Comparative Dynamics of NMDA- and AMPA-Glutamate Receptor N-Terminal Domains

Anindita Dutta, Indira H. Shrivastava, Madhav Sukumaran, Ingo H. Greger, and Ivet Bahar

INTRODUCTION

Ionotropic glutamate receptors (iGluRs) encompass a family of tetrameric glutamate-gated cation channels that mediate the majority of excitatory neurotransmission in the vertebrate central nervous system. Their operation underlies higher-order cognitive functions (Traynelis et al., 2010). Aberrant iGluR signaling is associated with several acute and chronic neurodegenerative diseases. The iGluR family harbors three major subfamilies: α-amino-3-hydroxy-5-methyl-4-isooazolepropionic acid receptors (AMPArs), kainate receptors (KRs), and N-methyl-D-aspartate receptors (NMDArs). Sequence similarity and structural data suggest a shared architecture between these subfamilies: an extracellular domain (ECD), a transmembrane domain (TMD), and an intracellular carboxyl-terminal domain that mediates trafficking and anchorage at synaptic sites (Traynelis et al., 2010; Dingledine, 2010). The ECD consists of a distal N-terminal domain (NTD) and ligand-binding domain (LBD).

The full-length structure of the GluA2 AMPAR (Figure 1A) provided a first view of the modular arrangement of the individual subunits of iGluRs (Sobolevsky et al., 2009). Both the NTDs and LBDs feature the clamshell-like bilobate fold belonging to the periplasmic binding protein (PBP)-like family (Quiocio and Ledvina, 1996). The function of the LBD is well characterized (Armstrong and Gouaux, 2000); L-glutamate docking to the cleft between the upper lobe (UL) and lower lobe (LL) of each subunit results in cleft closure, which is allosterically transmitted to the TMD to trigger channel activation. The function of the NTD remains to be established. In NMDArs, the NTD contributes to signaling as a key allosteric modulator of channel open probability (Gießen et al., 2009). However, the mechanism of NMDAR NTD allosterity is unclear as currently available crystal structures and with without the interlobe Zn<sup>2+</sup> ligand look almost identical (Karakas et al., 2009). NTD allosteric activity in non-NMDArs (i.e., AMPAR and KRs) remains poorly understood and is a matter of debate.

iGluR NTDs are organized in the tetrameric structure as dimeric entities, structurally homologous to metabotropic glutamate receptors (mGluRs) (Kunishima et al., 2000; Muto et al., 2007), and natriuretic peptide receptor (NPR) ligand-binding cores (He et al., 2006), which also operate as dimers. Recent structural data underscore a mechanistic basis for allosteric signaling via NMDAR NTDs, where the LLs are separated and thus free to move in response to ligand binding (Farina et al., 2011; Karakas et al., 2011). This arrangement mimics the ligand binding cores of mGluRs and NPRs but is different from AMPAR and KR NTD dimers that are mostly “zipped-up” across both lobes providing an extensive assembly interface (Kumar et al., 2009; Clayton et al., 2009; Jin et al., 2009; Rossmann et al., 2011), which likely restricts lobe motions. Accordingly, AMPA- and NMDA-R NTDs would have evolved different functions, with the AMPAR NTD mainly directing subunit assembly (Ayalon and Stern-Bach, 2001; Hansen et al., 2010; Rossmann et al., 2011) but with the NMDAR NTD additionally modulating ion channel function. This latter role has led to a surge in NMDAR modulator development, some of which have entered clinical trials (Mony et al., 2009; Karakas et al., 2011). Utilizing a combination of X-ray structural approaches and normal mode analysis (NMA) with elastic network models (ENMs), we showed recently...
that the NTDs of AMPARs do have access to well-defined collective fluctuations (Sukumaran et al., 2011). Together with biophysical measurements of NTD intrinsic fluctuations (Jensen et al., 2011), these recent results exemplify the dynamic capability of non-NMDAR NTDs and suggest a potential allosteric signaling capacity, which would provide a currently unexplored target for channel modulators.

To better understand this discrepancy between iGluR NTD activities, we set out to describe and compare NTD motions between the AMPAR and NMDAR subfamilies at multiple levels. We first solved the crystal structure of the remaining AMPAR NTD, GluA4, and conducted a comprehensive analysis of NTD dynamics across the entire subfamily of AMPARs using both NMA with anisotropic network model (ANM) and full-atomic molecular dynamics (MD) simulations; we also characterized the currently elusive dynamics of the NMDAR NTD, a powerful allosteric modulator. Combined with MD and ANM results, we determine the mechanisms of global and local motions favored by the iGluR NTD architecture, identify critical residues facilitating these motions, and reveal a mechanistic link between LL interface stability and cleft movements, which vary among subfamily members. Finally, upon comparison, we find that cooperative modes of motion intrinsically accessible to AMPAR NTD monomers are almost identical to those of NMDARs, albeit spatially more restricted upon dimerization. Significantly, AMPAR NTDs possess the ability to readily reconfigure into NMDAR NTD conformers. Together, these data reveal an unexpected parallel between AMPAR and NMDAR NTDs and provide a glimpse into the dynamic landscape of iGluR NTDs.

RESULTS

The GluA4 NTD at 2.2 Å

The NTD encompasses the most sequence-diverse part of the receptor, with the four AMPAR paralogs only sharing ~55% sequence similarity. In addition, the packing between NTD dimers is diverse (Rossmann et al., 2011). Hence, we first solved the structure of the remaining GluA4 NTD to facilitate a comprehensive analysis of this domain across the AMPAR subfamily.
**Table 1. Crystallographic Data Collection and Refinement Statistics**

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<td>RMSD from ideal bond angle (°)</td>
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**See also Figure 1.**

<sup>a</sup>Values in parentheses are for the highest resolution shell.

<sup>b</sup>Carbohydrate residues from N-glycosylation.

**Structure**

**Dynamics of iGluR N-Terminal Domains**

Between GluA1–A4 (Figure 1B), contacts across the LL interface of GluA4 are comparable to GluA2 (Protein Databank [PDB] ID code 3HSY), to GluA3 dimer BD (PDB ID code 3P3W; italicized letters indicate chain identifiers from PDB structures, i.e., protomers B and D from ID code 3P3W), and to GluA1 dimer AC (PDB ID code 3SAJ) (Yao et al., 2011). However, in GluA3 dimers CD (PDB ID code 3O21) and AC (PDB ID code 3P3W), the inter-LL distance is wider (Sukumaran et al., 2011) (Figure 1B). Also in GluA1 (dimer BD; PDB ID code 3SAJ), the LLs are packed more loosely partly because of rotations of Leu137 away from the dimer interface. At the equivalent position, GluA4-selective Tyr143 side chains stack across the interface, separated by ~3 Å (Figure S1D). Thus, UL interface contacts are mostly conserved in AMPAR NTDs, whereas LL packing is diverse and subunit selective. The looser LL contacts in GluA1 and GluA3 correlate with reduced NTD dimer stabilities (Rossmann et al., 2011).

To quantify contacts across the structurally variable LL interfaces, we calculated local atomic contact density (LD) indices, a measure for packing density across interfaces (Bahadur et al., 2004; Sukumaran et al., 2011) (Figure 1C). GluA4 exhibits interface characteristics similar to those of GluA2 (UL interface LD: 43.7, LL interface LD: 37.2). GluA1 shows high contact density in the UL (LDs of 40.8 and 44.4 in both dimers AC and BD), characteristic of tight, biologically relevant interfaces (Bahadur et al., 2004). However, the LL interface shows variability; GluA1 dimer AC is similar to those of GluA2 and GluA4, whereas BD is similar to GluA3 (CD; PDB ID code 3O21). Again, these structural features agree with measurements of AMPAR NTD homodimer stabilities by analytical ultracentrifugation (Rossmann et al., 2011), where GluA3 exhibited the weakest affinity (K<sub>d</sub> ~1 μM), followed by the intermediate GluA1 (K<sub>d</sub> ~100 nM), likely reflecting their LL separations and multiplicity of dimeric forms in crystal structures. GluA2 and GluA4 featured K<sub>d</sub> values between 2 and 10 nM, consistent with tighter LL packing (Zhao et al., 2012). Thus, the greatest structural variability between AMPAR NTDs is at the LL dimer interface; looser LL packing is expected to increase NTD interprotomer mobility.

**Comparative ANM Analysis Reveals Global Motions Shared by AMPAR NTDs**

To compare the structural dynamics between AMPAR NTDs and to contrast those to the allosterically active NMDAR NTDs, we first analyzed the collective motions of GluA1–A4 dimers using the ANM (Atiglan et al., 2001; Eyal et al., 2006). ANM is particularly suitable for a comparative assessment of global motions intrinsic to whole protein families (Bahar et al., 2010b). Global motions are those at the lowest frequency end of the mode spectrum, predicted by the ANM to be uniquely defined by the native fold. The lowest frequency mode, mode 1, represents a structural change (usually subunit/domain movements) along the softest/smallest ascent direction away from the original energy minimum (Bahar et al., 2010a).

ANM calculations performed for GluA1–A4 NTD dimers showed that the AMPAR subfamily members share a common mechanism of global reconfiguration along mode 1 with a correlation of 0.90 ± 0.04 (Figure 2; Table S2A): torsional counterrotation of the two protomers, as previously noted for GluA2.
and GluA3 (Sukumaran et al., 2011) (Movie S1), and extended here to the entire subfamily (Figures 2A and 2B). The four subunits exhibit similar global mode shapes (distribution of mobilities, Figure 2A); their motion amplitudes (peak heights), however, show variations, with GluA3 exhibiting higher mobilities (Table S3), particularly in the LL (residues 120–225; inset of Figure 2A). These data, that is, the flexibility between paralogs (Figure 2A), inset), directly correlate with experimental measurements of AMPAR NTD dimer K_d’s (Figure 1C) (Rossmann et al., 2001).

Intrinsic Ability of NMDAR to Sample Open and Closed Cleft Conformations Supported by ANM and MD
NMDAR NTDs allosterically modulate NMDAR ion channel function, triggered by small molecule ligands and Zn2+ ions that bind the dimer interface and cleft region between lobes, respectively. However, current NMDAR NTD structures are similar with regard to cleft-opening angle (Karakash et al., 2009) and proteromer conformation upon ligand binding (Karakash et al., 2011). Thus, the motions underlying NMDAR NTD allostery are unknown.

Our ANM analysis of the global dynamics of the NMDAR NTD, performed for the NR2B subunit, revealed a global twist of the LLs toward the dimer interface accompanied by cleft opening, whereas LL twist motion in the opposite direction induced cleft closure, highlighting the classic clamshell-like motion (Figure 3A; Movie S5). Full-atom molecular dynamic (MD) simulations (Table S4) performed for the same subunit in the apo (NMDA1) and Zn2+-bound (NMDA2) states also revealed an overall rigidification accompanied by cleft-angle closure in the presence of Zn2+ (Figure 3B, pink curve), whereas cleft opening was observed (blue curve) in the absence of Zn2+. The cleft angle was monitored based on the relative positions of L124 (UL), S149 (cleft), and I257 (LL) α-carbons. The apo form thus stabilizes a more open conformation by at least 17° compared to the Zn2+-bound form (Figure 3B, inset). Strikingly, the same type of structural change is predicted by the ANM mode 2 for NR2B (Figure 3A). Thus, both ANM and MD support a classic periplasmic-binding protein mode of ligand recognition for Zn2+ binding.

Similarity between the Intrinsic Dynamics of NMDAR and AMPAR NTD Protomers
Next, we compared global dynamics between AMPAR and NMDAR NTD monomers. Despite their distinctive structural features (Furukawa, 2012), the global modes between the two subfamilies are surprisingly preserved. Figure 4 illustrates the results for AMPAR GluA2 and GluA3 and NMDAR GluN1 and GluN2B subunits. Two dominant modes of motion are observed: counter-rotation between the two lobes (mode 1, panels A and B; Movies S2 and S3) and intralobe clamshell opening/closing (mode 2, panels C and D; Movies S4–S5). The global modes of all AMPAR and NMDAR NTD monomers exhibited a high level of similarity, with correlation coefficients varying in the range 0.73 ± 0.11, highlighting the universality of the observed motions despite stark differences in tertiary and quaternary packing (Table S2B). Especially, the clamshell-like motion seen in NR2B (Figure 3), which enables sampling of ligand unbound/bound conformations, is also preserved in AMPAR monomers.

Although global mode shapes are similar between GluN1, GluN2B, GluA2, and GluA3 monomers (Figures 4A–4C), the relative amplitudes are largest in GluN1 and smallest in GluA2. Table S3 shows an overall ranking of GluN1 > GluA3 > GluN2B > GluA1 > GluA4 (stiffest) based on mode 1 (see Experimental Procedures), and a similar trend is observed in mode 2. This analysis reveals an unexpected difference between GluN1 and GluN2B. Importantly, these modes of motions and intrinsic flexibility are largely conserved between NMDA- and AMPAR-NTDs, as discussed further below.

Effect of Dimeric Packing on the Intrinsic Dynamics of AMPAR and NMDAR NTD Monomers
Because AMPAR- but not NMDAR-NTDs assemble into stable homodimers (Clayton et al., 2009; Jin et al., 2009; Rossmann et al., 2011; Zhao et al., 2012), we next evaluated the changes in dynamics upon NTD dimerization, using a perturbation method (Zheng and Brooks, 2005; Ming and Wall, 2005b), which facilitates assessing the effect of environment on the dynamics of examined systems. Here, each monomer in the dimeric NTD
Figure 3. Intrinsic Ability of NMDAR NTD to Undergo Cleft Motions
(A) Deformation of NR2B subunit (PDB ID code 3JPW, pink) along ANM mode 2, leads to opening of the cleft (blue).
(B) The time evolution of the cleft angle observed in the MD runs NMDA1 (pink, in presence of Zn²⁺) and NMDA2 (blue, in absence of Zn²⁺). The cartoon in the inset is the superposition of 50 ns snapshots from NMDA1 and NMDA2. The histograms in the inset are of the distribution of the angles sampled by NR2B in the two simulations: the average angle is 138° in NMDA1 and 121° in NMDA2.

Structure
Dynamics of iGluR N-Terminal Domains

of AMPAR (GluA2 and GluA3 homodimers) and NMDAR (N1-N2B heterodimer) is taken as the system and is analyzed in the context of the other monomer, which represents its environment. The dynamics in the presence of the environment is then compared to that of the system in isolation (i.e., the intrinsic dynamics of the monomers presented above).

The maps in Figures 5A and 5B display the correlations between the top-ranking 40 ANM modes predicted for the isolated and dimeric forms of the monomers of GluA2 and GluN2B. Highest correlations are indicated by correlation cosines (see Experimental Procedures) close to ±1 (colored red/blue) and lack of correlation by values approaching zero (green). The observed high correlations along the diagonal indicate that the dynamic character of the monomer is maintained in the dimer, with minor alterations (and occasional swaps in the order of mode).

Although the shapes of the global modes are closely maintained, the amplitudes of the motions exhibit a dependence on dimerization. One would expect the amplitudes of fluctuations to be depressed by interprotomer interactions, especially at interface regions. This is the case for GluA2, GluA3, and GluN2B (Figures 5C and 5D; Figure S2C), where the protomer in the dimer exhibits lower mobility compared to the isolated monomer. The insets in Figures 5C and 5D show the ribbon diagrams of the GluA2 and GluN2B monomers, respectively, colored by the difference in mobility between the monomer in the dimeric system and the isolated monomer. The region that shows the largest suppression is χF in the LL followed by the UL interface, whereas UL and LL cores remain unchanged. Dimerization has almost no effect on the mobility of GluN1, that is, interprotomer interactions do not obstruct the conformational flexibility of this NTD (Figure S2D; Table S3). Notably, the suppression of mobility in the χF helix region may have implications on the allosteric capacity of AMPAR NTDs.

NMDAR and AMPAR NTDs Readily Reconfigure along a Single, Global Mode of Motion

The observed difference in the size of global motions between GluN1 and AMPAR NTD protomers are likely due to their differences in dimeric packing. We next determined whether dimer conformations are interconvertible between iGluR subfamilies. If high-energy barriers separate different dimeric forms and preclude structural rearrangement, their distinctive (nonconvertible) interprotomer packing would impact their dynamics, and the known allosteric capacity of the NMDAR NTDs could be attributed to higher conformational freedom. If, however, the different structures are alternative forms readily accessible via soft modes of motions, this would imply that the seemingly less mobile AMPAR NTDs (such as GluA2 and GluA4) can access conformers with known allosteric potential (i.e., NR2B).

To make a quantitative assessment of the ease of transition between different NTD dimers, we examined the overlap (see Experimental Procedures) between (1) structural difference, \( \Delta[R]_{S1 \rightarrow S2} = [R]_{S1} - [R]_{S2} \), that is required for the transition from dimeric conformer “S1” to conformer “S2” (based on PDB coordinates, after optimal superimposition of the endpoints), and (3) the soft modes of structural change favored by S1, as predicted by the ANM. A strikingly easy “conversion” between AMPAR and NMDAR NTD conformers is evidenced by the high overlap between \( \Delta[R]_{S1 \rightarrow S2} \) and mode 1 predicted for S1. Figures 6A and 6B illustrate the results for GluN1 → GluA2 and GluA3 → GluN1-N2B, respectively. The former provides evidence for the ease of transition from NMDAR (GluN1) homodimer to the GluA2 homodimer and the latter from the GluA3 homodimer to the GluN1/GluN2B heterodimer (Figures 6C and 6D; see also Movies S6 and S7). This analysis underscores the significance of global modes in providing access to functional conformers. For example, upon deforming GluA3 NTD along ANM mode 1 alone, the rmsd from the GluN1-N2B heterodimer decreases from 13.06 to 6.12 Å (more than 50%).

Results for other pairs of conformers between AMPAR and NMDA subfamilies are shown in Figure S3. Ninety percent cumulative overlap with the targeted deformation (red curve) is attainable with a small subset (e.g., 20–25) of soft modes (out of a total of ~1,800 ANM modes) in all cases, except for the GluA2 → GluN1 homodimer, in which the overlap is ~70%. Together with the ANM data (Figures 2, 3, 4, and 5), these results underscore an unexpected parallel between AMPAR and NMDAR NTD flexibility and dynamics.
All-Atom MD Simulations Indicate High Intra- and Inter-LL Mobilities in AMPAR NTDs

To obtain a better understanding of the molecular interactions that underlie iGluR NTD dynamics, we conducted all-atom MD simulations. Root-mean-square fluctuations (rmsfs) in residue positions (Figure S4) confirm that GluA3 exhibits the highest mobility among all AMPAR NTDs. This enhanced mobility is primarily mediated by LL helices and Inter-LL Mobilities in AMPAR NTDs.

By conducting all-atom MD simulations, we aimed to explore the dynamics of the AMPAR NTDs, particularly focusing on GluA3. The simulations revealed that GluA3 exhibits the highest mobility compared to other AMPAR NTDs. This behavior is due to the unique positive charge distribution in the GluA3 LL interface, which projects into the interface, destabilizing the dimer (Sukumaran et al., 2011). GluA3 also exhibits localized reorientations of the LL as a key structural determinant mediating intrinsic dynamics.

GluA3 NTD Protomers Undergo Clamshell-like Motions

To determine the influence of interface stability on classic PBP-like clamshell motions (Quiocho and Ledvina, 1996; Trakhanov et al., 2005), we examined the fluctuations in the interlobe cleft angle. We observe markedly larger angular fluctuations in GluA3 than in GluA2 (Figures 7D and 7E); GluA3 featured the widest opening of the interlobe cleft angle (ranging up to >130° in protein A). Interestingly, the two GluA3 protomers appear to undergo anticorrelated fluctuations, with protomer A closing and B opening with a periodicity of ~25 ns (Figure 7D). This motion is unique to GluA3 and not discernable in other AMPAR paralogs. Specifically, hydrophobic packing is disrupted as phenylalanine pairs (F56, F88) are separated and in some cases, irreversibly broken (Figure S5). This “acquired” UL instability further points to the unique behavior of the GluA3 NTD, potential coupling between the LL and UL interfaces in AMPAR NTDs, and the importance of the LL as a key structural determinant mediating intrinsic dynamics.

Effect of LL Residues on Interlobe Packing and Dynamics

The difference in GluA2 and GluA3 interface stability and residue fluctuations observed in MD simulations is also reflected in their dimer stabilities derived experimentally (Rossmann et al., 2011). In GluA2, hydrophobic residues contribute to the LL-LL contacts, whereas in GluA3 pairs of arginines (R163 and R184) project into the interface, destabilizing the dimer (Sukumaran et al., 2011). To gain further insight into the relationship between the interface stability and cleft dynamics, we analyzed...
two mutants generated in silico: L144D (GluA2), to destabilize the GluA2 interface via like-charge repulsion, and R163I (GluA3), to strengthen the labile GluA3 interface via hydrophobic contacts.

Introduction of like-charges into the GluA2 LL interface indeed led to destabilization as can be seen from the comparison of the LL-LL distances for the mutant L144D (teal curve in Figure 8A) and for the wild-type GluA2 (black curve). Notably, the extent of destabilization is comparable to that originally observed for GluA3: interlobe distance between the two substituted amino acids increases to more than 30 Å within tens of nanoseconds, whereas in wt GluA2, the equivalent interaction is maintained over a period of 100 ns. Conversely, introducing hydrophobic residues into the GluA3 LLs leads to a more stabilized interface (Figure 8C): the distance between the mutated R163I residues is maintained but is disrupted in wt GluA3 dimer at early stages of the simulation. Snapshot at t = 100 ns illustrates the disruption of the LL packing interface upon L144D mutation in GluA2 (Figure 8B) and strengthening in R163I (Figure 8D). These results thus demonstrate the stabilizing role of hydrophobic residues at the packing interface in GluA2, as well as the destabilizing role of buried arginines in GluA3. CM distances between the LLs (Figure S6A) further establish that the GluA2-L144D mutant weakens dimer contacts, whereas GluA3-R163I is stabilizing.

Analyses of the trajectories generated for the mutants show that LL stability is coupled to UL-LL dynamics: a salt bridge connecting the lobes of the clamshell (D98–K112; Figure S6B, expected to restrain cleft motions, is destabilized in the GluA2 mutant, whereas the R163I mutation stabilizes the equivalent salt bridge in GluA3 (D104–R141; Figure S6C). Moreover, stabilization of the LL in GluA3-R163I restricts clamshell motions as compared to the wt GluA3 dimer. Therefore, alterations of LL interface strength can propagate to the hinge region in both cases and has the capacity to alter lobe motions, that is, the perturbation of LL stability in both GluA2 and GluA3 has bidirectional effects that extend beyond local (LL) interactions.

**DISCUSSION**

In this study, we provide a series of insights into the dynamics of AMPA- and NMDA-receptor NTDs. First, we present the crystal structure of the GluA4 NTD, facilitating a comprehensive analysis of this sequence-diverse domain across the AMPAR subfamily. Second, we provide mechanistic insights into the intrinsic dynamics of GluN2B that facilitate ligand binding and offer a first glance into the motions driving GluN2B NTD allostery, whose modus operandi has not been elucidated to date. Third, we reveal that AMPAR- and NMDAR-NTD monomers share surprisingly similar global mode motions. These are restricted, but not abolished, upon dimerization in a subunit-dependent fashion, dictated by the evolutionary and structurally variable LL interface. Fourth, we show that AMPAR NTDs can readily reconfigure into NMDAR NTD conformers, a further indication of their unexpected similarity and their putative allosteric capacity. Fifth, we evaluate the dynamics of the AMPAR subfamily at atomic resolution, where, in accordance with experimental data, we find that GluA3 features the weakest LL dimer interface, which ruptures after ~5 ns of MD simulations, followed by GluA3 LL secondary structure elements (αF) flipping downward to the LBD. The unique LL packing of GluA3 also potentiates it to undergo classic PBP-like clamshell motions. Finally, we capture critical residues at the LL-LL interface that mediate interprotomer dynamics in AMPARs, consolidated by analyses of mutants designed to weaken or strengthen the LL interface.
The AMPAR NTDs studied here are stable homodimers in solution, with a highly conserved UL interface, which will maintain dimer stability. The LL, on the other hand, which potentially shares an interface with the LBD, may play a mediatory role in the allosteric regulation, also demonstrated by the recent NMDAR shares an interface with the LBD, may play a mediatory role in the maintain dimer stability. The LL, on the other hand, which potentially solution, with a highly conserved UL interface, which will main-

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Figure 6. Ease of Transition between Dimeric Conformers of NMDAR and AMPAR NTDs (A) Results are illustrated for the passage from GluN1 (NMDA) homodimeric conformer to GluA2 dimer conformer. The overlap (blue bars) repre-
sents the correlation cosine (see Experimental Procedures) for each of the top-ranking 80 ANM modes to the conformational change. The red curve represents the cumulative overlap, adding up the contribution of all modes starting from the low frequency end (mode 1). The dashed green curve displays the control, for random modes. The slowest mode predicted for GluN1 (PDB ID code 3Q41-A) yields an overlap of ~80%, indicating a strong predisposition of the GluN1 homodimer to assume the conformation of the GluA2 dimer.

(B) Two transitional end points (orange, yellow) and an intermediate structure reached by moving exclusively along mode 1 (green).

(C and D) Same as (A) and (B), for the change in the conformation of GluA3 homodimer (yellow) toward that of the heterodimer GluN1/GluN2B (PDB ID code 3QEL-CD, orange) along GluA3 ANM inter-

mediates (PDB ID code 3KG2) along with a deletion encompassing six residues (Sobolevsky et al., 2009). How this mutation affects domain packing and allosteric communication in iGluRs is a key open question.

Our analysis reveals the ability of individual protomers to undergo concerted clamshell opening/closing motions, which simultaneously affect interprotomer contacts. This supports a possible cooperative response of AMPAR NTDs upon ligand binding or interaction with protein partners (O’Brien et al., 1999) and a capacity to transmit signals toward the channel. We note that ligand interaction may not be restricted to the interlobe cleft but could target the LL dimer interface, as known from analogous cases (He et al., 2006; Mony et al., 2011), or the highly dynamic αH region.

The NTD of the NMDAR is known to modulate channel gating by binding Zn2+ ions and ifenprodil-like compounds, thereby sparking clinical interest in these domains. The twisted LL along with surface properties (Karakas et al., 2009; Stroebel et al., 2011) have been purported to be the primary reason why NTD-mediated modulation of the ion channel is seen in GluN2B and NMDAR heterodimers but have not been seen so far in non-NMDA recep-
tors. Structural dynamics analysis offers a different perspective, where the global motions accessible to the different NTD structures of the iGluR families overlap remarkably. Also, the global modes of dimeric AMPAR, NMDAR, and mGluR NTDs allow for facile transitions from one form to another, suggesting that the AMPAR NTDs may equally have allosteric signaling abilities.

Binding of ions and small molecules to the NTD are most likely facilitated by global motions in NMDARs. NTD clamshell motions have been implicated in facilitating an induced-fit binding mechanism (Karakas et al., 2011). Based on similarity of global motions
between AMPARs and NMDARs, the allosteric effect known to modulate NMDAR open probability should not be disregarded for non-NMDARs. This view is further strengthened by the small-molecule binding capacity in the GluA2 cleft reported previously (Sukumaran et al., 2011; Cameron et al., 2012) and the labile nature of the GluA3 NTDs observed here. The present analysis is a further step toward clarifying the putativeallosteric potential of AMPAR NTDs by highlighting their intrinsic ability to undergo motions comparable to NMDAR NTDs and their propensity to sample conformers observed in NMDARs. This opens an avenue of searching for molecules able to bind AMPAR NTDs, which may in turn play an important role in regulating gating of nonNMDA iGluRs.

EXPERIMENTAL PROCEDURES

Protein Crystallography

GluA4 NTD constructs (Ala1–Asp380; all residue numbers correspond to the sequence of the mature protein, after signal peptide cleavage) were designed, expressed, and purified as described previously (Rossmann et al., 2011). Crystalization was performed using the vapor-diffusion method (Benvenuti and Mangani, 2007). GluA4 NTD readily crystallized in 0.1 M sodium cacodylate and 0.7–1.0 M citrate (pH 5.0–7.0). Diffraction data were collected from beamline I03 at the Diamond Light Source (Oxford, UK). Data were processed using the iMOSFLM package (Battye et al., 2011). The structure was solved by molecular replacement with PHASER (McCoy et al., 2007), using a GluA2 NTD monomer (PDB ID code 3HSY, chain B) as a search probe. The model was alternately refined using REFMAC (Murshudov et al., 2011) and manually rebuilt in COOT (Emsley et al., 2010). MOLPROBITY (Davis et al., 2007) was used to validate model stereochemistry.

MD Simulations

The GROMACS program (Van Der Spoel et al., 2005) was used to generate MD trajectories for the systems listed in Table S4. The proteins were solvated with single point charge (Berendsen et al., 1981) water molecules, using GROMOS 43a1 force field (Lindahl et al., 2001). MD runs were performed at 310 K (by implementing Berendsen’s temperature coupling to protein and water molecules) and atmospheric pressure. Electroneutrality was achieved by adding counterions. Electrostatic interactions were treated with the particle mesh Ewald method (Darden and Pedersen, 1993) and the LINCS (Hess et al., 1997) algorithm was used to constrain the bond lengths, enabling an integration timestep of 2 fs. Each system was energy minimized using the steepest descent algorithm, followed by an equilibration of 2 ns, before the productive runs of 100 ns. During equilibration, backbone atoms were restrained by harmonic potentials, while side-chain atoms and water molecules were allowed to relax. Figure S7 displays the rmsds from the initial state averaged over all residues as a function of time for the four AMPAR NTD subtypes.

ANM Analysis of Collective Motions

NMA with ANM was performed as previously described (Atilgan et al., 2001; Eyal et al., 2006). In the ANM, the overall potential ($V_{\text{ANM}}$) is represented as the sum of harmonic potentials between interacting nodes. Those nodes within a certain cutoff distance (15 Å) are assumed to interact. The force constants (the second derivatives of the potential $V_{\text{ANM}}$) for the 3N×3N interactions (for N residues in 3D) are given by the elements of the Hessian matrix $H$ (a summation over interresidue pairwise potentials with uniform force constants). The method yields a unique set of collective modes for each
protein, represented by 3N-dimensional eigenvectors $u^m_k (1 \leq k \leq 3N-6)$ obtained by decomposing the Hessian matrix H. The eigenvalues $\lambda_k$ of H scale with the square frequencies and $\lambda_k^{-1/2}$ defines the weight of mode k such that the conformational change due to mode k scales with $\lambda_k^{-1/2} u^m_k$; the square displacements (Figures 2, 4, and 5) scale with the $\lambda_k^{-1}$.

**Overlap between Experimentally Observed Structural Changes and Soft Modes**

Consider two known structures S1 and S2 for the protein (family) of interest, represented each by conformational vectors $\{R_0\}_{S1}$ and $\{R_0\}_{S2}$. Their structural difference (after optimal alignment using STRAP; Gille and Frohme, 2001) is $\Delta(R_{0,S1} - R_{0,S2})$. To observe if S1 is predisposed to undergo this change, we evaluate the overlap or correlation cosine, $(d\cdot u^m)$ for a subset of soft modes $u^m_k$ accessible to S1. Here, $d$ is the unit vector along $\Delta(R_{0,S1} - R_{0,S2})$. The cumulative overlap achieved by a subset of $m$ modes is given by $CO(m) = \vert \sum_k (d\cdot u^m_k) \vert^{1/2}$, where the summation is performed over $1 \leq k \leq m$. The complete set of eigenvectors forms an orthonormal basis, that is, $CO(m) = 1$ for $m = 3N-6$.

**NMA of a Subsystem Coupled to an Environment**

The dynamics of a system (S) in the context of an environment (E) is evaluated by partitioning H into four submatrices (Zheng and Brooks, 2005; Ming and Wall, 2005a)

$$H = \begin{pmatrix} H_{SS} & H_{SE} \\ H_{ES} & H_{EE} \end{pmatrix},$$

where $H_{SS}$ refers to interactions within the system, $H_{SE}$ to those within the environment, and $H_{ES}$ (or $H_{SE}$) to the coupling between S and E. The resulting effective Hessian of the system is in the presence of the environment

$$H_{SS}^{SE} = H_{SS} - H_{SE}H_{EE}^{-1}H_{ES}.$$  

(A) and (B) in Figures 5 and 2 are obtained by evaluating the correlation cosine $u^m_k$ between the eigenvectors $u^m_k$ and $u^m_{il}$ for $i = 1-40$ corresponding to $H_{SS}$ and $H_{SS}^{SE}$, respectively.

**ACCESSION NUMBERS**

The Protein Data Bank accession number for the GluA4 structure reported in this paper is 4GPA.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes seven figures, four tables, and eight movies and can be found with this article online at http://dx.doi.org/10.1016/j.str.2012.08.012.

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