Mechanisms of AMPA receptor tetramerization

The transmembrane domain mediates tetramerization of α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) receptors

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ABSTRACT

AMPA receptors (AMPARs) mediate fast excitatory neurotransmission in the central nervous system. Functional AMPARs are tetrameric complexes with a highly modular structure, consisting of four evolutionarily distinct structural domains: an amino-terminal domain (ATD), a ligand-binding domain (LBD), a channel-forming transmembrane domain (TMD) and a carboxy-terminal domain (CTD). Here we show that the isolated TMD of the GluA1 AMPAR is fully capable of tetramerization. Additionally, removal of the extracellular domains from the receptor did not affect membrane topology or surface delivery. Furthermore, while the ATD and the CTD contributes positively to tetramerization, the LBD presents a barrier to the process by reducing the stability of the receptor complex. These experiments pinpoint the TMD as the “tetramerization domain” for AMPARs, with other domains playing modulatory roles. They also raise intriguing questions about the evolution of iGluRs as well as the mechanisms regulating the biogenesis of AMPAR complexes.

Ionotropic glutamate receptors (iGluRs), predominantly the α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) receptor subtype, mediate fast excitatory neurotransmission in the mammalian central nervous systems. Functional AMPARs are tetramers consisting of four identical (homomeric) or similar (heteromeric) subunits (1) arranged as a dimer-of-dimers (2-4). The process of AMPAR assembly determines the availability of AMPARs to be trafficked to the cell surface, and therefore represents a target for modulating glutamatergic signaling (5).

iGluR subunits are highly modular, being composed of four well-defined structural domains (6,7) possessing distinct evolutionary origins (8): an amino-terminal domain (ATD), a ligand-binding domain (LBD), a transmembrane (TMD) consisting of three hydrophobic segments (M1, M3 and M4) as well as a re-entrant loop (M2), and a carboxy-terminal (CTD) domain attached to the intracellular end of M4 (9,10). Previous studies have identified structural elements that contribute to the assembly of AMPARs in almost every domain (5,11,12). The ATD of AMPARs forms dimers in solution and drive the initial dimerization process (13-15). Alternative splicing in the LBD (flip/flop) influences subunit composition of AMPAR heteromers, potentially by affecting the tetramerization process (16). Residues in the TMD, especially those near the channel pore (e.g. the Q/R editing site), also impact tetramerization (3,17). Recent studies
pinpointed the M4 segment, which is unique to eukaryotic iGluRs, as a critical determinant of AMPAR tetramerization (18,19). Despite the large number of studies investigating individual factors influencing AMPAR assembly, the relative contribution of each structural domain is unclear. Here, we took advantage of the highly modular nature of the iGluR structure to address the assembly process by deleting entire domains from an AMPAR subunit, GluA1. We demonstrate that the TMD is the minimal structural unit required for AMPAR tetramerization, highlighting the central role of this domain to the assembly process. The three other domains modulate tetramerization of AMPARs but are not required. Surprisingly, while the ATD and CTD contribute positively to tetramerization, the LBD presents an unexpected hindrance to the process. Our findings hint at previously unknown mechanisms through which AMPAR biogenesis could be regulated. They may also help inspire novel therapeutic strategies that modulate glutamatergic signaling by targeting the assembly of AMPARs.

EXPERIMENTAL PROCEDURES

Construct design, mutagenesis and expression—All constructs were designed based on the “flip” variant of the rat GluA1 (Accession#P19490) subunit. Constructs lacking the amino-terminal domain (ΔATD) of GluA1 were tagged with hemagglutinin (HA) near the N-termini. The cDNA encoding the HA-GluA1-ΔATD construct was generated by introducing restriction sites (EcoRV and Ascl) at positions flanking the ATD (A1 and D376) and then excising the entire ATD-encoding region and replacing it with a segment encoding the HA tag. The GluA1-ΔCTD and HA-GluA1-ΔATD/ΔCTD constructs were created by substituting residue S814 with a stop codon in full-length GluA1 and HA-GluA1-ΔATD, respectively. The HA-GluA1-ΔECD construct was ordered from GeneScript and contains all LBD-TMD linkers as a stretch of 5 (GT)5 between the M3 and M4 segments to replace the S2 segment of LBD. The full amino acid sequence is as follows:

MPYIFAFFCTGFLGAVVGANFQYVDV
PDYAPNNIQIGGLFPNKQSKPGVFS
FLDPLAYEIWMCIYAFAYIGVSVVLFL
VSRFSPYEHXSEFEEGRDQTTSDQS

NEFGFNSLWFLGAFMQQGCDISP
RSLSGRIVGGVWFWTILIISSYANT
LAAFLTVERMVSPIEGTGTTGTTGTE
CGSKDSGSKDKTSALSLSNVAGYFYI
LIGGLGLAMLVALIECFYKSRSESK
RMKGFCLIPQSSINEAIRTSTLPNSG
AGASGGGSGENGRVVSQDFPKSMQ
SIPCMSHSSGMPLATGL

The M1, M2, M3 and M4 regions are marked as bold while the 5(GT) linker is italicized. The sequence encoding for EGFP was introduced after the CTD region of this construct to generate the HA-GluA1-ΔECD-EGFP construct. Residue S814 in the HA-GluA1-ΔECD construct was changed to a stop codon to create HA-GluA1-ΔECD/ΔCTD. Calculated molecular weights of these constructs are listed in Table 1. In all of our constructs the amino acid numbering is for the mature receptor (without the signal peptide). Point mutations were generated using PCR-based methods and confirmed by sequence analysis. Human embryonic kidney (HEK) 293 cells were transfected with the cDNAs using X-tremeGene 9 or X-tremeGene HP (Roche) (20).

Immunocytochemistry (ICC)—HEK 293 cells were plated in 24-well plates at a density of 0.5 x 10⁵ cells/well on coverslips pretreated with nitric acid and coated with poly-D-lysine (25 µg/ml in PBS). Cells were maintained in 10% FBS at 37°C and 95% O₂/5% CO₂. 36 ~ 48 hrs after transfection, surface expression was determined as previously described (18).

To determine surface expression of HA-tagged AMPAR constructs, primary antibodies labeling either the N-terminally tagged HA (Covance, MMS-101P, mouse monoclonal 16B12) or the GluA1 C-terminus (Millipore, rabbit polyclonal AB1504) dissolved in conditioned medium were added directly onto the coverslip and incubated in a humidified chamber at 37°C for 30 min. Cells were then fixed in 4% paraformaldehyde and blocked in 1% goat serum. Cells were rinsed with PBS and incubated with Alexa Fluor® 546 goat anti-mouse or Alexa Fluor® 488 goat anti-rabbit secondary antibodies (Invitrogen, Carlsbad, CA) at room temperature. For permeabilized conditions, the cells were fixed in 4% paraformaldehyde and permeabilized in 0.25% Triton X-100 before blocking and primary antibody incubation. Prepared coverslips were
examined using an upright Fluoview® FV1000 confocal microscope (Olympus).

Harvest of membrane proteins—Cells were plated on 15 x 60 mm tissue culture dishes at roughly 3 x 10⁵ cells/dish. Maintenance of culture and transfections were done as for ICC.

Whole-cell lysates (including membrane proteins) were harvested as previously described (21). Briefly, a solubilization buffer is prepared by dissolving 20 mM (1%) n-dodecyl-α-D-maltopyranoside (DDM) (Affymetrix, D310HA) in PBS containing a protease inhibitor cocktail (Thermo Scientific, 1861278) as well as 1 mM phenylmethylsulfonyl fluoride. Cells were rinsed in a washing buffer (PBS containing 0.5 mM N-ethylmaleimide) and then harvested in the solubilization buffer. The lysate was rotated for 1 hr at 4°C and centrifuged at 50,000 RPM on a Beckman TLA 120.2 rotor for 40 mins at 4°C. The supernatant containing solubilized membrane proteins was collected for blue native PAGE.

For assaying the detergent or thermal stability of various constructs, a lysis buffer (20 mM Tris in PBS) was used to harvest cells after rinsing in the aforementioned washing buffer. We homogenized the suspended cells by passing them through a gauge 25 needle for six times using a 1 ml syringe. The cell lysate was then centrifuged at 3,000 RPM on an Eppendorf 5417R microcentrifuge for 3 min at 4°C. The supernatant containing the membrane fraction was then centrifuged at 50,000 RPM on a Beckman TLA 120.2 rotor for 10 min at 4°C. The pellet was isolated and re-suspended in chilled PBS, and the suspension was centrifuged again. The final pellet containing the membrane fraction was re-suspended in an alternative solubilization buffer (20 mM Tris and 50 mM NaCl in PBS) and was sonicated 4 times, each for 15 sec. 6 mM (0.3%) of DDM along with various concentrations of SDS (0% ~ 0.2%) was added to the suspension. The membrane lysate was then rotated for 1 hr at 4°C and centrifuged at 50,000 RPM on a Beckman TLA 120.2 rotor for 20 mins at 4°C. The resulting supernatant containing only solubilized membrane proteins was collected for analysis. For thermal stability assays, the supernatant (solubilized with 20 mM DDM and no SDS) was incubated at a particular temperature (0°C ~ 40°C) for 20 min right before blue native PAGE.

Blue Native PAGE (BN-PAGE)—Membrane proteins contained in the supernatant were resolved using Blue Native-PAGE as previously described (21,22). Briefly, protein samples mixed with 1X NativePAGE sample buffer, 0.05% NativePAGE G-250 additive and supplemented with 10 mM DDM were loaded onto Novex 4-16% Bis/Tris gradient gels. NativeMark™ Unstained Protein Standard (Life Technologies) and horse spleen apoferritin (Sigma) were also loaded as molecular weight markers. Proteins were separated at constant voltage (115 V for 1 hr and then 215 V for 2 hrs) at 4°C. Gels were transferred overnight (14 ~ 16 hrs) at constant amperage (35 mA) to polyvinylidene fluoride (PVDF) membranes. Following transfer, membranes were briefly rinsed with methanol and proteins were fixed on the membrane by incubating in 8% acetic acid for 15 min. Presence of proteins was confirmed by Ponceau S staining. Membranes were then rinsed with water, rehydrated with methanol, and rinsed with TBS-T (0.05% Tween) to remove the stain. Membranes were blocked in 5% Milk-TBS and then incubated with either anti-GluA1 (Millipore, MAB2269, mouse monoclonal RH95) or anti-HA (Covance, MMS-101P, mouse monoclonal 16B12). After rinsing in TBS-T, membranes were incubated with HRP-conjugated anti-mouse IgG secondary antibodies (Santa Cruz Biotechnologies, sc-2030). Blots were developed using luminol reagent (Santa Cruz Biotechnologies, sc-2048) before exposure to chemiluminescence blue-sensitive film (Crystalgen). In certain instances, lanes from the same gel are presented in a different order from the original gel (indicated by a thin space between lanes) for clarity of presentation. Each experiment with a particular construct is repeated at least n=3 times to ensure reproducibility.

Surface biotinylation assay—Cells expressing constructs of interest were rinsed with PBS containing 100 µM CaCl₂ and 1 mM MgCl₂ (PBS/CM) 48 hours post-transfection. Cells were then incubated in either PBS/CM or 5mg/ml sulfo-NHS-LC-biotin (Thermo) for 45 min on ice. The reaction was terminated by incubation in 100 mM glycine. After rinsing with PBS/CM, whole-cell lysates were harvested as described above and subsequently rotated with strepavidin agarose beads (Thermo) top-to-end for 2 hrs at room temperature. Membrane proteins contained in the supernatant were resolved using Blue Native-PAGE as previously described (21,22). Briefly, protein samples mixed with 1X NativePAGE sample buffer, 0.05% NativePAGE G-250 additive and supplemented with 10 mM DDM were loaded onto Novex 4-16% Bis/Tris gradient gels. NativeMark™ Unstained Protein Standard (Life Technologies) and horse spleen apoferritin (Sigma) were also loaded as molecular weight markers. Proteins were separated at constant voltage (115 V for 1 hr and then 215 V for 2 hrs) at 4°C. Gels were transferred overnight (14 ~ 16 hrs) at constant amperage (35 mA) to polyvinylidene fluoride (PVDF) membranes. Following transfer, membranes were briefly rinsed with methanol and proteins were fixed on the membrane by incubating in 8% acetic acid for 15 min. Presence of proteins was confirmed by Ponceau S staining. Membranes were then rinsed with water, rehydrated with methanol, and rinsed with TBS-T (0.05% Tween) to remove the stain. Membranes were blocked in 5% Milk-TBS and then incubated with either anti-GluA1 (Millipore, MAB2269, mouse monoclonal RH95) or anti-HA (Covance, MMS-101P, mouse monoclonal 16B12). After rinsing in TBS-T, membranes were incubated with HRP-conjugated anti-mouse IgG secondary antibodies (Santa Cruz Biotechnologies, sc-2030). Blots were developed using luminol reagent (Santa Cruz Biotechnologies, sc-2048) before exposure to chemiluminescence blue-sensitive film (Crystalgen). In certain instances, lanes from the same gel are presented in a different order from the original gel (indicated by a thin space between lanes) for clarity of presentation.
temperature. Beads were rinsed three times with PBS/CM and bound proteins were eluted by heating at 95°C in SDS sample buffer containing 50 mM DTT. Whole-cell lysates ("W"), eluted proteins ("E") as well as unbound fractions ("U") were subjected to SDS-PAGE and proteins of interest are detected by immunoblotting using antibodies described above. The percentages of loading for these samples are 2%, 50% and 2%, respectively. GAPDH was selected as a cytosolic loading control and was detected using a mouse monoclonal anti-GAPDH antibody (Calbiochem, CB1001). Intensity of each band was quantified using ImageJ (see below) and surface expression was measured as the ratio between the eluted ("E") and the total protein ("W") bands.

Densitometry and quantification—The concentration of antibodies, duration of incubation as well as time of exposure were optimized to ensure that the intensities of major bands fall within the linear response range of the film (21). Developed films were scanned into .tiff format and analyzed using ImageJ. The oligomeric state of each well-resolved band was designated based on its position relative to the molecular weight markers. The mean intensity (I) as well as the area (A) of each band of interest was measured (21). The background mean intensity (I_b) was also measured on an area where there was no signal. The cumulative intensity (C) of each band was calculated as:

\[ C = (I - I_b) \times A \]

For each construct, the fraction of tetramer (T) was calculated from the cumulative intensities of each band (C_T, C_{Tn}, C_D and C_M) (21):

\[ T = \frac{C_T}{C_T + C_{Tn} + C_D + C_M} \]

where T is defined as the fraction of subunits that are assembled into tetrameric complexes, assuming a linear relationship between the number of antibodies bound to a complex and the number of subunits it contains.

It should be noted that the intensity of each band is not necessarily proportional to the total amount of protein the band contains, since it could not be guaranteed that every single subunit in a given oligomeric complex is bound with an antibody. The fraction of tetramer intensity is therefore treated only as a relative measurement of tetramerization efficiency and cannot be used to calculate absolute physical quantities such as free energy of oligomerization, as was done in previous studies (23).

Data analysis and statistics—Stability assays, the percentage tetramer at a particular temperature or SDS concentration (T_n) was normalized to the baseline percentage at 0°C or 0% SDS, respectively (T_0):

\[ \text{Norm}_T = \frac{T_n}{T_0} \]

For each temperature or SDS condition, Student’s t-tests are performed between the T_n of each construct and the corresponding T_n of full-length GluA1.

To assess the relative stability (RS) of a particular mutant, we calculated the ratio between its normalized T and that of a wild type sample run on the same gel, either at a selected temperature (37°C) or at a selected mid-point SDS concentration (0.11%):

\[ RS_{\text{mut}} = \frac{\text{Norm}_T_{\text{mut}}}{\text{Norm}_T_{\text{WT}}} \]

Paired Student’s t-tests were performed between the normalized T of each mutant and the normalized T of the corresponding wild type sample run on the same gel under the same condition.

Homology modeling and molecular dynamics simulation—The homology model of the GluA1 TMD is built based on the TMD of a full-length GluA2 crystal structure (PDB code: 3KG2) (6) in MODELLER (24) by generating 50 models and selecting the one with the lowest score. The sequences of GluA2 and GluA1 TMD were aligned using Clustal Omega (25). In order to reduce the system size and speed up simulations, the intracellular M1-M2 linker (Y545 to D562) of each subunit was replaced with a GGG linker; otherwise, the TMD construct used in modeling and simulation is the same as that used in the experiments.

The GluA1 TMD structural model was first energy-minimized in vacuum with its Cα atoms fixed in position for 5000 steps, then it was embedded into a pre-equilibrated POPC bilayer obtained from CHARMM-GUI (26). Lipid and
water molecules that overlapped with the protein were removed. The charge of the system was neutralized with NaCl at a salt concentration of about 150 mM. A second energy minimization was conducted with protein Cα atoms fixed in position for 5000 steps. MD simulation was then performed for 5 ns, in which only the C-α atoms of the transmembrane helices restrained using a harmonic potential with a gentle force constant of 1.0 kcal/(mol*Å²). Finally, a 71.7 ns unbiased MD simulation was performed in which no restraints were applied to the system.

System preparation was done in VMD (27). All simulations were performed using NAMD version 2.9 (28) with CHARMM27 protein force field and CHARMM36 lipid force field (29). The simulation parameters were the same as those for the previous simulations (30,31).

RESULTS

AMPAR GluA1 forms a tetramer in the absence of the ECD—Eukaryotic iGluRs consist of four structural domains: the amino-terminal domain (ATD), the ligand-binding domain (LBD), the transmembrane domain (TMD) and an intracellular carboxy-terminal domain (CTD). The extracellular domain (ECD), which consists of the ATD and the LBD, accounts for 70% ~ 80% of the molecular weight of an AMPAR complex (Fig 1A). To investigate the significance of ECD in AMPAR tetramer assembly, we used blue native PAGE (BN-PAGE) to assess the oligomeric states of GluA1 with or without ECD. Consistent with previously published results (17,19), full-length AMPAR subunits, in this case GluA1, expressed in HEK 293 cells readily formed homo-tetramers (Figure 1B, left lane). A small dimer fraction was also detected, in keeping with the notion that AMPARs are dimer-of-dimers (2-4). When the ECD was deleted (HA-GluA1-ΔECD), the remaining domains of GluA1, including the TMD and CTD, still formed a homo-tetramer as estimated from its molecular weight (Figure 1B, right lane). No dimer band was detected for the ECD-lacking construct, consistent with the pseudo-4-fold symmetry of the TMD (6) and the idea that the ATD is responsible for AMPAR dimerization (13).

To confirm that the single band observed with HA-GluA1-ΔECD represents a tetramer, we tagged the construct with EGFP at the C-terminus, which caused an up-shift of its molecular weight (Fig. 1C, far right lane). When HA-GluA1-ΔECD and HA-GluA1-ΔECD-EGFP were co-expressed at various ratios, a total of four incremental up-shifts were observed (Figure 1C, inner lanes), implying the incorporation of 1, 2, 3 and 4 EGFP-tagged subunits respectively. Thus, the ECD-lacking GluA1 maintains a tetrameric stoichiometry.

Surface trafficking, membrane topology and packing of pore-lining helices are maintained despite the lack of ECD—In full-length AMPARs, the N-terminus is extracellular while the C-terminus is cytoplasmic (Figure 1A). To test whether the tetramer formed by HA-GluA1-ΔECD maintained the same membrane topology, we performed immunocytochemistry (ICC) using two different primary antibodies: one against the N-terminal HA tag and the other against the CTD (Figure 2A, left panel). Under non-permeabilizing conditions, we observed cell surface labeling only by the anti-HA antibody but not the anti-CTD antibody (Figure 2A, right panel, left 3 columns). Under permeabilizing conditions, we observed labeling by both antibodies (Figure 2A, right panel, rightmost column). Thus, HA-GluA1-ΔECD not only is capable of surface trafficking, but also has a membrane topology just like full-length GluA1, with the N-terminus extracellular and the C-terminus cytoplasmic.

To test whether the ECD-lacking GluA1 tetramer still contained an ion channel resembling that of the intact receptor, we used the lurcher mutation (A618T), which disrupts tetramerization of full-length AMPARs (3) possibly due to its location near the tightly packed activation gate of the fully-formed tetrameric ion channel (6). Consistent with previous studies (3), A618T reduced the tetramer fraction for intact GluA1 (Figure 2B) from 0.83 ± 0.04, n = 7 to 0.36 ± 0.09, n = 5 (mean ± SEM, n = number of samples). Similarly, the same mutation in the ECD-lacking GluA1 also reduced the tetramer fraction (Figure 2C) from 1.00 ± 0.01, n = 5 to 0.70 ± 0.11, n = 5 (see Experimental Procedures for discussions on quantification). Thus, the general pattern of helical packing in the ion channel is retained even in the absence of the ECD. The reduction in tetramer fraction was accompanied by either an increase of dimers (Figure 2B) or the emergence of trimers.
and dimers (Figure 2C). No monomer band was observed. This might reflect that the positioning of A618 in a non-tetrameric oligomer is different from that in a fully formed tetramer, resulting in A618T having no disruptive effect. Alternatively, interactions within the ATD (in the case of the full-length receptor), TMD (e.g. those mediated by the M4 segment) and/or CTD might compete with the lurcher mutation, thereby preventing the further breakdown of dimers.

Figure 2A indicate that the ECD is not required for the surface expression of GluA1. To quantify the effect of ECD deletion on GluA1 trafficking, we performed cell surface biotinylation assay. Both full-length GluA1 and HA-GluA1-ΔECD could be immunoprecipitated by streptavidin from surface biotinylation, in contrast to the cytosolic enzyme GAPDH (Figure 2D). There was no significant difference between the immunoprecipitated fraction of GluA1 and HA-GluA1-ΔECD (Figure 2E). These results, combined with the ICC experiment, suggest that the ECD does not contribute significantly to the efficiency of GluA1 surface trafficking.

*The TMD mediates tetramerization while the CTD plays a modulatory role*—The ECD-lacking construct contains two domains, the TMD and the CTD. To pinpoint which one of these is responsible for mediating tetramerization, we removed the CTD from this construct (HA-GluA1-ΔECD/ΔCTD) (see Experimental Procedures). This highly truncated construct was still able to form an oligomer, whose estimated molecular weight was consistent with that of a tetramer (91.6 kDa) (Figure 3A, right lane). Given the homology between the TM region of iGluRs and K+ channels (8,32), which still tetramerize even in the absence their specialized tetramerization domains (33-35), it is highly likely that the oligomer formed by HA-GluA1-ΔECD/ΔCTD was also a tetramer. To probe the stoichiometry of this oligomer, we included 0.11% sodium dodecyl sulfate (SDS) in the solubilization buffer to disrupt inter-subunit interactions (see Experimental Procedures and the next section of Results). As a result, the single band seen in Figure 3A (right lane) and Figure 3B (left lane) was partially broken down into two bands, presumably representing the dimer and the monomer (Figure 3B, right lane).

Unfortunately, we could not further verify the stoichiometry of HA-GluA1-ΔECD/ΔCTD using the experiment shown in Figure 1C because attaching the EGFP tag directly to the N-terminus of the M1 helix severely impaired protein folding (data not shown). We therefore performed homology modeling of a tetramer formed by this construct based on the GluA2 structure 3KG2 (6). The model after energy minimization was almost identical to the ion channel domain of 3KG2 (Figure 3C, initial conformation), with an RMSD of 0.54 Å. We also performed a total of 76.7 ns of constrained and unconstrained molecular dynamics simulations with the model and observed minimal changes in its structure (RMSD < 3.5 Å) (Figure 3C, final conformation). Hence, the deletion of all extra-membrane domains (ECD & CTD) and the introduction of the GT linker (See Experimental Procedures) did not induce any notable structural changes in the TMD tetramer, lending further credence to our conclusion that the transmembrane segments alone, independent of other domains, are capable of tetramerizing.

In the absence of the ECD, the deletion of CTD caused no notable impairment to the tetramerization process (Figure 3A). To test for any interaction between the ECD and the CTD, we removed the CTD in either full-length GluA1 (GluA1-ΔCTD) or a construct that lacks the ATD (HA-GluA1-ΔATD/ΔCTD). Tetramerization of full-length GluA1 lacking the CTD still occurred (Figure 3D, right lane) but in contrast to the TMD construct (Figure 3A, right lane) the efficiency of tetramerization, assayed as the tetramer fraction, was significantly reduced from 0.83 ± 0.05, n = 7 to 0.39 ± 0.03, n = 7. Thus, the CTD positively modulates the efficiency of tetramer formation in full-length AMPARs. The ATD-lacking construct (HA-GluA1-ΔATD) still predominantly forms tetramers (Figure 3E, left lane), as published previously (3,36). The dimer band observed with full-length GluA1 is here replaced by a monomer, as expected from the idea that the ATD is responsible for the initial dimerization (13). When the CTD is removed in the absence of the ATD but still in the presence of the LBD, tetramerization was almost completely abolished (Figure 3E, right lane), reducing the tetramer fraction from 0.77 ± 0.03, n = 5 to 0.16 ± 0.02, n = 3. This result is again consistent with the idea that CTD positively modulates tetramerization, but also suggests that
the LBD presents a penalty to the tetramerization process, which we refer to here as the “LBD barrier”.

**ATD and LBD have opposite effects on tetrameric stability**—To further address the energetic contributions of different domains to the assembly process, we assessed their influence on tetrameric stability by measuring changes in the tetramer fraction in response to various concentrations of SDS. SDS is a harsh anionic detergent that tends to disrupt non-covalent interactions between proteins, while at the same time partially mimicking the membrane environments required for TM helical contact (37). To improve the dynamic range of the assay, we also lowered the concentration of DDM in the solubilization buffer from 20 mM to 6 mM (see Experimental Procedures).

At 0% SDS, full-length GluA1, HA-GluA1-ΔATD and HA-GluA1-ΔECD-EGFP largely remained as tetramers (Figure 4A). As SDS concentration increased, the intensities of the tetramer bands decreased for all three constructs (Figure 4A). At the same time, additional bands corresponding to lower-number oligomers (dimers and monomers) began to appear. Pre-existing bands representing these lower oligomers also increased in intensity. These outcomes support the idea that incubation in SDS disrupts already-formed tetramers, causing them to break down into lower-number oligomers. We normalized the tetramer fraction at each SDS concentration to that at 0%, and fitted a sigmoidal SDS response curve for each construct (Figure 4B). The curve for HA-GluA1-ΔATD (Figure 4B, gray triangle and gray dash line) was significantly left-shifted compared to that of the full-length GluA1 (Figure 4B, open diamond and black line), indicating that although the ATD is not required for tetramerization, it contributes to the stability of the tetramer once it is formed. Surprisingly, the SDS response curve of HA-GluA1-ΔECD-EGFP was right-shifted compared to that of HA-GluA1-ΔATD and was almost indistinguishable from that of full-length GluA1 (Figure 4B, black square and black dash line). Thus, consistent with results shown in Figure 3, the LBD seems to confer a penalty to tetrameric stability, effectively balancing out the stabilizing effect of the ATD.

As an alternative approach to measure tetramer stability, we incubated the membrane samples at progressively higher temperatures. Consistent with earlier studies (35) increasing temperatures led to progressively lower tetramer fractions for all three constructs (Figure 4C). The derived curves (Figure 4C) showed the exact same pattern as the SDS response curves (Figure 4B), with HA-GluA1-ΔECD-EGFP being almost indistinguishable from the full-length GluA1, and HA-GluA1-ΔATD being left-shifted. The concordance between the SDS and the thermal approaches indicate that the results from both assays reflect the structural stability of the tetramers in a general sense, instead of being a fortuitous outcome associated with either treatment in particular.

As an additional validation of the SDS approach, we assayed the effect of the lurcher mutation (A618T) on tetrameric stability. For convenience, we selected as a reference a SDS concentration (0.11%) roughly at the mid-point of the dynamic range on the SDS response curves of the full-length and ECD-lacking constructs (Figure 4B), where the tetramer fraction of these constructs would be most sensitive to mutations that affect their stability. We introduced A618T into both constructs and found that the mutation led to a significant decrease in normalized tetramer fraction at 0.11% SDS regardless of the presence of ECD (Figure 5A&B). This is consistent with the disruptive effect of A618T on tetramerization with or without the ECD (Figure 2B&C) and suggests that the effect of A618T resulted from reduced tetrameric stability.

Our stability assays on the ECD-lacking constructs exhibited a striking concordance with results shown in Figure 3D&E. Overall, these results indicate that the ATD and the LBD modulate tetramerization in opposite directions, at least partially by influencing the stability of an already formed tetramer. The CTD also contributes positively to tetramerization (Figure 3D&E), though we did not measure its contribution to tetrameric stability, because the GluA1-ΔCTD showed a low baseline tetramer fraction, which would have limited the dynamic range of our assay.

**The M4 segment is critical to tetramerization only in the presence of the LBD**—Previously we identified a face in the third transmembrane segment (M4) of AMPARs that is critical to tetramerization (18,19). This face,
dubbed “VLGAVE” after the amino acid residues comprising it, is closely aligned with the M1 and M3 segments of an adjacent subunit in the GluA2 structure (Figure 6A), suggesting that interactions between M4 and the other TM segments play a critical role in AMPAR tetramerization. In support of this interpretation, mutating the residues within the VLGAVE face to tryptophan (W), which presumably disrupts molecular interactions between M4 and the rest of the TMD, prevents the formation of tetramers (Figure 6B) (19). A similar outcome occurred for HA-GluA1-ΔATD (Figure 6C), indicating that the TM interactions mediated by the VLGAVE face are still critical for tetramerization in the absence of the ATD. In contrast, the same mutations had no effect on the tetramerization of the ECD lacking construct with or without the EGFP tag (Figure 6D), suggesting a less important role of M4 in tetramerization when the LBD is absent. Thus, and again consistent with earlier results (Figure 3-4), the LBD of GluA1 presents a barrier to receptor tetramerization that can be offset, at least in part, by interactions mediated by the M4 segment.

An alternative explanation for the results in Figure 5C is that in the absence of the LBD, the M4 helix gains more freedom of movement. As a result, inter-subunit interactions in the TMD are now mediated by a different set of residues instead of the VLGAVE face. We therefore tested the effect of a subset of M4 tryptophan substitutions both within and outside of the VLGAVE face (Figure 7, gray and black residues respectively) on the stability of the ECD lacking tetramer with the mid-point SDS concentration of 0.11% as a reference point, at which the assay would be most sensitive (see Figure 4B). All of the tested mutations within the VLGAVE face (L795W, G798W, V805W and E809W) significantly affected the stability of HA-GluA1-ΔECD-EGFP (Figure 7B). In contrast, mutations outside of the VLGAVE face (G800W and L807W), which did not disrupt tetramerization in either full-length GluA1 or GluA2 (data not shown), showed either negligible or very slight effects on tetrameric stability (Figure 7B). These results indicate that interactions mediated by the M4 VLGAVE face still occur in the absence of the LBD and contribute to the stability of the tetramer, despite not being necessary for its formation. Thus, in the full-length receptor, these interactions are at least partially responsible for offsetting the “LBD barrier”.

Restricting quaternary arrangement of the LBD enhances tetramerization—One possible mechanism underlying the “LBD barrier” involves steric clashes among the four subunits at the LBD level. These clashes could destabilize the quaternary arrangement of the subunits, thereby affecting the stability of the entire receptor complex. The LBD is arranged as a dimer-of-dimers. We therefore introduced a previously published disulfide bond located at the LBD dimer-dimer interface (Figure 8A), which is known to stabilize that interface and restrict the quaternary movements of the LBD (38). The homologous cysteine substitution in GluA1, A661C, formed a disulfide crosslink as evidenced by redox-sensitive dimers in SDS-PAGE (Figure 8B, upper and mid panels). To test the effect of the crosslink on GluA1 tetramerization, we used a mutation in the M4 VLGAVE face, G798A, which dramatically, but not completely, impeded tetramerization (Figure 8B, lower panel) (19). The dimer-dimer crosslink was able to fully rescue the tetramerization deficit caused by G798A (from 0.46 ± 0.02, n = 9 to 0.92 ± 0.06, n = 2) (Figure 8B, lower panel). Hence, the instability of the LBD quaternary arrangement is most likely the structural basis of the “LBD barrier” to tetramerization.

DISCUSSION

Our experiments directly demonstrate that the TMD acts as the tetramerization domain in AMPARs (Figure 3A). Neither the ECD, which encompasses the ATD and the LBD, nor the CTD was necessary for the tetrameric assembly of GluA1 (Figures 1 & 3). Interestingly, the VLGAVE face of the M4 segment critical to the tetramerization of full-length GluA1 (19) is not required in the absence of the ECD (Figure 6D), although it still contributes to tetrameric stability (Figure 7). Together, our results suggest that the ion channel core (M1-M2-M3) of AMPARs alone is probably capable of tetramerization, with the peripheral M4 helix acting as a stabilizing “barrel hoop”. This scenario is similar to that of the KcsA channel, whose ion channel domain (homologous to the M1-M3 region of iGluRs) is fully capable of forming a functional tetramer, with the C-terminal “tetramerization domain” providing additional
stability (35).

A surprising finding from our experiments was that the presence of the LBD reduces the tetramerization efficiency (Figure 3A&D). Indeed, multiple lines of evidence point to an energetic penalty introduced by the LBD to GluA1 tetrameric assembly, referred to here as the “LBD barrier” (Figure 3, 4 & 6). The “LBD barrier” can be offset by some combination of the ATD, the M4 transmembrane segment and the CTD (Figure 3, 4, & 7).

One possible mechanism underlying the LBD barrier is that steric hindrance among the subunits at the LBD level destabilizes the tetrameric complex. Quaternary arrangements of the LBD are highly flexible (39-41). During the tetramerization process, quaternary rearrangements at the LBD level could lead to transient increases in steric hindrances. Consistent with this hypothesis, a disulfide cross-link introduced at the interface between two LBD dimers (Figure 8A), which presumably prevents inter-dimer quaternary rearrangements (38), rescued the deficit in tetramerization caused by a mutation in the M4 VLGAVE face (Figure 8B). In contrast, a mutation that restricts intra-dimer rearrangements in the LBD (L483Y in GluA2) has been found to negatively impact AMPAR assembly and maturation (42). This discrepancy seems to suggest that the LBD barrier arises from transient steric clashes between two LBD dimers but not within the dimers. Nevertheless, further work is required to pinpoint the exact structural determinants underlying the steric hindrance that destabilizes the LBD tetramer in the full-length receptor.

An alternative mechanism might involve the binding of glutamate at the LBD cleft. Biogenesis and assembly of nicotinic acetylcholine receptors (nAChR) are enhanced by the binding of nicotine (43,44). Similarly, binding of glutamate, which exists in the endoplasmic reticulum at high concentrations (45), could influence AMPAR assembly and maturation (42). This discrepancy seems to hint at a “checks and balances” scenario among multiple domains during the biogenesis of a functional AMPAR complex, with the LBD impeding tetramerization and the other domains collaborating to overcome the “LBD barrier”. The biological significance of such a scenario is unclear. One possibility is that a sub-optimal efficiency of tetramerization provides an additional checkpoint for controlling the subunit composition of receptors available for surface expression. Another possibility is that the “LBD barrier” is an evolutionary by-product of the rapid gating kinetics characteristic of AMPARs. Consistent with this idea, the disulfide bond at the LBD dimer-dimer interface, in addition to enhancing tetramerization, also hampers gating efficiency (38). Furthermore, the prokaryotic GluR0, which lacks the ATD, the M4 segment as well as the CTD and is presumably more ancient than mammalian iGluRs, exhibit slow gating kinetics (51). We therefore speculate that the emergence of the ATD, the M4 and the CTD later in evolution conferred additional efficiency to receptor tetramerization, thus allowing the LBD to evolve structural features required for fast gating.

An intact and functional LBD capable of gating motions is essential for the maturation and forward trafficking of full-length AMPARs (52,53). Surprisingly, the HA-GluA1-ΔECD construct, which lacks the LBD and is therefore incapable of agonist-induced gating motions, was expressed to the cell surface instead of being trapped in the ER (Figure 2A, D&E). At least two possible explanations exist for this discrepancy: 1) Certain ER chaperones (e.g. BiP) might bind with this idea, the lurcher mutation, which confers constitutive activity to the ion channel, also affects tetrameric stability (Figure 2B&C; Figure 5). Interestingly, binding site mutations in the kainate receptor GluK2 that abolish its function actually increased the tetramer fraction (50). We therefore tested the tetrameric stability of the receptor using the SDS approach either with or without glutamate in the solubilization buffer (data not shown), but did not observe any difference, possibly due to limited sensitivity of the assay. Still, we cannot rule out that ligand binding and the gating motions it induces presents a barrier during tetramerization, but no longer poses a problem after the tetramer is formed.

Taken together, our observations seem to hint at a “checks and balances” scenario among multiple domains during the biogenesis of a functional AMPAR complex, with the LBD impeding tetramerization and the other domains collaborating to overcome the “LBD barrier”. The biological significance of such a scenario is unclear. One possibility is that a sub-optimal efficiency of tetramerization provides an additional checkpoint for controlling the subunit composition of receptors available for surface expression. Another possibility is that the “LBD barrier” is an evolutionary by-product of the rapid gating kinetics characteristic of AMPARs. Consistent with this idea, the disulfide bond at the LBD dimer-dimer interface, in addition to enhancing tetramerization, also hampers gating efficiency (38). Furthermore, the prokaryotic GluR0, which lacks the ATD, the M4 segment as well as the CTD and is presumably more ancient than mammalian iGluRs, exhibit slow gating kinetics (51). We therefore speculate that the emergence of the ATD, the M4 and the CTD later in evolution conferred additional efficiency to receptor tetramerization, thus allowing the LBD to evolve structural features required for fast gating.

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specifically to the LBD and block forward trafficking until they sense the proper gating motion within the LBD. Thus, the HA-GluA1-ΔECD construct, which lacks the LBD, would be exempt from this ER checkpoint; 2) cytoplasmic trafficking machineries (e.g. the COPII complex) might bind to either the TMD or the CTD and prevent the receptor from exiting the ER until they sense the proper conformational changes induced by LBD gating motions. The HA-GluA1-ΔECD construct might be more structurally flexible and therefore could take on those conformations without the help of the LBD. Again, much work is still needed to pinpoint the exact mechanism. Our findings have yielded novel insights into and provoked intriguing questions about the role of the LBD in AMPAR trafficking.

Based on our observations, we conclude that the TMD of AMPARs mediates its tetrameric assembly. The LBD constitutes a barrier to tetramer formation and stability, while the ATD, the M4 segment and the CTD counteract the “LBD barrier”. This “checks and balances” scenario might have important implications on the regulation of AMPAR subunit composition and forward trafficking. Finally, in the context of neurons, the biogenesis of AMPARs might be modulated by auxiliary subunits and ER chaperone (54,55) as well as other interacting partners.

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FOOTNOTES

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The abbreviations used are: AMPAR, α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptor; ATD, amino-terminal domain; LBD, ligand-binding domain; TMD, transmembrane domain; CTD, carboxy-terminal domain; ECD, extracellular domain; EGFP, enhanced green fluorescent protein; ICC, immunocytochemistry; DDM, n-dodecyl-α-D-maltopyranoside; BN-PAGE, blue native polyacrylamide gel electrophoresis; SDS, sodium dodecyl sulfate; DTT, dithiothreitol; ER, endoplasmic reticulum.

FIGURE LEGENDS

FIGURE 1. Assessing the oligomeric state of GluA1 with or without the ECD

A, Left: crystal structure of the AMPA receptor GluA2 (3KG2) (6). The extracellular domain (ECD), consisting of the amino-terminal domain (ATD) and the ligand-binding domain (LBD), is shaded gray. The dash-line area indicates location and estimated size (~26 Å) of the unresolved carboxy-terminal domain (CTD). Purple shade represents the lipid bilayer, with “in” and “out” indicating the intracellular and extracellular faces.

B, Blue native PAGE (BN-PAGE) of HA-GluA1 and HA-GluA1-ΔECD expressed in HEK 293 cells. Positions of molecular weight markers are labeled on the right (see Experimental Procedures). Oligomeric states of detected bands were estimated by molecular weight (see Table 1) and indicated as “T” and “T’” (tetramers for HA-GluA1 and HA-GluA1-ΔECD, respectively) or “D” (dimer for HA-GluA1).

C, BN-PAGE of HA-GluA1-ΔECD co-expressed with HA-GluA1-ΔECD-EGFP at various DNA ratios. Each construct formed a single tetramer band when expressed alone (leftmost and rightmost lanes). Four incremental upward shifts were observed above the HA-GluA1-ΔECD (open circles) tetramer band, each representing a distinct tetrameric stoichiometry with 1, 2, 3 or 4 EGFP-tagged subunit (green filled circles) incorporated, respectively.

FIGURE 2. Surface trafficking, membrane topology and helical packing of the ECD-lacking GluA1 construct

A, Left: presumed membrane topology of the HA-GluA1-ΔECD construct. An anti-HA antibody (red) and an anti-GluA1-CTD antibody (blue) are used in immunocytochemistry (ICC) to detect the N-terminus and the C-terminus of the construct, respectively. Right: ICC of HEK 293 cells expressing HA-GluA1-ΔECD under non-permeabilizing (left 3 columns) or permeabilizing (right column) conditions.

B & C, BN-PAGE of GluA1 and GluA1(A618T) (B) or HA-GluA1-ΔECD-EGFP and HA-GluA1(A618T)-ΔECD-EGFP (C). Estimated oligomeric states are indicated as “T” (tetramer), “Tri” (trimer), “D” (dimer) and “M” (monomer).

D, Surface biotinylation assay of GluA1 (left) or HA-GluA1-ΔECD (right) (see Experimental Procedures for details). Samples were immunoprecipitated (IP) with strepavidin agarose beads. Proteins of interest
were detected by immunoblot (IB) using either anti-GluA1 or anti-HA antibodies. Fractions are labeled as: W (total whole-cell), E (elution), -Ctrl (biotin-free control) and U (unbound). The percentage of each sample that was loaded onto the SDS-PAGE gel is indicated below. The cytosolic protein GAPDH is used as a loading control.

E, The ratio between the intensities of the “E” and “W” bands (elution/whole-cell) for each protein was used as a quantification of surface expression. “*” indicates p-value < 0.05 when compared to GAPDH in unpaired Student’s t-test.

FIGURE 3. Tetramerization of GluA1 with or without the CTD
A, BN-PAGE of HA-GluA1-ΔECD as well as the HA-GluA1-ΔECD/ΔCTD. For HA-GluA1-ΔECD/ΔCTD, a single band was detected (indicated as “T¨”) with an estimated molecular weight roughly consistent with that of a tetramer (91.6 kDa).
B, BN-PAGE of HA-GluA1-ΔECD/ΔCTD harvested either without SDS or with 0.11% SDS in the solubilization buffer.
C, Homology modeling and MD simulation of an ECD/CTD-free GluA1 tetramer based on 3KG2 (see Experimental Procedures). RMSD between the model and corresponding regions of 3KG2 over time is shown in purple (black dashed line shows averaged RMSD between 50 and 76.7 ns). Initial conformation of the model after energy minimization (point “1” in RMSD trace) was almost identical to 3KG2, while the final conformation after 76.7 ns of MD simulation (point “2” in trace) showed minimal changes (RMSD < 3.5Å).
D & E, BN-PAGE of GluA1 and GluA1-ΔCTD (D) or HA-GluA1-ΔATD and HA-GluA1-ΔATD/ΔCTD (E). Estimated oligomeric states are indicated as in Figure 2.

FIGURE 4. Resistance of GluA1 tetramers to SDS with or without the ECD
A, BN-PAGE of GluA1, HA-GluA1-ΔATD or HA-GluA1-ΔECD-EGFP with various concentrations of SDS added to solubilization buffer (also containing 6 mM DDM).
B, Tetramer fractions of GluA1 (open diamond, black solid line), HA-GluA1-ΔATD (gray triangle, gray dash line) and HA-GluA1-ΔECD-EGFP (black square, black dash line) were normalized to those at 0% SDS (see Experimental Procedures). Normalized fractions are plotted against SDS concentration and fitted with sigmoid functions. The mid-points of the sigmoid functions fitted for the three constructs are approximately 0.11%, 0.08% and 0.11%, respectively.
C, Tetramer fractions of the same three constructs [indicated as in (B)] at various temperatures are normalized to those at 0°C (see Experimental Procedures). Normalized tetramer fractions are plotted against temperature and fitted with sigmoid functions.

Figure 5. Effect of the “lurcher” mutation A618T on tetrameric stability in SDS
A, BN-PAGE of GluA1, GluA1(A618T), HA-GluA1-ΔECD-EGFP or HA-GluA1-ΔECD-EGFP(A618T) with either 0% or 0.11% SDS.
B, Normalized tetramer fractions of the constructs in (A) at 0.11% SDS. “*” indicates p-value < 0.05 in unpaired Student’s t-test.

FIGURE 6. The effect of M4 segment on tetramerization with or without the ECD
A, VLGAVE face (red) of the M4 segment in the GluA2 structure 3KG2 (6). Numbering is for GluA1.
B, BN-PAGE of GluA1 and mutants with tryptophan (W) substitutions in the VLGAVE face (19).
C&D, BN-PAGE of GluA1-ΔATD (C), HA-GluA1-ΔECD or HA-GluA1-ΔECD-EGFP (D) with W substitutions in the VLGAVE face.

FIGURE 7. Contribution of VLGAVE face to tetrameric stability in the absence of the ECD
A, Example BN-PAGE of HA-GluA1-ΔECD-EGFP and mutants with tryptophan (W) substitutions inside (gray) and outside of (black) the VLGAVE face.
B, Normalized tetramer fractions of HA-GluA1-ΔECD-EGFP with tryptophan substitutions relative to the wild type at 0.11% SDS. Dash line indicates normality. “*” indicates p-value < 0.05 in paired Student’s t-test (comparing to wild type assayed on the same gel) while “**” indicates p-value < 0.01.

FIGURE 8. Effect of a disulfide cross-link at the LBD dimer-dimer interface on tetramerization
A, LBD tetramer in the GluA2 structure 3KG2 (6) viewed from the N-terminus. Subunits are colored blue (A/C conformation) and red (B/D conformation). Yellow line indicates the location of the disulfide bond between substituted cysteine A661C (numbering for GluA1). Box below shows a zoomed-in image of the dimer-dimer interface with the disulfide bond.
B, Upper and middle: SDS-PAGE of GluA1, GluA1(A661C), GluA1(G798A) and GluA1(G798A/A661C) under non-reducing (upper) or reducing (middle) conditions. Lower: BN-PAGE of the same constructs.
**TABLES**

Table 1. Calculated molecular weights of constructs used in experiments

Molecular weight of each construct as well as the total molecular weights of the oligomeric complexes formed by that construct (in kDa) are calculated and listed below.

| Constructs                     | Monomer | Dimer   | Trimer  | Tetramer |
|-------------------------------|---------|---------|---------|----------|
| GluA1                         | 99.7 kDa| 199.4 kDa| 299.1 kDa| 398.8 kDa|
| HA-GluA1-ΔATD                 | 58.2 kDa| 116.4 kDa| 174.6 kDa| 232.8 kDa|
| GluA1-ΔCTD                    | 92.0 kDa| 184.0 kDa| 276.0 kDa| 368.0 kDa|
| HA-GluA1-ΔATD/ΔCTD            | 50.5 kDa| 101.0 kDa| 151.5 kDa| 202.0 kDa|
| HA-GluA1-ΔECD                 | 30.7 kDa| 61.4 kDa | 92.1 kDa | 122.8 kDa|
| HA-GluA1-ΔECD-EGFP            | 57.8 kDa| 115.6 kDa| 173.4 kDa| 231.2 kDa|
| HA-GluA1-ΔECD/ΔCTD            | 22.9 kDa| 45.8 kDa | 68.7 kDa | 91.6 kDa |
Figure 1

A

ECD
ATD
LBD
TMD
CTD

Out
In

B

ΔECD
ΔECD-EGFP

or

HA-GluA1-ΔECD
HA-GluA1-ΔECD-EGFP

C

1:0
0:1

ΔECD
ΔECD-EGFP

HA-GluA1-ΔECD
HA-GluA1-ΔECD-EGFP
Figure 2

A

Transfection: GluA1

W E U-Ctrl

2% 50% 50% 2%

IB: GluA1

IB: GAPDH

D

IP: streptavidin

Transfection: HA-GluA1-ΔECD

IB: GAPDH

IB: HA

E

Elution/whole-cell ratio

GAPDH

GluA1

HA-GluA1-ΔECD

ns
Figure 4

A

B

C

GluA1

SDS Concentration (%)

Norm. Tetramer Fraction

Temperature (˚C)

Norm. Tetramer Fraction

GluA1

GluA1

0% 0.05% 0.1% 0.15% 0.2%

0 10 20 30 40
Figure 5

A

B

Figure 5
Figure 6
Figure 7

A

B

Relative Stability

**

**
The transmembrane domain mediates tetramerization of α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) receptors
Quan Gan, Jian Dai, Huan-Xiang Zhou and Lonnie P. Wollmuth

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