectors [4,10,167]. GPCR crystal structures have shown structural conservation in the short IL1 chain, though high levels of variability have been observed within IL2 and IL3 suggesting dynamic and/or unstable conformations. Differences have been observed in IL2 in both the D₃R and ADRB receptor structures. In a crystal structure of D₃R (PDB: 3PBL) [6], where there were two copies of the protein from the same unit cell, IL2 had 2.5 turns of an α -helical conformation for chain A in contrast to a disordered loop lacking electron density for chain B (Figure 7A). ADRB₁ [69] and ADRB₂ [26], which have an overall percent identity of 80% and nearly identical IL2 sequences, have displayed an α -helical conformation in ADRB₁ and unstructured conformation in ADRB₂ (Figure 7B; PDB: ADRB₁-2VT4, ADRB₂-2RH1). The three dimensional structure of IL2 may be dependent upon the functional state of the protein, as well as interactions with intracellular partners. IL3 exhibits the greatest length variability amongst IL, ranging from five to hundreds of residues and has been implicated in G protein selectivity. IL3 has been observed to form a disordered conformation or, more often, has been replaced by a fusion partner for increased conformational homogeneity to enable crystal formation in several solved GPCR structures [10]. There are crystallographic structures of rhodopsin (PDB: 3CAP) [56],

ADRB₁ (PDB: 2YCW, 2YCX, 2YCY, 2YCZ) [71], and δ -opioid receptor (PDB: 4N6H) [132] where IL3 adopts an α -helical conformation resulting in extended TM5 and TM6 helices [10,167]. The IL region undergoes considerable conformational changes required for G protein interaction and the consequent initiation of the signal transduction cascade.

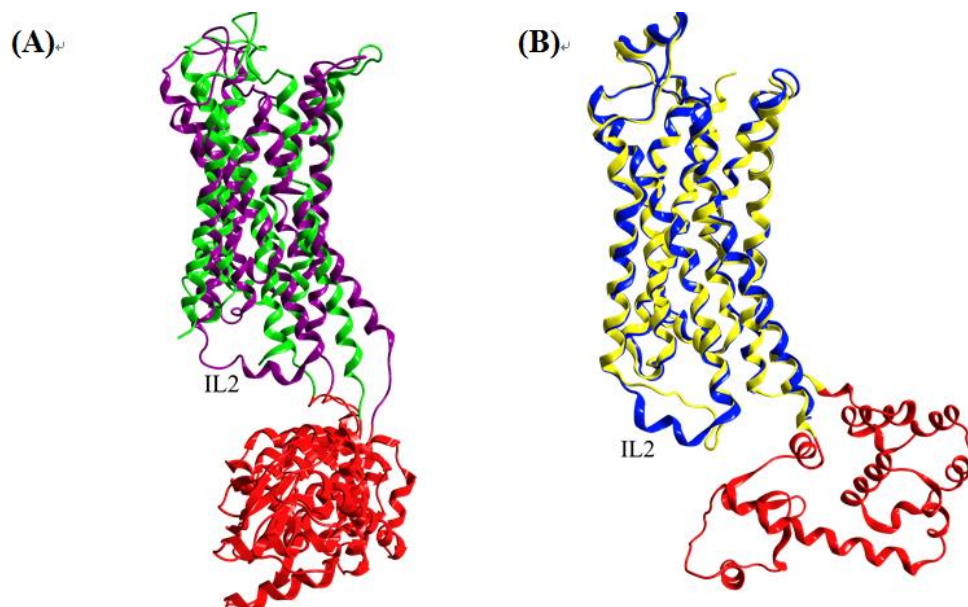


Figure 7. (A) Superposition of D₃R crystal structures (PDB: 3PBL) showing a 2.5 helical turn for chain A (purple) and no electron density for chain B (green) for IL2. The T4 lysozyme fusion partner replacing IL3 is shown in red for both chains. (B) Superposition of crystal structures comparing ADRB₁ (PDB: 2VT4 chain B; blue) and ADRB₂ (PDB: 2RH1 chain A; yellow) and T4 lysozyme in red) where IL2 exhibits an ordered structure in ADRB₁ and a disordered structure in ADRB₂.

6. Secretin and Adhesion (Class B)

Class B, the second largest class within the GPCR family, is comprised of the *Secretin* and *Adhesion* families. *Secretin* and *Adhesion* receptors have been classified together due to sequence similarities between their 7TM regions, although they have distinctions elsewhere that establish them as separate families. The *Secretin* family has an extracellular hormone-binding domain that interacts with peptide hormones [5]. The members of this family include the calcitonin/calcitonin-like receptors, the corticotropin-releasing hormone receptors, the glucagon receptor, the gastric inhibitory polypeptide receptor, the glucagon-like peptide receptors, the growth-hormone-releasing hormone receptor, the adenylyl cyclase activating polypeptide hormone receptor, the parathyroid hormone receptors, the secretin receptor, the vasoactive intestinal peptide receptors, and additional orphan receptors [3,5]. The *Secretin* family includes potential targets for drug development due to their involvement in central homeostatic functions. Members of this family have been connected to appetite regulation and type-2 diabetes [5].

The *Adhesion* family includes the epidermal growth factor receptors and lectomedin receptors, though a majority of the members are currently classified as orphan receptors [5]. This GPCR family exhibits a highly variable number of amino acids at the N-terminal region, ranging from 200–2800 in length, making this family phylogenetically and structurally different from the rest of the class B GPCR. The “*Adhesion*” family name is related to the N-terminal region that contains sequence motifs, such as the GPCR proteolysis (GPS) motif, which serve as intracellular autocatalytic processing sites that participate in cellular adhesion [3,5]. *Adhesion* family receptors bind extracellular matrix molecules rather than peptide hormones [5]. These receptors are believed to be involved in cell proliferation/migration, as well as immune system function via the mediation of leukocyte and neutrophil interactions. *Adhesion* receptors that contain long N-termini have become targets of monoclonal antibodies used as drugs candidates [5]. Numerous receptors from this family are localized in the central nervous system (CNS) [172] though their functional role in the CNS is not fully understood [5].

6.1. Structure

Within class B, sequence alignments show that *Adhesion* and *Secretin* families contain structural differences at the N-terminal region. *Adhesion* receptors contain distinctive O- and N-glycosylation sites, as well as EGF and GPS motifs [173]. *Secretin* family members have a 60–80 amino acid N-terminal domain [3] that include a hormone-binding (HRM) domain that is believed to have a key role in binding peptide hormones [168]. The Ballesteros-Weinstein numbering system somewhat extends to class B/*Adhesion/Secretin* where residues E3.50 and W4.50 are conserved [174]. The binding pockets of class B GPCR are broader and deeper inside the 7TM bundle compared to class A [151], as illustrated in Figure 6B, in order to accommodate endogenous peptide ligands. On the other hand, the crystal structure for GCGR (PDB: 5EE7) shows an allosteric binding site outside of the 7TM domain between TM6 and TM7 [143].

7. Glutamate (Class C)

The *Glutamate* (class C) family of receptors consists of the metabotropic glutamate receptors, GABA receptors, single calcium-sensing receptors, and sweet and umami taste receptors. The known endogenous ligands of this family are known to bind to the N-terminal region, but many allosteric ligands have been discovered to interact with TM3, TM5, TM6, and TM7 [175,176,177]. In addition, Ca^{2+} can bind to the extracellular region [178] and enhance the effects of glutamate in some glutamate receptors [179,180]. Many of the Ca^{2+} interacting residues are conserved within the “Venus flytrap” region of the *Glutamate* (class C) receptors [178] and have potential significance in drug design targeting depression learning, and memory [181].

7.1. Structure

In class C/*Glutamate* receptors, the 280–580 amino acid N-terminus [3] forms a cavity surrounded by two lobes which close upon ligand binding through a process known as the “Venus

flytrap” mechanism (VFTM) [5,182]. This mechanism involves a ligand binding-induced conformational change that results in the formation of disulfide bonds between the N-terminus and 7TM domain [168,183]. Receptors in this class lack the conserved residues defined by the Ballesteros-Weinstein numbering scheme seen extensively in Class A and to a lesser extent in Class B GPCR. Although the conserved ligand binding pocket of most class C receptors is located within the extracellular region, there are allosteric binding sites within the 7TM bundle [5] as seen in Figure 6C. These allosteric binding sites may be a way to achieve receptor specificity using allosteric modulators.

8. Frizzled/Taste2 (Class F)

The *Frizzled/Taste2* (class F) group of receptors includes frizzled and smoothed receptors (SMO) [3,5]. The frizzled receptors are known to bind secreted glycoproteins [184] while the SMO receptor functions in a ligand-independent manner through the SMO and sonic hedgehog (SHH) complex [185]. Frizzled receptors are involved in cell fate, proliferation, and polarity through association with secreted glycoproteins at the cysteine-rich N-terminus [3,5]. Although the cysteine-rich region is highly conserved in the frizzled receptor, there is evidence that there are additional binding sites located in the extracellular loops of the TM regions [186]. SMO receptors are structurally similar to frizzled receptors [5]. The cysteine-rich region found in the frizzled receptors, as well as the chemical properties of residues that bind secreted glycoproteins, are conserved in SMO [186,187]. Class F GPCR, though not well understood, have been linked to cancer development and are potential targets for cancer therapy [5].

8.1. Structure

Based on sequence analysis, class F/*Frizzled/Taste2* is the most highly conserved class within the GPCR superfamily [168]. These proteins have a distinctive N-terminus that spans 200–320 amino acids in length [3,5], in addition to a variable linker region between the EL and TM domains [5]. While the Ballesteros-Weinstein numbering scheme does not apply to this class, these proteins still share the common structure of the 7TM hydrophobic core [188]. There is limited structural information about this class of receptors since SMO is the only class F GPCR to be crystallized to date. The binding pocket is observed to be narrow and elongated in the currently available crystal structures (Figure 6D). In one SMO crystal structure (PDB: 5L7D), cholesterol is bound to the extracellular cysteine-rich domain (CRD) that is highly conserved in vertebrates. An oxysterol was observed to displace cholesterol and bind to the CRD groove leading to SMO activation and Hedgehog (Hh) signaling. Cholesterol has been proposed as an endogenous ligand that stabilizes the inactive SMO conformation [149].

9. Conclusions

Since the initial crystallization of rhodopsin in 2000, numerous technological advances have significantly impacted GPCR structure determination efforts, resulting in crystal structures of 42

individual receptors (to date). Currently more than 180 structures of GPCR have been solved and made available through the Protein Data Bank (Table 1). Of these, over 150 protein-ligand complexes are available (Table 2), representing the entire spectrum of ligand functions from inverse agonist to agonist. These data sets give insights into structural features and ligand recognition within this diverse protein superfamily. Specifically, the majority of crystallized ligand complexes with GPCR exhibit a ligand binding pocket near the extracellular end of the TM helical bundle, as shown in Figure 6. The TM helical bundle structure shows considerable structural similarity across the family (Figure 4), while EL2 shows the greatest structural diversity in the vicinity of the ligand binding pocket (Figure 5). Thus, differences in ligand selectivity between GPCR family members are driven both by sidechain differences in the TM helical bundle (rather than backbone conformational differences) as well as overall EL2 fold. GPCR have essential biological roles, and many have been confirmed to have value as therapeutic targets. Therefore solved, three-dimensional GPCR structures have tremendous potential to influence drug discovery. Structure-based drug discovery approaches can be applied directly to crystallized GPCR family members as therapeutic targets. Additionally, the crystallized GPCR structures exhibit close sequence identity to additional GPCR family members and can serve as templates for the development of reliable and predictive homology models. Validated models allow structure-based drug discovery approaches to be used against this even broader set of target GPCR members. Although efforts in GPCR crystal structure determination over the past two decades have been fruitful, vast amounts of work remain to characterize unrepresented family members that have lower sequence identities to currently crystallized GPCR family members. GPCR structural characterization will continue to be a rich research area in need of further advances and innovative approaches into the foreseeable future.

Acknowledgment

Research reported in this publication was supported by the National Institute of Mental Health of the National Institutes of Health under Award Number R15MH109034. The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health.

Conflict of Interest

The authors declare no conflict of interest in this paper.

References

1. Venter JC, Adams MD, Myers EW, et al. (2001) The sequence of the human genome. *Science* 291: 1304–1351.
2. Lander ES, Linton LM, Birren B, et al. (2001) Initial sequencing and analysis of the human genome. *Nature* 409: 860–921.

3. Fredriksson R, Lagerstrom MC, Lundin LG, et al. (2003) The G-protein-coupled receptors in the human genome form five main families. Phylogenetic analysis, paralogon groups, and fingerprints. *Mol Pharmacol* 63: 1256–1272.
4. Pierce KL, Premont RT, Lefkowitz RJ (2002) Seven-transmembrane receptors. *Nat Rev Mol Cell Biol* 3: 639–650.
5. Lagerstrom MC, Schioth HB (2008) Structural diversity of G protein-coupled receptors and significance for drug discovery. *Nat Rev Drug Discov* 7: 339–357.
6. Chien EY, Liu W, Zhao Q, et al. (2010) Structure of the human dopamine D3 receptor in complex with a D2/D3 selective antagonist. *Science* 330: 1091–1095.
7. Yang J, Villar VA, Armando I, et al. (2016) G protein-coupled receptor kinases: Crucial regulators of blood pressure. *J Am Heart Assoc* 5: e003519.
8. Bar-Shavit R, Maoz M, Kancharla A, et al. (2016) G protein-coupled receptors in cancer. *Int J Mol Sci* 17: 1320.
9. Flower DR (1999) Modelling G-protein-coupled receptors for drug design. *Biochim Biophys Acta* 1422: 207–234.
10. Katritch V, Cherezov V, Stevens RC (2012) Diversity and modularity of G protein-coupled receptor structures. *Trends Pharmacol Sci* 33: 17–27.
11. Gether U, Ballesteros JA, Seifert R, et al. (1997) Structural instability of a constitutively active G protein-coupled receptor. Agonist-independent activation due to conformational flexibility. *J Biol Chem* 272: 2587–2590.
12. Seifert R, Wenzel-Seifert K (2002) Constitutive activity of G-protein-coupled receptors: cause of disease and common property of wild-type receptors. *Naunyn Schmiedebergs Arch Pharmacol* 366: 381–416.
13. Nelson G, Hoon MA, Chandrashekar J, et al. (2001) Mammalian sweet taste receptors. *Cell* 106: 381–390.
14. Nelson G, Chandrashekar J, Hoon MA, et al. (2002) An amino-acid taste receptor. *Nature* 416: 199–202.
15. Milligan G (2004) G protein-coupled receptor dimerization: Function and ligand pharmacology. *Mol Pharmacol* 66.
16. Syrovatkina V, Alegre KO, Dey R, et al. (2016) Regulation, signaling, and physiological functions of G-proteins. *J Mol Biol* 428: 3850–3868.
17. Wess J (1997) G-protein-coupled receptors: molecular mechanisms involved in receptor activation and selectivity of G-protein recognition. *FASEB J* 11: 346–354.
18. Rasumussen SG, DeVree BT, Zou Y, et al. (2011) Crystal structure of the beta2 adrenergic receptor-Gs protein complex. *Nature* 477: 549–555.
19. Gilman AG (1987) G proteins: transducers of receptor-generated signals. *Annu Rev Biochem* 56: 615–649.
20. Nie J, Lewis DL (2001) Structural domains of the CB1 cannabinoid receptor that contribute to constitutive activity and G-protein sequestration. *J Neurosci* 21: 8758–8764.
21. Kobilka BK, Deupi X (2007) Conformational complexity of G-protein-coupled receptors. *Trends Pharmacol Sci* 28: 397–406.

22. Ross EM, Wilkie TM (2000) GTPase-activating proteins for heterotrimeric G proteins: regulators of G protein signaling (RGS) and RGS-like proteins. *Annu Rev Biochem* 69: 795–827.
23. Claing A, Laporte SA, Caron MG, et al. (2002) Endocytosis of G protein-coupled receptors: roles of G protein-coupled receptor kinases and beta-arrestin proteins. *Prog Neurobiol* 66: 61–79.
24. Okada T, Le TI, Fox BA, et al. (2000) X-Ray diffraction analysis of three-dimensional crystals of bovine rhodopsin obtained from mixed micelles. *J Struct Biol* 130: 73–80.
25. Palczewski K, Kumasaka T, Hori T, et al. (2000) Crystal structure of rhodopsin: A G protein-coupled receptor. *Science* 289: 739–745.
26. Cherezov V, Rosenbaum DM, Hanson MA, et al. (2007) High resolution crystal structure of an engineered human beta2-adrenergic G protein-couple receptor. *Science* 318: 1258–1265.
27. Rasmussen SG, Choi HJ, Rosenbaum DM, et al. (2007) Crystal structure of the human beta2 adrenergic G-protein-coupled receptor. *Nature* 450: 383–387.
28. Stroud RM (2011) New tools in membrane protein determination. *F1000 Biol Rep* 3: 8.
29. Ujwal R, Bowie JU (2011) Crystallizing membrane proteins using lipidic bicelles. *Methods* 55: 337–341.
30. Denisov IG, Sligar SG (2016) Nanodiscs for structural and functional studies of membrane proteins. *Nat Struct Mol Biol* 23: 481–486.
31. Xiang J, Chun E, Liu C, et al. (2016) Successful strategies to determine high-resolution structures of GPCRs. *Trends Pharmacol Sci* 37: 1055–1069.
32. Rawlings AE (2016) Membrane proteins: always an insoluble problem? *Biochem Soc Trans* 44: 790–795.
33. Neubig RR, Siderovski DP (2002) Regulators of G-protein signalling as new central nervous system drug targets. *Nat Rev Drug Discov* 1: 187–197.
34. Bayburt TH, Grinkova YV, Sligar SG (2002) Self-assembly of discoidal phospholipid bilayer nanoparticles with membrane scaffold proteins. *Nano Lett* 2: 853–856.
35. Delmar JA, Bolla JR, Su CC, et al. (2015) Crystallization of membrane proteins by vapor diffusion. *Methods Enzymol* 557: 363–392.
36. Jaakola VP, Griffith MT, Hanson MA, et al. (2008) The 2.6 angstrom crystal structure of a human A2A adenosine receptor bound to an antagonist. *Science* 322: 1211–1217.
37. Dore AS, Okrasa K, Patel JC, et al. (2014) Structure of class C GPCR metabotropic glutamate receptor 5 transmembrane domain. *Nature* 511: 557–562.
38. Thompson AA, Liu W, Chun E, et al. (2012) Structure of the nociceptin/orphanin FQ receptor in complex with a peptide mimetic. *Nature* 485: 395–399.
39. Wang C, Wu H, Katritch V, et al. (2013) Structure of the human smoothed receptor bound to an antitumour agent. *Nature* 497: 338–343.
40. Yin J, Babaoglu K, Brautigam CA, et al. (2016) Structure and ligand-binding mechanism of the human OX1 and OX2 orexin receptors. *Nat Struct Mol Biol* 23: 293–299.
41. Zhang K, Zhang J, Gao Z, et al. (2014) Structure of the human P2Y₁₂ receptor in complex with an antithrombotic drug. *Nature* 509.
42. Isberg V, Mordalski S, Munk C, et al. (2016) GPCRdb: an information system for G protein-coupled receptors. *Nucleic Acids Res*.

43. Berman HM, Westbrook J, Feng Z, et al. (2000) The protein data bank. *Nucl Acids Res* 28: 235–242.
44. Teller DC, Okada T, Behnke CA, et al. (2001) Advances in determination of a high-resolution three-dimensional structure of rhodopsin, a model of G-protein-coupled receptors (GPCRs). *Biochemistry* 40: 7761–7772.
45. Yeagle PL, Choi G, Albert AD (2001) Studies on the structure of the G-protein-coupled receptor rhodopsin including the putative G-protein binding site in unactivated and activated forms. *Biochemistry* 40: 11932–11937.
46. Okada T, Fujiyoshi Y, Silow M, et al. (2002) Functional role of internal water molecules in rhodopsin revealed by X-ray crystallography. *Proc Natl Acad Sci USA* 99: 5982–5987.
47. Choi G, Landin J, Galan JF, et al. (2002) Structural studies of metarhodopsin II, the activated form of the G-protein coupled receptor, rhodopsin. *Biochemistry* 41: 7318–7324.
48. Li J, Edwards PC, Burghammer M, et al. (2004) Structure of bovine rhodopsin in a trigonal crystal form. *J Mol Biol* 343: 1409–1438.
49. Okada T, Sugihara M, Bondar AN, et al. (2004) The retinal conformation and its environment in rhodopsin in light of a new 22 Å crystal structure. *J Mol Biol* 342: 571–583.
50. Nakamichi H, Okada T (2006) Crystallographic analysis of primary visual photochemistry. *Angew Chem Int Ed Engl* 45: 4270–4273.
51. Nakamichi H, Okada T (2006) Local peptide movement in the photoreaction intermediate of rhodopsin. *Proc Natl Acad Sci USA* 103: 12729–12734.
52. Salom D, Lodowski DT, Stenkamp RE, et al. (2006) Crystal structure of a photoactivated deprotonated intermediate of rhodopsin. *Proc Natl Acad Sci USA* 103: 16123–16128.
53. Standfuss J, Xie G, Edwards PC, et al. (2007) Crystal structure of a thermally stable rhodopsin mutant. *J Mol Biol* 372: 1179–1188.
54. Nakamichi H, Buss V, Okada T (2007) Photoisomerization mechanism of rhodopsin and 9-cis-rhodopsin revealed by x-ray crystallography. *Biophys J* 92: L106–L108.
55. Stenkamp RE (2008) Alternative models for two crystal structures of bovine rhodopsin. *Acta Crystallogr D* 64: 902–904.
56. Park JH, Scheerer P, Hofmann KP, et al. (2008) Crystal structure of the ligand-free G-protein-coupled receptor opsin. *Nature* 454: 183–187.
57. Scheerer P, Park JH, Hildebrand PW, et al. (2008) Crystal structure of opsin in its G-protein-interacting conformation. *Nature* 455: 497–502.
58. Standfuss J, Edwards PC, D'Antona A, et al. (2011) The structural basis of agonist-induced activation in constitutively active rhodopsin. *Nature* 471: 656–660.
59. Makino CL, Riley CK, Looney J, et al. (2010) Binding of more than one retinoid to visual opsins. *Biophys J* 99: 2366–2373.
60. Choe HW, Kim YJ, Park JH, et al. (2011) Crystal structure of metarhodopsin II. *Nature* 471: 651–655.
61. Deupi X, Edwards P, Singhal A, et al. (2012) Stabilized G protein binding site in the structure of constitutively active metarhodopsin-II. *Proc Natl Acad Sci USA* 109: 119–124.
62. Park JH, Morizumi T, Li Y, et al. (2013) Opsin, a structural model for olfactory receptors? *Angew Chem Int Ed Engl* 52: 11021–11024.

63. Singhal A, Ostermaier MK, Vishnivetskiy SA, et al. (2013) Insights into congenital stationary night blindness based on the structure of G90D rhodopsin. *EMBO Rep* 14: 520–526.
64. Szczepek M, Beyriere F, Hofmann KP, et al. (2014) Crystal structure of a common GPCR-binding interface for G protein and arrestin. *Nat Commun* 5: 4801.
65. Blankenship E, Vahedi-Faridi A, Lodowski DT (2015) The high-resolution structure of activated opsin reveals a conserved solvent network in the transmembrane region essential for activation. *Structure* 23: 2358–2364.
66. Kang Y, Zhou XE, Gao X, et al. (2015) Crystal structure of rhodopsin bound to arrestin by femtosecond X-ray laser. *Nature* 523: 561–567.
67. Singhal A, Guo Y, Matkovic M, et al. (2016) Structural role of the T94I rhodopsin mutation in congenital stationary night blindness. *EMBO Rep* 17: 1431–1440.
68. Gulati S, Jastrzebska B, Banerjee S, et al. (2017) Photocyclic behavior of rhodopsin induced by an atypical isomerization mechanism. *Proc Natl Acad Sci USA* 114: E2608–E2615.
69. Warne T, Serrano-Vega MJ, Baker JG, et al. (2008) Structure of a beta1-adrenergic G-protein-coupled receptor. *Nature* 454: 486–491.
70. Warne T, Moukhametzianov R, Baker JG, et al. (2011) The structural basis for agonist and partial agonist action on a beta(1)-adrenergic receptor. *Nature* 469: 241–244.
71. Moukhametzianov R, Warne T, Edwards PC, et al. (2011) Two distinct conformations of helix 6 observed in antagonist-bound structures of a beta-1 adrenergic receptor. *Proc Natl Acad Sci USA* 108: 8228–8232.
72. Christopher JA, Brown J, Dore AS, et al. (2013) Biophysical fragment screening of the beta1-adrenergic receptor: identification of high affinity arylpiperazine leads using structure-based drug design. *J Med Chem* 56: 3446–3455.
73. Warne T, Edwards PC, Leslie AG, et al. (2012) Crystal structures of a stabilized beta1-adrenoceptor bound to the biased agonists bucindolol and carvedilol. *Structure* 20: 841–849.
74. Miller-Gallacher JL, Nehme R, Warne T, et al. (2014) The 2.1 Å resolution structure of cyanopindolol-bound beta1-adrenoceptor identifies an intramembrane Na⁺ ion that stabilises the ligand-free receptor. *PLoS One* 9: e92727.
75. Huang J, Chen S, Zhang JJ, et al. (2013) Crystal structure of oligomeric beta1-adrenergic G protein-coupled receptors in ligand-free basal state. *Nat Struct Mol Biol* 20: 419–425.
76. Sato T, Baker J, Warne T, et al. (2015) Pharmacological analysis and structure determination of 7-Methylcyanopindolol-bound beta1-adrenergic receptor. *Mol Pharmacol* 88: 1024–1034.
77. Leslie AG, Warne T, Tate CG (2015) Ligand occupancy in crystal structure of beta1-adrenergic G protein-coupled receptor. *Nat Struct Mol Biol* 22: 941–942.
78. Hanson MA, Cherezov V, Griffith MT, et al. (2008) A specific cholesterol binding site is established by the 2.8 Å structure of the human beta2-adrenergic receptor. *Structure* 16: 897–905.
79. Bokoch MP, Zou Y, Rasmussen SG, et al. (2010) Ligand-specific regulation of the extracellular surface of a G-protein-coupled receptor. *Nature* 463: 108–112.
80. Wacker D, Fenalti G, Brown MA, et al. (2010) Conserved binding mode of human beta2 adrenergic receptor inverse agonists and antagonist revealed by X-ray crystallography. *J Am Chem Soc* 132: 11443–11445.

81. Rasmussen SG, Choi HJ, Fung JJ, et al. (2011) Structure of a nanobody-stabilized active state of the beta(2) adrenoceptor. *Nature* 469: 175–180.
82. Rosenbaum DM, Zhang C, Lyons JA, et al. (2011) Structure and function of an irreversible agonist-beta(2) adrenoceptor complex. *Nature* 469: 236–240.
83. Zou Y, Weis WI, Kobilka BK (2012) N-terminal T4 lysozyme fusion facilitates crystallization of a G protein coupled receptor. *PLoS One* 7.
84. Ring AM, Manglik A, Kruse AC, et al. (2013) Adrenaline-activated structure of beta2-adrenoceptor stabilized by an engineered nanobody. *Nature* 502: 575–579.
85. Weichert D, Kruse AC, Manglik A, et al. (2014) Covalent agonists for studying G protein-coupled receptor activation. *Proc Natl Acad Sci USA* 111: 10744–10748.
86. Huang CY, Olieric V, Ma P, et al. (2016) In meso in situ serial X-ray crystallography of soluble and membrane proteins at cryogenic temperatures. *Acta Crystallogr D* 72: 93–112.
87. Staus DP, Strachan RT, Manglik A, et al. (2016) Allosteric nanobodies reveal the dynamic range and diverse mechanisms of G-protein-coupled receptor activation. *Nature* 535: 448–452.
88. Shimamura T, Shiroishi M, Weyand S, et al. (2011) Structure of the human histamine H1 receptor complex with doxepin. *Nature* 475: 65–70.
89. Wang C, Jiang Y, Ma J, et al. (2013) Structural basis for molecular recognition at serotonin receptors. *Science* 340: 610–614.
90. Wacker D, Wang C, Katritch V, et al. (2013) Structural features for functional selectivity at serotonin receptors. *Science* 340: 615–619.
91. Liu W, Wacker D, Gati C, et al. (2013) Serial femtosecond crystallography of G protein-coupled receptors. *Science* 342: 1521–1524.
92. Wacker D, Wang S, McCorvy JD, et al. (2017) Crystal structure of an LSD-bound human serotonin receptor. *Cell* 168: 377–389.
93. Thal DM, Sun B, Feng D, et al. (2016) Crystal structures of the M1 and M4 muscarinic acetylcholine receptors. *Nature* 531: 335–340.
94. Haga K, Kruse AC, Asada H, et al. (2012) Structure of the human M2 muscarinic acetylcholine receptor bound to an antagonist. *Nature* 482: 547–551.
95. Kruse AC, Ring AM, Manglik A, et al. (2013) Activation and allosteric modulation of a muscarinic acetylcholine receptor. *Nature* 504: 101–106.
96. Kruse AC, Hu J, Pan AC, et al. (2012) Structure and dynamics of the M3 muscarinic acetylcholine receptor. *Nature* 482: 552–556.
97. Thorsen TS, Matt R, Weis WI, et al. (2014) Modified T4 lysozyme fusion proteins facilitate G protein-coupled receptor crystallogenesis. *Structure* 22: 1657–1664.
98. Glukhova A, Thal DM, Nguyen AT, et al. (2017) Structure of the adenosine A1 receptor reveals the basis for subtype selectivity. *Cell* 168: 867–877.
99. Dore AS, Robertson N, Errey JC, et al. (2011) Structure of the adenosine A(2A) receptor in complex with ZM241385 and the xanthines XAC and caffeine. *Structure* 19: 1283–1293.
100. Xu F, Wu H, Katritch V, et al. (2011) Structure of an agonist-bound human A2A adenosine receptor. *Science* 332: 322–327.
101. Lebon G, Warne T, Edwards PC, et al. (2011) Agonist-bound adenosine A2A receptor structures reveal common features of GPCR activation. *Nature* 474: 521–525.

102. Hino T, Arakawa T, Iwanari H, et al. (2012) G-protein-coupled receptor inactivation by an allosteric inverse-agonist antibody. *Nature* 482: 237–240.
103. Congreve M, Andrews SP, Dore AS, et al. (2012) Discovery of 1,2,4-triazine derivatives as adenosine A(2A) antagonists using structure based drug design. *J Med Chem* 55: 1898–1903.
104. Liu W, Chun E, Thompson AA, et al. (2012) Structural basis for allosteric regulation of GPCRs by sodium ions. *Science* 337: 232–236.
105. Lebon G, Edwards PC, Leslie AG, et al. (2015) Molecular determinants of CGS21680 binding to the human adenosine A2A receptor. *Mol Pharmacol* 87: 907–915.
106. Segala E, Guo D, Cheng RK, et al. (2016) Controlling the dissociation of ligands from the adenosine A2A receptor through modulation of salt bridge strength. *J Med Chem* 59: 6470–6479.
107. Batyuk A, Galli L, Ishchenko A, et al. (2016) Native phasing of x-ray free-electron laser data for a G protein-coupled receptor. *Sci Adv* 2: e1600292.
108. Carpenter B, Nehme R, Warne T, et al. (2016) Structure of the adenosine A(2A) receptor bound to an engineered G protein. *Nature* 536: 104–107.
109. Sun B, Bachhawat P, Chu ML, et al. (2017) Crystal structure of the adenosine A2A receptor bound to an antagonist reveals a potential allosteric pocket. *Proc Natl Acad Sci USA* 114: 2066–2071.
110. Hanson MA, Roth CB, Jo E, et al. (2012) Crystal structure of a lipid G protein-coupled receptor. *Science* 335: 851–855.
111. Chrencik JE, Roth CB, Terakado M, et al. (2015) Crystal structure of antagonist bound human lysophosphatidic acid receptor 1. *Cell* 161: 1633–1643.
112. Hua T, Vemuri K, Pu M, et al. (2016) Crystal structure of the human cannabinoid receptor CB1. *Cell* 167: 750–762.
113. Shao Z, Yin J, Chapman K, et al. (2016) High-resolution crystal structure of the human CB1 cannabinoid receptor. *Nature*.
114. White JF, Noinaj N, Shibata Y, et al. (2012) Structure of the agonist-bound neurotensin receptor. *Nature* 490: 508–513.
115. Egloff P, Hillenbrand M, Klenk C, et al. (2014) Structure of signaling-competent neurotensin receptor 1 obtained by directed evolution in *Escherichia coli*. *Proc Natl Acad Sci USA* 111: E655–E662.
116. Krumm BE, White JF, Shah P, et al. (2015) Structural prerequisites for G-protein activation by the neurotensin receptor. *Nat Commun* 6: 7895.
117. Krumm BE, Lee S, Bhattacharya S, et al. (2016) Structure and dynamics of a constitutively active neurotensin receptor. *Sci Rep* 6: 38564.
118. Yin J, Mobarec JC, Kolb P, et al. (2015) Crystal structure of the human OX2 orexin receptor bound to the insomnia drug suvorexant. *Nature* 519: 247–250.
119. Gayen A, Goswami SK, Mukhopadhyay C (2011) NMR evidence of GM1-induced conformational change of Substance P using isotropic bicelles. *Biochim Biophys Acta* 1808: 127–139.
120. Shihoya W, Nishizawa T, Okuta A, et al. (2016) Activation mechanism of endothelin ETB receptor by endothelin-1. *Nature* 537: 363–368.

121. Park SH, Das BB, Casagrande F, et al. (2012) Structure of the chemokine receptor CXCR1 in phospholipid bilayers. *Nature* 491: 779–783.
122. Zheng Y, Qin L, Zacarias NV, et al. (2016) Structure of CC chemokine receptor 2 with orthosteric and allosteric antagonists. *Nature* 540: 458–461.
123. Wu B, Chien EY, Mol CD, et al. (2010) Structures of the CXCR4 chemokine GPCR with small-molecule and cyclic peptide antagonists. *Science* 330: 1066–1071.
124. Qin L, Kufareva I, Holden LG, et al. (2015) Structural biology. Crystal structure of the chemokine receptor CXCR4 in complex with a viral chemokine. *Science* 347: 1117–1122.
125. Tan Q, Zhu Y, Li J, et al. (2013) Structure of the CCR5 chemokine receptor-HIV entry inhibitor maraviroc complex. *Science* 341: 1387–1390.
126. Oswald C, Rappas M, Kean J, et al. (2016) Intracellular allosteric antagonism of the CCR9 receptor. *Nature* 540: 462–465.
127. Miller RL, Thompson AA, Trapella C, et al. (2015) The importance of ligand-receptor conformational pairs in stabilization: Spotlight on the N/OFQ G protein-coupled receptor. *Structure* 23: 2291–2299.
128. Wu H, Wacker D, Mileni M, et al. (2012) Structure of the human kappa-opioid receptor in complex with JDTic. *Nature* 485: 327–332.
129. Manglik A, Kruse AC, Kobilka TS, et al. (2012) Crystal structure of the mu-opioid receptor bound to a morphinan antagonist. *Nature* 485: 321–326.
130. Huang W, Manglik A, Venkatakrisnan AJ, et al. (2015) Structural insights into micro-opioid receptor activation. *Nature* 524: 315–321.
131. Granier S, Manglik A, Kruse AC, et al. (2012) Structure of the delta-opioid receptor bound to naltrindole. *Nature* 485: 400–404.
132. Fenalti G, Giguere PM, Katritch V, et al. (2014) Molecular control of delta-opioid receptor signaling. *Nature* 506: 191–196.
133. Fenalti G, Zatspein NA, Betti C, et al. (2015) Structural basis for bifunctional peptide recognition at human delta-opioid receptor. *Nat Struct Mol Biol* 22: 265–268.
134. Zhang H, Unal H, Gati C, et al. (2015) Structure of the Angiotensin receptor revealed by serial femtosecond crystallography. *Cell* 161: 833–844.
135. Zhang H, Unal H, Desnoyer R, et al. (2015) Structural basis for ligand recognition and functional selectivity at angiotensin receptor. *J Biol Chem* 290: 29127–29139.
136. Zhang H, Han GW, Batyuk A, et al. (2017) Structural basis for selectivity and diversity in angiotensin II receptors. *Nature*.
137. Burg JS, Ingram JR, Venkatakrisnan AJ, et al. (2015) Structural biology. Structural basis for chemokine recognition and activation of a viral G protein-coupled receptor. *Science* 347: 1113–1117.
138. Zhang D, Gao ZG, Zhang K, et al. (2015) Two disparate ligand-binding sites in the human P2Y1 receptor. *Nature* 520: 317–321.
139. Zhang J, Zhang K, Gao ZG, et al. (2014) Agonist-bound structure of the human P2Y12 receptor. *Nature* 509: 119–122.
140. Zhang C, Srinivasan Y, Arlow DH, et al. (2012) High-resolution crystal structure of human protease-activated receptor 1. *Nature* 492: 387–392.

141. Srivastava A, Yano J, Hirozane Y, et al. (2014) High-resolution structure of the human GPR40 receptor bound to allosteric agonist TAK-875. *Nature* 513: 124–127.
142. Siu FY, He M, de Graaf C, et al. (2013) Structure of the human glucagon class B G-protein-coupled receptor. *Nature* 499: 444–449.
143. Jazayeri A, Dore AS, Lamb D, et al. (2016) Extra-helical binding site of a glucagon receptor antagonist. *Nature* 533: 274–277.
144. Hollenstein K, Kean J, Bortolato A, et al. (2013) Structure of class B GPCR corticotropin-releasing factor receptor 1. *Nature* 499: 438–443.
145. Dore AS, Bortolato A, Hollenstein K, et al. (2017) Decoding corticotropin-releasing factor receptor type 1 crystal structures. *Curr Mol Pharmacol*: Epub ahead of print, DOI: 10.2174/1874467210666170110114727.
146. Wu H, Wang C, Gregory KJ, et al. (2014) Structure of a class C GPCR metabotropic glutamate receptor 1 bound to an allosteric modulator. *Science* 344: 58–64.
147. Christopher JA, Aves SJ, Bennett KA, et al. (2015) Fragment and structure-based drug discovery for a class C GPCR: Discovery of the mGlu5 negative allosteric modulator HTL14242 (3-Chloro-5-[6-(5-fluoropyridin-2-yl)pyrimidin-4-yl]benzonitrile). *J Med Chem* 58: 6653–6664.
148. Wang C, Wu H, Evron T, et al. (2014) Structural basis for Smoothed receptor modulation and chemoresistance to anticancer drugs. *Nat Commun* 5: 4355.
149. Byrne EF, Sircar R, Miller PS, et al. (2016) Structural basis of Smoothed regulation by its extracellular domains. *Nature* 535: 517–522.
150. Lundstrom K (2006) Latest development in drug discovery on G protein-coupled receptors. *Curr Protein Pept Sci* 7: 465–470.
151. Shonberg J, Kling RC, Gmeiner P, et al. (2015) GPCR crystal structures: Medicinal chemistry in the pocket. *Bioorg Med Chem* 23: 3880–3906.
152. Attwood TK, Findlay JB (1993) Design of a discriminating fingerprint for G-protein-coupled receptors. *Protein Eng* 6: 167–176.
153. Chee MS, Satchwell SC, Preddie E, et al. (1990) Human cytomegalovirus encodes three G protein-coupled receptor homologues. *Nature* 344: 774–777.
154. Attwood TK, Findlay JB (1994) Fingerprinting G-protein-coupled receptors. *Protein Eng* 7: 195–203.
155. Strader CD, Sigal IS, Dixon RA (1989) Structural basis of beta-adrenergic receptor function. *FASEB J* 3: 1825–1832.
156. Liapakis G, Ballesteros JA, Papachristou S, et al. (2000) The forgotten serine. A critical role for Ser-2035.42 in ligand binding to and activation of the beta 2-adrenergic receptor. *J Biol Chem* 275: 37779–37788.
157. Swaminath G, Xiang Y, Lee TW, et al. (2004) Sequential binding of agonists to the beta2 adrenoceptor. Kinetic evidence for intermediate conformational states. *J Biol Chem* 279: 686–691.
158. Baldwin JM (1994) Structure and function of receptors coupled to G proteins. *Curr Opin Cell Biol* 6: 180–190.

159. Tyndall JD, Sandilya R (2005) GPCR agonists and antagonists in the clinic. *Med Chem* 1: 405–421.
160. Jacoby E, Bouhelal R, Gerspacher M, et al. (2006) The 7 TM G-protein-coupled receptor target family. *Chem Med Chem* 1: 761–782.
161. Spiss CK, Maze M (1985) Adrenoreceptors. *Anaesthetist* 34: 1–10.
162. Civelli O, Reinscheid RK, Zhang Y, et al. (2013) G protein-coupled receptor deorphanizations. *Annu Rev Pharmacol Toxicol* 53: 127–146.
163. Barst RJ, Langleben D, Frost A, et al. (2004) Sitaxsentan therapy for pulmonary arterial hypertension. *Am J Respir Crit Care Med* 169: 441–447.
164. Kotake T, Usami M, Akaza H, et al. (1999) Goserelin acetate with or without antiandrogen or estrogen in the treatment of patients with advanced prostate cancer: a multicenter, randomized, controlled trial in Japan. Zoladex Study Group. *Jpn J Clin Oncol* 29: 562–570.
165. Onuffer JJ, Horuk R (2002) Chemokines, chemokine receptors and small-molecule antagonists: recent developments. *Trends Pharmacol Sci* 23: 459–467.
166. Fatkenheuer G, Pozniak AL, Johnson MA, et al. (2005) Efficacy of short-term monotherapy with maraviroc, a new CCR5 antagonist, in patients infected with HIV-1. *Nat Med* 11: 1170–1172.
167. Lu M, Wu B (2016) Structural studies of G protein-coupled receptors. *IUBMB Life* 68: 894–903.
168. Strotmann R, Schrock K, Boselt I, et al. (2011) Evolution of GPCR: change and continuity. *Mol Cell Endocrinol* 331: 170–178.
169. Ballesteros JA, Weinstein H (1995) Integrated methods for the construction of three-dimensional models and computational probing of structure-function relations in G protein-coupled receptors. *Method Neurosci* 25: 366–428.
170. Zhang D, Weinstein H (1994) Polarity conserved positions in transmembrane domains of G-protein coupled receptors and bacteriorhodopsin. *FEBS Lett* 337: 207–212.
171. Katritch V, Fenalti G, Abola EE, et al. (2014) Allosteric sodium in class A GPCR signaling. *Trends Biochem Sci* 39: 233–244.
172. Bjarnadottir TK, Geirardsdottir K, Ingemansson M, et al. (2007) Identification of novel splice variants of Adhesion G protein-coupled receptors. *Gene* 387: 38–48.
173. Lin HH, Chang GW, Davies JQ, et al. (2004) Autocatalytic cleavage of the EMR2 receptor occurs at a conserved G protein-coupled receptor proteolytic site motif. *J Biol Chem* 279: 31823–31832.
174. Isberg V, Vroling B, Van DKR, et al. (2014) GPCRDB: an information system for G protein-coupled receptors. *Nucleic Acids Res* 42: D422–D425.
175. Gasparini F, Kuhn R, Pin JP (2002) Allosteric modulators of group I metabotropic glutamate receptors: novel subtype-selective ligands and therapeutic perspectives. *Curr Opin Pharmacol* 2: 43–49.
176. Malherbe P, Kratochwil N, Zenner MT, et al. (2003) Mutational analysis and molecular modeling of the binding pocket of the metabotropic glutamate 5 receptor negative modulator 2-methyl-6-(phenylethynyl)-pyridine. *Mol Pharmacol* 64: 823–832.

177. Litschig S, Gasparini F, Rueegg D, et al. (1999) CPCCOEt, a noncompetitive metabotropic glutamate receptor 1 antagonist, inhibits receptor signaling without affecting glutamate binding. *Mol Pharmacol* 55: 453–461.
178. Silve C, Petrel C, Leroy C, et al. (2005) Delineating a Ca²⁺ binding pocket within the venus flytrap module of the human calcium-sensing receptor. *J Biol Chem* 280: 37917–37923.
179. Hermans E, Challiss RA (2001) Structural, signalling and regulatory properties of the group I metabotropic glutamate receptors: prototypic family C G-protein-coupled receptors. *Biochem J* 359: 465–484.
180. Nakanishi S (1992) Molecular diversity of glutamate receptors and implications for brain function. *Science* 258: 597–603.
181. Riedel G, Platt B, Micheau J (2003) Glutamate receptor function in learning and memory. *Behav Brain Res* 140: 1–47.
182. Kunishima N, Shimada Y, Tsuji Y, et al. (2000) Structural basis of glutamate recognition by a dimeric metabotropic glutamate receptor. *Nature* 407: 971–977.
183. Niswender CM, Conn PJ (2010) Metabotropic glutamate receptors: physiology, pharmacology, and disease. *Annu Rev Pharmacol Toxicol* 50: 295–322.
184. Bhanot P, Brink M, Samos CH, et al. (1996) A new member of the frizzled family from *Drosophila* functions as a Wingless receptor. *Nature* 382: 225–230.
185. Murone M, Rosenthal A, de Sauvage FJ (1999) Sonic hedgehog signaling by the patched-smoothened receptor complex. *Curr Biol* 9: 76–84.
186. Chen CM, Strapps W, Tomlinson A, et al. (2004) Evidence that the cysteine-rich domain of *Drosophila* Frizzled family receptors is dispensable for transducing Wingless. *Proc Natl Acad Sci USA* 101: 15961–15966.
187. Nakano Y, Nystedt S, Shivdasani AA, et al. (2004) Functional domains and sub-cellular distribution of the Hedgehog transducing protein Smoothened in *Drosophila*. *Mech Dev* 121: 507–518.
188. Isberg V, de Graaf C, Bortolato A, et al. (2015) Generic GPCR residue numbers—Aligning topology maps minding the gaps. *Trends Pharmacol Sci* 36: 22–31.



AIMS Press

© 2017 Abby L. Parrill, et al., licensee AIMS Press. This is an open access article distributed under the terms of the Creative Commons Attribution License (<http://creativecommons.org/licenses/by/4.0>)