

The structural basis of function in Cys-loop receptors

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Abstract. Cys-loop receptors are membrane-spanning neurotransmitter-gated ion channels that are responsible for fast excitatory and inhibitory transmission in the peripheral and central nervous systems. The best studied members of the Cys-loop family are nACh, 5-HT₃, GABA_A and glycine receptors. All these receptors share a common structure of five subunits, pseudo-symmetrically arranged to form a rosette with a central ion-conducting pore. Some are cation selective (e.g. nACh and 5-HT₃) and some are anion selective (e.g. GABA_A and glycine). Each receptor has an extracellular domain (ECD) that contains the ligand-binding sites, a transmembrane domain (TMD) that allows ions to pass across the membrane, and an intracellular domain (ICD) that plays a role in channel conductance and receptor modulation. Cys-loop receptors are the targets for many currently used clinically relevant drugs (e.g. benzodiazepines and anaesthetics). Understanding the molecular mechanisms of these receptors could therefore provide the catalyst for further development in this field, as well as promoting the development of experimental techniques for other areas of neuroscience.

In this review, we present our current understanding of Cys-loop receptor structure and function. The ECD has been extensively studied. Research in this area has been stimulated in recent years by the publication of high-resolution structures of nACh receptors and related proteins, which have permitted the creation of many Cys loop receptor homology models of this region. Here, using the 5-HT₃ receptor as a typical member of the family, we describe how homology modelling and ligand docking can provide useful but not definitive information about ligand interactions. We briefly consider some of the many Cys-loop receptors modulators. We discuss the current understanding of the structure of the TMD, and how this links to the ECD to allow channel gating, and consider the roles of the ICD, whose structure is poorly understood. We also describe some of the current methods that are beginning to reveal the differences between different receptor states, and may ultimately show structural details of transitions between them.

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1. Introduction

Cys-loop ligand-gated ion channels (LGICs) are membrane-spanning proteins that are activated by neurotransmitters; they are responsible for fast excitatory and inhibitory neurotransmission in the central and peripheral nervous systems. Vertebrate members of this family include serotonin (5-HT₃), acetylcholine (nicotinic ACh or nACh), glycine (Gly), γ -aminobutyric acid (GABA_A, GABA_C) and zinc-activated (ZAC) receptors (R). There are also a range of invertebrate Cys-loop receptors gated by the same and other neurotransmitters (e.g. EXP-1, MOD-1, pHCl, HisCl, RDL, GluCl and SsCl), and related proteins have been identified in prokaryotes (e.g. ELIC and GLIC). Cys-loop receptors are the major targets for many active compounds, including anaesthetics, muscle relaxants, insecticides and a range of drugs that treat neurological disorders such as Alzheimer's, anxiety, epilepsy, learning, attention deficit and drug addiction. Methods such as high throughput screening (HTS) and fragment-based drug discovery (FBDD) use blind searches

of large compound libraries to find drug candidates, but a rational design of more effective drugs requires a detailed molecular knowledge of the sites at which they act. For Cys-loop receptors, this information lags behind that of many other proteins.

Cys-loop receptors derive their name from a 13-amino-acid loop within the extracellular domain (ECD) that is enclosed by a pair of disulphide-bonded Cys residues. Members of the family share a common structure, consisting of five pseudo-symmetrically arranged subunits surrounding a central ion-conducting pore (Fig. 1). Most receptors have more than one type of subunit, and these can combine in different combinations to yield a complex array of (usually) heteromeric receptor stoichiometries, with varying physiological and pharmacological properties. Each receptor family is selective for either cations or anions, and their activation can be either excitatory or inhibitory, depending on the distribution of ions at either side of the membrane, and the membrane potential of the cell. The structure of the subunits has been studied using a variety of biochemical techniques such as mutagenesis, photolabelling and cryo-electron microscopy, and more recently by X-ray crystallography. Each subunit can be functionally separated into three domains: The large ECD contains the ligand-binding site and is a major target for therapeutics. The transmembrane domain (TMD) consists of four membrane-spanning α -helices (M1–M4) that enable ions to cross the membrane and is the target for compounds such as alcohols, anaesthetics and steroids. The intracellular domain (ICD) is primarily formed by the large M3–M4 intracellular loop (~ 100 – 270 residues), and is responsible for receptor modulation, sorting and trafficking, and contains portals (openings) that allow ions access in and out of the pore and influence ion conductance. Recent studies have described homologous bacterial proteins that do not possess a Cys-loop or an ICD, and deletion studies in the 5-HT₃R and GABA_CR suggest that the ICD is not essential for the expression of vertebrate receptors (Bocquet *et al.* 2007; Jansen *et al.* 2008).

In summary, all Cys-loop receptors share homologous structures, and the basic mechanisms by which they function are also similar. In this review, we look at all members of the family, although concentrating largely on the 5-HT₃R, to explore the relationship between structure and function. The 5-HT₃R is a typical Cys-loop receptor, and has the advantage that it functions as a homomeric receptor, which simplifies the interpretation of experimental data. This protein has also been used extensively in homology modelling and ligand docking (Maksay *et al.* 2003; Reeves *et al.* 2003; Thompson *et al.* 2005; Yan & White, 2005). As this technique is becoming an accepted route to understanding the structural details of the proteins, we use the new homology models and docked ligands to explore the validity of these techniques to define specific molecular interactions with agonists and antagonists in the 5-HT₃ ligand-binding site.

For further reading, a number of recent reviews also cover some of the topics discussed here (Arias, 2006; Auerbach, 2010; Barnes *et al.* 2009; Chen, 2010; Corringer *et al.* 2010; Hogg *et al.* 2003; Lynch, 2004, 2009; Millar & Gotti, 2009; Peters *et al.* 2005, 2010; Webb & Lynch, 2007; Yakel, 2010).

2. Subunit stoichiometry

The stoichiometry of the neuromuscular nAChR was the first to be determined, and revealed that four different subunits formed a functional pentameric receptor with the stoichiometry $\alpha_2\beta\gamma\delta$ (Karlin *et al.* 1983). Determining the stoichiometry of other receptors has proved to be more problematic, as there are large numbers of subunits that could potentially contribute (e.g. 19 in the GABA_A receptor family), and it is becoming apparent that different arrangements

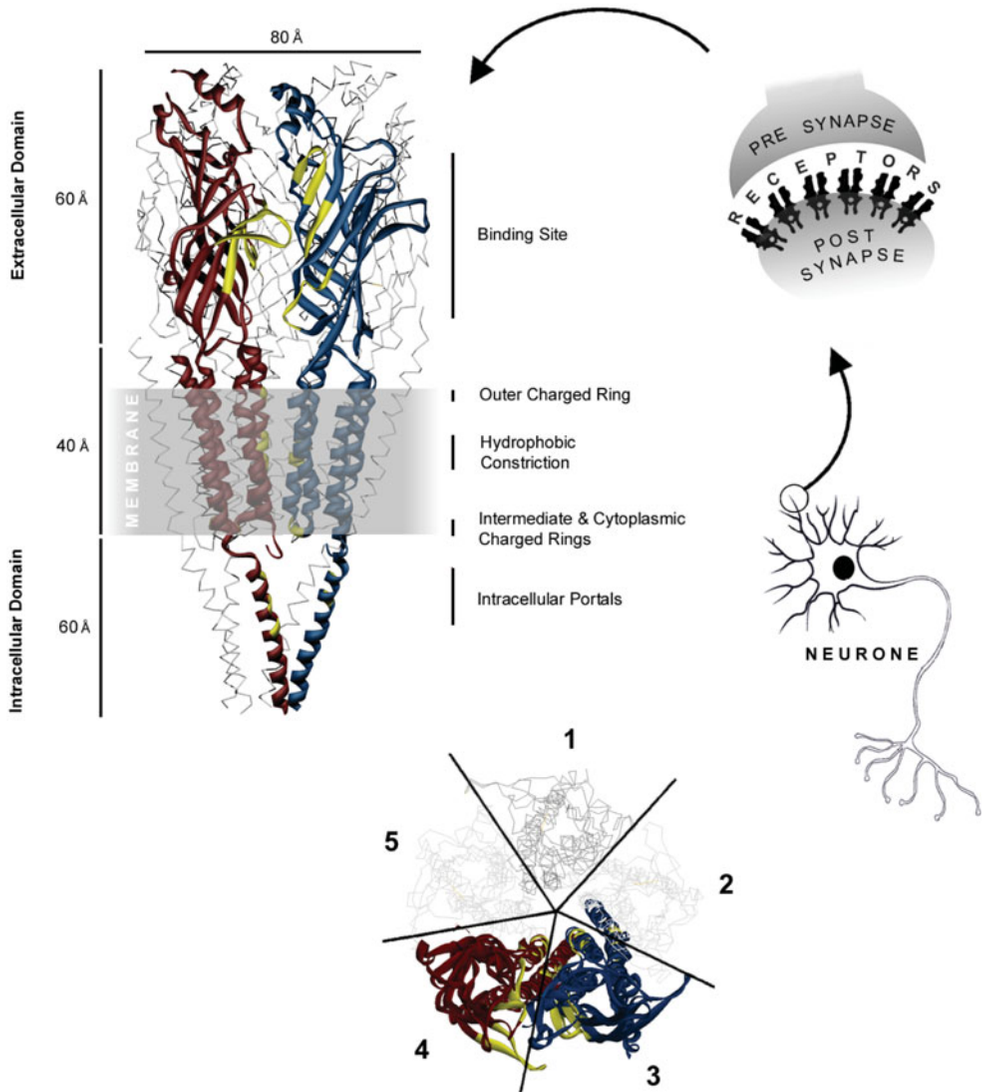


Fig. 1. Important functional components of the 5-HT₃ receptor, a typical member of the Cys-loop family of LGICs. The structure shown is a 5-HT₃ homology model based on a 4 Å-resolution structure of the nAChR (Miyazawa *et al.* 2003; PDB ID: 2BG9). The 5-HT₃ receptor, like the other members, consists of five subunits (1–5). The receptor is shown from above and from the side with two (red & blue) of the five subunits highlighted. Specific residues of interest are highlighted in yellow. The receptor is modular in nature and can be considered as having three main regions termed the ECD, TMD and ICD. The ECD contains the ligand-binding site that is formed by the convergence of six peptide loops located at the interface of two adjacent subunits (Noam *et al.* 2008; Thompson & Lummis, 2006). Three rings of charged amino acids (extracellular, intermediate and cytoplasmic) are found in the pore lining α -helices of the TMD (Gunthorpe & Lummis, 2001; Thompson & Lummis, 2003), and a hydrophobic constriction in the centre of the channel acts as the channel gate (Panicker *et al.* 2002).

may exist even with the same subunit types (Gotti *et al.* 2007; Millar & Gotti, 2009; Millar & Harkness, 2008; Olsen & Sieghart, 2009). For example, neuronal $\alpha 4\beta 2$ nAChR may be $\alpha_2\beta_3$ or $\alpha_3\beta_2$, which have differing pharmacologies (Moroni & Bermudez, 2006; Moroni *et al.* 2006). Studies indicate, however, that only a limited number of the possible stoichiometries are

found *in vivo*, possibly because of cell-specific expression and/or interactions between subunit interfaces that form during receptor assembly (e.g. neuronal GABA_A receptors are predominantly $\alpha 1_2\beta 2_2\gamma 2$). Receptor types may also be restricted to specific regions of the body (e.g. GABA_C receptors are largely restricted to retinal bipolar cells; Cutting *et al.* 1991; Enz & Cutting, 1999). Some Cys-loop receptors have considerably fewer potential stoichiometries. For example, there are only four known isoforms of the GlyR α -subunit ($\alpha 1$ – $\alpha 4$) and a single β -subunit, with the probable stoichiometry of $\alpha 1_3\beta_2$ or $\alpha 1_4\beta$ (Lynch, 2009; Webb & Lynch, 2007), and only a single subunit has been described for vertebrate Zn²⁺-activated receptors (Davies *et al.* 2002). Invertebrate receptors may also have multiple subunits; several glutamate-gated and pHCl receptor subunits have been reported, although currently there are only two known histamine-gated receptor subunits (HisCl1 and HisCl2SsCl) and single SsCl and MOD-1 receptor subunits (Cully *et al.* 1996, 1994; Mounsey *et al.* 2007; Ranganathan *et al.* 2000; Zheng *et al.* 2002). Prokaryotic receptors discovered to date also only have single subunits, but as many of these have only recently been described, the diversity of their subunits types may grow with further investigation (Bocquet *et al.* 2007; Hilf & Dutzler, 2008; Nury *et al.* 2009).

The 5-HT₃R is an example of a Cys-loop receptor with relatively few subunits; five have been identified to date (A–E), although like many other receptors, some of these demonstrate a further level of complexity that results from different splice-variations and differing post-translational modifications (Bruss *et al.* 2000; Tzvetkov *et al.* 2007; Werner *et al.* 1994). For example, there are long and short forms of the mouse 5-HT₃A subunit that differ by six amino acids, and there are three translational variants of the human 5-HT₃B subunit (Fig. 2). Only 5-HT₃A subunits can form functional homomeric 5-HT₃Rs, and appear to be obligatory in heteromeric receptors (Holbrook *et al.* 2009; Niesler *et al.* 2007). Of the heteromeric receptors, only 5-HT₃AB receptors have been extensively characterized and, compared to homomeric 5-HT₃A, 5-HT₃AB receptors differ in their EC₅₀, Hill slope, desensitization kinetics, shape of current–voltage relationship, and most noticeably, a much larger single-channel conductance (~ 16 pS in 5-HT₃AB compared to < 1 pS in 5-HT₃A; Davies *et al.* 1999; Dubin *et al.* 1999). However, the pharmacology of 5-HT₃A and 5-HT₃AB receptors is almost identical, suggesting that they contain a common binding site (an A–A interface), a hypothesis supported by a recent study of mouse 5-HT₃AB receptors (Brady *et al.* 2001; Lochner & Lummis, 2010), but conflicting with the BABBA arrangement determined using atomic force microscopy (Barrera *et al.* 2005). The subunit types and stoichiometry of 5-HT₃Rs have been recently reviewed (Barnes *et al.* 2009; Jensen *et al.* 2008).

3. The ECD

3.1 Structure

Recent X-ray crystal structures of the nAChR ECD have revealed molecular details of residues that contribute to the ligand-binding domain, but such studies of whole receptors, or even ECD pentamers are proving difficult to obtain (Dellisanti *et al.* 2007). Therefore, most molecular details of Cys-loop receptors have been extrapolated from 4 Å resolution cryo-electron microscopy images of the nAChR, or from higher-resolution images of related acetylcholine binding proteins (AChBPs) and bacterial receptors. AChBPs are homologous to the ECD of nACh ($\sim 25\%$ amino-acid sequence identity) and other Cys-loop receptors (15–20% identity). The original AChBP structure was determined at 2.7 Å resolution in 2001 (Brejci *et al.* 2001), and

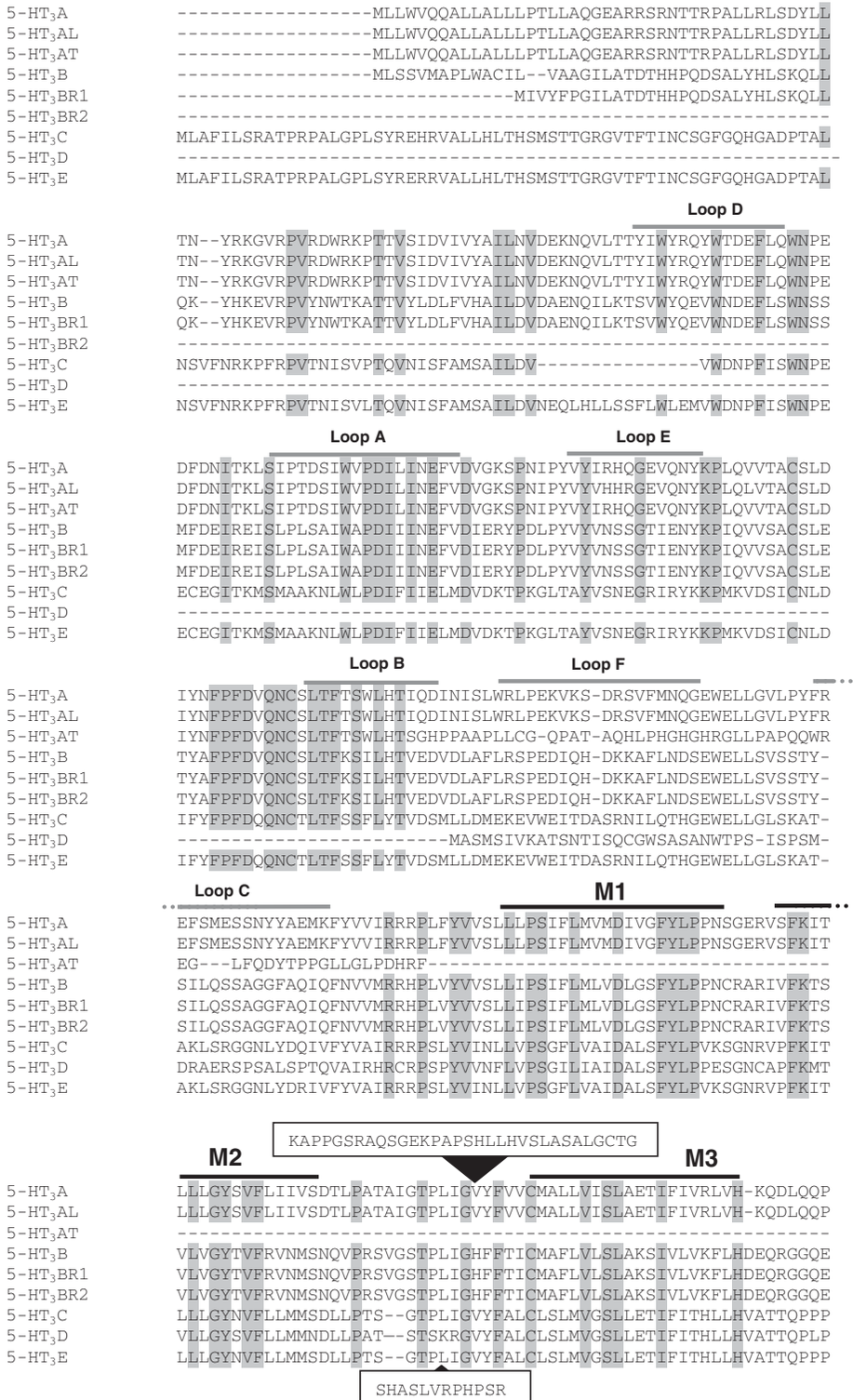


Fig. 2. (Cont.)

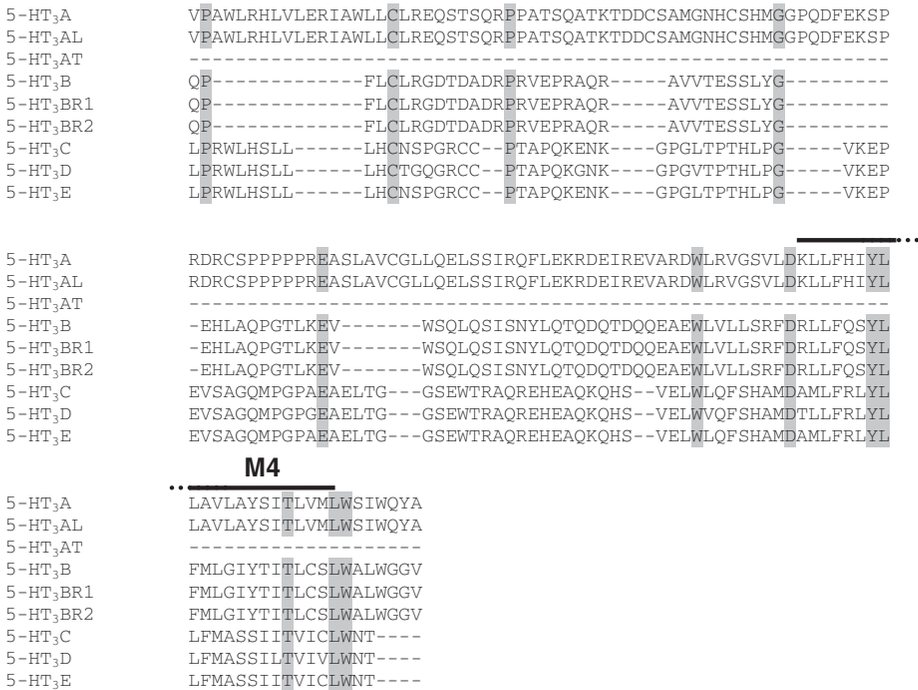


Fig. 2. Alignment of human 5-HT₃ receptor subunits. The binding loops (A–F) and transmembrane (M1–M4) regions are highlighted by horizontal lines above the text. Conserved residues are highlighted with a grey background. The human alternative long (5-HT₃AL), truncated (5-HT₃AT), Brain1 (5-HT₃BR1) and Brain2 (5-HT₃BR2) forms are shown. The boxes show the additional residues found in 5-HT₃AL (32 amino acids) and 5-HT₃D (12 amino acids) variants. Accession numbers for the alignment are as follows: 5-HT_{3A} P46098, 5-HT_{3B} O95264 and 5-HT_{3C} Q6V706. 5-HT_{3D} and 5HT_{3E} were taken from Niesler *et al.* (2003). 5-HT₃AL and 5-HT₃AT were taken from Bruss *et al.* (2000). 5-HT₃BR1 and 5-HT₃BR2 were taken from Tzvetkov *et al.* (2007).

since this time other AChBP structures have been reported (e.g. Celie *et al.* 2004, 2005b; Hansen & Taylor, 2007; Hibbs *et al.* 2009). The similarity between AChBP and the ECD of Cys-loop receptors was confirmed when the structures of an nACh subunit monomer and subsequently homologous prokaryotic receptors were determined (Bocquet *et al.* 2009; Dellisanti *et al.* 2007; Hilf & Dutzler, 2008; Nury *et al.* 2009). With this similarity established, we can be more confident that studies that utilized the AChBP structure to make predictions within Cys-loop receptors were broadly correct. A review of prokaryotic receptors can be found in Corringer *et al.* (2010).

3.2 The ligand-binding site

Early biochemical and labelling studies indicated that Cys-loop receptor ligand-binding sites were constituted by three non-contiguous regions from the ECDs of two contributing subunits. With the advent of the AChBP crystal structure, it was confirmed that the binding site was at the interface between two adjacent subunits (Brejc *et al.* 2001; Celie *et al.* 2004, 2005a, 2005b). The two adjacent subunits are termed the principal and complementary subunits, and the binding site is formed by three peptide loops (loops A–C) from the principal subunit, and three β -sheets

(loops D–F) from the complementary subunit; as this terminology was introduced before crystallographic studies revealed the secondary structure, these regions are not all ‘loops’ (Fig. 3).

The exact location of the loop region varies subtly with different subunits of different receptors; the locations that we have shown in Fig. 3 are therefore only approximate. Only one or a few residues within each loop may face into the binding pocket, with residues in the remainder of the loop probably maintaining the structure of the pocket and/or participating in the conformational changes that result in channel opening. Evidence from AChBP structures (discussed later in section 7.5) suggests that binding of different ligands results in different movements of the binding pocket; the ECD generally contracts around agonists, but adopts a more open structure with antagonists. It has been known for some time that antagonists and agonists may interact with different binding pocket residues, and one ligand may interact with more or less residues than another (e.g. a large nAChR antagonist such as α -bungarotoxin (α -BTX) interacts with a much larger repertoire of residues than a small antagonist such as methyllycaconitine). What is perhaps more surprising is that agonists do not need to interact with the same residues to activate the receptor. For example, 5-HT forms a critical hydrogen bond with Glu129 in the 5-HT₃R, but 5-FT, which still activates the receptor (albeit as a partial agonist), does not appear to interact at all with this residue (Bower *et al.* 2008).

For the majority of Cys-loop receptors, at least two binding sites are required for channel activation, and at muscle nAChRs, the principal subunits at both these sites are the α 1 subtype; for some neuronal receptors, it appears that the two principal subunits differ within a single receptor, for instance α 4 and α 6 (Champiaux *et al.* 2003; Rayes *et al.* 2009). Questions about cooperative binding are often finessed with the statement that the open state of the channel is more likely to be associated with the presence of at least two bound agonists, although certain mutant receptors have Hill slopes near unity, allowing for the possibility of opening with just a single bound agonist. In some mutant receptors that are highly agonist-sensitive, there is also constitutive activation in the total absence of agonist, as though the open state is rather more stable than normal (e.g. Bhattacharya *et al.* 2004).

Cys-loop receptor-binding sites all contain a number of aromatic residues (Table 1). For many of the Cys-loop receptors, a cation– π interaction has been described between the natural ligand and one of these aromatic residues. This type of interaction has been observed in a variety of proteins using high-resolution structural data, but for Cys-loop receptors cation– π interactions have only been identified using unnatural amino acid mutagenesis (Dougherty, 2008). For this technique, a series of electron-withdrawing or electron-donating groups are substituted onto the side chains of aromatic residues, subtly altering the energy of the cation– π interaction. If the EC₅₀ varies monotonically with the calculated strength of the interaction, this is evidence for the presence of a cation– π interaction. In all Cys-loop receptors examined to date, when a cation– π interaction is found, the ligand interacts with only one aromatic side chain in the binding pocket; in some other proteins, the efficient stabilization of this bond relies upon interactions with several aromatic rings, and the optimal orientation of the cationic centre is normal to the planes passing through the centroids of these rings (Schärer *et al.* 2005).

Different aromatic side chains (Trp, Phe or Tyr) make a cation– π interaction in different Cys-loop receptors. Each of these is located in one of the three loops on the principal subunit (Table 1); as yet, no cation– π interactions have been found in the complementary subunit. In the 5-HT₃R, the contributing residue is a loop B Trp (Beene *et al.* 2002, 2004), while in the MOD-1 receptor (also activated by 5-HT) it is a Trp in loop C (W226; Mu *et al.* 2003). Therefore, even in receptors activated by the same ligand, the residue involved in the cation– π interaction can differ.

Table 1. *Aromatic residues in the binding sites of different Cys-loop receptors. Residues that contribute to cation- π interactions are shown in bold.* ^{1,3}Beene et al. (2002), ²Xiu et al. (2009), ⁴Mu et al. (2003), ⁵Padgett et al. (2007), ⁶Lummis et al. (2005a), ⁷Pless et al. (2008). See Dougherty (2008) for further information. Note that loop D is in the complementary face, and residues from this face have not been found to form cation- π interactions to date

| Receptor | Ligand | Loop A | Loop B | Loop C | Loop D | |
|---|---------------|------------|-------------|--------|-------------|------|
| nACh ($\alpha 1\beta\gamma\delta$) ¹ | ACh | Y93 | W149 | Y190 | Y198 | W55 |
| nACh ($\alpha 4\beta 2$) ² | ACh, Nicotine | Y98 | W154 | Y195 | Y202 | W55 |
| 5-HT ₃ ³ | 5-HT | E129 | W183 | F226 | Y234 | W90 |
| MOD-1 ⁴ | 5-HT | C120 | Y180 | Y221 | W226 | F83 |
| GABA _A ⁵ | GABA | Y97 | Y157 | F200 | Y205 | F65 |
| GABA _C ⁶ | GABA | F138 | Y198 | Y241 | Y247 | Y102 |
| Gly ⁷ | Gly | F99 | F159 | Y202 | F207 | F63 |

Similarly in the GABA_C receptor, GABA has a cation- π interaction with a loop B Tyr residue, but in the GABA_A receptor, GABA has a cation- π interaction at a Tyr on loop A. Several exogenous or synthetic agonists can also make cation- π interactions (e.g. epibatidine; Cashin et al. 2005) but it is not essential; for example, nicotine can make a cation- π interaction at the neuronal $\alpha 4\beta 2$ nAChR, but does *not* make a cation- π interaction at the muscle nAChR (Beene et al. 2002; Xiu et al. 2009). These data provide an explanation for the low potency of nicotine at muscle nAChRs (and an understanding of why smoking does not cause severe muscle contractions), and also demonstrates the importance of understanding the molecular interactions when designing receptor-specific drugs. These data also highlight the problem that even with good structural information, docking a ligand into a protein may not always be accurate, and experimental data are essential to allow the correct solution to be selected from possible options.

3.2.1 Ligand binding; *in silico* predictions from the 5-HT₃R

In silico predictions of ligand binding require either a high-resolution structure or a homology model, and the template used for the latter will determine its accuracy. To show how differing starting templates can introduce conformational variability, Fig. 4 overlays two 5-HT₃ homology models that were created using similar AChBP structures containing the same bound ligand (HEPES; PDB ID's 119B and 1UX2). The overlay shows that the backbones closely mirror each other, but there are considerable differences in the orientations of side chains. Other starting templates that contain different bound ligands produce further variation, and if we use these for *in silico* docking, the positions of the side chains can have a significant impact on the final orientation of the ligand. Nevertheless, homology models have been produced for many receptors, and a range of ligands docked into their binding sites (e.g. Abdel-Halim et al. 2008; Cromer et al. 2002; Le Novere et al. 2002; Maksay et al. 2003; Reeves et al. 2003; Schapira et al. 2002; Trudell & Bertaccini, 2004; Yan & White, 2005). The ability of 5-HT₃R to form homomeric receptors means that they are a relatively simple system for molecular modelling, and they have the considerable advantage that the experimental determination of the effects of amino acid substitutions on the properties of the receptor is straightforward. In the following section, this receptor is used as a model system to illustrate some of the pros and cons of *in silico* techniques.

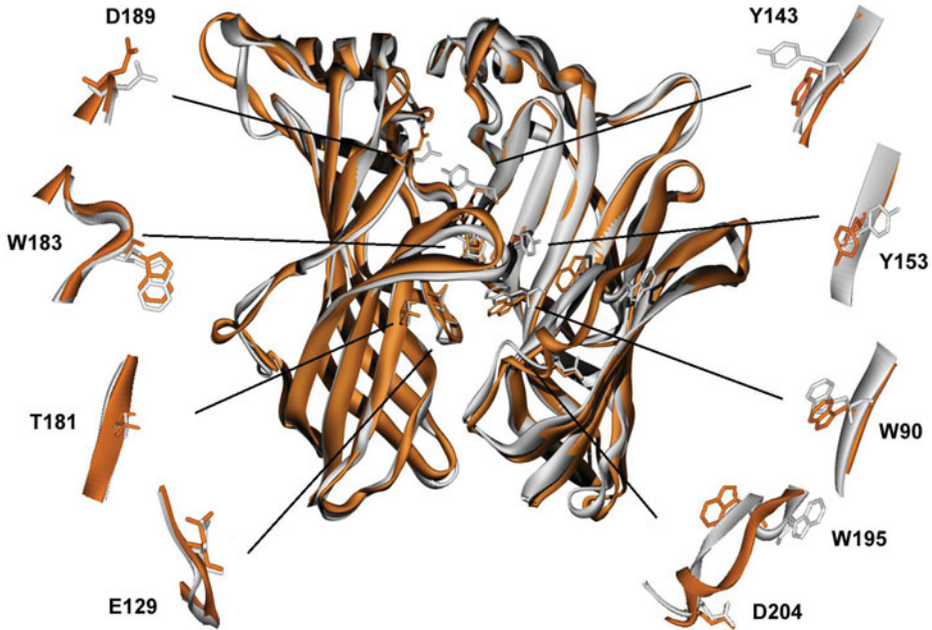


Fig. 4. An overlay of two 5-HT₃ receptor homology models that were based on HEPES bound AChBP structures (PDB ID: 119B white and 1UX2 orange). Some residues that have been shown to be important for granisetron (a selective 5-HT₃ antagonist) binding are highlighted and emphasize that some regions e.g. close to W195, have large differences in the orientation of their side chains. The relative positions of the models were compiled by Swiss-PdbViewer ‘magic fit’ using loop B as a reference point.

In silico docking of ligands can be performed using a variety of software tools. One of the most widely used and well regarded is GOLD (The Cambridge Crystallographic Data Centre, Cambridge UK), which places a ligand into the protein and then improves the fit by iteratively moving the ligand into the most energetically favourable orientation (Olsen *et al.* 2004a). To explore its accuracy, we determined whether GOLD could adequately locate binding sites and correctly position ligands in them by removing ligands from their original protein structures and re-docking them. Figure 5 shows the 10 predicted ligand orientations for nicotine and carbamylcholine in their original AChBP crystal structures, and in two other randomly selected structures. In each instance, the software correctly located the ligand within the receptor, although there are some subtle differences in their precise orientations.

Granisetron is a selective, competitive antagonist of 5-HT₃Rs. Nuclear magnetic resonance (NMR) and crystallography studies of granisetron show that the azabicyclic of granisetron adopts a boat–chair configuration, and the carbonyl linker is relatively immobile (i.e. rigid), with a dihedral angle of 180° (Fludzinski *et al.* 1987; Roe & Kuntz, 1995; Schmidt & Peroutka, 1989; Vernakar *et al.* 2010). Using this structure, we docked granisetron into a range of 5-HT₃R homology models, the templates of which were the 18 currently available AChBP, nACh and prokaryotic receptor structures (Tables 2 and 3). We have used a flexible ligand (non-constrained bond angles) and a rigid ligand (constrained bond angles), and the tables show the additional variability that is introduced by altering the flexibility of the ligand. A comparison of the results shows that granisetron is located in broadly similar locations in the binding pockets, although the predicted orientations differ (Fig. 6). Flexible (Table 2) and rigid (Table 3) ligand docking

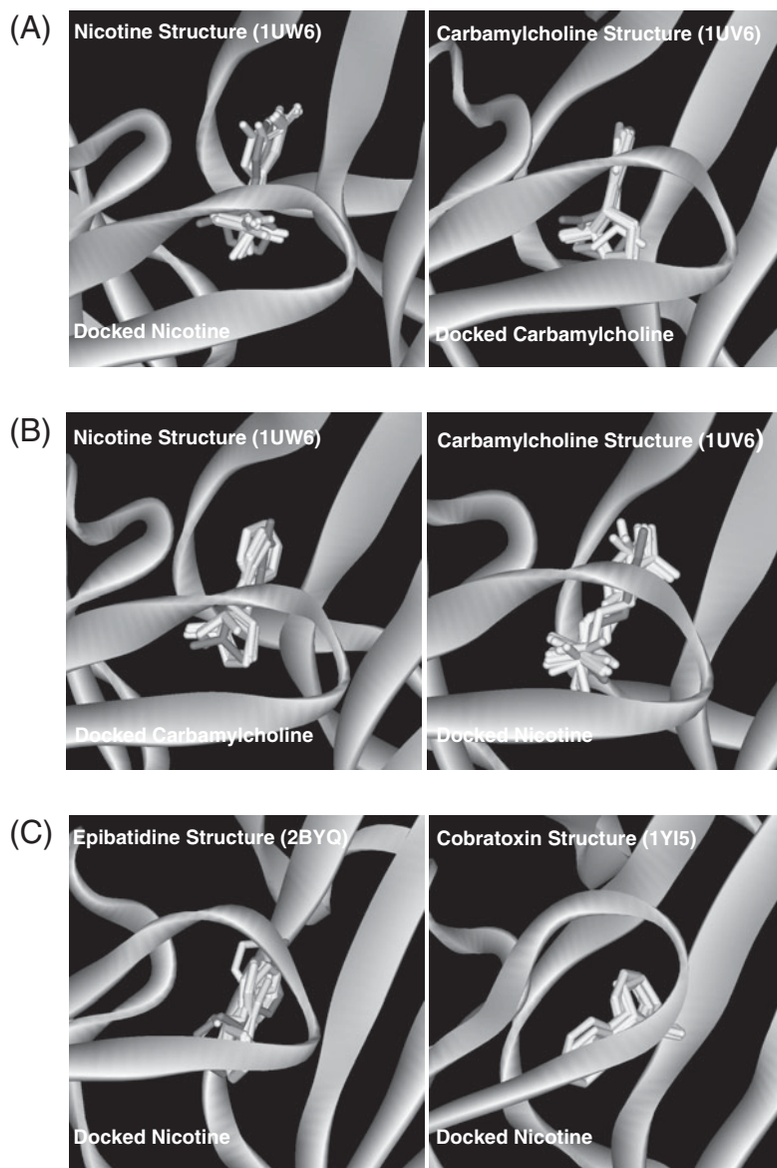


Fig. 5. A test for the accuracy of computational ligand docking. (A) Nicotine and carbamylcholine re-docked into their respective AChBP structures. (B) Nicotine docked into the carbamylcholine structure and vice versa. (C) Nicotine docked into two other AChBP structures. In each panel, the original ligand molecule is shown in grey and 10 docking solutions are shown in white; the ligands are clearly positioned on top of one another within the binding site.

generated eight and five categories of potential ligand orientations, respectively. Rigid docking increased the incidence of granisetron being placed outside the binding site (described as others); docking errors may be responsible, although some locations may represent local energy minima within the binding and unbinding routes, as previously suggested (Joshi *et al.* 2006; Maksay *et al.* 2003; Thompson *et al.* 2006a; Zhang *et al.* 2006).

Table 2. Flexible ligand docking of granisetron into the 5-HT₃ receptor-binding site produces docked clusters that can be categorized into one of eight groups (A1–E). Docking solutions that place granisetron outside the binding site are described as others. The data were created using the boat-chair configuration of granisetron docked into a series of 5-HT₃ homology models that were generated from the 18 currently available PDB templates that show homology to Cys-loop receptors. The protein sequence of the murine 5-HT₃ receptor (accession number: Q6J1J7) was co-aligned with each of the template sequences using FUGUE, which takes into account secondary structures (Shi et al. 2001). Using Modeller 6v2, five homology models were generated from each PDB template (Sali & Blundell, 1993), and the best model selected using Ramachandran plot analysis. The protonated form of granisetron was constructed in Chem3D Ultra 7.0 (CambridgeSoft, Cambridge, UK) based on the crystal structure of a related indazole carboxamide (Cambridge Structural Database; reference code FIZXUH) and docked with GOLD v3.0 (The Cambridge Crystallographic Data Centre, Cambridge, UK), into a binding site that was defined as being within 20 Å of the α-carbon of W183. To dock ligands, 10 genetic algorithm runs were carried out for each homology model, with a population size of 50 and the maximum number of generations set to 27 000. For each homology model, the 10 docking solutions were categorized according to the ligand orientation, and the number of solutions in each category are shown (see Fig. 6 legend for descriptions)

| Starting Template for the 5-HT ₃ R homology model | | | Number of ligand orientations in each 5-HT ₃ R homology model | | | | | | | | |
|--|------------------|------|--|----|----|----|----|----|---|----|-------|
| Protein | Ligand type | PDB | A1 | A2 | B1 | B2 | C1 | C2 | D | E | Other |
| Aplysia AChBP | – | 2BYN | | 3 | | | | | 7 | | |
| Aplysia AChBP | HEPES | 2BR7 | | 7 | | | | | | 3 | |
| Limnaea AChBP | HEPES | 1I9B | | 9 | | | | | | | 1 |
| Limnaea AChBP | HEPES | 1UX2 | | 1 | | 6 | 2 | 1 | | | |
| Aplysia AChBP | Epibatidine | 2BYQ | 3 | 4 | | | 1 | | | 1 | 1 |
| Limnaea AChBP | Carbomylcholine | 1UV6 | | | 1 | | | 1 | | 8 | |
| Limnaea AChBP | Nicotine | 1UW6 | | 10 | | | | | | | |
| Aplysia AChBP | Methylcaconitine | 2BYR | | | | | | | 1 | 7 | 2 |
| Aplysia AChBP | Lobeline | 2BYS | 5 | 4 | | | | | | 1 | |
| <i>Bulinus truncatus</i> AChBP | CAPS | 2BJO | | | | | | | | 10 | |
| <i>Torpedo marmorata</i> nAChR | – | 2BG9 | 3 | 1 | | 1 | | | | | 5 |
| Limnaea AChBP | α-Cobratoxin | 1YI5 | | 4 | | | | | | 5 | 1 |
| Aplysia AChBP | α-Cobratoxin | 2BYP | | | 3 | 7 | | | | | |
| Aplysia AChBP | α-Conotoxin | 2C9T | | | 2 | 8 | | | | | |
| Erwinia LGIC | – | 2VL0 | | 2 | | 3 | | 5 | | | |
| Gleobacter LGIC | – | 3EAM | | | | | | | | | 10 |
| Aplysia AChBP | Cocaine | 2PGZ | | | 10 | | | | | | |
| Aplysia AChBP | Tropisetron | 2WNC | | 1 | 2 | 1 | 1 | 4 | | | 1 |
| Total number of orientations in each category | | | 11 | 46 | 18 | 26 | 4 | 11 | 8 | 35 | 21 |

An alternative method for orientating a ligand uses a protein in which a structurally similar ligand with a common pharmacophore has been co-crystallized (Fig. 7). To predict interacting amino acids, the new ligand can be pasted into the model. This method can result in steric clashes between the ligand and receptor, but these can be minimized with the software. It must be stressed, however, that all these *in silico* methods only estimate the possible orientations of amino-acid side chains and docked ligands. They do nevertheless provide testable hypotheses that can be validated by experimentation.

Table 3. Rigid ligand docking of granisetron into the 5-HT₃ receptor binding site produces docked clusters which can be categorized into one of five groups. Methods used can be found in the legend of Table 2

| Starting template for the 5-HT ₃ R homology model | | | Number of ligand orientations in each 5-HT ₃ R homology model | | | | | |
|--|----------------------|-------|--|----|----|----|----|-------|
| Protein | Ligand Type | PDB | A1 | A2 | B2 | C1 | C2 | Other |
| Aplysia AChBP | – | 2BYN | | | 10 | | | |
| Aplysia AChBP | HEPES | 2BR7 | 5 | 2 | | | | 3 |
| Limnaea AChBP | HEPES | 1I9B | 5 | 2 | | | | 3 |
| Limnaea AChBP | HEPES | 1UX2 | | 4 | 2 | | 4 | |
| Aplysia AChBP | Epibatidine | 2BYQ | 5 | | 5 | | | |
| Limnaea AChBP | Carbomylcholine | 1UV6 | | | | 8 | 2 | |
| Limnaea AChBP | Nicotine | 1UW6 | | 3 | 1 | | | 6 |
| Aplysia AChBP | Methylcaconitine | 2BYR | | | | 7 | | 3 |
| Aplysia AChBP | Lobeline | 2BYS | 1 | 1 | 8 | | | |
| <i>B. truncatus</i> AChBP | CAPS | 2BJO | 3 | | | | | 7 |
| <i>T. marmorata</i> nAChR | – | 2BG9 | | 1 | | | | 9 |
| Limnaea AChBP | α -Cobratoxin | 1YI5 | 1 | 2 | 1 | | | 6 |
| Aplysia AChBP | α -Cobratoxin | 2BYP | | 4 | | | 6 | |
| Aplysia AChBP | α -Conotoxin | 2C9 T | | | 6 | | | 4 |
| Erwinia LGIC | – | 2VL0 | | | | | | 10 |
| Gleobacter LGIC | – | 3EAM | | | | | | 10 |
| Aplysia AChBP | Cocaine | 2PGZ | | | | 10 | | |
| Aplysia AChBP | Tropisetron | 2WNC | | 5 | 5 | | | |
| Total number of orientation in each category | | | 20 | 24 | 38 | 25 | 12 | 61 |

3.2.2 Ligand binding: experimental evidence for the 5-HT₃R *in silico* predictions

The two methods (flexible and rigid docking) produced in total eight distinct categories (or clusters) of ligand orientations (see Fig. 6), with, for example, 26 and 38 poses respectively in the orientation B2. This places granisetron with the azabicyclic rings between W183 and Y234, and the indazole ring towards loop E. Orientation A2 is the total most common (46 and 24 poses, respectively) and has more interactions with residues that have been identified as important in the 5-HT₃R binding site (Joshi *et al.* 2006; Thompson *et al.* 2005). The orientation that is best supported by the experimental evidence, however, is A1, where the orientation of granisetron is reversed so that the indazole ring is located between W183 and Y234 with the azabicyclic ring orientated towards the transmembrane region, between residues E129 and W90. Double-mutant cycle analysis shows that the azabicyclic ring of granisetron is close to W90 and the indazole ring is orientated away from the membrane (Yan & White, 2005), and this orientation is also supported by experimental evidence as described by both Joshi *et al.* (2006) and Thompson *et al.* (2005). In both orientations A1 and A2, there is an interaction with W183, a residue that is important for both agonist and antagonist binding (Beene *et al.* 2002; Spier & Lummis, 2000), and with Y234, which also contributes to the binding site; substitutions of Y234 to Ala or Ser severely compromise granisetron binding, although Y234F mutants have similar binding affinities to wild-type receptors (Spier & Lummis, 2000; Suryanarayanan *et al.* 2005). An Ala mutation at the adjacent S233 residue also abolishes binding, which may be due to its altering the

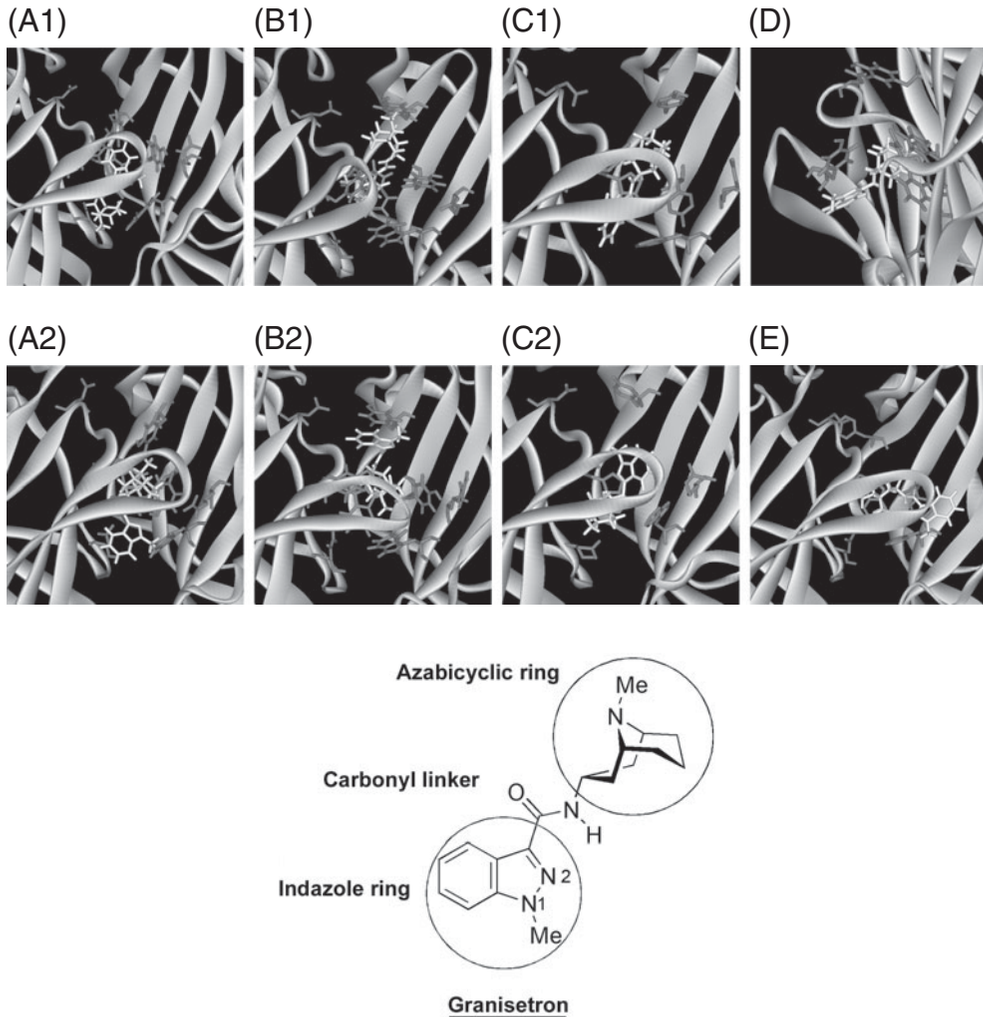


Fig. 6. Examples of the eight main categories of docked poses found in the 320 homology models generated for this study. Categories were largely based on the proximity of granisetron atoms to W183, and the orientation of the azabicyclic and indazole rings. The number of docked poses that fell into each of these categories can be seen in Tables 2 and 3. In brief, the descriptions of these clusters are as follows: (A1) Indazole ring close to W183 and the azabicyclic ring orientated towards the membrane. (A2) Same as A1, but with the azabicyclic and indazole rings reversed. (B1) Indazole ring close to W183 and the azabicyclic ring orientated towards Y143 in loop E. (B2) Same as B1, but with the azabicyclic and indazole rings reversed. (C1) Carbonyl linker close to W183 and the azabicyclic ring orientated away from the membrane. (C2) Same as (C1), but with the indazole ring orientated away from the membrane. (D) Either the azabicyclic or indazole rings close to W183 and the opposite end of the ligand orientated towards loop C. (E) Granisetron lies horizontally across the back of loop C. (Other) A number of unique positions located throughout the ECD.

location of the adjacent Y234 residue (Suryanarayanan *et al.* 2005). Mutation of both E129 and W90 strongly affect granisetron binding regardless of the amino acid used, showing that they are both essential; E129 hydrogen bonds with 5-HT, and may similarly interact with granisetron (Price *et al.* 2008; Spier & Lummiss, 2000; Sullivan *et al.* 2006; Yan *et al.* 1999), and W90 may

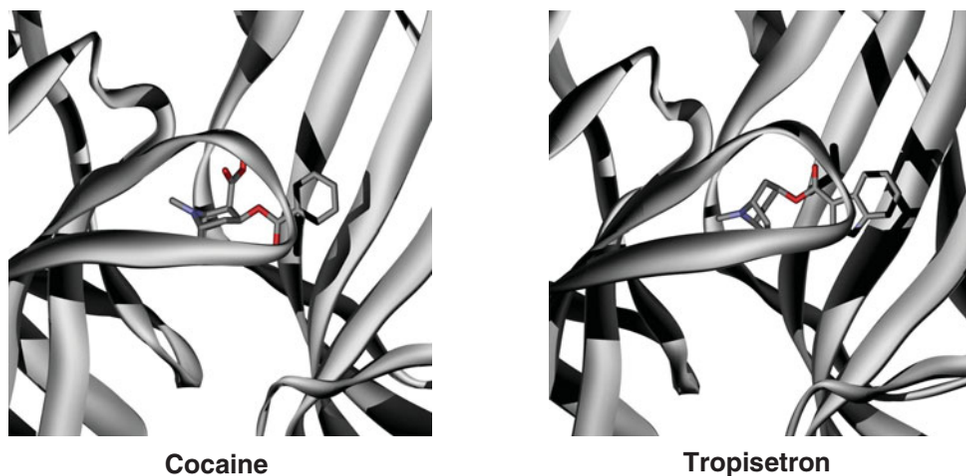


Fig. 7. AChBP crystal structures showing the orientations of cocaine (PDB ID: 2PGZ at 1.76 Å) and tropisetron (2WNC at 2.2 Å). The orientations are most similar to category E described in Fig. 6.

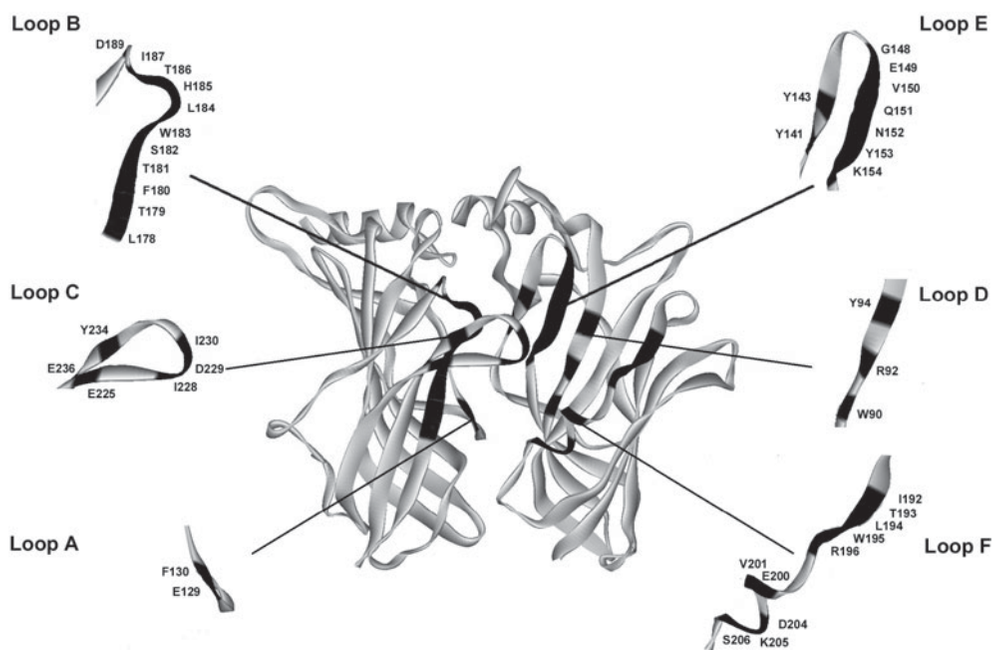


Fig. 8. Binding site substitutions that cause significant changes in the binding affinity of granisetron at the 5-HT₃ receptor. Residues have been superimposed upon a homology model of the 5-HT₃ receptor that was generated using 119B. The data were taken from Beene *et al.* (2004), Boess *et al.* (1997), Joshi *et al.* (2006), Price *et al.* (2008), Schreiter *et al.* (2003), Spier & Lummis (2000), Thompson *et al.* (2005, 2006b, 2008), Venkataraman *et al.* (2002a, b), Yan *et al.* (1999), Yan & White (2005) and Sullivan *et al.* (2006).

stabilize the structure of region by a T-type interaction with Tyr234 (Gallivan & Dougherty, 1999).

Residues that have an impact on granisetron binding are shown in Fig. 8. These include a number of residues centred around W195 and D204 in loop F. Whether the residues in

loop F are directly involved in ligand binding is difficult to determine from the homology models as this region is poorly resolved in the crystal structures, and some residues may interact with adjacent β -sheets rather than with the ligand itself (Spier & Lummis, 2000; Thompson *et al.* 2006b). Loop E residues G148 and V150 have been shown to abolish ligand binding when mutated to Ala and there are also effects at residues L178, F180, Q188, D189, I190 and N191 in loop B (Joshi *et al.* 2006; Thompson *et al.* 2008; Venkataraman *et al.* 2002b). As many of these are at some distance from the binding site, and some are on opposite sides of β -sheets, it is unlikely that they directly interact with the ligand; their effects may be due to intramolecular interactions that are critical for the structure of the binding site (Thompson *et al.* 2006b, 2008). Some of these residues have also been implicated in the binding/unbinding pathway of the ligand, while others may contribute to the subunit interface, or be involved in the transduction of binding energy into channel opening (Joshi *et al.* 2006; Thompson *et al.* 2006a).

3.2.3 Ligand binding: summary

Our docking results show a wide range of ligand orientations, highlighting the potential problem of developing theories based solely on *in silico* predictions. We can, however, use this information to design experiments to probe the accuracy of the predictions. For the 5-HT₃R, experimental data best support the predicted orientations of granisetron and 5-HT shown in Fig. 9, which are not the most common docking solutions, but are in general agreement with structure–activity relationships (Bower *et al.* 2008; Maksay *et al.* 2003; Reeves *et al.* 2003; Schmidt & Peroutka, 1989). It must also be considered that it may be possible for ligands to adopt multiple orientations. For example, molecular dynamic studies examining GABA binding to the GABA_CR show that GABA appears to ‘flip’ from one orientation to another during the simulation, although there is currently only experimental data to support one of the orientations (Melis *et al.* 2008), and *in silico* predictions in the 5-HT₃R have concluded that there are two possible orientations for both *m*CPBG and granisetron (Joshi *et al.* 2006; Schulte *et al.* 2006).

Comprehensive reports can be found elsewhere on the binding sites of 5-HT₃ (Schulte *et al.* 2006; Thompson & Lummis, 2006), nACh (Romanelli *et al.* 2007), Gly (Lynch, 2004) and GABA receptors (Abdel-Halim *et al.* 2008; Huang *et al.* 2006; Korpi *et al.* 2002; Sedelnikova *et al.* 2005).

3.3 Allosteric modulation

Cys-loop receptors are allosteric proteins, but they themselves are also subject to allosteric modulation by a wide range of organic and inorganic substances (Changeux *et al.* 1984). Some of these substances occur endogenously and may reinforce or attenuate the natural response under physiological conditions, but, given the central importance of Cys-loop receptors in the nervous system and neurological disorders, it is not surprising that some synthetic receptor modulators are widely used potent and effective drugs, such as the benzodiazepines, which act at GABA_A receptors. We will not attempt to describe the effects of all of these modulators, but will briefly describe some examples to give an indication of the diversity of compounds and the range of studies being undertaken to understand their mechanisms of action. Further information can be obtained from the following reviews for GABA_A (Huang *et al.* 2006; Olsen *et al.* 2004b), Gly

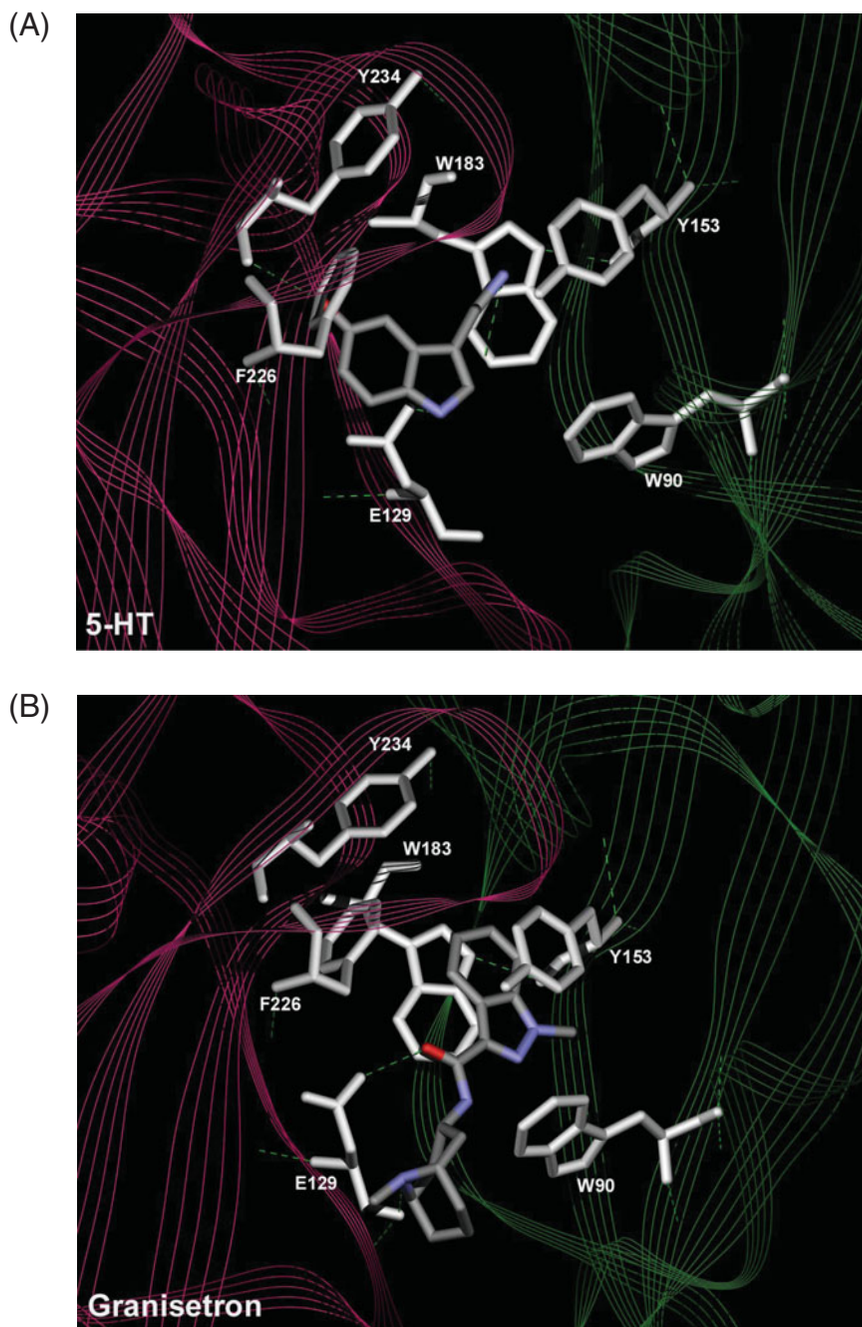


Fig. 9. Preferred orientations for 5-HT and granisetron docked into a homology model of the 5-HT₃ receptor binding site. Both orientations provide the best fit for the experimental data. The ligands can potentially interact with W183, are influenced by a range of aromatic residues that are orientated with their π -rings normal to the ligand, and have critical hydrogen bonds interactions with E129. See text for more details.

(Hawthorne & Lynch, 2006; Lynch, 2004, 2009), nACh (Faghieh *et al.* 2008; Arias & Bouzat, 2006) and 5-HT₃ receptors (Reeves & Lummis, 2002).

3.3.1 Ions as modulators

Receptors in the Cys-loop family can be significantly affected by physiologically relevant ions such as calcium, magnesium and zinc. The effects of these ions vary according to the receptor type and subunit composition. For example, in the $\alpha 7$ nAChR, these cations potentiate responses, while 5-HT₃R responses are typically reduced (Brown *et al.* 1998; Hu & Lovinger, 2005; Hubbard & Lummis, 2000; Niemeyer & Lummis, 2001; Thompson & Lummis, 2008a). Ion-binding sites in many Cys-loop receptors have been identified in the channel (Bertrand *et al.* 1993; Eddins *et al.* 2002a, 2002b; Gill *et al.* 1995; Hu & Lovinger, 2005; Livesey *et al.* 2008; Niemeyer & Lummis, 2001; Noam *et al.* 2008; Quirk *et al.* 2004; Thompson & Lummis, 2008a; Van Hooft & Wadman, 2003), but there are also binding sites in other regions of these proteins. A specific binding site for calcium, for example, has been identified in the ECD of the $\alpha 7$ nACh (Galzi *et al.* 1996), and insertion of this region into the 5-HT₃R results in an enhancement of the 5-HT-induced response. This region coincides with residues that have been shown to bind Ca²⁺ in AChBP (Brejc *et al.* 2001). Zinc-binding sites have been located at subunit interfaces in nAChR and GlyR (Hsiao *et al.* 2006; Nevin *et al.* 2003), while in GABA_A receptors, zinc binds to both the ECD and the pore (Dunne *et al.* 2002; Fisher & Macdonald, 1998; Fisher, 2002; Horenstein & Akabas, 1998; Hosie *et al.* 2003). Binding of these ions is likely to have important physiological consequences although these are not yet fully understood. In the GABA_A receptor, for example, sensitivity to zinc changes with the onset of epilepsy (Kapur & Macdonald, 1997), an effect that has been genetically linked to a mutation within the M2 region of the GABA_A $\gamma 2$ subunit (Baulac *et al.* 2001).

3.3.2 Benzodiazepines

Benzodiazepines are an important class of therapeutic compounds that modulate GABA_A receptors by binding at the α - γ subunit interface (Olsen & Sieghart, 2009). Differences in the pharmacological profiles of different α - and γ -subunit subtypes have enabled the identification of amino-acid residues that are involved in benzodiazepine binding. For example, $\alpha 1$ -His102 directly interacts with flunitrazepam and diazepam (Berezhnoy *et al.* 2004; McKernan *et al.* 1998; Tan *et al.* 2007), while $\alpha 1$ -Tyr160, $\alpha 1$ -Tyr210 (Amin *et al.* 1997) and $\gamma 2$ -Phe77 (Buhr *et al.* 1997a) form part of the aromatic-binding site for benzodiazepines. Residues $\alpha 1$ -Thr206, $\alpha 1$ -Glu209, $\alpha 1$ -Tyr162, $\alpha 1$ -Thr207, $\gamma 2$ -Tyr58, $\gamma 2$ -Ala79, $\gamma 2$ -Met130 and $\gamma 2$ -Thr142 contribute to benzodiazepine selectivity and efficacy (Buhr & Sigel, 1997; Buhr *et al.* 1997a, 1997b; Kucken *et al.* 2000; Mihic *et al.* 1994; Sigel & Buhr, 1997; Teissere & Czajkowski, 2001).

The mechanisms that communicate conformational changes between the GABA- and benzodiazepine-binding sites are less well understood. Mutations in loop F of the $\gamma 2$ subunit do not change the binding affinity of benzodiazepines or the agonist response, but decrease potentiation, indicating that this region may be involved in telegraphing the modulatory behaviour to other areas of the receptor (Hanson & Czajkowski, 2008; Padgett & Lummis, 2008). Other regions of the protein are also probably involved, including the $\beta 10$ sheet of the ECD (see Fig. 3), and residues in M1, M2 and the M2–M3 loop (Boileau & Czajkowski, 1999; Jones-Davis

et al. 2005). Further reading on benzodiazepines can be found in Olsen & Sieghart (2009), Rudolph *et al.* (2001) and Sigel (2002).

3.3.3 Alcohols and anaesthetics

A wide range of alcohols and anaesthetics modulate Cys-loop receptor function, and their behaviours are mostly mediated via interactions with the TMD (Arias & Bhumireddy, 2005; Hawthorne & Lynch, 2006; Huang *et al.* 2006; Sessoms-Sikes *et al.* 2003). Effects of these compounds vary according to the receptor type, subunit composition, and the nature and concentration of compound being used. For example, long *n*-alkanols and anaesthetics are inhibitory at nAChRs, but ethanol is potentiating at low concentrations (Zuo *et al.* 2002), while 5-HT₃Rs are potentiated and inhibited depending upon the alcohol or anaesthetic (e.g. Machu & Harris, 1994; Suzuki *et al.* 2002; Zhang *et al.* 1997). Other examples include $\alpha 4\beta 2$ nAChRs, which are sensitive to the anaesthetics isoflurane and propofol, and $\alpha\beta\gamma\delta$ nACh and $\alpha 7$ nAChRs which are not (Flood *et al.* 1997; Violet *et al.* 1997). There are many other examples, and there are excellent reviews on this subject by Arias & Bhumireddy (2005), Urban *et al.* (2006) and Yamakura *et al.* (2001).

3.3.4 Ivermectin – a commercially important modulator of invertebrate GluCl receptors

Ivermectin, a macrocyclic lactone produced by bacteria, is the world's largest-selling veterinary drug, and has also largely eradicated 'river blindness' resulting from nematode infections in sub-Saharan Africa. Ivermectin, its avermectin analogues and the milbemycins are probably allosteric potentiators of invertebrate GluCl channels at submicromolar concentrations (Vassilatis *et al.* 1997). These heteromultimeric channels, found in several invertebrate phyla, are homologous to vertebrate GlyR (and slightly less so to GABA_A receptors).

We know little about the binding site for ivermectin, but because the GluCl channels resemble other Cys-loop receptors, it is certain that the GluCl β subunit carries the principal-binding site. The GluCl β Tyr182 residue aligns with the loop B cation- π residues: Trp of the nAChR, Trp of the 5-HT₃R and (probably most similar) Phe of the GlyR (see Table 1). Substitutions to several other residues at this position abolish the responses to both glutamate and ivermectin. However, the GluCl β -Y182F mutation decreases the maximal glutamate response by ~6-fold, without changing the ivermectin response (Li *et al.* 2002). This is evidence that the binding sites for glutamate and IVM do not overlap. Twenty other mutations were studied in the β -subunit ECD; none preserved glutamate sensitivity while abolishing ivermectin sensitivity (Li *et al.* 2002). Therefore, we cannot say where ivermectin binds to GluCl channels.

GlyR are also activated by ivermectin, but are ~1000-fold less sensitive (Shan *et al.* 2001), suggesting that ivermectin may act differently on GlyR. Nevertheless, several GlyR-binding site mutations abolish glycine but not ivermectin sensitivity, supporting the idea that the agonist and ivermectin sites do not overlap (Shan *et al.* 2001). Also supporting the idea that ivermectin binds at a non-agonist site, voltage-clamp fluorometry established that the 19' residue near the top of M2 changes its environment when the channel is opened by all agonists but not when opened by ivermectin (Pless *et al.* 2007).

A mystery associated with ivermectin is its very low reversibility, which vitiates concentration-response experiments. Ivermectin effects take > 8 h to wash out and this lower limit could actually be governed by a synthesis of new receptors (Slimko *et al.* 2002). The most appropriate experiments show that GluCl channels are half-activated by ivermectin during a 1 nM puff lasting

several seconds (Slimko & Lester, 2003). In unpublished experiments (HAL lab), hypersensitive nAChR mutants with comparably low EC₅₀ values show washout time constants of several minutes; therefore, simple high affinity does not explain the long washout times for ivermectin at GluCl channels. An unnatural Pro substitute in the M2–M3 linker of 5-HT₃Rs produces an apparently irreversible activation (Lummiss *et al.* 2005b) with an EC₅₀ of 20 nM, and may represent a good analogy to the action of ivermectin. Further comments on ivermectin can be found in sections 3.3.5 and 7.5.

3.3.5 $\alpha 7$ nACh receptor allosteric modulators

Among the nAChRs, the $\alpha 7$ nAChR has received recent attention as a target for allosteric activators (Bertrand *et al.* 2008; Hogg & Bertrand, 2007). Potent positive allosteric modulators include NS-1738, 4-naphthalene-1-yl-3*a*,4,5,9*b*-tetrahydro-3-*H*-cyclopenta[*d*]quinoline-8-sulfonic acid amide (TQS), PNU-120596, *N*-(4-chlorophenyl)- α -[[[4-chloro-phenyl]amino] methylene]-3-methyl-5-isoxazoleacet-amide ('compound 6'; Ng *et al.* 2007), LY-2087101 (Broad *et al.* 2006) and galanthamine (Lopes *et al.* 2007); these act at concentrations $\leq 10 \mu\text{M}$. TQS and PNU-120596, but not NS1738, have the property that they either reactivate desensitized receptors and/or significantly retard desensitization, properties that are also shared by ivermectin at GlyR (Gronlien *et al.* 2007). However, according to results from $\alpha 7/5$ -HT₃ chimeras, NS-1738 and PNU-12059 bind at different sites. The ECD of the $\alpha 7$ nAChR is required for NS-1738 action, and the ECD in combination with the M2–M3 linker of the $\alpha 7$ nAChR are required for agonist-independent activity in the presence of NS-1738 (Bertrand *et al.* 2008). In contrast, the entire TMD of the $\alpha 7$ nAChR is required for allosteric modulation by PNU-12059 (Bertrand *et al.* 2008).

4. The TMD

4.1 Structure

A range of experimental techniques show that the TMD is composed of four membrane spanning α -helices (M1–M4) from each subunit; each receptor therefore has 20 such α -helices within the membrane (Fig. 10). The α -helical nature of these regions, which was originally inferred from hydrophathy plots (Noda *et al.* 1982), was verified in the nAChR by photolabelling (Blanton & Cohen, 1994), two-dimensional ¹H-NMR spectroscopy (Lugovskoy *et al.* 1998), Fourier transform infrared (FTIR) spectroscopy (Baenziger & Methot, 1995; Corbin *et al.* 1998; Görne-Tschelnokow *et al.* 1994; Methot *et al.* 1994) and mutagenesis (e.g. Tamamizu *et al.* 2000). The best available structural information comes from 4 Å resolution cryo-electron microscopy images of the nAChR (Miyazawa *et al.* 2003); higher-resolution structures from related prokaryotic receptors have been solved, but it is not yet clear how representative these are of vertebrate Cys-loop receptors (Bocquet *et al.* 2009; Hilf & Dutzler, 2008; Nury *et al.* 2009). The 4 Å nAChR images show that α -helices from each subunit are arranged symmetrically, forming an inner ring of M2 helices that line the central pore, and an outer ring composed of M1, M3 and M4 that shields the inner ring from the lipid environment (De Planque *et al.* 2004; Miyazawa *et al.* 2003). On the extracellular side, the transmembrane helices are spread apart, but gather together as they cross the membrane towards the intracellular side (Goren *et al.* 2004; Miyazawa *et al.* 2003; Panicker *et al.* 2002). Overlaying electron densities of subunits in the resting state reveals that M1,

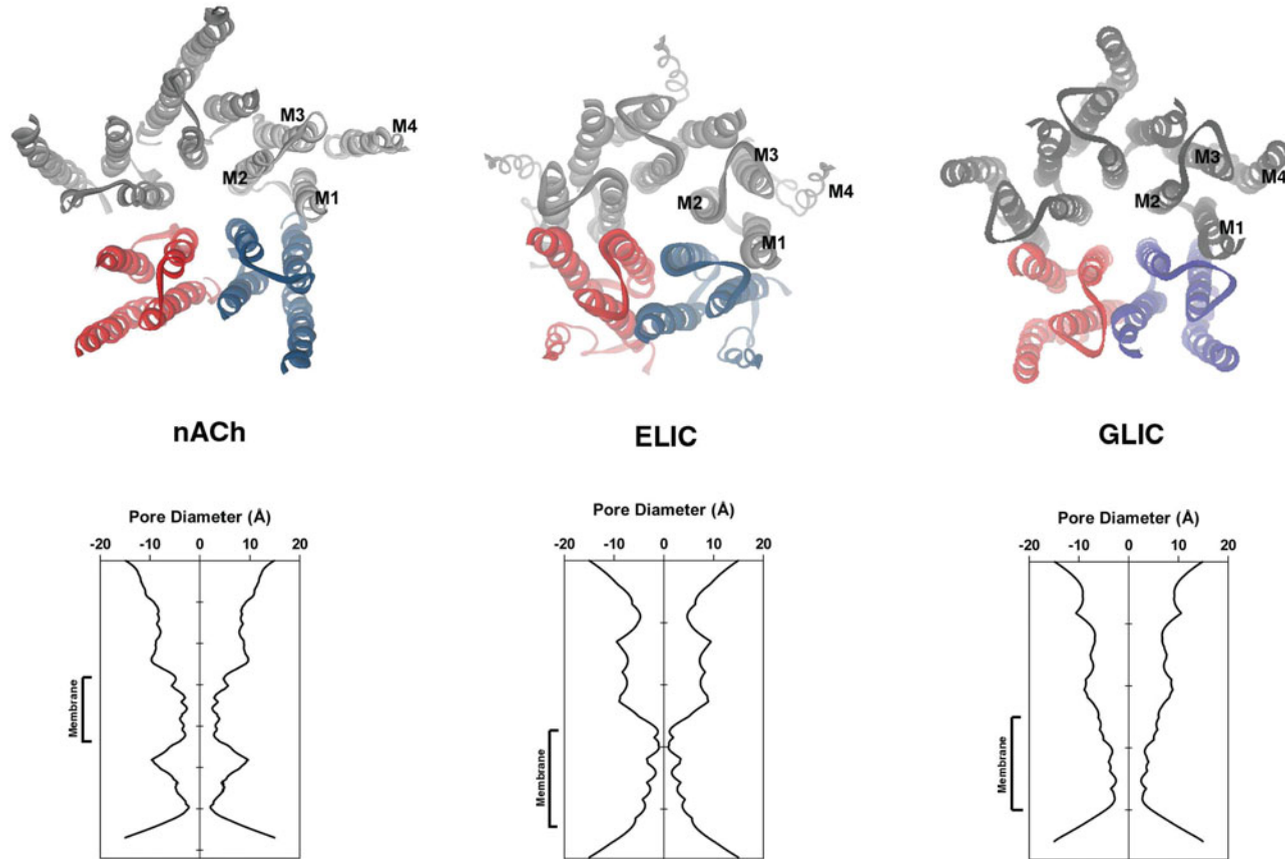


Fig. 10. Comparisons of TMDs from open and closed receptors. Top panel: structures of nAChR (closed; Miyazawa *et al.* 2003; PDB ID: 2BG9), ELIC (closed; Hilf & Dutzler, 2008; PDB ID: 2VL0) and GLIC (open; Bocquet *et al.* 2007; PDB ID: 3EAM) are shown. Two (red and blue) of the five subunits are highlighted. M2 lines the central pore, and residues that face this water accessible surface are shown in Fig. 11. Lower panels: pore diameter as calculated by HOLE, with a 15 Å cut-off to find the ends of the pores (Smart *et al.* 1993). Each tick on the vertical axis is 25 Å. The nAChR pore appears longer, because the structure also contains part of the ICD.

M2 and M3 have precise positioning within the structure while the location of M4 is more relaxed, particularly at its C-terminus. A review of pore structures can be found in Absalom *et al.* (2009).

4.2 M1 and the M1–M2 loop

M1 forms part of the outer ring that is in contact with the lipid environment and may also contact M2. Mutations in M1 have been shown to produce receptors that have altered desensitization, changes in EC_{50} or are non-functional (Akabas & Karlin, 1995; Bianchi *et al.* 2001; Dang *et al.* 2000; Engblom *et al.* 2002; England *et al.* 1999; Greenfield *et al.* 2002; Lobitz *et al.* 2001; Lobo *et al.* 2004; Spitzmaul *et al.* 2004; Zhang & Karlin, 1997). M1 may therefore be a structural element involved in transmitting movement of the ligand-bound ECD into M2, possibly through direct interactions with the M2 helix following activation (Unwin *et al.* 2002). Indeed, some of the roles of specific residues that contribute to this activity are beginning to emerge. For example, the highly conserved proline residue in the centre of M1 has been shown to be critical due to its lack of ability to act as a hydrogen bond donor (Dang *et al.* 2000), and may permit M2 to transiently alter its position upon channel activation. Recent experiments introducing ionizable side chains into M1 revealed the current response is reduced 25–50% by protonation at any of five α -helically spaced M1 side-chains, suggesting M1 is not completely shielded from the channel axis by M2. The data also show that the side chains closest to the axis in the open state are also those closest in the cryo-electron microscopy studies, revealing that M1 moves little, or may not move at all, between the open and closed states (Cymes & Grosman, 2008; Miyazawa *et al.* 2003).

There is evidence that the region that links M1 with the ECD is an important functional element involved in the gating process. Mutations at the extracellular end of the 5-HT₃R (R222, Hu *et al.* 2003), GlyR (R218, Castaldo *et al.* 2004), GABA_AR (K215, Kash *et al.* 2004) and nAChR (several residues, Zhang & Karlin, 1997) have been shown to effect receptor gating. The other end of M1 may form part of the intracellular mass that lies at the cytoplasmic face of the pore. Evidence from SCAM has indicated that the intracellular end of M1 and the M1–M2 linker lie along the path of the permeating ions, and these regions contain residues responsible for anion/cation selectivity (Filippova *et al.* 2004; also see section 4.4). Coordination of cadmium ions in 5-HT₃R Cys mutants and the use of negatively and positively charged thiol reactive MTS reagents have demonstrated that residues in the M1–M2 loop are accessible (Panicker *et al.* 2002, 2004).

4.3 M2 lines the channel pore and acts as the channel gate

A great number of experiments over many years have shown that the residues in M2 line the channel pore and M2 is an α -helix. In particular, substituting cysteine, histidine, lysine or arginine residues into M2 has revealed water-accessible pore lining residues that have a periodicity consistent with an α -helical conformation (Akabas *et al.* 1994; Cymes *et al.* 2005; Reeves *et al.* 2001; Xu & Akabas, 1996; Zhang & Karlin, 1998). Structural data from cryo-electron microscopy indicate that M2 is ~ 40 Å long and extends beyond the embrace of the lipid environment (Bachy *et al.* 1993; Miyazawa *et al.* 2003). These data also show that within the limits of the membrane the M2 helices tilt radially towards the centre of the pore until they reach residues 6'–9', a region that is considered to be the channel gate (Fig. 10). This is consistent with earlier

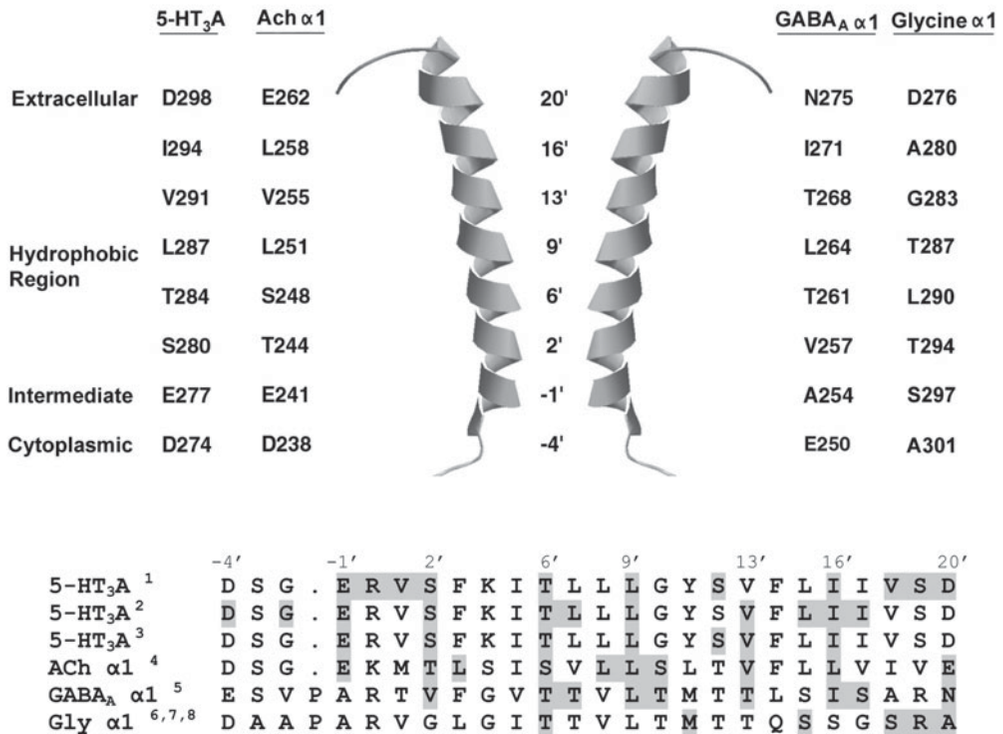


Fig. 11. M2 channel lining residues in different Cys-loop receptors. Pore-lining residues are shown next to the M2 α -helix (taken from the nAChR structure; PDB ID: 1oed). To allow for comparisons between M2 residues of different Cys-loop receptors, a prime (') notation is used: residues are assigned numbers according to their position relative to a charged M2 residue that is conserved in Cys-loop receptors. Residues that are accessible to modification are highlighted: ¹Kaneez & White (2004), ²Reeves *et al.* (2001), ³Panicker *et al.* (2002), ⁴Akabas *et al.* (1994), ⁵Xu & Akabas (1996), ⁶Shan *et al.* (2002), ⁷Lynch *et al.* (2001) and ⁸Lobo *et al.* (2004). Accession numbers for the sequence alignment are: Mouse 5-HT_{3A}, Q6J1J7; Electric Ray ACh α 1, P02710; Human GABA_A α 1 P14867; Human Gly α 1, P23414.

studies that showed the binding site for the open-channel blocker QX-222 was located at the region between 6' and 10' of the nAChR, and suggested that the open channel tapered to its narrowest point just below 6' (Charnet *et al.* 1990; Leonard *et al.* 1988).

The centre of the pore coincides with a conserved collar of hydrophobic side chains; in the nAChR α 1 subunit this includes Leu251, Val255 and Leu258 (Figs 1, 2, 10 and 11). Five symmetrically placed M2 helices from each of the five subunits create a hydrophobic region that is 3 Å at its narrowest and less than 3.5 Å over a distance of approximately 8 Å in the closed state, and has been referred to as a hydrophobic girdle (Miyazawa *et al.* 2003). Ion permeabilities suggest that the diameter of the open channel is between 7.4 Å and 8.4 Å for cation channels and between 5.2 Å and 6.2 Å for anion channels (Brown *et al.* 1998; Cohen *et al.* 1992; Fatima-Shad & Barry, 1993; Rundstrom *et al.* 1994; Wang & Imoto, 1992). Originally, the channel gate was predicted to be close to the cytoplasmic end of M2 (Wilson & Karlin, 1998; Wilson *et al.* 2000). This conclusion was supported by data from applying thiol reactive compounds in the closed or open state of the receptor, but the results were limited as channels may spontaneously open, modifying reaction rates can vary in the presence and absence of agonist and some of the modifying reagents may be small enough to pass the closed gate (Bali & Akabas, 2007; Tikhonov

et al. 2004; Wilson & Karlin, 2001). It is now widely accepted that the gate is located in the centre of the channel. This constricted region shares structural similarities to the gate of other membrane-permeable channels which are also occluded by a narrow hydrophobic region (Chang *et al.* 1998), and represents the only obstruction within the channel that provides an energetic barrier to ion permeation (Beckstein *et al.* 2001; Hummer *et al.* 2001).

4.4 M2 and ion selectivity

The residues that line the ion-accessible inner face of the channel are predominantly non-polar except for rings of charged amino acids (Figs 1 and 11) (Akabas *et al.* 1992, 1994; Panicker *et al.* 2002; Reeves *et al.* 2001; Xu & Akabas, 1993; Xu *et al.* 1995; Zhang & Karlin, 1998). Initially, Konno *et al.* (1991) reported that the three rings of charged amino acids (referred to as extracellular, intermediate and cytoplasmic rings) in the nAChR M2 region were responsible for ion selectivity, with the intermediate ring exerting the strongest influence (Fig. 11). The mechanisms of charge selectivity were later evaluated by substituting M2 residues of the $\alpha 7$ nAChR with the corresponding residues from Gly $\alpha 1$ (Galzi *et al.* 1992). As the number of mutations was gradually reduced, the smallest number of residues required to reverse ion selectivity was found to be valine to threonine (V251T or V9'T), neutralization of a glutamate (E237A or E-1'A) and the insertion of a proline (P236 or P-2') in the M1–M2 loop. Homologous changes also alter ion selectivity in the 5-HT₃ (Gunthorpe & Lummis, 2001; Thompson & Lummis, 2003), MOD-1 (Menard *et al.* 2005), GABA ρ (Carland *et al.* 2004; Wotring *et al.* 2003), GABA_A (Jensen *et al.* 2002, 2005a) and Gly $\alpha 1$ receptors (Keramidas *et al.* 2000). The contribution of each of the three mutations was studied in more detail in the $\alpha 7$ nAChR, and showed that the residue at the $-1'$ position is the most critical (Corringer *et al.* 1999a). At this position in cationic receptors glutamate predominates, while the $-1'$ residue in anionic receptors is uncharged (Keramidas *et al.* 2002; Wotring *et al.* 2003). In the $\alpha 7$ nAChR V9' was found not to be directly involved in charge selectivity, and the effect of the Pro insertion was the result of localized structural modifications at the intracellular end of M2 (Corringer *et al.* 1999a). Indeed, in the $\alpha 7$ nAChR, functional anionic channels could be generated by inserting a Pro at any of the positions 234, 236 or 237, although was dependent on being accompanied by the E237A and V251T mutations. The structural importance of the Pro was further illustrated by changes in functional properties of the mutant receptors, including differences in activation, desensitization, EC₅₀, Hill co-efficient and spontaneous channel openings (Corringer *et al.* 1999a; Wotring *et al.* 2003). At the invertebrate GluCl receptor, charge selectivity remains unaltered in similar Pro mutants and a structural role was also concluded (Sunesen *et al.* 2006). Further complications were presented in a study by Wotring & Weiss (2008), who also showed that the introduction of Glu residues within an eight amino-acid stretch ($-2'$ to $5'$) of GABA $\rho 1$ produces varied permeability ratios depending upon the location of the substitution. However, in native receptors the character of the $-1'$ residue is conserved across the Cys-loop family, indicating that this position is critical. Consequently, the ring of charge at the $-1'$ position is now universally regarded as an essential component of charge selectivity within the Cys-loop family, with other residues in the region playing some roles in some receptors.

Mutations in the 5-HT₃R close to the extracellular ring of charge have also been implicated in charge selectivity (Thompson & Lummis, 2003). When the 19' residue is changed from a serine to an arginine and combined with the $-1'$ Glu to Ala mutation, the receptor is predominantly anion selective. Importantly, unlike the triple mutants described above, these 5-HT₃R mutants

do not display concomitant changes in the biophysical properties of the channel, suggesting that these data more accurately reflect the residues directly involved in ion selectivity. Neutralization of R19' in the GlyR, however, does not alter ion selectivity when expressed in conjuncture with A1'E and P2' Δ mutations, although it does modify conductance and rectification (Keramidas *et al.* 2002; Moorhouse *et al.* 2002). These data indicate that there are structural differences between the cation- and anion-selective receptors (also see section 4.5).

As functional receptors can be formed from different combinations of subunits (which have different amino acids lining their pores), there can be large differences in the permeation of certain ions. One of these is Ca^{2+} , and nAChRs have a wide range of Ca^{2+} permeabilities (Arias, 2006; Cens *et al.* 1997; Gerzanich *et al.* 1998; Livesey *et al.* 2008; Noam *et al.* 2008; Tapia *et al.* 2007; Vernino *et al.* 1992) determined primarily by the residues located at the intermediate ($-1'$) and extracellular ($20'$) rings (Bertrand *et al.* 1993; Galzi *et al.* 1992; Hu & Lovinger, 2005; Livesey *et al.* 2008). For example, reduced Ca^{2+} conductance in $(\alpha 4)_2(\beta 2)_3$ nAChRs compared to $(\alpha 4)_3(\beta 2)_2$ nAChRs is a consequence of only $\beta 2$, but not $\alpha 4$ subunits having acidic residues at their $-1'$ positions (Tapia *et al.* 2007). Such data can be extrapolated to other receptors: 5-HT₃ABR have lower Ca^{2+} permeability than 5-HT₃AR, which may be the consequence of $20'$ residue being Asp and Asn in A and B subunits, respectively. Consistent with this, a D20'A substitution reduces Ca^{2+} permeability, as does the replacement of the adjacent R19' with Ser (Livesey *et al.* 2008; Thompson & Lummis, 2003). Recent studies suggest that the ICD may also play a role in Ca^{2+} permeability as substitutions of charged residues in this region can have a major effect on Ca^{2+} permeability (Livesey *et al.* 2008; Thompson & Lummis, 2003). In both nAChR and 5-HT₃Rs, Ca^{2+} binding sites have also been reported in the ECD (see section 3.3.1).

Comprehensive reviews on ion selectivity in the Cys-loop family of receptors can be found in Jensen *et al.* (2005b), Keramidas *et al.* (2004), Peters *et al.* (2010) and Sine *et al.* (2010).

4.5 The M2–M3 loop

The M2–M3 loop forms part of the interface that links the ECD with the TMD, and it has a critical role in transmitting the energy of binding into channel opening (discussed further in section 6). Studies have shown that mutations in this region disrupt activation in nACh, 5-HT₃, GABA and Gly receptors (Campos-Caro *et al.* 1996; Deane & Lummis, 2001; Grosman *et al.* 2000a, 2000b; Kusama *et al.* 1994; Lewis *et al.* 1998; Lynch *et al.* 1997; O'Shea & Harrison, 2000; Rajendra *et al.* 1995; Rovira *et al.* 1998, 1999; Saul *et al.* 1999; Sigel *et al.* 1999). The structure of this loop has been examined by a range of techniques, including NMR and electron microscopy, and the data suggest that there are differences between cation- and anion-selective receptors. In the nAChR, the M2 helix extends two rings above the membrane (i.e. up to the $23'$ residue), while in the GlyR, the helix terminates at the $15'$ residue (Ma *et al.* 2005). The loop moves during receptor activation; in the Gly $\alpha 1$ receptor, SCAM studies reveal that all the residues within the M2–M3 region are accessible to modification, and surface accessibility increases when the receptor is activated (Bera *et al.* 2002; Lynch *et al.* 2001). Specific residues in this loop play particular roles, for example, in the 5-HT₃R a *cis*–*trans* isomerization of the Pro at the apex of this loop (Pro308, P8') can trigger channel opening (Lummis *et al.* 2005b). While the same mechanism seems not to activate the nAChR, the equivalent proline functionally couples to flanking Val residues extending from the $\beta 1$ – $\beta 2$ and Cys-loops, and together these regions form a critical part of the transduction pathway (Lee *et al.* 2008). A conserved proline within the Cys-loop has also

been identified as a candidate for channel activation (Limapichat *et al.* 2010). This topic is also discussed in section 6.

4.6 M3 and M4 helices

Structural data show that M3 and M4 are α -helical and shield M2 from the lipid bilayer, although there are water-accessible clefts that lie between the TMD α -helices (Miyazawa *et al.* 2003; Fig. 10). SCAM studies on the M3 segment of GABA_A receptors show that in the absence of GABA, only those residues towards the extracellular side of the membrane are accessible. Activation allows modifying reagents to approach residues located closer to the centre of the M3 α -helix, as water-permeable clefts between adjacent α -helices widen as a consequence of conformational changes in M2 (Miyazawa *et al.* 2003; Wang *et al.* 1999; Williams & Akabas, 1999). The outer face of the M3 helix is in close contact with the membrane and is inaccessible to these modifying agents (Blanton & Cohen, 1992, 1994; Blanton *et al.* 1998). Recent nAChR experiments introduced ionizable side chains into M3 to reveal relative distance from the channel's axis, similar to the experiments performed in M1 and M2 (Cymes *et al.* 2005). The extent of block was <40% for only five side chains, limiting the precision of the data, but was not inconsistent with the α -helical pattern and orientation of the closed state found in structural studies. M3, like M1, apparently rotates little between the closed and open states.

Given the location and apparent roles of M3 and M4, it is surprising that mutations can have significant effects on receptor function, but such mutations cause changes in both the whole-cell current (Cruz-Martin *et al.* 2001; Guzman *et al.* 2003; Lasalde *et al.* 1996; Williams & Akabas, 1999; Wu *et al.* 2010) and single-channel kinetics of nAChRs and 5-HT₃Rs (Bouzat *et al.* 2000, 2002; Corradi *et al.* 2009; De Rosa *et al.* 2002; Lee *et al.* 1994; Navedo *et al.* 2004; Ortiz-Miranda *et al.* 1997; Tamamizu *et al.* 1999, 2000; Wang *et al.* 1999). The effect of these mutations can be additive both in terms of the contribution from each subunit (Bouzat *et al.* 1998; De Rosa *et al.* 2002) and within the same α -helix (Lasalde *et al.* 1996), although this observation is not supported by all studies (Cruz-Martin *et al.* 2001). M4 may also detect the lipid environment and influence the functional properties of the receptor (daCosta & Baenziger, 2009).

These regions also influence receptor expression: the number and characteristics of C-terminal residues in M4 are critical for the expression of 5-HT₃R on the cell surface (Butler *et al.* 2009) and the expression of non-assembling receptors that contain only the ECD and M1–M3 helices can be rescued by co-expression with M4 (Haeger *et al.* 2010; Villmann *et al.* 2009). Thus, the M3 and M4 regions are an integral part of the receptor, and have a function that extends beyond simply shielding M2 from the membrane.

5. The ICD

5.1 Structure

The structure of the ICD is unresolved apart from a single α -helix that is located in the M3–M4 loop of each subunits. Electron microscopy-derived images at 9 Å resolution show that the ICD adopts a 'hanging-basket'-type structure with openings or 'portals' that are the likely site of ion entry and exit to/from the channel (Hales *et al.* 2006; Unwin, 1993). One side of each portal is contributed by the α -helix described above. This amphipathic α -helix was

originally identified by sequence analysis many years ago and was considered to line the pore (Finer-Moore & Stroud, 1984; Miyazawa *et al.* 1999). More recent studies have shown that it contributes significantly to the channel conductance in both nAChRs and 5-HT₃Rs (Hales *et al.* 2006; Peters *et al.* 2005). Interestingly, homologous receptors in bacteria do not have an extended loop region between M3 and M4, which has led to experiments where this region was deleted in 5-HT₃Rs and GABA_CRs, demonstrating that these receptors maintained most of the characteristics of the parent receptor (Bocquet *et al.* 2007; Hilf & Dutzler, 2008; Jansen *et al.* 2008). Thus, these regions are not essential for function or expression.

5.2 Channel conductance

Studies that demonstrated a role of the M3–M4 loop in channel conductance were originally performed in the 5-HT₃R, and more recently extended to nAChRs and GlyRs (Carland *et al.* 2009; Deeb *et al.* 2007; Hales *et al.* 2006; Kelley *et al.* 2003; Livesey *et al.* 2008; Peters *et al.* 2004). In the 5-HT₃R, A-subunits can form functional homomeric channels with a conductance < 1 pS, but when combined with B-subunits receptors display a much larger conductance (9–17 pS; Brown *et al.* 1998; Davies *et al.* 1999; Derkach *et al.* 1989; Hussy *et al.* 1994). By replacing parts of the A-subunit sequence with homologous regions from the B-subunit, Kelley *et al.* (2003) identified three amino acids that govern the differences between the low conductance of the homomeric receptor and the higher conductance of the heteromeric receptor, and which align with a polar stripe of residues identified by Finer-Moore & Stroud (1984). Electrophysiological data support the suggestion that the charged groups line portals within the sides of the ICD, and influence the ion flux between the cytoplasm and the inner vestibule at the base of the pore (Miyazawa *et al.* 1999; Unwin, 2000). As the widest region of the portals resembles the diameter of a hydrated permeant ion, they provide an explanation for the homomeric 5-HT₃ channel having a much smaller unitary conductance than most nAChRs, despite their similar ionic permeabilities and very similar M2 sequences (Brown *et al.* 1998; Lambert *et al.* 1989; Malone *et al.* 1991; Mochizuki *et al.* 2000; Yakel *et al.* 1990; Yang, 1990). Additional studies have shown that the conductance of the channel can be dynamically altered by sulphhydryl modifying reagents (Deeb *et al.* 2007) and the permeability of divalent cations is also altered by mutations in this region (Livesey *et al.* 2008). A peptide that mimics this region at GABA_ARs similarly modulates conductance at inside-out patches (Everitt *et al.* 2009). Charged residues at this location have also been proposed to interact with phosphate groups on intracellular proteins, regulating both channel conductance and ion selectivity (Livesey *et al.* 2008; Noam *et al.* 2008).

5.3 Intracellular modulation

Interactions of proteins and ions with the M3–M4 loop of Cys-loop receptors can modulate receptor activity, assembly, targeting and trafficking (e.g. Akk & Steinbach, 2000; Bouzat *et al.* 1994; Boyd *et al.* 2002; Melzer *et al.* 2010; Yu & Hall, 1994). Some interactions are highly specific to different subunits of different receptors, such as gephyrin that specifically targets GlyR to postsynaptic synapses, while others are more general; phosphorylation of the M3–M4 loop, for example, is linked to changes in channel behaviour in nACh, 5-HT₃, GABA_A and Gly receptors (Filippova *et al.* 2000; Hubbard *et al.* 2000; Lankiewicz *et al.* 2000; McDonald & Moss, 1994; McDonald & Moss, 1997; Moss *et al.* 1992, 1996; Nishizaki & Ikeuchi, 1995; Nishizaki & Sumikawa, 1998; Ruiz-Gomez *et al.* 1991; Sedelnikova & Weiss, 2002; Vaello *et al.* 1994;

Van Hooft & Vijverberg, 1995; Wecker *et al.* 2001). A comprehensive series of reviews by Connolly (2008), Gaimarri *et al.* (2007), Kneussel & Loebrich (2007), Millar (2008), Millar & Harkness (2008) and Sarto-Jackson & Sieghart (2008) cover the assembly and trafficking of nACh, 5-HT₃, GABA_A and Gly receptors.

A variety of kinases including casein kinase II, tyrosine kinase, protein kinase A (PKA) and protein kinase C (PKC) phosphorylate different residues with differing effects. For example, PKA phosphorylates Ser409 in the 5-HT₃R causing an increase in the rate of desensitization (Coultrap and Machu, 2002; Hubbard *et al.* 2000; Lankiewicz *et al.* 2000; Sun *et al.* 2003; Yakel *et al.* 1990), while PKC actions on this receptor regulate the probability of certain conductance states (Coultrap & Machu, 2002; Van Hooft & Vijverberg, 1995) and rapidly increase surface expression (Emerit *et al.* 2002; Grailhe *et al.* 2004; Ilegems *et al.* 2004; Sun *et al.* 2003). Other processes of post-translational modulation such as protein glycosylation and palmitoylation have also been described, but the exact roles of these processes in the regulation of receptor assembly, targeting and trafficking are not yet fully determined (e.g. Boyd *et al.* 2002; Drisdell *et al.* 2004; Green *et al.* 1995).

6. Molecular basis of Cys-loop receptor activation

Binding of an agonist to its receptor causes movements of the ECD that are transduced to the M2 helices and lead to the opening of the pore (e.g. Grosman *et al.* 2000b; Lee *et al.* 2009; Unwin *et al.* 2002). In the heteromeric nAChR, this movement is initiated within the α -subunits, which undergo rotations, although several recent studies also describe movements that precede rotation (Horenstein *et al.* 2001; Lape *et al.* 2008; Pless & Lynch, 2009; Unwin *et al.* 2002). The structural integrity of the ECD is important as weakening backbone hydrogen bonds in the β 7, β 9 and β 10 sheets abolish receptor function, while photochemically cleaving the backbone between loops A and E has similar effects on GABA activation, but not on activation by pentobarbital (Gleitsman *et al.* 2009; Hanek *et al.* 2010). The movement of the ECD, mediated by the M2–M3 linker at the extracellular side of the TMD, destabilizes the hydrophobic ‘girdle’ in the channel, which moves away from the centre of the pore into space that resides between the inner and outer rings, opening the channel (Law *et al.* 2000; Miyazawa *et al.* 2003). It is often claimed that Miyazawa *et al.* (2003) found a ‘rotation’ in the M2 regions, but only rotations in the ECD were identified, along with two alternative structures of the M2 helices: straight in the closed state and kinked in the open state. The cytoplasmic ends of M2 remain relatively static during these events (Panicker *et al.* 2002, 2004). In support of this hypothesis, mutations of the gate residues affect ion permeation, cause increased sensitivity to channel opening, slow desensitization of macroscopic currents, increase closing events and/or increase channel open times (Chang & Weiss, 1998; Filatov & White, 1995; Labarca *et al.* 1995). A collective movement of all the M2 helices is likely, as they maintain their five-fold symmetry in both the closed and open states (Unwin, 1995), and the gate residue α L251 effects pore opening independent of the nAChR subunit type mutated (Filatov & White, 1995; Labarca *et al.* 1995). Gating of Cys-loop receptors can occur in the absence of bound ligand, but at very low frequency (e.g. Jackson, 1984, 1986; Hu & Peoples, 2008). The binding of ligand increases the probability of opening and maximizes as the quantity of bound ligands rises to at least two (Beato *et al.* 2004; Corradi *et al.* 2009). As the channel opening rate can be quicker than the dissociation rate of the ligand, several openings can occur during a single ligand occupancy.

Electron microscopy studies (Miyazawa *et al.* 2003; Unwin *et al.* 2002) have indicated that the $\beta 1$ – $\beta 2$ and $\beta 8$ – $\beta 9$ loops and the $\beta 10$ strand in the ECD are closely associated with residues in the M2–M3 linker, providing a direct link between the ECD and TMD (Lee *et al.* 2008; Unwin, 1995; Unwin *et al.* 2002). Comparing the structures of the ELIC (apparent closed conformation) and GLIC (apparent open conformation) prokaryotic receptors also shows distinct differences in this region (Bocquet *et al.* 2009; Hilf & Dutzler, 2008). In support of the structural data, experiments show that coupling of binding to gating in a chimaeric AChBP (ECD)/5-HT₃ (ICD) receptor could only be achieved when these three amino-acid loops from the ECD of the 5-HT₃R were substituted into the corresponding regions of AChBP. This indicates that compatibility between the two regions is necessary (Bouzat *et al.* 2004), although subsequent studies which could not repeat these functional data led these authors to conclude that AChBP is in the desensitized form (Grutter *et al.* 2005b). Specific residues in the loops at the ECD/TMD interface have been identified as playing roles in the transduction process; in particular, a salt-bridge between the ECD and TMD regions has been identified in nACh (Lee & Sine, 2005) and GABA_C (Price *et al.* 2007; Wang *et al.* 2007) receptors. However, there is no such salt bridge between the equivalent residues in GABA_AR (Kash *et al.* 2003, 2004), 5-HT₃ (Price *et al.* 2007) or GlyR (Schofield *et al.* 2003), although it is clear that charged residues are important. A good explanation for these results is that there is a global electrostatic attraction between the two regions (Dougherty, 2008; Xiu *et al.* 2005). A series of recent reviews have summarized current knowledge of the changes that induce channel opening (Bartos *et al.* 2009; Cederholm *et al.* 2009; Chang *et al.* 2009; Gay & Yakel, 2007).

7. Time-resolved structural information

It has long been a goal of biophysicists to understand the conformational changes in an ion channel at sufficient time resolution (a) to link each structure to a functionally defined state, and possibly (b) to visualize the transition states as well. This goal remains elusive for two reasons. (1) Some Cys-loop receptors desensitize within a few ms after opening, and therefore, structural measurements must have sufficient temporal resolution to distinguish between active and desensitized states. (2) We do not yet understand the kinetic bases of equilibrium side chain parameters such as polarity, polarizability, volume and (in the case of Pro), backbone *cis*–*trans* isomerization, or how such properties govern the kinetics of changes in the secondary and tertiary structures of proteins. Therefore, results from site-directed mutagenesis alone cannot usually be interpreted in kinetic terms.

7.1 Time-resolved cryo-electron microscopy

Conceptually, the clearest approach is Unwin's pioneering experiment that obtained cryo-electron microscopy data at 9 Å resolution from *Torpedo* nAChR-rich membranes, both in the closed state, and ~5 ms after ACh was sprayed onto the membrane (Unwin, 1995). Unwin identified rotations in the ECD and two alternative structures of the M2 helices; straight in the closed state and kinked in the open state. These measurements may eventually be brought to the 4 Å resolution of Unwin's later cryo-electron studies (Unwin, 2005), but are currently insufficiently resolved to identify the orientation of individual amino-acid side chains.

7.2 Time-resolved mass spectrometry

As detailed in section 4, state-dependent cross-linking and SCAM experiments reveal that residues change their distance from each other, or their water-accessibility, as the channel opens, closes and desensitizes. A complementary technique uses photoaffinity labelling during periods when the Cys-loop receptor is in the open, closed or desensitized state. Using this technique, the membrane was kept intact to allow voltage control manipulations; oocytes expressing nAChR were exposed to constant ACh during a voltage-clamp experiment. The receptors, which have voltage-dependent gating, were switched from open to closed by voltage jumps. The oocyte was exposed to the lamp during 500 ms epochs that coincided with either the open or closed states (Leite *et al.* 2003). Because of the photoprobe concentrates in the membrane, mass spectrometry then revealed regions whose exposure to the membrane changed during the state transitions. In the open state, there was specific probe incorporation within the ECD in the $\beta 8$ – $\beta 9$ loop. In the closed state, probe incorporation occurred within the Cys-loop, and these findings agree well with present concepts about the gating interface. In the closed state, there was also probe incorporation in the M3–M4 region, emphasizing that this region too may move relative to the membrane during gating (Akk & Steinbach, 2000; Bouzat *et al.* 1994).

7.3 Light-flash relaxations

Other strategies produce a perturbation at a structurally defined location, and then ask how the receptor relaxes with a new equilibrium. The kinetics of the relaxation then reveal the speed of a conformational change that propagates from the perturbed residue to the channel gate(s). Unfortunately, the location of voltage dependence is unknown in those Cys-loop receptors (such as the muscle nAChR) that display voltage-dependent kinetics. However, light-flash relaxations are more informative, because they originate at known locations. In one example, the photoisomerizable nicotinic agonist Bis-Q was photoisomerized from the active, *trans* configuration to the non-agonist *cis* configuration while the Bis-Q molecule was bound to the binding site. Channels closed completely within $<100 \mu\text{s}$ showing that dissociation of the agonist and channel closure are linked to this time scale (Nass *et al.* 1978; Sheridan & Lester, 1982). The photon cross-section for this closure corresponds to two Bis-Q molecules per channel, showing that this conclusion applies to either of the two bound agonists. Information of this sort can be obtained only indirectly from other kinetic studies.

In another example, the unnatural side chain Tyr-ONB, or caged Tyr, was introduced in place of the Tyr residues in the nAChR α subunit at loop A (Tyr93), near the Cys-loop (Tyr127) or in loop C (Tyr198). The mutant receptor did not respond to ACh. Flash decaging, in the presence of ACh, produced conductance increases that covered a wide range of time scales (1 ms to 10 s), with at least two phases in each case (Miller *et al.* 1998). The faster phase (τ_1) was governed by the time course of the flash (~ 1 ms); this is important, because it implies that changing any of these side chains to the native Tyr activates the gate within ~ 1 ms. The time constant of the slower phase (τ_2) was considerably slower: τ_2 at Tyr127 (13 ms) was faster than for Tyr198 (41 ms) or for Tyr93 (820 ms). We now understand that Tyr127 is closer to the channel gate than are the other two side chains, and we also know that Tyr198 is on a loop that moves when agonist binds in AChBP (Zheng & Zagotta, 2003). On the other hand, in most models, Tyr93 does not participate directly in conformational changes that open the channel. Thus, a structural

change in a side chain influences the conductance more rapidly when it directly participates in the gating pathway.

7.4 Rate-equilibrium free energy relationships

Rate-equilibrium free energy relationships (REFERs) provide a more general way to specify the order at which various residues participate in the conformational changes that open and close the gate (Edelstein *et al.* 1996; Grosman *et al.* 2000b). In this analysis, one performs a kinetic experiment that can distinguish the rate of channel opening from that of channel closing. This can be accomplished with macroscopic measurements, but is most simply done by measuring single-channel kinetics at agonist concentrations so high that they saturate the binding step. The kinetic and equilibrium data are compared for a set of mutations at a residue under investigation. If the data are well behaved, they yield a parameter $0 < \Phi < 1$, which indicates the proximity of the transition state controlled by the residue of interest. Values of Φ near 0 and 1 indicate that a transition state is nearest to the open or closed state, respectively. Strikingly, the initial investigation showed that progression of Φ , from near 1 to near 0, approximated the physical position of residues from extracellular (near the binding site) to intracellular (near the channel gate), respectively (Chakrapani *et al.* 2003; Grosman *et al.* 2000b). These data led to the conclusion that opening proceeds in a conformational wave, from the binding site to the channel gate. Evidently, the Φ values imply both mechanistic order and temporal order.

More recent analyses have shown clusters of Φ values for neighbouring residues, giving rise to the idea that domains of the receptor move together and allowing the ordering of conformational changes that involve various domains. Thus, a recent analysis concludes that the residues at the top of α -M2 region move at about the same time as the binding site (Bafna *et al.* 2008). Later in the conformational wave, the M2 regions have several distinct steps at Φ values between 0.64 and 0.31 (Purohit *et al.* 2007).

7.5 Voltage-clamp fluorometry

Voltage-clamp fluorometry provides another way to identify changes in the local structure. Fluorescence changes (ΔF) that differ from conductance changes are the more interesting, whether these differences occur along the axis of time, agonist concentration, blocker concentration or agonist efficacy. In the usual experiment, an introduced Cys residue is derivatized with a Cys-reactive fluorophore whose fluorescence is highly dependent on local polarity. The typical probe, tetramethylrhodamine, increases its fluorescence by factors approaching 100 when its environment becomes less polar. We do not yet fully understand the photophysical nature of the relation between conformational changes and fluorescence, but can simply say that a ΔF means a local change in the environment. Indeed, several Cys-loop experiments show that the sign (positive or negative) of ΔF varies with position for closely spaced residues (Muroi *et al.* 2006; Dibas *et al.*, HAL lab). We call this the 'sign caution'. One must also rule out a direct interaction between the tethered fluorophore and the ligand under test, as well as making sure that the ligand does not have a fluorescent signature of its own.

Following the lead of experiments on voltage-gated channels and neurotransmitter transporters, voltage-clamp fluorometry measurements usually involve a voltage-clamped oocyte, because the large measureable membrane area provides better signal to noise ratios than experiments on an individual voltage-clamped mammalian cell. The excised inside-out patch procedure, which has proven to be useful for cyclic nucleotide-gated channels (Zheng and Zagotta, 2003), is less

useful for channels gated by extracellular ligands and therefore has not been employed. There are also indications that covalently bound fluorophores can sense structural changes in the GABA_AR that presumably originate at binding interfaces, and then propagate to a non-binding subunit that has the fluorescent label (Muroi *et al.* 2006).

Concentration–response relationships are possible when the maximal $\Delta F/F$ (e.g. signal/background) exceeds $\sim 2\%$. The data for GABA_A and GABA_C receptors show that fluorophores tethered to loops A, C and E undergo agonist-induced conformational changes that change the fluorescence of bound fluorophores; these fluorescence changes have the same concentration–response relationship as the conductance (Chang & Weiss, 2002). Thus, we have another indication that part of the ECD moves as the channel opens; but the small $\Delta F/F$ prevented experiments that would compare the kinetics of the fluorescence changes and conductance changes. Interestingly, fluorophores in loops A (Chang & Weiss, 2002) and E (Chang & Weiss, 2002; Muroi *et al.* 2006) experience antagonist-induced fluorescence decrease, opposite to the agonist-induced changes. This difference eliminates concerns about the ‘sign caution’. On the other hand, a fluorophore in loop C exhibits similar changes whether the binding site is occupied by agonists or antagonists (Chang & Weiss, 2002). These findings about antagonists could not have been obtained from mutagenesis alone. Because of the ‘sign caution’, the results do not conflict with recent conclusions, mostly based on AChBP, that agonists allow loop C to collapse on the agonist, while antagonists tend to push loop C away from the other loops (Gao *et al.* 2006; Hansen *et al.* 2005). Picrotoxin, a pore blocker, does not induce ΔF by itself (Muroi *et al.* 2006), but partially blocked ΔF for a loop E position, fully blocked ΔF for a loop A position, and failed to block ΔF for a loop C position (Chang & Weiss, 2002). Pore block is a complex kinetic and equilibrium phenomenon and detailed concentration–response experiments for both agonist and blocker would be required to resolve the question of whether a fluorophore in the ECD senses a different environment when the channel is blocked by picrotoxin (Lester, 1992).

The largest $\Delta F/F$ (10–20%) has been measured for the environment-sensitive fluorophore, tetramethylrhodamine (TMR) tethered at the extracellular end (typically close to the channel lining residue 19') of the M2 region at the muscle nAChR (Dahan *et al.* 2004) or GlyR (Pless *et al.* 2007). These signals enable spectrally resolved studies on the fluorescence, verifying the expectation (from studies in solution) that the emission spectrum shifts towards the blue as the emission increases. The original report concerned the muscle nAChR β -subunit containing TMR tethered to the 19'Cys mutation; but signals almost as large have now also been found for the γ 19' and δ 19' positions as well (Dibas *et al.*, unpublished results). The concentration–response relations for agonist-induced ΔF are shifted well to the left of those for agonist-induced current, implying that conformational changes occur at concentrations too low to open the channel. Further experiments led to the conclusion that β 19' positions events that closely follow agonist binding at the $\alpha\delta$ interface. In this case, $\Delta F/F$ was sufficiently large to allow kinetic studies, down to a time resolution of ~ 5 ms. The kinetic studies showed that one or more desensitized states of the nAChR retain the fluorescence increase, consistent with the idea that most desensitized receptors have agonist bound. The question arose, does ΔF arise solely from one or more desensitized states? The answer is no, because when the receptors were activated by a flash-induced increase of the agonist *trans*-Bis-Q, ΔF was complete with 20 ms, one to two orders of magnitude faster than desensitization (Dahan *et al.* 2004).

Experiments with tetramethylrhodamine tethered to a GlyR 19' Cys residue show that large $\Delta F/F$ has the same Gly concentration–response relation as the Gly-induced conductance

(Pless *et al.* 2007). But β -alanine and taurine produced robust ΔF without appreciable activation. Propafol converted taurine and β -alanine to full agonists, yet failed to produce a Gly-like blue-shifted emission spectrum. Thus, as in the nAChR, TMR at 19' reports a conformational change associated with a binding event in the ECD that occurs in the absence of channel opening. On the other hand, ivermectin, suspected of acting in the TMD, activates the channel without inducing ΔF . Strychnine, which competes at the binding site, blocks ΔF ; picrotoxin, which blocks the channel, does not reduce ΔF .

Voltage-clamp fluorometry has justified its promise as a procedure that can reveal conformational changes induced by ligands, separable from those associated with channel opening (Pless & Lynch, 2008). Cys-loop receptors apparently have wondrously complex conformational states and flexibility. Unfortunately, voltage-clamp fluorometry has told us little about the specific nature of the additional conformations.

7.6 Total internal reflection fluorescence

Some of Axelrod's pioneering experiments with total internal reflection microscopy (TIRF) were performed on muscle nAChRs labelled with antibodies (Wang & Axelrod, 1994). Since then, many investigators have appreciated that TIRF is an optimal technique for resolving membrane-associate proteins. Recent TIRF experiments with single-molecule resolution have utilized channels that incorporate fluorescent proteins to count subunits in various ion channels (Ulbrich & Isacoff, 2007). In a further refinement of the channel- and subunit-counting theme, it is now possible to label nAChRs with fluorescent unnatural amino acids, which vastly decreases the possible structural perturbation produced by the fluorophore (Pantoja *et al.* 2009). It is also possible to count individual subunits with fluorescent ligands, either α -BTX (Pantoja *et al.* 2009) or a Cy3-derivative of carbamoylcholine (Fujimoto *et al.* 2008). Within a given individual cell, there is good agreement among the estimates for the number of channels from electrophysiology, fused fluorescent proteins, the fluorescent unnatural side chain and fluorescent α -BTX (Pantoja *et al.* 2009). It may also be possible to develop high-throughput assays for 5-HT₃Rs with TIRF detection of fluorescent ligands (Hovius *et al.* 1999).

The 'optical patch clamp' is an entirely different use of single-molecule TIRF microscopy (Demuro & Parker, 2005). Cells are injected with a dye whose fluorescence increases when it binds Ca²⁺. When an nAChR channel opens, the resulting transient cytoplasmic microdomain of increased fluorescence is sufficiently close to the membrane to be visualized by TIRF. The increased fluorescence has a square-wave time course (at a temporal resolution of ~2 ms) and exhibits all the expected kinetics, pharmacology, dose-dependence and voltage dependence expected from single nAChR channels. The technique simultaneously images and resolves the opening of hundreds of channels. It is especially encouraging that muscle nAChRs and normal Ringer solutions were used in the experiment, because the muscle nAChR has a relatively lower Ca²⁺ permeability than most nACh and 5-HT₃Rs, implying that nearly all receptors could be studied with the 'optical patch clamp'.

8. Conclusions

Members of the Cys-loop of LGICs display considerable structural and functional homology. In this review, we have seen how evidence from structural studies can often be applied across the

whole family of receptors. Functional conservation cannot be better demonstrated than by the creation of chimaeric receptors that combine varying regions of the different family members to create new receptors that possess the functional properties of both receptors (i.e. Eiselé *et al.* 1993; Galzi *et al.* 1996; Grutter *et al.* 2005a, b). Considering the level of sequence and structural similarity between members, it is not surprising that there is also cross-talk by agonists (e.g. Nakazawa *et al.* 1995; Macor *et al.* 2001) and antagonists within the group (e.g. Ballestro *et al.* 2005; Broad *et al.* 2002; Drisdell *et al.* 2008; Gurley & Lanthorn, 1998; Macor *et al.* 2001; Thompson & Lummis, 2008b). There has already been considerable research on these receptors, and the more recent identification of new members (e.g. Histamine-gated (Beg & Jorgensen, 2003; Bocquet *et al.* 2007; Davies *et al.* 2002; Zheng *et al.* 2002), EXP-1 (Beg & Jorgensen, 2003), Zinc-activated (Davies *et al.* 2002), ELIC (Hilf & Dutzler, 2008), proton-gated (Bocquet *et al.* 2007), glutamate-gated (Cully *et al.* 1994), MOD-1 (Ranganathan *et al.* 2000) and SsCl (Mounsey *et al.* 2007)) suggests that others still remain undiscovered. A combination of traditional experimental methods and some of the more recent developments described in section 7 will provide us with further insights, many of which will be widely applicable to the whole Cys-loop receptor family.

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