Crystal structure of a human GABA<sub>A</sub> receptor

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Summary

Type-A γ-aminobutyric acid receptors (GABA<sub>A</sub>Rs) are the principal mediators of rapid inhibitory synaptic transmission in the human brain. A decline in GABA<sub>A</sub>R signalling triggers hyperactive neurological disorders such as insomnia, anxiety and epilepsy. Here we present the first three-dimensional structure of a GABA<sub>A</sub>R, the human β<sub>3</sub> homopentamer, at 3 Å resolution. This structure reveals architectural elements unique to eukaryotic Cys-loop receptors, explains the mechanistic consequences of multiple human disease mutations and shows a surprising structural role for a conserved N-linked glycan. The receptor was crystallised bound to a previously unknown agonist, benzamidine, opening a new avenue for the rational design of GABA<sub>A</sub>R modulators. The channel region forms a closed gate at the base of the pore, representative of a desensitised state. These results offer new insights into the signalling mechanisms of pentameric ligand-gated ion channels and enhance current understanding of GABAergic neurotransmission.

Introduction

In response to binding the neurotransmitter GABA, released at inhibitory synapses, GABA<sub>A</sub>R chloride channels open and depress neuronal excitability in the adult central nervous system<sup>1</sup>. GABA<sub>A</sub>Rs belong to a superfamily of pentameric ligand-gated ion channels (pLGICs) known as the Cys-loop receptors, that include the cation-selective nicotinic acetylcholine receptors (nAChRs) and serotonin type-3 receptors (5HT<sub>3</sub>Rs), as well as anion-selective glycine receptors (GlyRs)<sup>2</sup>. These molecules assemble as pentamers from a variety of subunits. In human GABA<sub>A</sub>Rs these subunits are encoded by nineteen different genes: α<sub>1</sub>-6, β<sub>1</sub>-3, γ<sub>1</sub>-3, δ, e, θ, π and ρ1-3<sup>3</sup>. Most physiological heteromeric formats are thought to include two α, two β and one other, most frequently a γ subunit<sup>4</sup>. β<sub>3</sub> subunits also efficiently assemble into functional homomeric channels, and although they have yet to be identified as discrete populations in the brain, they serve as meaningful models for the heteromeric receptors<sup>5</sup>. Each subunit contributes an extracellular domain of
200-250 amino acids, an α-helical M1-M4 transmembrane bundle and an M3-M4 intracellular loop of 85-255 residues. Neurotransmitter molecules bind at extracellular pockets between subunits to induce a conformational switch that crosses to the transmembrane region to open the ion channel. GABA\textsubscript{A}Rs are the targets of a wide range of drugs including benzodiazepines, used in treatment of epilepsy, insomnia, anxiety and panic disorder, and the intravenous general anaesthetics propofol and etomidate. GABA\textsubscript{A}Rs also mediate alcohol inebriation and are targets for endogenous modulators such as neurosteroids.

In the absence of GABA\textsubscript{A}Rs structural information, insights relied on analogies with related proteins. The soluble acetylcholine binding protein (AChBP) provided the first high-resolution model for the extracellular region. Ground-breaking electron microscopic studies gradually led to an atomic model (at 4Å) of a complete heteromeric nAChR from the Torpedo marmorata electric organ, and a general framework for molecular understanding of pLGICs. Subsequently, crystal structures of two bacterial homologues, ELIC and GLIC, as well as the first structure of an anion-selective Cys-loop receptor, the Caenorhabditis elegans glutamate-gated chloride channel α homopentamer, were reported, providing insights into potential mechanisms of interaction with orthosteric ligands and allosteric modulators. Nevertheless, due to limited sequence identities, these models alone cannot adequately explain GABA\textsubscript{A}R receptor assembly, ligand binding and modulation, its gating mechanism, or the consequences of numerous human mutations linked to epilepsy and insomnia. Aiming to address these unknowns, here we report the crystal structure of a human GABA\textsubscript{A}R, the β3 homopentamer.

**Architecture of the GABA\textsubscript{A}R β3 homopentamer**

Crystallisation of human GABA\textsubscript{A}R β3 required truncation of the intracellular loop between transmembrane helices 3 and 4 (M3-M4). Residues from Gly308 to Asn421 were substituted by a linker sequence, SQPARAA, to give the construct GABA\textsubscript{A}R-β3\textsubscript{cryst} (residue numbering used here corresponds to the mature isoform 1, i.e. Gln26 in UniProt entry P28472 is Gln1 in GABA\textsubscript{A}R-β3\textsubscript{cryst}). GABA\textsubscript{A}R-β3\textsubscript{cryst} solubilised in detergent retained the ability to bind agonists (GABA and histamine), channel blockers (fipronil and picrotoxin) and the anaesthetic etomidate (Extended Data Fig. 1a-e). Furthermore, in patch-clamped HEK293T cells expressing GABA\textsubscript{A}R-β3\textsubscript{cryst}, application of the GABA\textsubscript{A}R β3 agonists histamine or propofol induced inward currents that were inhibited by the channel blockers fipronil and picrotoxin (Extended Data Fig. 1f, g).

We crystallized and determined the structure of GABA\textsubscript{A}R-β3\textsubscript{cryst} at 3Å resolution (Extended Data Table 1). Viewed perpendicular to the central five-fold pseudo-symmetry axis, the receptor approximates a cylinder 110 Å in height, with diameter ranging 60 to 80 Å (Fig. 1a), spanning the plasma membrane and protruding ~65 Å into the extracellular space. Viewed along the same pseudo-symmetry axis from the extracellular side, the pentamer has a toroidal (doughnut-like) profile, surrounded by fifteen (three per subunit) N-linked glycans (Fig. 1b). Each extracellular domain (ECD) comprises an N-terminal α-helix (α1) followed by ten β-strands folded into a curled β-sandwich, topologically similar to other family members characterised to date (Fig. 1c, Extended Data Figs. 2 and 3). A second α-helix...
(α2), between β-strands 3 and 4, is located under the α1 helix (Fig. 1c). Four additional helices (M1-M4) from each subunit come together to create the pentameric transmembrane domain (TMD), with M2 segments lining a pore that tapers as it traverses towards the intracellular side of the membrane (Fig. 1b-d).

On the extracellular side, water and solutes access the pore from a vestibule surrounded by the ECDs and from lateral tunnels situated between the ECDs that are lined by an excess of negatively charged groups, suggesting putative cation modulation sites (Fig. 1c, e and Extended Data Fig. 4a, d, g, h).

A positively-charged ring halfway down the vestibule hosts putative anion binding sites at each inter-subunit interface (Extended Data Fig. 4d), revealed by peaks (visible to an ~6σ level) in $F_o-F_c$ electron density maps calculated following refinement without anions modelled (Extended Data Fig. 5a). Attempts to confirm the nature of these peaks by soaking heavy atom anions (e.g. iodide) into the crystals and analysis of anomalous electron density maps were inconclusive. However, placement of chloride ions followed by refinement satisfactorily accounts for these electron density features (Extended Data Fig. 5a-c). This positively-charged ring overlaps with a vestibule ion selectivity filter previously proposed for Cys-loop receptors20. Furthermore, these putative chloride sites are in spatial proximity (but structurally non-equivalent) to the “anion site 1” reported in the bacterial channel GLIC21, and thus chloride ions might be important stabilizers of pLGIC assembly.

At the TMD level, two large non-overlapping pockets are located near residues previously inferred to bind the intravenous anaesthetics etomidate and propofol5,9 (Extended Data Fig. 6a-c). The putative propofol binding pocket is structurally distinct from the one identified in the bacterial channel GLIC18. The binding and transduction modes of propofol on GABA$_A$R-β3$_{cryst}$ and GLIC are therefore unrelated, as are its relative potencies. Propofol potentiates and activates GABA$_A$Rs but inhibits GLIC5,18.

**Determinants of assembly**

The nineteen different GABA$_A$R subunits obey stringent rules for assembly specificity4, the molecular determinants of which are poorly understood. Comparing inter-subunit interfaces from pLGIC structures available to date reveals striking differences in the geometry and thermodynamics of complex formation, with GABA$_A$R-β3$_{cryst}$ subunits forming the most extensive, energetically favourable interactions. These occur in particular between ECDs through patchworks of hydrogen bonds, salt bridges and van der Waals contacts (Fig. 2 a-c and Extended Data Fig. 7a-e). Within the upper portions of the ECDs, between the α1 helices of adjacent subunits, the side chains of Arg26 and Asp17 form a salt-bridge network extending to Asp24 and Lys13 (Fig. 2b), an interaction specific to β-β and α-β subunits in human GABA$_A$Rs (Extended Data Fig. 3). Between the α2 helices, the inter-subunit interface is stabilised by a network of hydrogen bonds and salt bridges surrounding Arg86, specific to the GABA$_A$R-β and GlyR subunits (Fig. 2b and Extended Data Fig. 3). The loop connecting the α1 helix with the β1 strand also bridges this interface and mutations in it (or its proximity) are associated with childhood absence epilepsy and febrile seizures22,23 (Extended Data Fig. 3). A GABA$_A$R-β3 Gly7Arg mutation that excludes γ2 subunits from
heteromeric GABA\(_A\)Rs\(^{22}\) opposes and is likely to perturb the \(\alpha_1-\beta_1\) loop conformation (Fig. 2b). A GABA\(_A\)R-\(\gamma_2\) Arg43Gln mutation (equivalent to Arg28Gln in GABA\(_A\)R-\(\beta_3\cryst\)) that excludes \(\gamma_2\) subunits\(^{23}\) resides within the loop and is predicted to disrupt the backbone organisation of the \(\alpha_1-\beta_1\) loop (Fig. 2b).

Inter-subunit contacts between the central portions of ECDs involve the \(\beta_4\), \(\beta_5\), \(\beta_5'\) and \(\beta_6\) strands and flanking loops. The \(\beta_5-\beta_5'\) loop is extended in GABA\(_A\)R-\(\beta_3\cryst\) and GluCl\(_\alpha\) compared to nAChRs and AChBP (Extended Data Fig. 2d), and protrudes into the neighbouring subunit (Fig. 2c). The \(\beta_5-\beta_5'\) loop His107, strictly conserved in GABA\(_A\) and Gly receptors, and main chain amino groups in positions 104 and 108 coordinate putative chloride anions across this boundary (within the positively-charged ring lining the vestibule discussed earlier; Fig. 2c and Extended Data Fig. 5c). Further interface contacts include a salt bridge network centred on Asp101 and Arg129, and a potential cation-\(\pi\) interaction between Lys103 and Phe105, both likely to influence assembly specificity based on residue conservation patterns (Fig. 2c and Extended Data Fig. 3). Mutations within this ECD interface are linked to epileptic encephalopathies, for example Asn85Asp\(^{24}\) which will remodel interactions with basic residues adjoining the \(\beta_5-\beta_5'\) loop (Fig. 2c).

**Neurotransmitter pocket and binding mode of a novel agonist**

The neurotransmitter-binding pocket of GABA\(_A\)Rs is located between ECDs. It comprises the \(\beta_4\) strand and adjacent residues (Asp95-Leu99), part of the \(\beta_7-\beta_8\) loop (Glu155-Tyr159) and the \(\beta_9-\beta_{10}\) loop (Phe200-Tyr205) from the “principal” (“P”) face, also known as loops A-C, respectively. The “complementary” (“C”) face in GABA\(_A\)R-\(\beta_3\cryst\) comprises a segment of the \(\beta_2\) strand (Tyr62-Gln64) and the \(\beta_6\) strand (Leu125-Arg129), also known as loops D and E respectively, which come from equivalent motifs in \(\alpha\) subunits in heteromers (Fig. 3a-c and Extended Data Fig. 3)\(^{25}\).

The \(\beta_9-\beta_{10}\) loop adopts a “closed” conformation over the site, consistent with an agonist-bound conformation\(^{17,26-28}\) (Extended Data Fig. 8). Closure is stabilised by salt-bridges between the side chains of Arg207, Glu153 and Glu155 (Fig. 3b,c), residues previously implicated in GABA binding and activation\(^{29-31}\), and for which analogous interactions are implicated in binding and activation in nAChRs\(^{12,32}\).

We observed large positive peaks in the \(F_\sigma-F_c\) electron density map in all five neurotransmitter-binding sites, that are accounted for by benzamidine molecules, an additive that helped us obtain high-resolution diffracting crystals (Extended Data Fig. 5d-f). To our knowledge, benzamidine or its derivatives have not previously been reported to act as GABA\(_A\)R ligands. We recorded GABA\(_A\)R-\(\beta_3\cryst\) currents in HEK293S-GnTI\(^{-}\) cells, and found benzamidine to behave as an agonist capable of inducing desensitisation (\(EC_{50} = 61 \pm 12 \mu M; n = 4\); Fig. 3d, e). Thermostabilisation of GABA\(_A\)R-\(\beta_3\cryst\) in detergent micelles by benzamidine revealed a similar sensitivity to histamine (benzamidine \(EC_{50} = 370 \pm 180 \mu M;\) histamine \(EC_{50} = 400 \pm 150 \mu M; n = 3\); Fig. 3d and Extended Data Fig. 1c). Classically, benzamidine is known as a highly potent serine-protease inhibitor, derivatives of which are in clinical trials for prevention of blood clotting\(^{33}\). However, given that GABA\(_A\)R-\(\beta_3\) agonists such as histamine act instead as potentiators of heteromeric GABA\(_A\)Rs,
characterization of benzamidine derivatives as positive modulators may offer new opportunities in drug development.

The benzamidine benzyl ring stacks between side chains Phe200 and Tyr62, while its amidinium group hydrogen bonds with the Glu155 side-chain and backbone carbonyls of Ser156 and Tyr157, and participates in putative cation-π interactions with the Tyr157 and Tyr205 aromatic rings (Fig. 3b,c). This binding mode is reminiscent of the “principal” face one observed in GluClα and AChBP (Extended Data Fig. 8a, b). Two epileptic encephalopathies are linked to mutations in this region, β3 subunit Asp95Asn and Glu155Gly24. Loss of Asp95, which hydrogen bonds Ser156 and Tyr157, will likely destabilise the conformation of the β7-β8 loop, while Glu155Gly will impair binding of amino-group containing ligands (Fig. 3b,c and Supplementary Discussion).

The impact of N-linked glycosylation

Two of the three N-linked glycosylation sites present in each subunit (Asn8, only visible in chain A, and Asn80) are occupied by residual GlcNAc moieties following endoglycosidase F1 treatment. The third site, Asn149 on the β7 strand (Fig. 4), is conserved in almost all GABAAR-β, nAChR and 5HT3R subunits (Extended Data Fig. 3) and was resistant to enzymatic digestion (Extended Data Fig. 5g-i). This glycan extends along the β9 and β10 strands that support the agonist binding “loop C”, providing further contacts between these units and the β7 strand, which may facilitate the ECD-TMD signal transduction. Substitution of Asn149 in GABAAR-β2 reduces sensitivity to GABA34, and in GlyRs, Zn2+ coordination between the same three β-strands potentiates agonist sensitivity35. Furthermore, mutation of Arg192 at the core of this network of glycan interactions is associated with chronic insomnia and increases the inactivation rate of GABAARs36. Finally, the equivalent glycan in nAChR α1 also contacts the outer face of the ECD, and its enzymatic cleavage decreases nAChR function37.

Structural coupling between the extracellular and transmembrane domains

For a neurotransmitter binding event to transmit to the channel gate, a signal must be transduced across the ECD-TMD interface. The inner and outer β-sheets of each GABAAR-β3cryst ECD come together at the base through a conserved salt-bridge between β1-β2 loop Glu52 and β10-strand Arg21614,38 (Extended Data Fig. 7e). Beneath this, in each subunit the ECD-TMD interface consists of two clusters of interactions: an array of polar contacts linking the outer portion of the M2-M3 loop with the β6-β7 (“Cys”) loop (Fig. 5a); and van der Waals contacts between residues in the inner portion of the M2-M3 loop near the pore, the β1-β2 loop and the β6-β7 loop (Fig. 5b). Central to both clusters is Pro144 in the β6-β7 loop, conserved in all pLGICs (Extended Data Fig. 3) and adopting a cis conformation, which orients the neighbouring Tyr143 backbone carbonyl downwards to hydrogen bond with M3 helix backbone amino groups (Fig. 5a). These contacts ensure that the GABAAR-β3cryst ECD-TMD interface is tightly structurally coupled, with a solvent inaccessible surface area of 689 Å2, 100 Å2 larger than observed in ELIC, but comparable to those observed in GluClα, GLIC, and the three nAChR structures solved in a membrane context by electron microscopy (Extended Data Fig. 9a). Nevertheless, despite similarly strong
interfaces, structural alignments of GABA\textsubscript{AR-β\textsubscript{3 cryst}} with other pLGICs reveal differences in their ECD-TMD relative orientations, where rotations within a ~20° range likely reflect the multiplicity of states (resting, activated and desensitised) that these receptors occupy\textsuperscript{39} (Extended Data Fig. 9b-f).

Mutations linked to epileptic encephalopathies highlight the importance of the ECD-TMD region\textsuperscript{24,40}. One such mutation, GABA\textsubscript{AR-β3 Tyr277Cys} will reduce hydrogen bond connectivity between the M2-M3 loop and the β6-β7 loop and disrupt a stacking interaction with the Arg141 guanidinium group (Fig. 5a). Another mutation, GABA\textsubscript{AR-β1 Phe246Ser} (β3 Phe 221) is located at the top of M1 and will disrupt hydrophobic association with the critical Tyr143/Pro144 motif at the apex of the β6-β7 loop (Fig. 5b). A Lys289Met mutation reported in GABA\textsubscript{AR-γ2} (Lys274 in GABA\textsubscript{AR-β3 cryst}) affects a residue whose side chain, very well ordered in electron density maps, reaches across a neighbouring subunit (Fig. 5b and Extended Data Fig. 7e). This residue is conserved in GlyRs, for which mutations are linked to the rare genetic startle disorder hyperekplexia, and where detailed kinetic analysis of GlyR\textsubscript{α1} Lys276Glu has revealed a significantly slower gating\textsuperscript{41}. Thus, based on the GABA\textsubscript{AR-β3 cryst} structure, it appears that this lysine facilitates the coordination of inter-subunit motions.

**Channel structure and the desensitisation mechanism**

The GABA\textsubscript{AR-β3 cryst} pore is lined by five M2 helices (Fig. 1a-d), supporting “rings” of residues between positions −5′ and 20′ (Fig. 6a). In GABA\textsubscript{AR-β3 cryst} the M2 helices taper inwards from the 13′ Thr263 down to the intracellular border where the −2′ Ala248 side chains define the narrowest point, just 3.15 Å in diameter (Fig. 6a,b). This pore is too narrow to permit the passage of chloride anions (with a Pauling radius of 1.8 Å) and therefore delineates a closed gate. However, this geometry fundamentally differs from closed structures reported to date, ELIC\textsuperscript{15} and Torpedo nAChR\textsuperscript{14}, which exhibit almost vertical M2 helices and have closed gates in the extracellular portion of the pore (9′ up to 20′) formed by bulky hydrophobic side chains (Fig. 6b). In GABA\textsubscript{AR-β3 cryst} none of the M2 hydrophobic rings (1′ Val, 3′ Leu, 5′ Ile, 8′ Val, 9′ Leu, 11′ Met, 14′ Ile and 18′ Leu) line the pore. Instead, its pore trajectory more closely resembles the open conformations of GLIC\textsuperscript{16} and GluCl\textsubscript{α17}, that are narrowest at the intracellular border (Fig. 6b and Extended Data Fig. 9g).

The pore-lining residues in GABA\textsubscript{AR-β3 cryst} (Fig. 6a) overlap with the equivalent α-subunit residues in the GABA\textsubscript{AR} identified by cysteine accessibility studies and with the open nAChR pore residues identified by protonation of introduced charged residues\textsuperscript{42,43}. However, the 9′ Leu residues in GABA\textsubscript{AR-β3 cryst} (conserved across the mammalian Cys-loop receptors) are rotated out of the pore, with side chains placed between neighbouring M2 helices. This rotation is not observed in the open-to-closed M2 flexion motion in nAChR\textsuperscript{44}. Irrespective, rotation of M2 cannot account for the closed state of GABA\textsubscript{AR-β3 cryst} because superposition of each TMD individually onto open pore GluCl\textsubscript{α} TMDs to remove its influence reveals that the GABA\textsubscript{AR-β3 cryst} pore would remain closed (Fig. 6c-f). Thus, the closed pore of GABA\textsubscript{AR-β3 cryst} is accounted for by a unique conformation of its M2 helix (Fig. 6a and Extended Data Fig. 9g).
The expanded extracellular portion of the pore in GABA\(_{\text{AR-}}\beta_3\) crystallised is stabilised by two rings of salt-bridges. The first one involves 17' His267 and 20' Glu270 from adjacent M2 helices (Extended Data Fig. 7e). His267 lines both the pore and an inter-subunit cavity, coordinating Zn\(^{2+}\) for inhibition or propofol for potentiation\(^5,45\) (Extended Data Fig. 6b,c and 7e). The second ring is an intra-subunit interaction between the side chains of 19' Arg269 and M3 Asp282, which retracts M2 against M3 (Extended Data Fig. 7e). This ring is conserved in GABA\(_{\text{AR}}\) \(\alpha\)-subunits and GlyR \(\alpha\)-subunits, and substitutions of GlyR \(\alpha\)1 19' Arg are the most frequent cause of the human genetic startle disorder, hyperekplexia, with disease mutations 19' Leu and 19' Gln decoupling agonist binding from gating\(^46\).

The contracted intracellular portion of the pore is confined by the conformation of Tyr299 side chains, which press downwards upon the back of the M2 gate, compressing it shut (Fig. 6g). Aromatic Phe or Tyr residues occupy this position across all human GABA\(_{\text{AR}}\) and GlyR subtypes and in \(C.\) elegans GluCl\(_{\alpha}\). Strikingly however, in the GluCl\(_{\alpha}\) open pore the equivalent Phe294 side chain points upwards, preventing such compression (Fig. 6g).

During the gating process, conformation switching of side chains at this key position might account for local desensitisation, and drugs that potentiate function by blocking desensitisation\(^47-49\) might do so by disabling this aromatic switch. A desensitised state of GABA\(_{\text{AR}}\)\(-\beta_3\) crystallised is in agreement with our electrophysiological recordings of benzamidine-induced desensitising currents measured in HEK cells at saturating concentrations (10 mM), which approach those used in crystallisation (33 mM; Fig. 3e). Furthermore, in heteromeric GABA\(_{\text{AR}}\)s, swapping the \(\beta\)-subunit intracellular border with the equivalent nAChR residues ablates desensitisation\(^50\).

### Conclusion

Here we present the first X-ray structure of a GABA\(_{\text{AR}}\), the human \(\beta_3\) homopentamer, co-cryssalised with a novel agonist, benzamidine. GABA\(_{\text{AR}}\)\(-\beta_3\) crystallised has a closed \(\beta_9\)-\(\beta_{10}\) loop, being in an agonist-bound state, but the pore is shut, consistent with a desensitised conformation. To our knowledge, this is the first time when a pLGIC desensitised state has been described crystallographically. These results shed new light on the conformational transitions that occur across pLGICs and provide a rational basis for understanding how human disease mutations that cause epilepsy, insomnia and hyperekplexia impact upon GABA\(_{\text{AR}}\) assembly, glycosylation, agonist binding, as well as the signal transduction and gating processes.

### Methods

#### Construct design, optimization and small-scale screening

A synthetic cDNA construct encoding the full length human GABA\(_{\text{AR}}\) \(\beta_3\) subunit, based on GenBank accession number M82919, was codon-optimised for expression in mammalian cells. Approximately one-hundred construct variants were subsequently generated by PCR, to evaluate the consequences of N-linked glycosylation sites removal, mutation of Cysteine residues, N-terminal and C-terminal truncations, truncations in the intracellular loop connecting the transmembrane helices 3 and 4 (M3-M4) and introduction of chimeric domains (such as T4 lysozyme) into predicted flexible loops. These constructs were cloned...
into the pHSec vector, making use of the secretion signal sequence provided by the plasmid. To facilitate small scale screening of protein expression, solubilisation, purification and stabilization conditions, constructs were tagged N-terminally with monoVenus and C-terminally with a nine amino acid sequence derived from bovine rhodopsin (TETSQVAPA) that is recognised by the Rho-1D4 monoclonal antibody (University of British Columbia). Small-scale expression trials were performed by transient transfection in adherent HEK293T cell cultures, as previously described. Expression levels of recombinant GABA<sub>AR</sub>-β3 variants were evaluated by western blotting using Rho-1D4 as a primary antibody and the efficiency of their cell surface trafficking was monitored by wide-field fluorescence microscopy.

Suitable constructs underwent high-throughput solubilisation screening by FSEC, using a broad panel of detergents. For this and all subsequent steps, small-scale expression was performed in HEK293S-GnTI<sup>−</sup> cells, to reduce N-linked glycosylation heterogeneity. Cells were transiently transfected using lipofectamine (Invitrogen) in an adherent format and 48-72 h later were re-suspended and solubilized in a 10 mM HEPES pH 7.2, 300 mM NaCl buffer supplemented with a 1:100 (v/v) dilution of mammalian protease inhibitor solution (Sigma-Aldrich) and 1 % detergent, for 2 h at 4 °C. Insoluble material was removed by centrifugation (10,000g, 15 min) and the supernatant incubated for 2 h at 4 °C with purified Rho-1D4 antibody coupled to CNBr-activated sepharose beads (GE Healthcare). Resin-bound samples were washed with 10 mM HEPES pH 7.2, 300 mM NaCl buffer containing detergent at 3 x CMC and receptor constructs were eluted overnight in the same buffer supplemented with 500 μM TETSQVAPA peptide (Genscript). For size-exclusion chromatography (SEC), samples were loaded onto a Superdex 200 3.2/300 column (GE Healthcare) equilibrated in 10 mM HEPES pH 7.2, 300 mM NaCl, 0.02 % n-Dodecyl-β-D-maltopyranoside (DDM, Anatrace), attached to a high-performance liquid chromatography system with automated micro-volume loader and in-line fluorescence detection (Shimadzu).

Large-scale protein expression and purification

In preliminary large-scale expression trials, six-litre batches of HEK293S-GnTI<sup>−</sup> cells were grown in suspension to densities of 2 x 10<sup>6</sup> cells ml<sup>−1</sup> in Protein Expression Media (PEM, Invitrogen) supplemented with L-glutamine, non-essential amino-acids (Gibco) and 1% foetal calf serum (Sigma). For transient transfection, cells from 1 litre cultures were collected by centrifugation (200 g for 5 mins) and resuspended in 150 ml Freestyle medium (Invitrogen) containing 3 mg PEI Max (Polysciences) and 1 mg plasmid DNA, followed by a 3-6 h shaker-incubation. Subsequently, culture media were topped up to 1 litre with PEM containing 4 mM valproic acid. Typically, 40-70 % transfection efficiencies were achieved, as assessed by control transfections with a monoVenus-expressing plasmid. 48-72 h post-transfection cell pellets were collected, snap-frozen in liquid N2 and stored at −80 °C.

FSEC screening indicated that decyl maltose neopentyl glycol (DMNG, Anatrace) is the most suitable detergent for handling GABA<sub>AR</sub>-β3 constructs, and that protein stability is enhanced by the addition of cholesterol hemisuccinate (CHS, Anatrace). Therefore, the solubilisation buffer contained 10 mM HEPES pH 7.2, 300 mM NaCl, 1% DMNG and 0.05% CHS. Proteins were affinity-purified on Rho-1D4 antibody-coupled beads as
described above, concentrated by ultrafiltration to 1-2 mg/ml using 100-kDa cut-off membranes (Millipore) and applied to a Superdex 200 10/300 gel filtration column (GE Healthcare) equilibrated in 10 mM HEPES pH 7.2, 150 mM NaCl, 0.007 % (w/v) DMNG, 0.0006 % (w/v) CHS.

The GABA<sub>A</sub>R-β<sub>3</sub>cryst construct

Residue numbers correspond to the Uniprot entry P28472. Throughout the manuscript, in agreement with the established customs within the ion channel community and to facilitate comparisons between our results and others’, we adopted a numbering scheme whereby Gln26 becomes Gln1, as the first residue of the mature protein (after cleavage of the secretion signal sequence).

The most suitable construct in terms of yield and monodispersity spanned the full length of the mature GABA<sub>A</sub>R β3 isoform 1 (from Gln26 to Asn473) but with the intracellular M3-M4 loop (Gly333–Asn446) replaced with the short amino acid sequence SQPARAA. This is the construct we refer to as GABA<sub>A</sub>R-β<sub>3</sub>cryst, because it eventually led to the crystal structure described in this manuscript. The N-terminal region is native (the N-terminal monO Venus tags was not present on the construct used for large scale expression), however we maintained the C-terminal TETSQVAPA tag for affinity-purification.

Aiming to obtain the highest possible protein yields, a HEK293S-GnTI<sup>−</sup> cell line stably expressing GABA<sub>A</sub>R-β<sub>3</sub>cryst was generated as described previously. Pure protein yields from this cell line were typically ~0.02 mg/g cells (typically 10 g cells was obtained from 1 litre cell-media), roughly twice that of transient transfection.

Crystallization and data collection

The homopentameric GABA<sub>A</sub>R-β<sub>3</sub>cryst contains 15 N-linked glycosylation sites, bringing a considerable extra volume, flexibility and potential heterogeneity. Therefore, prior to crystallization, purified GABA<sub>A</sub>R-β<sub>3</sub>cryst samples were concentrated to 3 mg/ml and incubated with 0.01 mg/ml endoglycosidase F1 for 2h at room temperature. Sitting drop vapour diffusion crystallization trials were performed in 96-well Greiner Crystal Quick X plates, using 200nl protein plus 100nl reservoir dispensed by a Cartesian Technologies robot. Plates were maintained at 6.5 °C in a Formulatrix storage and imaging system. Initial crystals grew within 1-7 days in a range of conditions, however they were poorly diffracting (no better than ~13Å). Subsequent rounds of optimization by additive screening led to well diffracting crystals (~3Å), grown in 11.5 % PEG 4000, 100 mM NaCl, 100 mM Li<sub>2</sub>SO<sub>4</sub>, 100 mM N-(2-Acetamido)iminodiacetic acid pH 6.5, 2 % (w/v) benzamidine. Crystals were cryoprotected by soaking in reservoir solution supplemented with 20 % glycerol, followed by flash-freezing in liquid nitrogen. Diffraction images of 0.2° oscillation were collected at the Diamond Light Source beamline I03, on a Pilatus 6M-F detector, and indexed, integrated and scaled using xia. Data collection statistics are shown in Supplementary Table 1.
Structure determination, refinement and analysis

The GABA\(_\text{AR-}{\beta}3\text{cryst}\) structure was solved by molecular replacement using the *C. elegans* glutamate-gated chloride channel \(\alpha\) (GluCl\(_\alpha\)66, PDB accession code 3RHW) as a search model in Phaser\(^{67}\). An initial round of automated model building, structure refinement and density modification was performed using Phenix AutoBuild\(^{68}\) followed by iterative steps of manual model building in Coot\(^{69}\) and refinement in Buster\(^{70}\). During the refinement/building process it became clear that the N-terminal region of one GABA\(_\text{AR-}{\beta}3\text{cryst}\) monomer (chain A) adopted a distinct, well-ordered, conformation because of its involvement in crystal contacts. As a result, the strict five-fold non-crystallographic symmetry (NCS) restraints strategy was replaced at later stages by a local structural similarity restraints NCS approach, to allow pruning of genuine differences among matching chains from the NCS relation\(^{71}\). The final model contains one GABA\(_\text{AR-}{\beta}3\text{cryst}\) homopentamer per asymmetric unit. The complete polypeptide chains could be built, except the C-terminal TETSQVAPA purification tag and the first nine N-terminal residues (QSVNDPGNM) in chains B, C, D and E. Furthermore, clear electron density is visible for benzamidine molecules, one of which occupies every orthosteric ligand binding site, as well as 11 out of the 15 N-linked glycosylation sites, the remaining four being located in the N-terminal disordered regions of chains B-E. Glycans attached to Asn 149 in each chain were protected from endoglycosidase F1 cleavage due to extensive interactions with the protein core, underlying their important structural role. Stereochemical properties of the model were assessed in Coot\(^{69}\) and Molprobity\(^{72}\). Protein geometry analysis revealed no Ramachandran outliers, with 96.98\% residues in favoured regions and 3.02\% residues in allowed regions. Molprobity clash score after adding hydrogens is 5.74 (100\textsuperscript{th} percentile) and the overall Molprobity score is 1.85 (100\textsuperscript{th} percentile).

Sequence and structural alignments were performed in ClustalW\(^{73}\) and SHP\(^{74}\), respectively. Protein interfaces were analysed using the PDBePISA web server at the European Bioinformatics Institute (http://www.ebi.ac.uk/pdbe/prot_int/pistart.html)\(^{75}\) and residue conservation was mapped onto the crystal structure using ProtSkin\(^{76}\). Electrostatic surface potential calculations were performed using the APBS Tools plug-in in PyMOL\(^{77}\) and pore/tunnel dimensions were analysed using the Caver 3.0 software for a probe radius of 1.4 Å\(^{78}\). Structural figures were prepared with the PyMOL Molecular Graphics System, Version 1.6, Schrödinger, LLC.

Thermostability binding experiments

The thermostability of detergent-solubilised constructs was determined by heating them over a range of temperatures for equal time periods and then measuring the reduction in the intensity of the monodisperse SEC profile for each sample\(^{79}\). With increasing temperature, an increased proportion of protein is denatured, aggregates and therefore lost from the monodisperse peak when the protein is subsequently run on SEC. Protein stability was evaluated by plotting the decay in peak UV absorbance against increasing temperature, for example to obtain a 50 % melting temperature (\(T_m\)). Purified GABA\(_\text{AR-}{\beta}3\text{cryst}\) protein at 0.05 mg/ml (20 nM) in 150 mM NaCl, 10 mM HEPES p\(H\) 7.2, 0.007 % DMNG (w/v), 0.0006 % CHS (w/v) was separated into 50 μl aliquots in PCR tubes, and heated at a range of temperatures from 30-80 °C for 1 hour. Samples were run on a high-performance liquid
chromatography system with automated micro-volume loader (Shimadzu) through a Superdex 200 3.2/300 column (GE Healthcare) maintained in 300 mM NaCl, 10 mM HEPES, 0.007 % DMNG (w/v), 0.0006 % CHS (w/v). Monodisperse peak reduction with increasing temperature was measured relative to an unheated control sample maintained at 4 °C.

Importantly, because some drugs thermostabilise detergent-solubilised proteins upon binding\textsuperscript{79}, the above thermostability assay also offers an efficient strategy to measure protein sensitivity to drugs in the detergent-solubilised environment, and therefore to probe the effects of detergent on protein structural integrity, and to evaluate the structural integrity of engineered constructs. Purified GABA\textsubscript{A}R-β\textsubscript{3}\textsubscript{cryst} was aliquoted into PCR tubes, supplemented with ligand at a range of concentrations and heated at 66 °C (the temperature at which the monodisperse peak was reduced by 70 %) for 1 hour. Afterwards samples were run on a high-performance liquid chromatography system with automated micro-volume loader (Shimadzu) through a Superdex 200 3.2/300 column (GE Healthcare) maintained in 300 mM NaCl, 10 mM HEPES, 0.007 % DMNG (w/v), 0.0006 % CHS (w/v). Drug dose-response curves were generated by plotting UV absorbance against drug concentration.

**Fluorescence displacement assay**

Fluorescence emission spectra from 490-560 nm for the fluorescent ligand 1-aminoanthracene (1-AMA) at 5 μM, in the presence of 0.3 μM GABA\textsubscript{A}R-β\textsubscript{3}\textsubscript{cryst} and increasing concentrations of the anaesthetic etomidate, were measured using a Varian Cary Eclipse Fluorescence spectrophotometer, with excitation at 400 nm (5 nm slit width).

**Electrophysiology**

Electrophysiological recordings were made from HEK293S-GnTI\textsuperscript{−} cells stably expressing GABA\textsubscript{A}R-β\textsubscript{3}\textsubscript{cryst} using a Nanion Port-a-Patch (Nanion)\textsuperscript{80} or from HEK293T cells transiently transfected with GABA\textsubscript{A}R-β\textsubscript{3}\textsubscript{cryst} using a traditional patch-clamp set-up. The internal pipette solution contained (mM): 140 KCl, 2 MgCl\textsubscript{2}, 1 CaCl\textsubscript{2}, 10 HEPES, 11 EGTA, and 2 ATP, pH 7.2 (≈ 300 mOsm). The external saline solution contained (mM): 140 NaCl, 4.7 KCl, 1.2 MgCl\textsubscript{2}, 2.5 CaCl\textsubscript{2}, 10 HEPES, and 11 D-Glucose, pH 7.4 (≈ 300 mOsm). An Axopatch 200B amplifier (Axon instruments) was used to record whole-cell currents voltage-clamped at −40 mV. Recording chips with pore resistances of 3-5 MΩ were used for Nanion recordings, or patch electrodes were pulled from thick-walled borosilicate glass (Harvard Electronics). For Nanion recordings, 20 μl of a HEK293S-GnTI\textsuperscript{−} cell suspension in external saline solution was manually pipetted over the chip pore and the Nanion suction protocol was applied to obtain a giga-seal and subsequent whole-cell configuration. Five microliters of agonist drug (histamine or benzamidine) were manually pipetted at 5-fold final concentration (50 mM) into the 20 μl saline bath. Subsequently 6.25 μl of agonist drug + channel blocker (fipronil or picrotoxin) was manually pipetted at 5-fold concentration (50 mM for agonist, 50 μM for blocker) into the 25 μl saline bath. For the traditional patch-clamp set-up, drug-solutions were applied using a rapid solution changer (model RSC-200; Biological Science Instruments). Maximal whole-cell agonist evoked currents were in the range 0.2 – 1 nA. Data was recorded in several minute acquisition
epochs into pClamp 10.0 software on a computer via a Digidata 1322A (Axon instruments) sampling at 200 μs intervals.

**Extended Data**

**Extended Data Figure 1. Ligand binding to GABA\(_{A}\)R-β3\(_{\text{cryst}}\) in detergent micelles and in HEK293T cells**

a, Overlays of size-exclusion profiles for equal amounts of purified GABA\(_{A}\)R-β3\(_{\text{cryst}}\) that were pre-heated at different temperatures for 1 hour in order to evaluate protein stability. b, Overlays of size-exclusion profiles of purified GABA\(_{A}\)R-β3\(_{\text{cryst}}\) heated at 66 °C (70 % decay temperature) for 1 hour in the presence of increasing doses of histamine. Histamine binding protects (stabilises) the protein in accordance with its affinity, giving a dose-response profile. c, Profiles of GABA\(_{A}\)R-β3\(_{\text{cryst}}\) thermostabilisation in detergent micelles by four ligands. Values in brackets represent the EC\(_{50}\) of thermostabilisation. The assay was used to evaluate two channel blockers, the insecticide fipronil (26 ± 5 nM) and the convulsant picrotoxin (900 ± 480 nM), and two neurotransmitter agonists, GABA (2.3 ± 0.2 mM) and histamine (400 ± 150 μM). NOTE: the GABA\(_{A}\)R-β3\(_{\text{cryst}}\) low sensitivity to GABA is in keeping with that observed for full-length homomeric GABA\(_{A}\)R-β3 receptors, which

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are also less sensitive than αβ and αβγ heteromeric GABA\textsubscript{R}s\textsuperscript{81}. \textbf{d, e} Displacement of bound 1-aminoanthracene (1-AMA) from GABA\textsubscript{A}R-β3\textsubscript{cryst} by the anaesthetic etomidate. The fluorescent ligand 1-AMA (5 μM) in the presence of 0.3 μM GABA\textsubscript{A}R-β3\textsubscript{cryst} experienced an increase in fluorescence (due to binding in the hydrophobic anaesthetic pocket\textsuperscript{82}) that was displaced by increasing concentrations of etomidate; 50 % maximal displacement occurs at 7.1 ± 1.1 μM, n = 5. Peak heights of single traces were measured as the average intensities of peak points between 519 and 533 nm. Equivalent doses of etomidate with the fluorescent ligand 1-AMA (5 μM) in the absence of GABA\textsubscript{A}R-β3\textsubscript{cryst} did not displace (reduce) fluorescent signal (not shown). \textbf{f-g}, HEK293T whole-cell patch-clamp recordings of the natural ligand histamine (\textbf{f}) activating an inward current through GABA\textsubscript{A}R-β3\textsubscript{cryst}, subsequently blocked by the channel-blocker picrotoxin (fipronil also blocked histamine currents, not shown) and of the anaesthetic propofol activating an inward current through GABA\textsubscript{A}R-β3\textsubscript{cryst} (\textbf{g}). All error bars are s.e.m.
Extended Data Figure 2. Structural alignment of pLGIC extracellular domains and AChBP

The top panel shows structures aligned to the GABA<sub>A</sub>-β<sub>3</sub>cryst ECD, viewed perpendicular to the five-fold pseudo-symmetry axis, from inside the extracellular vestibule. Colour coding: GABA<sub>A</sub>-β<sub>3</sub>cryst in red (PDB 4COF); GluClα in green (PDB 3RIF); ELIC in yellow (PDB 2VL0); GLIC in cyan (PDB 4HFI); AChBP in dark blue (PDB 1UX2); nAChR in violet (PDB 2BG9). Four variable loop regions of functional significance for neurotransmitter binding, signal transduction and receptor assembly have been individually rotated to optimize viewing of the disparities between these structural elements.

- **a.** Loop C (β9-β10), capping the neurotransmitter binding site.
- **b.** Loop β6-β7 (Cys-loop in eukaryotic structures), important for ECD-TMD coupling and signal transduction.
- **c.** Loop [β1-β2], important for ECD-TMD coupling and signal transduction.
- **d.** Loop β5-β5′, important for subunit assembly at the ECD level.
- **e.** Table showing parameters of structural alignment between the GABA<sub>A</sub>-β<sub>3</sub>cryst extracellular (ECD) and transmembrane (TMD) domains.

\[
\begin{array}{|c|c|c|c|c|}
\hline
\text{Reference structure} & \text{Moving structure} & \text{Equivalent Co positions} & \text{Identical residues at equivalent positions} & \text{rmsd of equivalent Co coordinates (Å)} \\
\hline
\text{(chain/residue range/PDB ID)} & \text{(chain/residue range/PDB ID)} & \text{(A)} & \text{(B)} & \text{(C)} \\
\hline
\text{GABA<sub>A</sub>-β<sub>3</sub>cryst ECD} & \text{GluCl ECD (A/1-217/4COF)} & 198 & 73 (26.9%) & 1.38 \\
 & \text{ELIC ECD (A/11-199/2VL0)} & 163 & 42 (25.8%) & 1.55 \\
 & \text{GLIC ECD (A/15-193/4HFI)} & 176 & 37 (21.1%) & 1.78 \\
 & \text{AChBP (A/1-205/1UX2)} & 185 & 39 (21.1%) & 2.14 \\
 & \text{nAChR closed ECD (A/1-210/2BG9)} & 180 & 37 (20.6%) & 2.16 \\
 & \text{nAChR open ECD (A/1-210/4AQS)} & 185 & 37 (20%) & 2.31 \\
 & \text{nAChR closed ECD (A/1-210/4AQS)} & 182 & 33 (18.1%) & 2.30 \\
\hline
\text{GABA<sub>A</sub>-β<sub>3</sub>cryst TMD} & \text{GluCl TMD (A/213-340/3RIF)} & 123 & 52 (42.3%) & 0.97 \\
 & \text{ELIC TMD (A/194-315/4HFI)} & 121 & 38 (21.4%) & 1.62 \\
 & \text{GLIC TMD (A/200-316/2VL0)} & 111 & 26 (25.2%) & 2.14 \\
 & \text{nAChR closed TMD (A/211-437/2BG9)} & 110 & 11 (10%) & 2.27 \\
 & \text{nAChR open TMD (A/211-437/4AQS)} & 111 & 11 (10.1%) & 2.41 \\
 & \text{nAChR closed TMD (A/211-437/4AQS)} & 109 & 11 (10.1%) & 2.45 \\
\hline
\end{array}
\]

* The amino acid sequences in the M2 and M3 helices of the nAChR open and closed models (PDB IDs 4AQS, 2BG9 and 4AQS) appear to be out of register by 4 residues. As a result, the number of identical residues occupying equivalent positions in nAChR and GABA<sub>A</sub>-β<sub>3</sub>cryst is lower than expected from sequence alignment (22%).

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respectively, and equivalent regions in pLGICs as well as the acetylcholine binding protein (AChBP) from *Lymnaea stagnalis*. Superpositions were performed using the SHP programme (see Methods for details).

Extended Data Figure 3. Sequence alignment of GABA\(_\text{A}\)R-\(\beta^3\)cryst with representative human Cys-loop receptor family members and the other pLGICs crystallized to date (ELIC, GLIC and GluCl\(\alpha\))

Residue conservation is indicated by black/grey highlights. Residues involved in inter-subunit salt-bridges are highlighted in red/blue, sites of N-linked glycosylation are highlighted in orange. Sequence block-highlights indicate the classically-defined neurotransmitter-binding loops (A-F, in light cyan), as well as key loops discussed in the manuscript: \(\beta1-\beta2\) in purple; \(\beta6-\beta7\) (the Cys-loop) in dark green; \(\beta8'-\beta9\) loop in light red; M2-M3 loop in yellow; M3-M4 loop in mustard. Dots above the sequence mark residues linked to human diseases (red), binding of anaesthetics (violet), interactions with agonist...
benzamidine (green) and Tyr299, whose side chain conformation appears to contribute to the control of channel desensitisation (cyan). Orange hexagons indicate N-linked glycans observed in the GABA<sub>A</sub>R-β<sub>3</sub>cryst structure. C-terminal residues on dark-blue background represent affinity purification tags. Secondary structure element colouring corresponds to Fig. 1c. The GABA<sub>A</sub>R-β<sub>3</sub>cryst residue numbering, shown above the sequence, matches the mature isoform 1 (UniProt entry P28472 Gln26 becoming Gln1). Other sequences are from the following Uniprot entries: GBRB2, P47870; GBRB1, P18505; GLRA1, P23415; GLRB, P48167; GBRA1, P14867; GABRG1, Q8N1C3; ACHA1, P02708-2; ACHB1, P11230; ACHD, Q07001; ACHG, P07510; ACHA7, P36544; 5HT3A, P46098; ELIC, P0C7BY7; GLIC, Q7NDN8. GluClα sequence is from PDB ID: 3RIF. To keep the alignment as compact as possible, the following regions of poor conservation were removed: secretion signal sequences and cytoplasmic M3-M4 loop (as annotated in Uniprot) except for bacterial channels and GluClα; residues 184-192 (GLGPDGQGH) from ACHB1; residues 188-199 (KENRTYPVEWII) from ACHD; residues 187-194 (GQTIEWIF) from ACHG.
Extended Data Figure 4. Solvent-accessible surfaces of GABA\textsubscript{A}R-\beta3\textsubscript{cryst} coloured by electrostatic potential

a, Outside view of the receptor, perpendicular to the 5-fold pseudo-symmetry axis. The exit point of an ECD side tunnel is indicated by a dotted circle (transversal sections in g-h are at this level). b, View from the extracellular side, along the 5-fold pseudo-symmetry axis. c, View from the intracellular side, along the 5-fold pseudo-symmetry axis. The positively charged region surrounding the central pore originates from the dipoles of the M2 helices. d, Longitudinal cross-section (interior cartoon coloured grey except for pore-lining helices in deep teal), showing electrostatic surface potential inside the pore and in the extracellular vestibule. Arrowheads indicate positions of the transverse cross-sections. A chloride ion bound within the positively charged vestibule belt is shown as a green sphere. The green asterisk marks the exit of an inter-subunit side-tunnel. e, f, Transverse section at level of the neurotransmitter binding site (negatively charged), observed from above and underneath. g,
h, Transverse section at level of the ECD tunnels, negatively charged. i, j, Transverse section at level of the anaesthetic (etomidate) binding site, positively charged.

Extended Data Figure 5. Crystallographic quality control for non-protein elements in the GABA\textsubscript{A}R-β3\textsubscript{cryst} structure

a-c, The anion binding site between ECD interfaces (corresponding to main text Fig. 2c). a, SigmaA-weighted 2F\textsubscript{o}−F\textsubscript{c} (blue, contoured at 1.5σ) and F\textsubscript{o}−F\textsubscript{c} (green/red contoured at +3σ/−3σ) electron density maps following autoBUSTER refinement in the absence of chloride. b, The same electron density maps and contour levels following refinement in the presence of chloride. c, Final model, showing the chloride coordination sphere. d-f, Equivalent panels to the ones described above, for the benzamidine ligand bound to the neurotransmitter pocket. g-i, Equivalent panels to the ones described above, for the N-linked glycan at site 3 (Asn149) except that the contour level of the 2F\textsubscript{o}−F\textsubscript{c} maps is 1σ. Dotted lines in f and i indicate contacts within hydrogen-bonding distance.
Extended Data Figure 6. Binding cavities for intravenous anaesthetics

a, Side view of a GABA$_A$R-β$_3$ crystal surface representation. Dotted lines indicate the planes of the transverse sections shown in b and c. b, Transverse section through the pentamer at the level of 17′ His (His267), previously found to bind photolabelled propofol$^{29}$. c, Close-up of tilted transverse section indicated in (a), revealing the putative anaesthetic binding pockets in GABA$_A$R-β$_3$ crystal in agreement with previous mutagenesis and photolabelling studies with etomidate$^{28}$ and propofol$^{29}$ analogues. P: propofol binding site. E: etomidate binding site.
Extended Data Figure 7. Assembly interfaces between GABA$_A$R-$\beta_3$$_{\text{cryst}}$ subunits

a, Top and side (from the vestibule) view of two GABA$_A$R-$\beta_3$$_{\text{cryst}}$ neighbouring subunits, highlighting the nature of inter-subunit contacts between the “principal” (P) face of one subunit and “complementary” (C) face of the next (box indicates the region enlarged in e). Residues involved in salt-bridges are coloured purple and red, those forming putative hydrogen-bonds in cyan and residues forming van der Waals contacts are in orange.

b, Analysis of the inter-subunit interfaces between the ECDs and the TMDs. Values shown correspond to the most extensive inter-subunit interface in each PDB entry. The ECD was defined from the N-terminus up to one residue C-terminal of the conserved Arg at the end of $\beta$10 strands in all pLGICs (Arg 216 in GABA$_A$R-$\beta_3$$_{\text{cryst}}$). The TMD was defined as all residues beyond this point. c, “Open book” view of the inter-subunit interfaces (subunits were rotated 126° outwards around their long axis, relative to their side orientation in a), with surfaces coloured by the nature of interactions. Dotted lines delineate the trajectory of
an inter-subunit side-tunnel. d, “Open book” view (as above) of the inter-subunit interfaces with surface shaded by degree of conservation among GABA_A and GlyR family members, revealing that key determinants of specificity are located largely in the ECDs where conservation is lower (Extended Data Fig. 7d). e, Top down view at the ECD-TMD interface level, showing key interactions within a single ECD (small oval) and between subunits (large oval). Grey dashed lines indicate putative hydrogen-bonds and salt-bridges. Boxed residues mark positions of disease mutations discussed in main text.

Extended Data Figure 8. Comparison of the β9-β10 (loop C) conformation, the agonist-binding site and the ligand orientation in GABA_A-β3_cryst and equivalent GluClα and AChBP regions. GABA_A-β3_cryst is shown in grey (with blue strands) and its agonist benzamidine in green. Selected N and O atoms are in blue and red, respectively. a, Structural alignment of GABA_A-β3_cryst and GluClα (PDB ID: 3RIF, red backbone, with its agonist glutamate in orange); b, Structural alignment of GABA_A-β3_cryst and AChBP (PDB ID: 1UV6, red backbone) with agonist carbamylcholine bound (orange backbone); c, Structural alignment of GABA_A-β3_cryst and AChBP (PDB ID: 2BYN, red backbone) in apo form; d, Structural alignment of GABA_A-β3_cryst and AChBP (PDB ID: 2C9T, red backbone) in an inhibitor toxin bound form (the toxin was excluded for clarity). Key binding residues from loop B and loop C are presented to highlight interactions with nitrogen atoms. In both a and b, the β9-
β10 strand of GABA\textsubscript{A}R-β3\textsubscript{cryst} adopts a similar conformation to the closed loop from GluCl\textsubscript{α}-glutamate or AChBP-carbamylcholine. In both c and d, where AChBPs lack agonists, loop C is in an extended, open conformation different from its equivalent in GABA\textsubscript{A}R-β3\textsubscript{cryst}. Structural alignments were performed using SHP (see Methods).

Extended Data Figure 9. Analysis of ECD-TMD interfaces and pore-lining M2 helices

a, Comparative analysis of ECD-TMD interfaces in pLGIC structures reported to date. Values correspond to chain A in each PDB entry. The ECD/TMD boundaries were set between the two residues C-terminal from the Arg that ends the β10 strands in all pLGICs (Arg 216 in GABA\textsubscript{A}R-β3\textsubscript{cryst}, see sequence alignment in Extended Data Fig. 3). b-f, GABA\textsubscript{A}R-β3\textsubscript{cryst} is shown in red (except the M2 helix, in teal). Its ECD was structurally aligned with equivalent regions of GluCl\textsubscript{α} (PDB ID: 3RIF, in b), GLIC (PDB ID: 4HFI, in c), ELIC (PDB ID: 2VLO, in d), nAChR open (PDB ID: 4AQ9, in e) and nAChR closed.
Alignments reveal relative variations in the ECD-TMD orientation. Structural alignments were performed using SHP (see Methods). Comparative analysis of M2 helix curvature, based on individual Cα positions, and pore diameter in GABA_{A}R-β_{3}cryst (desensitised conformation) versus GluCl (PDB ID: 3RIF, open conformation). Residues whose side chains line the pore are highlighted in bold. Pore diameters were calculated at the level of Cα atoms, using Caver (see Methods). Legend: †Calculated as the difference between total accessible surface areas of isolated and interfacing structures, divided by two, using PISA. ‡Indicates the solvation free energy gain upon formation of the interface. The value is calculated as difference in total solvation energies of isolated and interfacing structures, using PISA. *N residues: number of residues involved in ECD-TMD interactions. §NH/NSB/NvdW: number of putative H-bonds, salt bridges and additional van der Waals interactions that contribute to the ECD-TMD interface. ¶Rotation of TMD relative to the equivalent region in GABA_{A}R-β_{3}cryst, around the inter-domain “effective hinge axis”, following superposition of the A-chain ECDs (calculated using DynDom, http://fizz cmp.uea.ac.uk/dyndom). ±The apparent register shift in the M2 and M3 helices (and connecting M2-M3 loop, which forms a large part of the ECD-TMD interface) in currently available nAChR models may impact on the values shown.

Extended Data Table 1
Crystallographic data collection and structure refinement statistics

| Data collection | GABA_{A}R-β_{3}cryst |
|-----------------|----------------------|
| Space group     | C2                   |
| Cell dimensions |                      |
| a, b, c (Å)     | 174.1, 108.9, 207.4  |
| α, β, γ (°)     | 90.0, 107.4, 90.0    |
| Wavelength (Å)  | 0.976                |
| Resolution (Å)  | 98.96-2.97 (3.05-2.97) |
| R_{merge}       | 0.077 (0.833)        |
| R_{pim}         | 0.034 (0.362)        |
| I/σI            | 13.5 (2.0)           |
| Completeness (%)| 99.8 (99.8)          |
| Redundancy      | 6.8 (7.1)            |
| Wilson B factor (Å²) | 74.07       |

| Refinement      |                      |
| Resolution (Å)  | 40.0-2.97 (3.05-2.97) |
| No. reflections | 76328 (5599)          |
| R_{work}        | 0.207 (0.237)         |
| R_{free}        | 0.227 (0.278)         |
| No. atoms       | 13972                |
| Protein         | 13643                |
| N-linked glycans| 279                  |
This structure was determined from one crystal.
Numbers in parentheses refer to the highest resolution shell.
$R_{free}$ was calculated as per $R_{work}$ for a 5% subset of reflections that was not used in the crystallographic refinement.
Ramachandran statistics and Molprobity scores are included in the Methods section.

Supplementary Material
Refer to Web version on PubMed Central for supplementary material.

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**Figure 1. Architecture of GABA<sub>AR</sub>-β<sub>3</sub>cryst**

**a**, GABA<sub>AR</sub>-β<sub>3</sub>cryst viewed parallel to the plasma membrane (α-helices red, except the pore-lining M2 shown in teal; β-strands blue; loops grey). N-linked glycans shown in orange “ball-and-stick” representation. **b**, View from the extracellular space (synaptic cleft) down the five-fold pseudo-symmetry axis, with a single subunit coloured in grey. **c**, Two subunits, rainbow coloured from blue N-terminus to red C-terminus, illustrating secondary structure nomenclature. A water-filled ECD vestibule and TMD pore shown in light green (diameter indicated periodically) runs through the five-fold pseudosymmetry axis of the pentamer.
joined by lateral tunnels coming from between each of the subunit ECDs (two only shown for clarity, in grey). d, The pentameric transmembrane region, to illustrate the arrangement of helices M1-M4 and the M2-M3 loop (yellow). e, View of a lateral tunnel running between subunits into the central vestibule.
Figure 2. Assembly interactions in GABA<sub>AR</sub>-β<sub>3</sub>cryst

a, Top-down view of the GABA<sub>AR</sub>-β<sub>3</sub>cryst pentamer and side-on view from the vestibule of two neighbouring subunits, highlighting the nature of inter-subunit contacts between the “principal” face of one subunit and “complementary” face (residues marked by “c-”) of the next. Salt-bridging residues are coloured purple and red, those forming putative hydrogen-bonds in cyan, and residues forming van der Waals contacts in orange.

b, The upper ECD close-up shows the inter-subunit α1-β1 loop (upper dotted oval), the inter-subunit salt-bridges connecting α1 helices (middle dotted oval) and inter-subunit α2-α2 interactions (lower dotted oval). Boxed residue labels correspond to disease mutations discussed in main text.

c, The ECD anion-binding site and surrounding inter-subunit interface (chloride shown as a green sphere). Grey dashed lines indicate putative salt-bridges and hydrogen-bonds, green dashes indicate chloride coordination.
Figure 3. Neurotransmitter pocket occupied by the agonist benzamidine

a, Benzamidine (green/blue spheres) bound in the neurotransmitter pocket. Of note, the β8-β8′ loop, known as loop F (Arg169-Ala174; running into the red loop) in heteromeric GABA_ARs does not contribute to the GABA_AR β3_cryst orthosteric site. b, c, Benzamidine binding mode. “Complementary” face residues marked by “c-“. Grey dashed lines indicate putative hydrogen-bonds, salt-bridges and cation-π interactions; blue dashed lines indicate the coordination sphere of benzamidine nitrogen atoms. Boxed residue labels indicate disease mutations. d, Benzamidine dose-response curves determined by patch-clamp of GABA_AR-β3_cryst expressed in HEKS-GnTI- cells (solid line) and by thermostabilisation of GABA_AR-β3_cryst in detergent micelles (dashed line; error bars are s.e.m.). e, Electrophysiological response to 10 mM benzamidine and block by 10 μM channel blocker fipronil (an alternative blocker, picrotoxin, also blocked GABA_AR-β3_cryst currents – Extended Data Figure 1d).
Figure 4. A conserved glycosylation site interacts with β9-β10 loop residues
Close-up of the N-linked glycosylation site 3, attached to the β7-strand that supports the β6-β7 (Cys) loop (Cys-bridge shown in black spheres) and its interactions with surrounding residues from the β9-β10 agonist binding loop (indicated by dashed orange lines; grey dashed lines highlight putative salt-bridges). Boxed residue labels correspond to disease mutations discussed in main text.
Figure 5. Structural coupling at the ECD-TMD interface

a, Side-on view of the ECD-TMD interface, rotated 135° in b. a, Putative hydrogen bonds (indicated by grey dashed lines) between residues of the β6-β7 (Cys) loop, the outer portion of the M2-M2 loop and the top of M3 and M4 helices. b, Hydrophobic packing in the ECD-TMD interface on the pore side, involving residues from the β6-β7 loop, the β1-β2 loop, the inner portion of the M2-M3 loop and the N-terminus of the M1 helix from a neighbouring subunit. Boxed residue labels correspond to disease mutations discussed in main text.
**Figure 6. Structure of the ion channel in a desensitised state**

a, Two GABA<sub>A</sub>R-β<sub>3</sub>cryst M2 helices (teal), with side chains of pore-lining residues in stick representation. An equivalent GluCl<sub>α</sub> (3RIF) M2 helix, in orange, illustrates its distinct flexure. 

b, Pore diameter of GABA<sub>A</sub>R-β<sub>3</sub>cryst (teal) and related structures: open GluCl<sub>α</sub>, open nAChR (4AQ9), closed nAChR (2BG9), open GLIC (4HFI) and closed ELIC (2VL0).

c, Chain A superposition of pentameric GABA<sub>A</sub>R-β<sub>3</sub>cryst (red/teal) over GluCl<sub>α</sub> (orange), revealing the relative rotation of transmembrane regions.

d, The pore constriction at −2<sup>′</sup>A248 in GABA<sub>A</sub>R-β<sub>3</sub>cryst compared to GluCl<sub>α</sub> at −2<sup>′</sup>P243, using alignment in c. 

e, Superposition of individual GABA<sub>A</sub>-β<sub>3</sub>cryst subunit TMDs over GluCl<sub>α</sub> removes the relative rotation, but the pore remains shut.

f, g, GABA<sub>A</sub>R-β<sub>3</sub>cryst (red/teal) showing Tyr299
“pressing” M2 to constrict the channel. In GluClα (orange) Phe294 points upwards, enabling an open pore conformation.