Mechanism of NACHO-mediated assembly of pentameric ligand-gated ion channels

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Pentameric ligand-gated ion channels (pLGICs) such as γ -aminobutyric acid receptors (GABA_ARs) and nicotinic acetylcholine receptors (nAChRs) are cell surface proteins of crucial importance for animal physiology. The molecular mechanisms that control their assembly at the endoplasmic reticulum (ER) remain unknown. Here, we identified and determined the cryo-EM structure of an assembly intermediate containing two α 1 subunits of GABA_AR each bound to an ER-resident membrane protein NACHO. The structure showed how NACHO shields the principal (+) transmembrane interface of α 1 subunits containing an immature extracellular conformation. Crosslinking and structure-prediction revealed an adjacent surface on NACHO for β 2 subunit interactions to guide stepwise oligimerisation. Mutations of either subunit-interacting surface on NACHO also impaired the formation of homopentameric α 7 nAChRs, pointing to a generic framework for pLGIC assembly.

Roughly half of all integral membrane proteins are part of stable multi-subunit complexes (1). In eukaryotes, membrane protein complexes are assembled at the ER before trafficking to their intended sites of function (2). The assembly reaction poses several mechanistic obstacles for the cell. The subunits must be produced in roughly the appropriate stoichiometry to satisfy all physiologically meaningful arrangements; they must find each other in a large and crowded ER membrane environment; excess subunits and partial assemblies must be recognised and degraded; and inappropriate interactions should be minimised. How these obstacles are overcome to enable efficient membrane protein assembly is unknown.

Moreover, should subunit folding precede oligomerisation, individual subunits would expose interfaces for interaction with their partners, or for lining ion channel pores, prior to assembly. This poses major problems within a membrane environment, where partially hydrophilic surfaces that can be energetically disfavoured and act as recognition elements for quality control ubiquitin ligases (3, 4). It is thought that chaperone-like assembly factors might temporarily shield such interaction surfaces to minimise off-pathway outcomes (5-8). However, the identities and mechanism of putative intramembrane assembly factors for most protein complexes remain unknown.

The pLGICs provide an ideal system to investigate oligomeric assembly for multiple reasons. First, their structures, functions, and expression properties have been characterised extensively owing to their exceptional biological importance (9, 10). Second, their assembly can be mediated solely by their membrane domain (11-13), simplifying the problem and allowing us to focus on intramembrane factors. Third, the transmembrane domain (TMD) of each subunit is a relatively simple four-helix bundle (with the helices named M1 through M4), thereby facilitating biochemical analysis (fig. S1, A to B). Finally, some pLGIC subunits cannot form homopentamers. This allows stalling of the assembly pathway and provides a route to

biochemically identify directly-interacting candidate assembly factors whose interaction with subunits of assembling homomeric channels might be too transient to capture effectively.

NACHO is a candidate biogenesis factor for GABA_A receptors

We sought to identify interactors for the membrane domain of a membrane-inserted but notyet-assembled GABA_AR α 1 subunit (GABRA1). A version of α 1 lacking its extracellular domain (ECD) was verified to insert correctly in an in vitro translation reaction containing ER-derived microsomes from either canine pancreas or HEK293 cells (fig. S1, C to D). HEK293 cells are not known to express endogenous GABA_AR subunits but can support the production of multiple receptor subtypes (*14–16*), suggesting that any putative factors involved in their assembly are present in these cells. The newly-translated ECD-lacking α 1 (hereafter mini- α 1) was affinity purified under non-denaturing conditions and the recovered proteins were identified by mass spectrometry. As a specificity control, we also analysed interaction partners of a thermostabilised variant of β 1-adrenergic receptor (β 1AR Δ CL3)(*17*), a monomeric G protein-coupled receptor.

Mini- α 1 co-purified with several proteins previously implicated in protein biogenesis, ER quality control, membrane traffic, and membrane proteins of unclear function (Fig. 1A, table S1). Amongst interaction partners, we were intrigued by NACHO (also called TMEM35A), an ER membrane protein originally identified by its ability to enhance surface expression of a subset of nicotinic acetylcholine receptors (nAChRs) (*18, 19*). The fact that nAChRs are pLGICs (*20*), combined with our currently opaque understanding of the NACHO mechanism (*21, 22*), motivated us to investigate this interaction further.

Immunoprecipitation (IP) of newly inserted full length α 1 and mini- α 1, but not β 1AR, coprecipitated endogenous NACHO from both canine-derived and HEK293-derived microsomes (Fig. 1B; fig. S1E). Similarly, native IP of endogenous NACHO recovered newly inserted α 1 and mini- α 1, but not β 1AR (Fig. 1C). Co-IP analysis in cultured cells detected NACHO interactions with the α 1, β 2, and β 3 subunits of GABA_AR and the α 7 subunit of nAChR (fig. S2A). Direct comparison of the GABA_AR α 1 versus nAChR α 7 interactions showed that the latter recovered substantially less NACHO (fig. S2B) despite NACHO being more strongly required for functional α 7 surface expression (*19*). This can be explained because the partners of GABA_AR α 1 are unavailable for assembly whereas nAChR α 7 assembles into homopentamers with only a transient unassembled state.

Knockdown of NACHO reduced surface expression of both homopentameric α 7 nAChR and GABA_AR (with subunits α 1, β 3, and γ 2) in stably expressing HEK293S cells (Fig. 1, D to F). By contrast, NACHO knockdown had little or no impact on the expression of other classes of membrane proteins (Fig. 1F). In over-expression experiments, NACHO modestly increased surface expression of co-transfected GABA_AR α 1- β 2 (fig. S2C). By contrast, NACHO markedly stimulated α 7 nAChR surface expression (fig. S2D) as expected from earlier work (*19*).

The magnitude of reduced GABA_AR surface expression upon NACHO knockdown was similar to that seen with loss of EMC (Fig. 1F), a membrane protein insertase needed for GABRA1 biogenesis (23). A well-established EMC substrate (squalene synthase) (24) was not affected by NACHO (Fig. 1F), arguing against NACHO influencing EMC function. GABA_AR surface expression was not dependent on Asterix, a subunit of a general chaperone termed the PAT complex (5, 8, 25). Thus, NACHO interacts with the unassembled α 1 GABA_AR and α 7 nAChR (likely via its membrane domain) at the ER and modestly or markedly facilitates surface expression of functional receptor.

Although NACHO was thought to be neuron-specific (19, 20), public transcriptomic data indicate that NACHO is also expressed in tissues outside the nervous system (fig. S3), consistent with its presence in microsomes derived from both canine pancreas and HEK293 cells (Fig. 1B, fig. S1E). This discrepancy might be due to loss of NACHO in some sub-lines of HEK293 cells (fig. S1F), explaining how it was found in a gain-of-function screen for stimulators of α 7 nAChR expression (18). The widespread expression of NACHO, together

with its apparent interaction with a GABA_AR subunit, is consistent with the broad tissue distribution of the pLGIC family (fig. S3). Although earlier work had suggested a neuron- and nAChR-specific function (19–21), the functional effect on GABA_AR surface expression (Fig. 1F), together with a pre-assembly interaction with the α 1 subunit (Fig. 1, A to C), point to a broader role of NACHO that extends to GABA_A receptors and potentially other pLGICs.

NACHO interacts with the plus interface of folded $\alpha 1$

In principle, the α 1-NACHO interaction could rely on a linear segment of the yet-to-be folded α 1 chain or a folding-dependent surface. To distinguish these possibilities, we tested the NACHO interaction with α 1 mutants in which residues within its four-helix TMD were sequentially replaced by tryptophan. Helix-packing mutants that would disrupt the M1-M2 (C234W and V238W) or M1-M3 (T262W) interactions strongly diminished co-IP with NACHO, whereas a mutant that should disrupt the M4-M3 interaction (S396W) had only a modest effect (Fig. 2A). This suggests that the NACHO interaction might involve a folding-dependent surface from the M1-M2-M3 subdomain.

This hypothesis was tested using stalled $\alpha 1$ insertion intermediates in which zero, one, two or three transmembrane helices had emerged from the ribosome. Affinity purification of these ribosome-nascent chain complexes via an N-terminal tag on $\alpha 1$ recovered the ribosome and Sec61 translocation channel with each intermediate, whereas NACHO was recovered only when the first three helices had been inserted (Fig. 2B). Importantly, a matched intermediate with three transmembrane helices of Rhodopsin did not recover NACHO, instead co-purifying with the PAT complex (5), a recently characterised general intramembrane chaperone(8). Thus, NACHO can initially be recruited to nascent $\alpha 1$ co-translationally once M3 has inserted into the ER. The ability of NACHO to engage its substrate co-translationally would explain earlier findings that NACHO co-immunoprecipitates with the oligosaccharyl transferase complex (OST), the translocon-associated protein complex (TRAP), and Calnexin (22), all of which are part of the native ribosome-translocon machinery at the ER (26–28).

The NACHO- α 1 interaction could either be direct or mediated by an intermediary chaperone such as Calnexin, as proposed for the NACHO interaction with nAChR subunits (*21, 22*). To distinguish these possibilities and detect proteins in direct physical proximity, we replaced residues at various surface positions in the α 1 TMD with the UV-activated crosslinking amino acid benzoyl-phenylalanine (Bpa). Consistent with the mass spectrometry results (Fig. 1A), a diverse set of interactions were seen with different surfaces of unassembled α 1 (fig. S4). Native IPs via endogenous NACHO identified six proximal α 1 positions: 252, 260, 274, 294, 301 and 309 (Fig. 2C). These residues decorate the so-called plus interface of the α 1 subunit TMD (fig. S1B), formed by the M2 and M3 helices, that will ultimately abut the minus interface of either β , γ , δ , or ϵ subunits in pentameric GABA_AR arrangements.

To identify the region of NACHO involved in this interaction, we performed analogous experiments in which NACHO contained a photo-crosslinker at various predicted surface sites based on the high-confidence AlphaFold2-predicted structure (*29*) (fig. S5, A to B). In this experiment, tagged NACHO variants containing an amber stop codon at different positions were expressed in HEK293 cells co-expressing amber-suppression factors for site-specific incorporation of the UV-activated crosslinking amino acid AbK. These cells were then semi-permeabilised to allow insertion of radiolabelled mini- α 1 into the ER by in vitro translation. Following UV irradiation, non-denaturing IPs via NACHO verified that each AbK-containing NACHO variant was still competent for interaction with α 1, albeit with reduced efficiency in a few cases. Of these NACHO variants, four positions (55, 99, 106, and 126) physically crosslinked α 1. Three of these sites mapped to a single surface on the predicted NACHO model, formed by helices 5 (H5) and 6 (H6). The fourth crosslinking site is exposed to the ER lumen.

We conclude that in intact membranes, NACHO directly engages the plus interface of $\alpha 1$ (Fig. 2E). This interface can only form once M3 has been membrane-inserted, explaining why this is the point of initial NACHO recruitment (Fig. 2B). Because this interaction is lost when the $\alpha 1$ folding is disrupted (Fig. 2A), we posit that the M1-M2-M3 subdomain begins folding co-translationally to generate a surface that is recognised by NACHO. The $\alpha 1$ interface shielded by NACHO might otherwise be a target for quality control factors. As expected, the $\alpha 1$ -interacting surface of NACHO has several conserved patches (fig. S5C). However, intriguingly, the most conserved NACHO surface does not engage $\alpha 1$. As demonstrated later, it can interact with a GABA_AR β subunit.

Structure of a NACHO-α1 assembly intermediate

The fact that $\alpha 1$ TMD and NACHO interact directly suggests that this putative assembly intermediate might be amenable to structural analysis. Assuming that the $\alpha 1$ subunit domains fold sequentially (i.e. ECD precedes TMD), we co-expressed, affinity-purified and reconstituted in lipidic nanodiscs the full-length human $\alpha 1$ and NACHO. The resulting particles were analysed by cryogenic electron microscopy (cryo-EM; fig. S6). A map at 3.6 Å global resolution was obtained for a hetero-tetrameric complex containing two molecules each of $\alpha 1$ and NACHO (Fig. 3A; fig. S7). This map enabled de novo and template-based modelling of the membrane and extracellular domains, respectively (Figure 3, B to E, table S4).

The NACHO- α 1 interface occludes ~ 1000 Å² on each side and involves the H5 and H6 helices of NACHO and the M2 and M3 helices of α 1 (Fig. 3F). This interaction shields the partially hydrophilic surface of the α 1 TMD principal (or "plus") interface that, in fully assembled GABA_ARs, would contact the TMD of complementary (or "minus") subunits (fig. S8A). The high conservation of "plus" TMD interfaces across the GABA_A receptor family (fig. S9, A to B) suggests that NACHO could in principle engage other subunits similarly. Consistent with this idea, the NACHO surface that engages α 1 is also conserved. Intriguingly, the equivalent region of the NACHO paralog TMEM35B diverges at multiple sites (fig. S9, C to D). Although the function of TMEM35B is not known, its roughly complementary expression pattern to NACHO (fig. S3) suggests that these two paralogs might engage subunits belonging to different classes of pLGICs.

The key role of M3 and the absence of M4 in the NACHO-binding interface explains the timing of co-translational NACHO recruitment to nascent α 1. The interactions observed are also consistent with the site-specific photo-crosslinking results from both α 1 and NACHO mutants. Furthermore, the structure explains why mutations that perturb the M1-M2-M3 bundle, but not M4, impair the α 1-NACHO interaction. Thus, the structure of over-expressed proteins reflects the interactions in native ER membranes between nascent α 1 and endogenous NACHO.

Whereas the α 1 plus interface is shielded by NACHO, its minus interface is occupied by the same interface of another α 1, whose plus interface is bound to another NACHO (Fig. 3A; fig. S8B). In this configuration, M1 and M2 of one subunit are juxtaposed to M2 and M1, respectively, of the other. In addition to shielding each subunit's minus interface, portions of the channel-lining M2 helix are partially shielded and would probably be sterically inaccessible to quality control factors (*30*). Thus, the NACHO- α 1- α 1-NACHO structure explains how each of the α 1 surfaces that are ultimately buried in the final receptor are temporarily shielded, exposing a primarily hydrophobic surface to the surrounding membrane (fig. S8C). Shielding these surfaces and minimizing hydrophobic mismatch with the relatively thin ER membrane would help obscure α 1 from quality control factors that recognise exposed hydrophilicity within or exposed hydrophobicity outside the membrane.

Interestingly, the NACHO-engaged α 1 is tilted within the membrane compared to its more upright position in the GABA_AR (Fig. 3C). By favouring this tilt, NACHO may help minimise the hydrophobic mismatch of the α 1 TMD during its residence in the ER, which is thought to have a thinner membrane than downstream compartments of the secretory pathway (*31*). Because

the ER membrane retains proteins with short TMDs (32, 33), a tilted α 1 induced by its interaction with NACHO may contribute to ER retention of unassembled subunits.

The α 1 ECD adopts a pre-folded conformation in the NACHO- α 1 complex.

Comparison of the ECD in the NACHO-engaged α 1 subunit with that observed in the fullyassembled pentamer reveals that the former adopts a pre-folded, 'immature' conformation (Fig. 4, A to C; fig. S10). Specifically, the β 5-5' hairpin, which engages both adjacent subunits in the fully assembled pentamer, is shifted toward the β 5' strand (Fig. 4, B to C). Consequently, if this conformation were observed in the fully assembled pentamer, the β 5 strand would be unable to reach its partner subunit at the α 1⁻ interface, while the β 5' strand would clash with the subunit on the α 1⁺ side (fig. S10C).

Additionally, the position of the N111-linked vestibule glycan is displaced compared to its location in the pentameric receptor (Fig. 4D). The vestibule N111-linked glycans are strictly conserved in all GABA_A alpha subunit subtypes and provide an additional level of receptor stoichiometry control by ensuring that no more than two alpha subunits (identical or not) can be incorporated into a pentamer owing to steric clashes (*14*, *15*, *34–37*). Accordingly, recombinant alpha subunits where this glycosylation site is mutated can form homopentamers (9). Within the NACHO-bound context reported here, the N111 glycans also seem to act as "spacers" that help maintain the 2 subunits in a relative orientation compatible with the subsequent incorporation of non-alpha subunits (Fig. 3C and Fig. 4D). These observations suggest that the ECD adopts its mature conformation once the subunit engages both neighbouring subunits, and that this conformational transition is part of the assembly process.

NACHO engages $\alpha 1$ and $\beta 2$ subunits via adjacent functional surfaces

The α 1-NACHO interaction seen in our structure and by crosslinking likely represents an assembly intermediate given that receptor biogenesis is impaired in the absence of NACHO and this interaction can only occur prior to assembly of α 1 with either β or γ subunits. If so, how would additional GABA_AR subunits be recruited to this intermediate given the occlusion of both minus and plus interfaces? One possibility is if NACHO uses a surface other than its α 1-interacting domain to recruit a non- α subunit. Indeed, one of the exposed surfaces of NACHO in the NACHO- α 1 complex is highly conserved (fig. S5C).

To test this idea, we explored a potential direct physical interaction between NACHO and the GABA_AR β 2 subunit, chosen because it cannot form homomeric interactions. Using semi-permeabilized HEK293 cells expressing NACHO with Abk at several sites, we found that newly inserted β 2 photo-crosslinked to NACHO. The two sites on NACHO that crosslink to β 2 (21 and 26) do not crosslink to α 1, whereas the α 1-interacting sites do not crosslink to β 2 (Fig. 5A). The region of β 2 interaction is near the highly conserved surface of NACHO that is available for interaction in the α 1-NACHO structure. AlphaFold2 predictions (*38*, *39*) further support the idea of GABA_A receptor subunits interacting with two adjacent non-overlapping surfaces of NACHO (fig. S11). First, co-folding NACHO with the α 1 subunit suggests that α 1 can bind to two distinct interfaces: one closely matches the experimentally predicted here to bind β 2 (fig. S11, A and B). Second, co-folding NACHO with the β 3 subunit predicted that β 3 can also bind to the highly conserved putative β 2-binding surface on NACHO. Subunit binding at this site on NACHO would not clash with any part of the NACHO- α 1- α 1-NACHO structure (fig. S11, B to C), suggesting a role for this binding site in facilitating subunit oligomerisation.

The functional relevance of the two subunit-interacting surfaces on NACHO was tested by mutagenesis combined with surface expression of homopentameric α 7 nAChR as a readout for successful assembly (Fig. 5B). α 7 nAChR was chosen because its assembly is strictly dependent on NACHO, allowing more facile testing of mutants. Alanine mutations of key pairs of residues on either the α -binding surface or β -binding surface partially or substantially reduced nAChR surface expression, whereas alanine mutations on the non-conserved non-

binding surface had no effect. In all cases, the mutant protein was expressed at comparable levels to wild type NACHO. Thus, both the α - and β -interacting surfaces of NACHO are crucial for its ability to facilitate nAChR assembly and surface expression.

Discussion

We have identified NACHO as an intramembrane assembly factor for GABA_AR and α 7 nAChR. In the case of GABA_AR, NACHO functions by binding and shielding the plus interface of the α 1 subunit, whose minus interface can potentially be shielded by homodimerization with another α 1-NACHO complex. An adjacent surface of NACHO binds the β 2 subunit, presumably for its recruitment to α 1. In the case of α 7 nAChR, the same two surfaces of NACHO would bind and bring together two subunits of α 7 to facilitate their subsequent assembly. This work sheds light on the general problem of membrane protein complex assembly, about which very little is understood in any system. Furthermore, the α 1-NACHO structure provides one of the very few examples of how membrane protein subunits are temporarily stabilized prior to their assembly. Although additional factors will likely be involved in pentameric ion channel assembly, our findings suggest the following working framework onto which future findings can be added (Fig. 5C).

As a GABA_AR α subunit is co-translationally inserted into the ER, the individual TMDs would begin interacting with each other. Once M3 is in the membrane, the M1-M2-M3 bundle can presumably form, generating the M2-M3 interface to which NACHO binds. Importantly, this occurs at the ribosome-Sec61 complex, whose surrounding area excludes most membrane proteins due to steric hinderance by the ribosome and other translocon components (8, 26, 40). NACHO's minimal protrusion from the membrane would permit access, thereby providing it priority over quality control factors (such as ubiquitin ligases) with large cytosolic domains (30). Upon completion of translation, the NACHO- α complex would be released and can engage another NACHO- α complex to generate the structure we have observed. It is plausible that the minus interface is temporarily shielded by a yet-unidentified factor, as hinted by crosslinks from these positions (fig. S6A), prior to forming this putative heterotetramer intermediate.

Recruitment of a β subunit to the NACHO- α - α -NACHO complex would be facilitated by the NACHO- β interaction, positioning it between the two α subunits. It is attractive to posit that once recruited, the ECDs could interact with each other, taking advantage of their flexible connections to the membrane domain. The "immature" ECD conformation reported here, in which the β 5-5' hairpin extends toward the β 5' strand, may facilitate initial contact with the ECD of the incoming β subunit, followed by full engagement. Indeed, ECD interactions are thought to be favoured by membrane tethering to drive assembly of some pentameric ion channels (41–43).

Once the ECDs interact, the membrane domains would be at very high local concentrations, facilitating displacement of NACHO from α 's plus interface in favour of an interaction with β 's minus interface. Similarly, β 's plus interface would favour interaction with the minus interface of the other α , thereby generating an α - β - α -NACHO complex. The vacant slot between α and NACHO would present an ideal site for another β subunit due to the exposed β -interacting site on NACHO and minus interface on α . Importantly, the α - β ECD interaction would shift α subunit's β 5-5' hairpin into its position observed in the fully assembled receptor, enabling it to 'capture' the incoming β subunit at the α ⁻ interface. The β - α - β - α complex could now accept the final subunit (e.g., γ 2) upon dissociation of the remaining NACHO. This model should be considered speculative, but plausible and consistent with our biochemical, structural and mutational analyses.

Our findings rationalise some but contradict other previous claims regarding NACHO function. The main similarity with earlier work is the conclusion that NACHO facilitates expression of functional nAChRs (19–21). However, these earlier studies proposed that NACHO acts indirectly via other putative substrate-interacting factors such Calnexin and the

OST and TRAP complexes (22). Our findings now suggest that NACHO acts directly on ion channel subunits. Furthermore, the proposal that NACHO is nAChR- and neuron-specific (19, 20) seems to have been premature, with more recent expression studies showing that NACHO is widely expressed in many tissues, including the ER of HEK293 cells and pancreas as shown here. Our studies provide a molecular and structural foundation from which the assembly principles of the pentameric receptor family can now be dissected in mechanistic depth.

Methods

Plasmids, GeneBlocks, and antibodies

Constructs for *in vitro* translation (IVT) in rabbit reticulocyte lysate were cloned from existing plasmids in the Hegde and the Aricescu lab into a pSP64-based vector or ordered as gene blocks (from Integrated DNA Technologies) containing a 5' SP6 promoter for transcription (44, 45) and are described in table S2. The NACHO constructs for photo-crosslinking and flow cytometry experiments were sub-cloned from the cDNA construct of full-length human NACHO (Genscript, Ohu24486). Antibodies were either from commercial sources or were custom antibodies that have been described previously (46) as detailed in table S3. For structural analysis, synthetic cDNA constructs encoding the full-length human GABAAR a1 subunit, based on Uniprot ID P14867, and full-length human NACHO (TMEM35A, Uniprot ID Q53FP2) were codon-optimized for expression in mammalian cells. To facilitate protein production and purification, the native signal peptide of the GABA_AR α 1 subunit was replaced with that of chicken RPTPo (MGILPSPGMPALLSLVSLLSVLLMGCVA), followed by a Twin-Strep affinity tag, a GGS linker, an Agel restriction site and the mature GABA_AR α1 sequence (Uniprot residues 28-456).. For the NACHO construct used for structural analysis, a C-terminal GGSGGSGGS linker was added, followed by the Rho-1D4 affinity tag (TETSQVAPA). Both constructs were cloned into the lentiviral expression vector pHR (47).

Cell culture

Cells expressing GABA_A (N)–FLAG– α 1 β 3 γ 2L–(C)–(GGS)3GK–1D4 have been described previously (*48*). Briefly, HEK293S-TetR-Blasticidin cells were transfected with a 2:2:1 ratio of Flag–GABAAR α 1/pcDNA4/TO–Zeocin, hGABAAR β 3/pcDNA3.1/TO–Hygro1, and hGABAAR γ 2L(GGS)3GK-1D4/pACMV/TO–G418. Cells were selected in antibiotics (Zeocin, Hygromycin, G418, and Blasticidin) and clones were expanded. One clone was selected for high level expression and the inducible expression of functional GABA receptors was confirmed by RT-PCR, western blotting, agonist binding, whole-cell patch-clamp physiology and flow cytometry.

CRISPR-Cas9-mediated disruption of NACHO was performed using pSPCas9(BB)-2A-Puro (PX459) plasmid (Addgene) encompassing the gRNA 5'-GGCCACAATAGTTACGGTTC-3'. Transfected cells were selected for 48 h with 1 μ g/ml puromycin. Remaining cells were sorted into 96-well plates at 1 cell/well concentration to select for single-cell colonies. Single colonies were expanded and screened for successful gene disruption by sequencing and western blots using TMEM35A antibodies.

HEK293 Flp-In TRex cell lines with various stably expressed doxycycline-inducible reporters have previously been described (24, 49). These reporter cell lines were grown in DMEM was supplemented with tetracycline-free FCS (Biosera) and 15 μ g/ml blasticidin and 100 μ g/ml hygromycin.

Flow cytometry analysis

For knockdown experiments in reporter cell lines, siRNAs were transfected using the Lipofectamine RNAiMAX reagent according to manufacturer's instructions (Thermo Fisher Scientific). After 72 hours, reporter expression was induced with 0.1 µg/mL doxycycline in DMEM supplemented with 10% fetal calf serum for 6 h prior to analysis by flow cytometry. GABAAR-expressing cells were collected in ice-cold PBS, washed, and resuspended in PBS with 1:100 PE-conjugated Rat anti-DYKDDDDK (to label surface α1) for 1 hour. Cells were washed once, resuspended in PBS, and passed through 70-µm prior to analysis using Beckton Dickinson LSRII with ex488, em585/42. GFP-P2A-RFP tagged reporter cells were collected by trypsinization, washed in PBS, and passed through 70-µm prior to analysis using Beckton Dickinson LSRII with ex488, em525/50 (GFP) or ex561, em612/20 (RFP). Data was collected with FACSDiva (BD Biosciences) and subsequently analyzed with FlowJo to exclude dead cells and debris, based on forward-scatter and side-scatter profiles.

In experiments where NACHO was transfected in GABAAR-expressing ∆NACHO cells (figures 1d, 4b), 100 ng of NACHO-expressing plasmid was combined with with 900 ng of empty vector in Opti-MEM media (ThermoFisher) and transfections were performed in 6-well plates with TransIT-293 (Mirus Bio) according to manufacturer's instructions. NACHO constructs lacked a Tet operator, such that expression began immediately after transfection. 16 hours later, GABAAR expression was induced with 0.1 ug/mL doxycycline and cells were prepared for flow cytometry as described above.

Preparation of semi-permeabilised cells

Semi-permeabilised (SP) cells were prepared by modification of earlier protocols(8) as follows. All steps of SP-cell preparation were performed at 0-4°C on cells at ~70% confluency, typically from a 10 cm dish. After removing the growth media, the cells were washed once with ice-cold PBS, collected by gentle pipetting in 1 ml PBS, and counted using ScepterTM 2.0 Cell Counter (Merck Millipore) with the 60 μ M sensor (Merck Millipore, PHCC60050). The cells were recovered by centrifugation for 2 min at 5000 rpm in a microcentrifuge, washed once with ice-cold PBS, then resuspended in 1 ml of 1X "physiologic salt buffer" [PSB: 50 mM HEPES-KOH, pH 7.5, 100 mM KOAc, 2.5 mM Mg(OAc)₂] supplemented with 0.01% digitonin. Following a 10 min incubation on ice, the cells were collected by centrifugation, washed twice with 1X PSB, then resuspended in 0.5X PSB to a concentration of 4 × 10⁷ cells/ml. The SP cells were used immediately without freezing at a final concentration in translation reactions of 4 × 10⁶ cells/ml.

In vitro translation

All in vitro transcription reactions used PCR-generated templates containing the SP6 promoter (44, 45). The transcription reactions were for 1 hour at 37°C. The resulting transcript was used without further purification and was diluted 1:20 in the IVT reaction, which was carried out in rabbit reticulocyte lysate (RRL) as described earlier (44, 45). Where indicated in the figure legends, the reaction was supplemented with either canine rough microsomes (RMs) prepared and used according to the method of Walter and Blobel (50), SP cells prepared as above, or RMs prepared from HEK293-Expi cells as described previously (25). Labelling of nascent proteins was achieved by including ³⁵S-methionine (500 µCi/ml). Site-specific incorporation of the photo-crosslinkable amino acid benzoyl-phenylalanine (BPA) was achieved via amber suppression as described previously (51). In brief, amber codon(s) were suppressed by supplementing translation reactions with 0.1 mM BPA, 5 µM B. Stearothermophilus tRNA^{Tyr} with a CUA anti-codon, and 0.25 µM BPA-tRNA synthetase. All translation reactions were incubated for 30 min at 32°C, then halted by transferring the samples to ice. All further steps were performed at 0-4°C, unless stated otherwise. Prior to SDS-PAGE analysis, the tRNA on RNCs was removed by adjusting the sample to 50 µg/ml RNaseA, 10 mM EDTA, 0.05 % SDS and incubating 10-15 min at room temperature.

Affinity purification of RNCs

Biochemical analysis of proteins associated with defined RNC intermediates (Fig. 2B) was done by immunoblotting of products affinity purified via an epitope tag on the nascent chain as described(8). In short, microsomes from the IVT reactions were first recovered by centrifugation at 4°C in the TLA55 rotor (Beckman) for 20 min at 55,000 rpm. The pellet was washed three times with 1XRNC buffer [50 mM HEPES-KOH, pH 7.5, 200 mM KOAc, 5 mM $Mg(OAc)_2$] then resuspended in one-fourth the volume of the original translation reaction. The resuspended microsomes were diluted 8-fold in solubilization buffer (1XRNC buffer supplemented with 1.5% digitonin) and incubated for 10-30 min on ice. Insoluble material was sedimented for 15 min at 20,000 × g at 4°C in a microcentrifuge and the supernatant was transferred to 20 µl Streptactin sepharose (IBA Lifesciences) that had been equilibrated in 1XRNC buffer supplemented with 0.25% digitonin (wash buffer). After 2 h with gentle end-over-end rotation at 4°C, the beads were washed three times with wash buffer, then

transferred to a new tube. Elution was with 50 mM biotin in wash buffer on ice for 1 h. The eluates were analysed by immunoblotting with the antibodies indicated in the figures.

Photo-crosslinking via probes in NACHO

Site-specific NACHO interactions (Fig. 2D and Fig. 4A) were analysed in SP cells derived from NACHO KO cells reconstituted with exogenous NACHO variants containing BPA installed at defined sites by amber suppression. For reconstitution, the plasmid encoding NACHO was co-transfected with plasmids encoding amber suppression components (amber suppressor tRNA and the appropriate synthetase for charging with AbK) as described before (*52*). The cells were grown in the presence of AbK for 48 h prior to harvesting and preparation of SP cells as described above. The reconstituted resuspended SP cells were used for in vitro translation of the desired ³⁵S-labelled substrate after which the SP cells were isolated by centrifugation and transferred to 384-well plates for UV irradiation as described above. The samples were subjected to native IPs using anti-NACHO antibodies and analyzed by by SDS-PAGE and autoradiography.

Photo-crosslinking via probes in the substrate

In experiments shown in Fig. 2C and fig. S4, photo-crosslinking utilized probes in the substrate. The ³⁵S-methionine labelled substrate containing BPA was generated in the presence of RMs as described above. RMs were isolated by centrifugation, resuspended in PSB, and UV-irradiated. The samples were either analyzed directly, subjected to native IPs using anti-NACHO antibodies or denaturing IPs using anti-NACHO or anti-FLAG antibodies (against the substrate) as indicated in the figure legends.

Protease protection assays

Proteinase K (PK) protection assays to assess the topology of different integral membrane proteins was done directly following the translation reaction as described before (*45*, *49*). In brief, translation reactions performed in the absence or presence of RMs were put on ice, then divided into aliquots and adjusted to 0.5 mg/ml PK without or with 1% Triton X-100 as indicated in the figure. After 1 h on ice, 5 mM of freshly-prepared PMSF in DMSO was added from a 250 mM stock and incubated for 2-5 min on ice to stop the reaction. The entire reaction volume was transferred to 10 volumes of boiling 1% SDS, 100 mM Tris-HCl, pH 8.0. The samples were then analysed by SDS-PAGE and autoradiography either directly or after denaturing immunoprecipitation as described below.

Immunoprecipitations

Denaturing IPs were performed on samples denatured in SDS-PAGE sample buffer by heating for 10 minutes at 95°C. After cooling, the samples were diluted 10-fold in denaturing IP buffer [50 mM HEPES pH 7.5, 100 mM NaCl, 2.5 mM Mg(OAc)₂, 1% Triton X-100] and incubated for 2-3 hours at 4°C with either 5 µl of anti-FLAG-M2 affinity resin (Sigma-Aldrich), Streptactin sepharose (IBA Lifesciences), or CaptivA Protein A sepharose (Repligen) plus the desired antibody. The resin was washed three times with 0.5 ml each of denaturing IP buffer and eluted in SDS-PAGE sample buffer by heating to 95°C. Native IPs were done by first solubilizing the samples on ice in 50 mM HEPES pH 7.5, 200 mM NaCl, 2.5 mM Mg(OAc)₂, 1% Digitonin, removing insoluble material by centrifugation at 4°C for 10 min at maximum speed in a microcentrifuge, then diluting samples 10-fold in native native IP buffer [50 mM HEPES pH 7.5, 200 mM NaCl, 2.5 mM Mg(OAc)2, 0.1% Digitonin]. The samples were then incubated for 2-3 hours at 4°C with either 5 µl of anti-FLAG-M2 affinity resin (Sigma-Aldrich), Streptactin sepharose (IBA Lifesciences), or CaptivA Protein A sepharose (Repligen) plus the desired antibody. The resin was washed three times with 0.5 ml each of native IP buffer, the beads were transferred to a fresh tube, all residual wash buffer removed, and eluted in SDS-PAGE sample buffer by heating to 95°C.

Mass spectrometry

Translation reactions containing transcripts coding for the desired protein (or no transcript as a control) were subjected to affinity purification of via the FLAG tag as described above, but without the elution step. Proteins samples bound to anti-FLAG beads were reduced with 5 mM DTT and alkylated with 10 mM iodoacetamide in the dark, at room temperature. Proteins were digested on-bead with 0.15 ug trypsin (Promega) over night at 25°C. The samples were centrifuged at 10,000 x g for 5 min and supernatants were transferred to a clean tube. Beads were washed once with 30% acetonitrile (MeCN) and 0.5% formic acid (FA) and the wash solution was combined with the supernatant. The peptide mixtures were desalted using homemade C18 (3M Empore) stage tips contained 1 μ l of Poros Oligo R3 (Thermo Fisher Scientific) resin. Bound peptides were eluted from the stage tip with 30-80% MeCN and partially dried down in a Speed Vac (Savant).

Peptide mixtures were analysed by LC-MS/MS using a fully automated Ultimate 3000 RSLC nano System (Thermo Fisher Scientific) coupled online to a Q Exactive Plus hybrid quadrupole-Orbitrap mass spectrometer (Thermo Fisher Scientific). Peptides were trapped by a 100 μ m x 2 cm PepMap100 C18 nano trap column (Thermo Fisher Scientific) and separated on a 75 μ m × 25 cm, nanoEase C18 T3 column (Waters) using a binary gradient consisting of buffer A (2% MeCN, 0.1% FA) and buffer B (80% MeCN, 0.1% FA) at a flow rate of 300 nl/min. Eluted peptides were introduced directly via a nanoFlex ion source into the mass spectrometer. MS1 spectra were acquired at a resolution of 70K, mass range of 380–1600 m/z, automatic gain control target of 1 x 10⁶, maximum injection time of 100 ms and dynamic exclusion of 40 s. MS2 analysis was carried out at a resolution of 17.5K, automatic gain control target of 5 x 10⁴, maximum injection time of 108 ms, normalized collision energy of 27 % and isolation window of 1.5 m/z.

MS raw files were searched against the *Homo sapiens* reviewed UniProt Fasta database (downloaded Dec. 2020) using MaxQuant (53) with the integrated Andromeda search engine (v.1.6.6.0). The database search included tryptic peptides with maximum of two missed cleavage, cystine carbamidomethylation as a fixed modification, and methionine oxidation and acetylation of the protein N-terminal as variable modifications. The MaxQuant output file, proteinGroups.txt, was then processed with Perseus (v. 1.6.6.0) software. The complete data plotted in Fig. 1A is provided in table S1.

Protein production and purification

For large-scale protein production, 1 L of suspension Expi293 cells (ThermoFisher #A14527) was grown to a density of 2×10^6 cells ml⁻¹ in Expi293 expression media (Gibco #A1435101) at 37 °C, 160 r.p.m., and 8% CO₂. Next, 1.1 mg of a 1:1 (w/w) mixture of the GABA_AR α 1 subunit and NACHO DNA and 3 mg of Polyethylenimine "Max" (Polysciences #24765) were dissolved separately in 30 mL Expi293 expression media, then mixed and incubated for 15 min at room temperature. The DNA-PEI mixture was then added to the suspension cells. After 24h, cells were harvested by centrifugation at 3000 *g* for 15 min at 4 °C and snap-frozen in liquid N₂.

Protein purification and nanodisc reconstitution was done as previously described (14, 15, 54). Frozen cell pellets were resuspended on ice in buffer A (50 mM HEPES pH 7.5, 300 mM NaCl) supplemented with 1% (v/v) mammalian protease inhibitor cocktail (Sigma-Aldrich). Cells were lysed by 1% (w/v) Lauryl Maltose Neopentyl Glycol (LMNG, Anatrace) for 1 h at 4 °C then centrifuged for 20 min at 12,000g (4 °C) (15). The supernatant was incubated with 1D4 affinity resin rotating slowly for 1 h at 4 °C (16). The resin was recovered by centrifugation (500*g*, 5 min) then washed with buffer B [buffer A supplemented with 0.1% (w/v) LMNG and 0.01% BBE (w/v)].

While attached to 1D4 resin, receptors were incubated with phosphatidylcholine (POPC, Avanti) and bovine brain lipid (BBL) extract (type I, Folch fraction I, Sigma-Aldrich) mixture (POPC:BBL = 85:15) for 30 min at 4 °C. Excess lipids were removed by pipetting after allowing the beads to settle, then samples were mixed with 100 uL (5 mg/ml) of MSP 2N2 and incubated for 30 min at 4 °C (*54*). The detergent was removed by incubating the resin with 20

mg Biobeads for 90 min at 4 °C, followed by washing with 20-30 bed volumes of buffer A. Receptor samples were eluted with buffer A supplemented with 2 mM 1D4 peptide (TETSQVAPA).

Cryo-EM sample preparation and data collection

A 3.5 µl volume of sample was applied to glow-discharged (PELCO easiGlow, 30 mA for 30 s) gold R1.2/1.3 300 mesh UltraAuFoil grids (*55*) (Quantifoil). The excess liquid was blotted for 4 s prior to plunge-freezing into liquid ethane using a Leica EM GP2 plunger (Leica Microsystems; 95% humidity, 14 °C). Grids were stored in liquid nitrogen prior to data collection. Cryo-EM data were collected on Titan Krios G3 microscopes at the University of Cambridge Department of Biochemistry EM facility (BiocEM) in electron counting mode at 300 kV. The microscope was equipped with a Gatan K3 camera and Gatan BioQuantum energy filter. Before data acquisition, two-fold astigmatism was corrected and beam tilt was adjusted to the coma-free axis using the autoCTF program (Thermo Fisher Scientific). The data were acquired automatically using EPU software (Thermo Fisher Scientific) in super-resolution mode and on-the-fly binning by 2. Detailed data acquisition parameters for all datasets are given in table S4.

Cryo-EM image processing

Image processing pipeline is shown in fig. S6. Gain-uncorrected K3 movies in tiff format were motion- and gain-corrected using RELION's implementation of the MotionCor2 algorithm (56), with frames grouped to yield a total fluence corresponding to ~1 e⁻/Å² per frame. Contrast transfer function (CTF) estimation was performed with CTFFIND-4.1.13 (57) using the sums of power spectra from combined fractions corresponding to an accumulated fluence of 4 e⁻/Å². Micrographs whose estimated resolution from CTFFIND was worse than 10 Å were removed. Particles were picked using a re-trained BoxNet2D neural network in Warp (58) then re-extracted in RELION with a pixel size 1.46575 Å and 256 pix² box size. Extracted particles were imported into cryoSPARC (v3.3.2) (59), subjected to 2D classification, then good classes (as shown in fig. S6C) selected to generate ab-initio models without applying symmetry. Heterogeneous refinement was then performed with all particles using the outputs from ab-initio job as references.

Next, multiple iterations of heterogeneous refinement, followed by homogeneous and nonuniform refinement of best classes were used to prune the set of good particles. All refinements at this stage were performed without imposing symmetry. The final set of particles was subjected to non-uniform refinement in cryoSPARC with C2 symmetry, then converted into STAR format using csparc2star from UCSF PyEM suite (*60*) and imported into RELION (v4.0.0) (*61*, *62*). First, a 3D auto-refinement was performed with C2 symmetry, local searches only (1.8°), local signal-to-noise filtering using SIDESPLITTER (*63*) (implemented in RELION), while limiting the maximum number of poses and translations to consider to 1000 and a minimum angular sampling set to 1°. The reference used in refinement was the output of the last non-uniform refinement in cryoSPARC.

Next, three steps of CTF refinement were performed: first refining magnification anisotropy; then refining optical aberrations (up to the 4th order); and finally refining per-particle defocus and per-micrograph astigmatism (64). A round of 3D auto-refinement was performed with the same parameters as above, with the most recent map as the reference. The particles were then subjected to non-uniform refinement in cryoSPARC with no symmetry imposed to check for deviations from the C2 symmetry, which did not result in further improvement of the map. A final round of non-uniform refinement was performed with C2 symmetry imposed. Local resolution plots were generated with RELION (version 4.0.0). Orientation distributions were analysed by cryoEF (65). All renderings of maps and models were done in ChimeraX (66).

Atomic model building and refinement

The initial model for the GABA_AR α 1 subunit was derived from PDB ID 7QNE (14). Starting model for NACHO was downloaded from the AlphaFold2 database (29). Iterative rounds of model building and refinement were performed in Coot v0.9.4 (67), Servalcat (68), REFMAC

v5.8.0258 (69) and Phenix v1.19.2 and dev-5430-0000 (70). The extracellular domains of the α 1 subunits were refined with additional Geman-McClure restraints in Coot, with the distance alpha value set to 0.001. Models were validated using MOLPROBITY v4.2 (71). Model building and refinement parameters and statistics are provided in table S4.

Computational analysis of the α 1-Nacho interface

Residue contacts for the α 1-Nacho complex and the fully-assembled α 1 β 3 γ 2 hetero-pentamer (PDB ID 7QNE) were calculated with the Protein Contact Atlas (72). A contact between a pair of residues is considered to exist if the distance between any two atoms from the residue pair is smaller than the sum of their van der Waals radii plus a cut-off distance of 1 Å (ref. (73)). Contact fingerprints (fig. S8) were generated by summing the per-residue number of contacts a given α 1+ residue makes with Nacho, β 3-, or γ 2- (ref. (73)). The contact fingerprint similarity score is a dot product between pairs of contact fingerprints for α 1+/Nacho, α 1+/ β 3- and α 1+/ γ 2- interfaces. Sequence alignments were generated with Clustal Omega and conservation scores calculated with bio3d (v2.4.3) (74) using the "blosum62" substitution matrix. Buried surface area (Fig. 3F) was calculated using the PDBePISA (75). For visualisation purposes, any residue with a ratio of buried surface area to accessible surface area greater than 0.3 was considered as buried. Custom scripts in R (v4.1.2) were employed for all analyses. Renderings were generated in PyMOL (v2.5.5) and graphs using the pheatmap (v1.0.12) or ggplot2 (v3.4.2) packages in R.

Gene expression analysis

Median gene-level TPM (transcripts per kilobase million) by tissue data were obtained from the GTEx Portal on 22 November 2021 at 18:00 GMT (download link: https://storage.googleapis.com/gtex_analysis_v8/rna_seq_data/GTEx_Analysis_2017-06-05_v8_RNASeQCv1.1.9_gene_median_tpm.gct.gz) (76, 77). Visualization was done using custom R scripts (R version 4.1.2) and the pheatmap package (v1.0.12).

Data availability: Data are available in the main article, supplementary materials, or public repositories. Atomic coordinates for the NACHO-α1 complex will be deposited in the Protein Data Bank; the cryo-EM density map in the Electron Microscopy Data Bank; and the raw movies in the Electron Microscopy Public Image Archive. Meanwhile, the atomic model, maps, and the validation report can be downloaded from:

ftp://ftp.mrc-lmb.cam.ac.uk/pub/knayde/nacho/

Author Contributions: YH identified candidate assembly factors and performed most of the biochemical and functional analyses on GABA_ARs; AS determined the cryo-EM structure of the NACHO- α 1 complex, built the initial model, and performed gene-expression and conservation analyses. RMJ verified key biochemical results, contributed to NACHO structure-function studies, and extended the findings to nAChRs; LS analysed co-translational interactions between α 1 and NACHO; SY-PC performed mass spectrometry; KN, TM, SWH and DYC contributed to cryo-EM data collection and processing, model building together with AS and ARA, refinement, and analysis of the NACHO- α 1 structure; RSH conceived the project and developed an overall plan with ARA, both of whom provided project management and supervision. YH and RSH wrote the initial draft, with key contributions from AS and ARA. All authors contributed to manuscript editing.

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Main Figures:



Fig. 1. NACHO facilitates GABA_A receptor biogenesis.

(A) Interaction partners of in vitro translated membrane domain of the α 1 subunit (termed mini- α 1) of the GABA_A receptor (GABRA1) versus a well-folded thermostable β 1AR variant (β 1AR Δ CL3) identified by affinity-capture mass spectrometry (Supplementary Table 1). Mini- α 1specific membrane protein interactors of uncertain function are indicated in cyan. (B) The indicated proteins were synthesized in reticulocyte lysate supplemented with microsomes derived from Expi293 cells, subjected to anti-FLAG immunoprecipitation, and analysed by immunoblotting for either NACHO (top) or the FLAG epitope (bottom). TST and 3F denote the twin-strep tag and 3xFLAG tag, respectively. (C) The indicated proteins were synthesized in reticulocyte lysate supplemented with ³⁵S-methionine and microsomes derived from Expi293 cells, subjected to either anti-FLAG or anti-NACHO non-denaturing immunoprecipitation, and visualised by autoradiography. (D) Topology diagram of the tagged reporters for pentameric ligand-gated ion channel subunits, the single-pass membrane proteins ASGR1 and SQS, and the multipass membrane protein TRAM2. '2A' denotes the viral-derived 2A ribosome skipping sequence. (E) HEK293 cells stably expressing doxycycline-inducible FLAG-tagged α 7 subunit of nAChR (CHRNA7) were treated with the indicated siRNAs for 72 hours, induced for 6 hours

with doxycycline (or left uninduced), then surface labelled with PE-labelled anti-FLAG antibody and analysed by flow cytometry. **(F)** Experiment similar to panel E but with cells stably expressing the indicated reporter proteins under a doxycycline-inducible promoter. Four different siRNAs were tested as indicated. The GABA_A receptor comprises the $\alpha 1\beta 3\gamma 2$ subunits and was detected by surface staining using PE-labelled anti-FLAG antibody (which detects the $\alpha 1$ subunit). The other reporters were analyzed directly for fluorescence signal.



Fig. 2. Nascent α 1 engages NACHO in the membrane.

(A) The indicated mini- α 1 mutant was translated in reticulocyte lysate supplemented with ³⁵Smethionine and microsomes derived from Expi293 cells. The products were divided in two, subjected to anti-FLAG (top) and anti-NACHO (bottom) immunoprecipitation, and visualised by autoradiography. (B) The α 1 subunit of GABA_A receptor containing a twin-strep tag (TST) at the N-terminus (downstream of the signal peptide) and truncated at the indicated distances (in amino acids) downstream of the first TMD was translated in reticulocyte lysate supplemented with microsomes derived from Expi293 cells. The resulting ribosome-nascent chain complexes (RNCs) were affinity purified via the TST and analysed by immunoblotting for the indicated proteins relative to serial dilutions of Expi293 microsomes. A mock translation reaction (neg.) and one containing Rhodopsin (Rho) truncated 40 amino acids downstream of its third TMD were analysed in parallel. (C) Mini- α 1 variants lacking (WT) or containing an amber codon at the indicated positions was translated in reticulocyte lysate supplemented with ³⁵S-methionine, microsomes derived from Expi293 cells, and amber suppression reagents for incorporation of the photo-crosslinking amino acid Bpa. The reactions were

irradiated with UV light, subjected to native or denaturing anti-NACHO IP, and the products visualised by autoradiography. The position of mini- α 1 and its crosslink to NACHO (x NACHO) are indicated. **(D)** Mini- α 1 was translated in reticulocyte lysate supplemented with ³⁵S-methionine and semi-permeabilized cells containing NACHO variants with the photocrosslinking amino acid AbK at the indicated positions. After UV irradiation, the samples were subjected to anti-NACHO native IP and visualized by autoradiography. **(E)** The residues of mini- α 1 and NACHO observed to crosslink with NACHO and mini- α 1, respectively, are shown on structural models of each protein. The right diagram shows the proposed mode of interaction between the membrane domain of α 1 and NACHO.



Fig. 3. Structure of the α 1-NACHO complex.

(A-D) Overview of the cryo-EM density (A) and model (B-D) for the α 1-NACHO structure. NACHO contains six helices (H1 to H6, of which H2 and H3 are in the ER lumen, with the remaining helices acting as TMDs). The four TMDs of α 1 are termed M1 through M4. (E) Diagram showing the organisation of TMDs in the α 1-NACHO structure and in a fully assembled GABA_A receptor. Key surfaces of α 1 are indicated as lines. (F) The interaction surfaces of α 1 and NACHO in the α 1-NACHO structure. (G) The NACHO-engaged α 1 is tilted by 6° relative to its orientation in the pentameric receptor.



Fig. 4. Maturation of the GABA_AR α 1 subunit β 5-5' hairpin during assembly.

(A) Superposition of $\alpha 1$ as observed in the $\alpha 1$ -NACHO structure and in a fully assembled GABA_A receptor, aligned on the ECDs. The box highlights the region zoomed in on panels (I-J). (B-C), Conformation of the $\beta 5$ -5' hairpin in the NACHO-engaged $\alpha 1$ (red) compared to its conformation in the $\alpha 1$ subunit of the pentameric receptor (grey). (D) Structural rearrangements in the $\alpha 1$ ECDs upon maturation from the $\alpha 1$ -NACHO assembly intermediate state to fully assembled channel state, as viewed down the symmetry axis of the $\alpha 1$ -NACHO complex.



Fig. 5. Dual mode of NACHO-substrate engagement.

(A) Mini- α 1 (top) or mini- β 2 (bottom) was translated in reticulocyte lysate supplemented with ³⁵S-methionine and semi-permeabilized cells containing NACHO variants with the photocrosslinking amino acid AbK at the indicated positions. After UV irradiation, the samples were subjected to anti-NACHO native IP and visualized by autoradiography. The position of mini- α 1, - β 2, and their crosslinks to NACHO (x NACHO) are indicated. The diagram below the autoradiograph shows the putative interaction surfaces on NACHO for the membrane domains of α 1 and β 2. Below the diagram is shown a structural model of the α 1-NACHO complex indicating the mutation sites analyzed in panel B. (B) Flow cytometry assay for α 7 nAChR surface expression as in fig. S2D with either wild type NACHO or the indicated mutants. The dashed line indicates the mode of α 7 nAChR expression seen with wild type NACHO. (C) Model for the role of NACHO in assembly of a pentameric GABA_A receptor.

Supplementary Figures:



Figure S1. Characterisation of α 1 and NACHO in vitro.

(A) Diagram of proteins used in Fig. 1a-c. Glycosylation sites, tags, and sites of proteinase K (PK) accessibility are indicated. (B) Schematic of the membrane domain of $\alpha 1$ in isolation (left) and within the context of an assembled GABA_A receptor (right). (C) Assay of membrane insertion and topology for $\alpha 1$, mini- $\alpha 1$, and $\beta 1AR\Delta CL3$. Each protein was translated in reticulocyte lysate containing 35S-methionine with ER-derived rough microsomes (RM) from HEK293 cells where indicated. The translation products were treated with proteinase K (PK) without or with detergent (superscript d) as indicated, and the samples analysed directly (top - total IVT) or after immunoprecipitation via the N-terminus (middle) or C-terminus (bottom). Red asterisks indicate ubiquitination, downward blue arrowheads indicate glycosylated products, and upward red arrowheads indicate protease-protected fragments. (D) Assay as in panel c but using microsomes derived from canine pancreas. Note that in this system, a small proportion of $\alpha 1$ is glycosylated at a cryptic site in its lumenal domain, resulting in two glycosylated products. (E) The indicated proteins were synthesized in reticulocyte lysate supplemented with RM derived from canine pancreas, subjected to anti-FLAG immunoprecipitation, and analysed by immunoblotting for NACHO. TST and 3F denote the twin-strep tag and 3xFLAG tag, respectively. (F) Immunoblotting for NACHO in microsomes, semi-permeabilized cells (SP cells) or total cell lysate derived from adherent HEK293 cells and a suspension-adapted sub-line termed Expi293 (two independent samples are analysed). Two exposures are shown. NACHO is expressed at much lower levels in HEK293 cells than in Expi293 cells. Immunoblotting for CCDC47, a resident ER protein of the multipass translocon, serves as a loading control.



Figure S2. Interaction and flow cytometry analysis of GABA_AR and nAChR.

(A, B) The indicated HA-tagged receptor subunit was co-expressed by transient transfection with untagged NACHO in HEK293 cells and subjected to non-denaturing anti-HA immunoprecipitation (IP). The input and IP samples were analyzed by immunoblotting for either the HA tag or NACHO. (C, D) HEK293 cells were transfected with the indicated subunit(s) and analyzed for surface-expression of the HA epitope tag.



Figure S3. Tissue expression of NACHO and pentameric ion channels.

Heat map of mRNA levels from the Genotype-Tissue Expression (GTEx) database for the indicated genes (along the x-axis) in the indicated tissues (along the y-axis).



Figure S4. Photo-crosslinking analysis of mini-α1.

(A) A structural model of mini- α 1 is shown with spheres indicating the sites of incorporation of the UV-activated photo-crosslinking amino acid Bpa (left). Between four to eight positions were sampled in each of the four TMDs (M1 to M4). (B) FLAG-tagged mini- α 1 variants lacking (WT) or containing an amber codon at the indicated positions was translated in reticulocyte lysate supplemented with ³⁵S-methionine, microsomes derived from Expi293 cells, and amber suppression reagents for incorporation of Bpa. The reactions were irradiated with UV light where indicated, subjected to denaturing anti-FLAG IP to recover mini- α 1, and the products visualised by autoradiography. Two exposures are shown. The position of glycosylated (and hence, inserted) mini- α 1 is indicated. Crosslinks of mini- α 1 to various products are indicated by asterisks, with different colours indicating different interaction partners. The teal asterisk proved to be the crosslink to NACHO (see Fig. 2c). (C) Crosslinking experiment similar to panel b. Total IVT products and denaturing anti-NACHO IPs are shown in the top and bottom panels, respectively. Relative to the strong NACHO crosslink from position 309, crosslinks from positions 252 and 260 were weaker; no crosslink to NACHO was seen from position 298.



Figure S5. Structure prediction and conservation of NACHO.

(A) AlphaFold2 model of human NACHO (TMEM35A) coloured by pLDDT. (B) The structural model of NACHO is shown with spheres indicating the sites of incorporation of the UV-activated photo-crosslinking amino acid AbK (right). (C) The structural model of NACHO coloured by conservation.



Figure S6. Cryo-EM data collection and processing.

(A) Portion of a typical cryo-EM micrograph of purified α 1-NACHO complex reconstituted into nanodiscs. (B) Flowchart of the processing pipeline for structure determination of the α 1-NACHO complex. (C) Manually selected 2D class averages of the α 1-NACHO complex.



Figure S7. Quality of cryo-EM maps and models.

(A) Fourier shell correlation (FSC) curves of the α 1-NACHO structure. (B) Particle orientation distribution with efficiency *E* (calculated by cryoEF) of the α 1-NACHO dataset. (C) Cryo-EM map coloured by local resolution.



Figure S8. Analysis of NACHO- α 1 and α 1- α 1 interactions.

(A) The surfaces of $\alpha 1$ that interact with gamma2 (yellow), NACHO (teal) or both (pink) are depicted at left. The right shows a close-up of the $\alpha 1$ -NACHO interface. (B) The surfaces of $\alpha 1$ that interact with beta3 (blue), another $\alpha 1$ (red) or both (purple) are depicted at left. The right shows a close-up of the $\alpha 1$ - $\alpha 1$ interface. (C) The NACHO- $\alpha 1$ - $\alpha 1$ -NACHO structure coloured by hydrophobicity. The approximate position of the membrane is indicated.



Figure S9. Conservation of the NACHO- α 1 interaction surfaces.

(A) The surface residues on α 1 that contact NACHO are coloured by similarity across other GABA_A receptor subunits. (B) Relative similarity of α 1 residues that contact NACHO. (C) The surface residues of NACHO that contact α 1 are coloured by similarity with TMEM35B. (D) NACHO residues that contact α 1 are compared to the corresponding residues from TMEM35B.



Figure S10. Conformational changes in the β 5-5' hairpin of the α 1 subunit during receptor assembly.

(A) Schematic representation of the β 5-5' hairpins in the NACHO- α 1 complex (left) and the fully assembled pentamer (right). In the pentameric receptor, the β 5-5' hairpin of each subunit interacts with both neighbouring subunits. (B) Cross-section of the α 1 β 3 γ 2 GABA_AR (PDB: 7QNE) depicting the positions of the α 1 subunit β 5-5' hairpins. For clarity, α 1 subunit are represented only by a schematic. (C) Putative conformational changes of the α 1 subunit β 5-5' hairpins during assembly and ECD maturation. The "extended" conformation of the hairpin seen in the NACHO- α 1 complex may facilitate initial contact with the incoming β 3-E subunit.



Figure S11. Putative mechanism of β subunit incorporation based on AlphaFold predictions.

(A) AlphaFold-Multimer prediction of the α 1-NACHO complex closely matches the experimentally derived model. (B) AlphaFold predicts an alternative binding mode for α 1-NACHO, in which α 1 interacts with the interface on NACHO that we experimentally identified here as the binding site for the β subunit. (C) Proposed model of α 1 β 3 γ 2 GABA_AR assembly. The first step is based on the experimental model presented here, while the final step is the previously reported experimentally determined α 1 β 3 γ 2 GABA_AR structure (PDB: 7QNE). The second step - involving incorporation of the first β subunit - is based on AlphaFold predictions for the NACHO- β 3 complex, aligned with the experimentally derived α 1-NACHO structure using NACHO as the template. No clashes are observed between the transmembrane or extracellular domains in this model. The remaining steps are modelled by aligning subunits according to their interactions in the pentameric receptor.

Supplementary Tables:

Supplementary Table 2: Constructs used in this study

	Internal ref.	Figure			
	Base constructs				
Mini-α1-3xFLAG	SP64-based vector containing epitope-tagged mini- α 1 (residues xx-yy of human GABRA1). The N-terminal tail contains an HA epitope and glycosylation site; the C-terminal tail contains a 3X-FLAG epitope. The stop codon was changed to TAA (ochre).	YH027	1a, 1b, 1c, 2a, 2c, 2d, EDF1c, EDF1e		
Mini-α1-TST	Same as Mini- α 1-3xFLAG except the FLAG tag is replaced by the twin-strep tag (TST). The stop codon was changed to TAA (ochre).	YH029	1b, EDF1e		
β1AR∆CL3-FLAG	SP64-based vector containing the thermostabilised turkey β 1AR in which the third cytosolic loop has been shortened. Characterised in Chitwood et al., 2018, Cell, 175(6):1507-1519.	YH063	1a, 1b, 1c, EDF1c, EDF1e		
α1-3xFLAG	SP64-based vector containing epitope-tagged full length human GABRA1. The C-terminal tail is appended with a 3XFLAG tag.	YH090	1b, 1c, EDF1c, EDF1e		
TST-α1	SP64-based vector containing full length human GABRA1 with the twin-strep tag (TST) inserted downstream of the signal peptide.	-	2b		
TST-Rho-Ext	Gene block coding for Rhodopsin extended at the N- terminus with a signal sequence, twin-strep tag (TST) and translocated soluble domain. Described in Smalinskaite et al., 2022, Nature, 611(7934):161-166.	MK16 (gBlock)	2b		
Mini-β2-3xFLAG	SP64-based vector containing epitope-tagged mini-β2 (residues xx-yy of human GABRB2). The N-terminal tail contains an HA epitope and glycosylation site; the C-terminal tail contains a 3X-FLAG epitope.	YH116	4a		
NACHO-TST	TMEM35A (NACHO) in the pCDNA5.1 vector with a C-terminal twin-step tag (TST).	YH147	1d, 4b		
NACHO-TST (102A,103A)	TMEM35A (NACHO) in the pCDNA5.1 vector with a C-terminal twin-step tag (TST).	YH169	4b		
NACHO-TST (12A,120A,121A)	TMEM35A (NACHO) in the pCDNA5.1 vector with a C-terminal twin-step tag (TST).	YH172	4b		
NACHO-FLAG	pCDNA3.1-based vector expressing NACHO tagged at the C-terminus with FLAG.	-	2d, 4a		
CHRNA7-3xFLAG	pCDNA5.1 vector containing human CHRNA7 containing a C-terminal 3xFLAG tag.	YH123	EDF2c		
pHR-TST-α1	Codon-optimised human GABA _A R α 1 residues 28- 456 preceded by the chicken RPTP σ secretion signal peptide (MGILPSPGMPALLSLVSLLSVLLMGCVA), secretion leader sequence (ETG), the twin-strep tag (TST), and GGS linker.	-	3, EDF6		
pHR-NACHO-1D4	Codon-optimised human TMEM35A (NACHO) ACHO-1D4 appended with a linker (GGSGGSGGS) and Rho-1D4 at the C-terminus.		3, EDF6		
Mini-α1 folding mutants					
Mini-α1-3xFLAG (C234W)	Backbone is same as Mini-α1-3xFLAG.	YH136	2a		
Mini-α1-3xFLAG (V238W)	Backbone is same as Mini-α1-3xFLAG.	YH137	2a		
Mini-α1-3xFLAG (T262W)	Backbone is same as Mini-α1-3xFLAG.	YH138	2a		
Mini-α1-3xFLAG (S396W) Backbone is same as Mini-α1-3xFLAG.		YH143	2a		
Mini-α1 amber mutants for photo-crosslinking					
Mını-α1-3xFLAG (Y225amber)	Backbone is same as Mini- α 1-3xFLAG.	YH078	EDF4		

Mini-α1-3xFLAG (I228amber)	Backbone is same as Mini-α1-3xFLAG.	YH079	EDF4
Mini-α1-3xFLAG (C234amber)	Backbone is same as Mini-α1-3xFLAG.	YH039	EDF4
Mini-α1-3xFLAG (L240amber)	Backbone is same as Mini-α1-3xFLAG.	YH080	EDF4
Mini-α1-3xFLAG (L247amber)	Backbone is same as Mini-α1-3xFLAG.	YH081	EDF4
Mini-α1-3xFLAG (V252amber)	Backbone is same as Mini-α1-3xFLAG.	YH131	EDF4
Mini-α1-3xFLAG (V257amber)	Backbone is same as Mini-α1-3xFLAG.	YH012	EDF4
Mini-α1-3xFLAG (F258amber)	Backbone is same as Mini- α 1-3xFLAG.	YH013	EDF4
Mini-α1-3xFLAG (V260amber)	Backbone is same as Mini- α 1-3xFLAG.	YH132	EDF4
Mini-α1-3xFLAG (L264amber)	Backbone is same as Mini- α 1-3xFLAG.	YH014	EDF4
Mini-α1-3xFLAG (L269amber)	Backbone is same as Mini- α 1-3xFLAG.	YH015	EDF4
Mini-α1-3xFLAG (I271amber)	Backbone is same as Mini- α 1-3xFLAG.	YH016	EDF4
Mini-α1-3xFLAG (R274amber)	Backbone is same as Mini- α 1-3xFLAG.	YH082	2c, EDF4
Mini-α1-3xFLAG (Y294amber)	Backbone is same as Mini- α 1-3xFLAG.	YH083	2c, EDF4
Mini-α1-3xFLAG (F298amber)	Backbone is same as Mini- α 1-3xFLAG.	YH133	EDF4
Mini-α1-3xFLAG (L301amber)	Backbone is same as Mini- α 1-3xFLAG.	YH084	2c, EDF4
Mini-α1-3xFLAG (L309amber)	Backbone is same as Mini- α 1-3xFLAG.	YH085	2c, EDF4
Mini-α1-3xFLAG (I398amber)	Backbone is same as Mini- α 1-3xFLAG.	YH086	EDF4
Mini-α1-3xFLAG (F404amber)	Backbone is same as Mini- α 1-3xFLAG.	YH087	EDF4
Mini-α1-3xFLAG (L409amber)	Backbone is same as Mini- α 1-3xFLAG.	YH088	EDF4
Mini-α1-3xFLAG (Y411amber)	Backbone is same as Mini- α 1-3xFLAG.	YH089	EDF4
	NACHO amber mutants for photo-crosslinking		
NACHO- FLAG(L18amber)	pCDNA3.1-based vector expressing NACHO tagged at the C-terminus with FLAG.	YH153	2d, 4a
NACHO- FLAG(V21amber)	pCDNA3.1-based vector expressing NACHO tagged at the C-terminus with FLAG.	YH105	2d, 4a
NACHO- FLAG(S38amber)	pCDNA3.1-based vector expressing NACHO tagged at the C-terminus with FLAG.	YH107	2d
NACHO- FLAG(Y47amber)	pCDNA3.1-based vector expressing NACHO tagged at the C-terminus with FLAG.	YH108	2d, 4a
NACHO- FLAG(K55amber)	pCDNA3.1-based vector expressing NACHO tagged at the C-terminus with FLAG.	YH109	2d, 4a
NACHO- FLAG(L71amber)	pCDNA3.1-based vector expressing NACHO tagged at the C-terminus with FLAG.	YH110	2d
NACHO- FLAG(L99amber)	pCDNA3.1-based vector expressing NACHO tagged at the C-terminus with FLAG.	YH111	2d
NACHO- FLAG(Q106amber)	pCDNA3.1-based vector expressing NACHO tagged at the C-terminus with FLAG.	YH112	2d, 4a
NACHO- FLAG(H117amber)	pCDNA3.1-based vector expressing NACHO tagged at the C-terminus with FLAG.	YH113	2d
NACHO- FLAG(T126amber)	pCDNA3.1-based vector expressing NACHO tagged at the C-terminus with FLAG.	YH156	2d
NACHO- FLAG(R138amber)	pCDNA3.1-based vector expressing NACHO tagged at the C-terminus with FLAG.	YH158	2d

Antibody	Source	Catalog No.	RRID	Dilution for blotting	Dilution for IP	Dilution for flow cytometry
Rb anti-TMEM35 (NACHO)	Proteintech	24786-1-AP	AB_2879723	1:1000	1:500	-
Rb anti-RPL8	Abcam	ab169538	AB_2714187	1:10000- 1:20000	-	-
Rb anti-CCDC47	Bethyl	A305-100A	AB_2631495	1:10000	-	-
Rb anti-Asterix	Invitrogen	PA5-66788	AB_2665196	1:2000	-	-
Rb anti-Sec61β	Fons et al. (2003) <i>J.</i> <i>Cell Biol.</i> 160 :529-539.	-	-	1:5000	-	-
Mo anti-FLAG-HRP	Sigma	A8592	AB_439702	1:5000	-	-
Rb anti-Strep tag	Abcam	ab76949	AB_1524455	1:5000	-	-
Anti-GABAA Receptor α1	Merck- Millipore	06-868	AB_310272	-	1:500	-
Rat PE anti- DYKDDDDK	Biolegend	637310	AB_2563148	-	-	1:100

Supplementary Table 3: Antibodies used in this study

Data collection and	
processing	BiocEM Krios
Microscope	G3
Electron Gun	XFEG
Detector	K3
Magnification	81k
Energy filter slit width (eV)	20
Voltage (kV)	300
Flux on detector (e/pix/s)	20.44-20.52
Electron exposure on sample	44.96-45.14
(e ⁻ /Å ²)	
Objective aperture (µm)	100
Target defocus range (um)	1-2.6
Calibrated pixel size (Å)	1.066
Symmetry imposed	C2
Number of collected movies	23.057
Initial particle images (no)	10 530 693
Final particle images (no.)	304 241
Map resolution at ESC=0 143	36
	0.0
EMPIAR code	11691
	11001
Refinement	
Model resolution (Å)	3.5
FSC threshold	0.143
Model composition	0.110
Non-hydrogen atoms	7942
Protein residues	03 <u>1</u>
Nucleotide bases	0
Ligande	0 172
B factors $(Å^2)$	472
Brotoin	145 60
Pioleini Nucleatide	140.02 Not oppliaable
ligand	
Ligano Des deviations	177.70
R.m.s. deviations	0.000
Bond lengths (A)	0.002
Bond angles (°)	0.408
Validation	4.05
MolProbity score	1.25
Clash score	4.83
Poor rotamers (%)	0.00
Ramachandran plot	
Favored (%)	98.26
Allowed (%)	1.74
Disallowed (%)	0.00
PDB code	8QM2

Table S4: Cryo-EM data collection and refinement of the α 1-NACHO model

B-factors (ADPs) were calculated with Phenix. Other values are calculated by MolProbity using the Phenix package (version dev-5430-000).