

1 **Non-selective cation permeation in an AMPA-type** 2 **glutamate receptor**

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9 **Abstract**

10 Fast excitatory synaptic transmission in the central nervous system relies on the AMPA-type glutamate
11 receptor (AMPA). This receptor incorporates a non-selective cation channel which is opened by the
12 binding of glutamate. Although the open pore structure has recently become available from cryo-electron
13 microscopy (Cryo-EM), the molecular mechanisms governing cation permeability in AMPA receptors are
14 not understood. Here, we combined microsecond molecular dynamics (MD) simulations on a putative
15 open state structure of GluA2 with electrophysiology on cloned channels to elucidate ion permeation
16 mechanisms. Na⁺, K⁺ and Cs⁺ permeated at physiological rates, consistent with a structure that
17 represents a true open state. A single major ion binding site for Na⁺ and K⁺ in the pore represents the
18 simplest selectivity filter (SF) structure for any tetrameric cation channel of known structure. The
19 minimal SF comprised only Q586 and Q587, and other residues on the cytoplasmic side formed a cone-
20 shaped void that lacked major interactions with ions. We observed Cl⁻ invasion of the upper pore,
21 explaining anion permeation in the edited form of GluA2. A permissive architecture of the SF
22 accommodated different alkali metals in distinct solvation states to allow rapid, non-selective cation
23 permeation, and co-permeation by water. Simulations suggested Cs⁺ uses two equally populated ion
24 binding sites in the filter and we confirmed with electrophysiology of GluA2 that Cs⁺ is more permeant
25 than Na⁺, consistent with serial binding sites preferentially driving selectivity.

26 **Significance Statement**

27 AMPA-type glutamate receptors (AMPA) are key actors in neurotransmission, making the final step in
28 a relay of excitability from one brain cell to another. The receptor contains an integral ion channel,
29 which, when opened by neurotransmitter binding, permits sodium and other cations to cross the cell
30 membrane. We investigated permeation of sodium, potassium and caesium in an AMPAR at the atomistic
31 level using a computational molecular dynamics approach on a structure with the ion channel pore in a
32 presumably open state. We determined that the region selecting between cations is the simplest of any
33 channel of this type. Distinct from ion channels that select single ion species, cations are never fully
34 dehydrated and have only one major ion binding site in the filter. Simulations suggested two similar
35 binding sites for caesium, and studies of AMPARs in mammalian cell membranes showed that this makes
36 caesium more permeant than sodium.

37 **Introduction**

38 Glutamate receptor ion channels are found at synapses throughout the vertebrate nervous
39 system, where they convert sub-millisecond glutamate signals into cation currents. Advances
40 in structural biology have provided molecular scale maps of their ion pores, permitting
41 comparison with a burgeoning menagerie of structures from related ion channels. It has been
42 comparatively difficult to obtain candidate open pore structures of glutamate receptors, with
43 the notable exceptions being from single particle cryo-electron microscopy (Cryo-EM) of
44 complexes between GluA2 and Stargazin (1, 2). However, it is unclear from simple inspection
45 of these structures whether a) the ion pore is conductive, or b) it is open to its fullest extent,
46 or to the highest conductance level. However, a substantial body of electrophysiological work
47 provides good benchmarks for how these structures should behave. For example, canonical
48 measurements of channel conductance and reversal potentials show that alkali earth cations
49 from sodium (Na^+) up to caesium (Cs^+) should permeate GluA1 (3) and unedited GluA2 (4)
50 approximately equally well and the single channel conductance of the full level of GluA2 (Q)
51 should be considerable (~ 30 pS, ref. 5).

52 Most computational work on ion permeation through channels has been focused on simple,
53 selective potassium (K^+) channels like KcsA (6, 7), being the first reported crystal structures of
54 ion channels (8). Thanks to their minimal sequences, these channels demand little
55 computational overhead. Their key structural features are two membrane spanning helices and
56 a reentrant loop forming a narrow selectivity filter (SF) for permeant ions. This core motif
57 defines a superfamily of tetrameric and pseudotetrameric channels that encompass selective,
58 semi-selective and non-selective cation conductances. In common with many eukaryotic
59 channels, the AMPA type glutamate receptor has a pore domain whose gating state is controlled
60 by substantially larger domains outside the membrane (amino terminal domain, ATD and
61 ligand binding domain, LBD, Figure 1A) which account for about 75% of the protein mass. The
62 large size presents a challenge for conventional molecular dynamics (MD) simulations, with
63 the AMPA receptor being about 6 times bigger than KcsA.

64 It remains to be seen to what extent key features of ion permeation elucidated in prokaryotic
65 channels (selectivity, discrete sites, desolvated ions and block by divalent ions) are widely
66 applicable in channels with more substantial architectures. Recent experimental and MD work
67 on non-selective prokaryotic channels like NaK and NaK-CNG suggests that ion permeation in
68 these channels differs substantially from classical K^+ selective ion permeation (9–11). The SF
69 of non-selective cation channels is much more flexible, with fewer ion binding sites leading to
70 distinct conduction mechanisms and hydration states for Na^+ and K^+ when passing through the

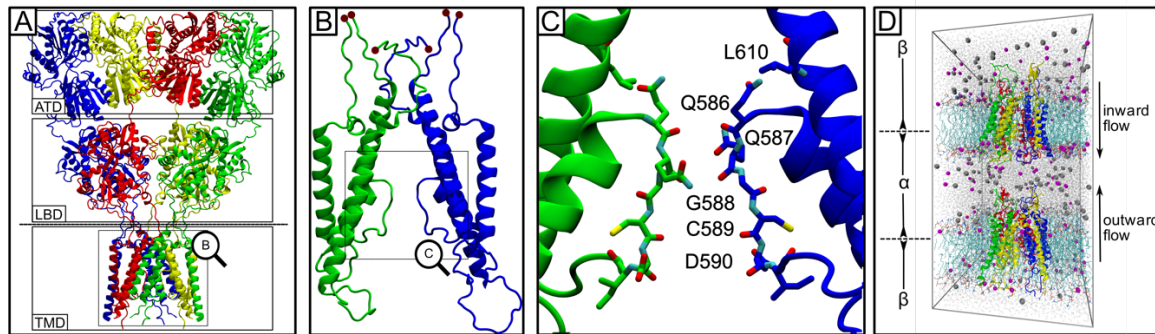


Figure 1. AMPA receptor and the simulation setup (A) The activated open state of the AMPA receptor from Cryo-EM (PDB ID: 5WEO) (1) with Stargazin molecules removed. The receptor is composed of amino terminal, ligand binding and transmembrane domains (ATD, LBD and TMD). (B) The TMD and linkers to the LBD layer were included the MD simulations. The sites where the linkers were truncated and physically restrained (see Methods) are marked with red balls. Two out of the four subunits are shown. (C) The selectivity filter region of the AMPA receptor pore, with key residues labelled. Again, only two diagonally-opposed subunits are drawn. (D) The computational electrophysiology setup was composed of two tetrameric AMPA channels, each embedded in a separate POPC lipid bilayer, solvated with water molecules and ions. A small cation imbalance between the two compartments α and β was maintained during the simulations. The resulting gradient gave a transmembrane potential to drive ion permeation.

71 filter. In the AMPA receptor, in one activated open Cryo-EM structure of GluA2 (1), density for
72 a presumptive hydrated sodium ion was observed adjacent to the unedited Q586 residue. In
73 the structure of edited GluA2 (Q586R) with Stargazin (2), ions were absent. The closed state
74 structure of a GluA1:A2 heteromer featured strong density of unknown identity adjacent to
75 C589 (12). Whether the paucity and heterogeneity of resolved ions is due to a lack of order in
76 the filter region, the lack of detail in the coulomb potential density or a true deficit of ions
77 remains unclear. However, observations in these AMPA receptor structures are in marked
78 contrast to the Cryo-EM structures of several other non-selective cation channels such as
79 hyperpolarization-activated cyclic nucleotide-gated (HCN) (13) and cyclic nucleotide-gated
80 (CNG) channels (14) where two or three bound ions were visible. Further context comes from
81 canonical K^+ -selective channels which feature up to 4 ions in a row (15) and crystal structures
82 of Ca^{2+} -selective TRP channels where two Ca^{2+} ions were readily resolved (16).

83 Here we used MD-based computational electrophysiology to examine ion permeation through
84 the mammalian AMPA receptor ion channel. We determined that the published structure is
85 stably open, suitably detailed for MD simulations and probably represent a native fully-open
86 state. We identified a minimal selectivity filter consisting of a single ion binding site that does
87 not dehydrate ions. In simulations, Cs^+ co-opted a secondary binding site. Consistent with
88 multiple sites promoting ion selectivity, electrophysiology of AMPARs in HEK cells showed that
89 Cs^+ is more permeant than Na^+ .

90 Results

91 *Monovalent cation permeation*

92 Our starting structure for MD simulations was the AMPAR channel in its activated conformation
93 (PDB ID: 5WEO) (1), which we embedded into a palmitoylcholine (POPC)
94 lipid bilayer (Figure 1D). To reduce the number of atoms in the simulations, we removed the
95 extracellular ATD and LBD and the four Stargazin molecules decorating the periphery of the
96 channel. To retain the open channel conformation, we held the transmembrane domain (TMD)
97 of AMPAR open by physically restraining the end of the truncated linkers (Figure 1B, see
98 Methods for details). This approach, while facile, was highly reproducible, and the channel did
99 not close spontaneously during any simulation run (Figure 2A). Simulations of ion conduction
100 were performed with the computational electrophysiology method (17, 18). In this setup, the
101 simulation box contains two membranes that define two compartments and a voltage
102 difference across each membrane is created by an ion gradient that is maintained alchemically
103 during the simulations (Figure 1D).

104 Classical electrophysiology experiments suggest AMPARs are broadly non-selective over
105 different alkali metal cations, although most measurements were not made on GluA2 (3, 19).
106 We began by performing a series of ion permeation simulations with the principal biological
107 ion species, Na⁺ and K⁺. To ensure the robustness of our observations, we undertook
108 simulations at different temperatures, with different force fields and different driving forces on
109 the ions (see Discussion and Methods). The conductances of the AMPA channel pore were
110 directly calculated from the ion permeations observed in these simulations. We counted cations
111 that traversed the entire narrow region of the pore domain delineated by the ⁵⁸⁶QQGCDI⁵⁹¹
112 sequence, summarized in Table 1 and Figure 2B. Outside of this region, ion movements lacked
113 order and made only rare interactions with the channel leading to a largely flat energy
114 landscape (Figure S1). In all trajectories, the channel pore remained conductive for Na⁺ or K⁺
115 during the entire 500 ns run, as exemplified by the ion track plots in Figure 2A. In a typical
116 500 ns trajectory with a transmembrane potential of 450 mV, we recorded over 40 K⁺
117 permeation events, corresponding to a conductance of 32 pS.

118 Although K⁺ permeated readily in the simulations with AMBER99sb force field (20), we observed
119 substantially fewer permeations of Na⁺ under the same simulation condition. The simulated
120 conductances of both K⁺ and Na⁺ were higher in the CHARMM36 force field (21), allowing direct
121 observation of Na⁺ permeation at close to physiological rates (Table 1, Figure 2B). However,
122 the permeation of K⁺ in the CHARMM36 forcefield approaches implausibly high rates, albeit

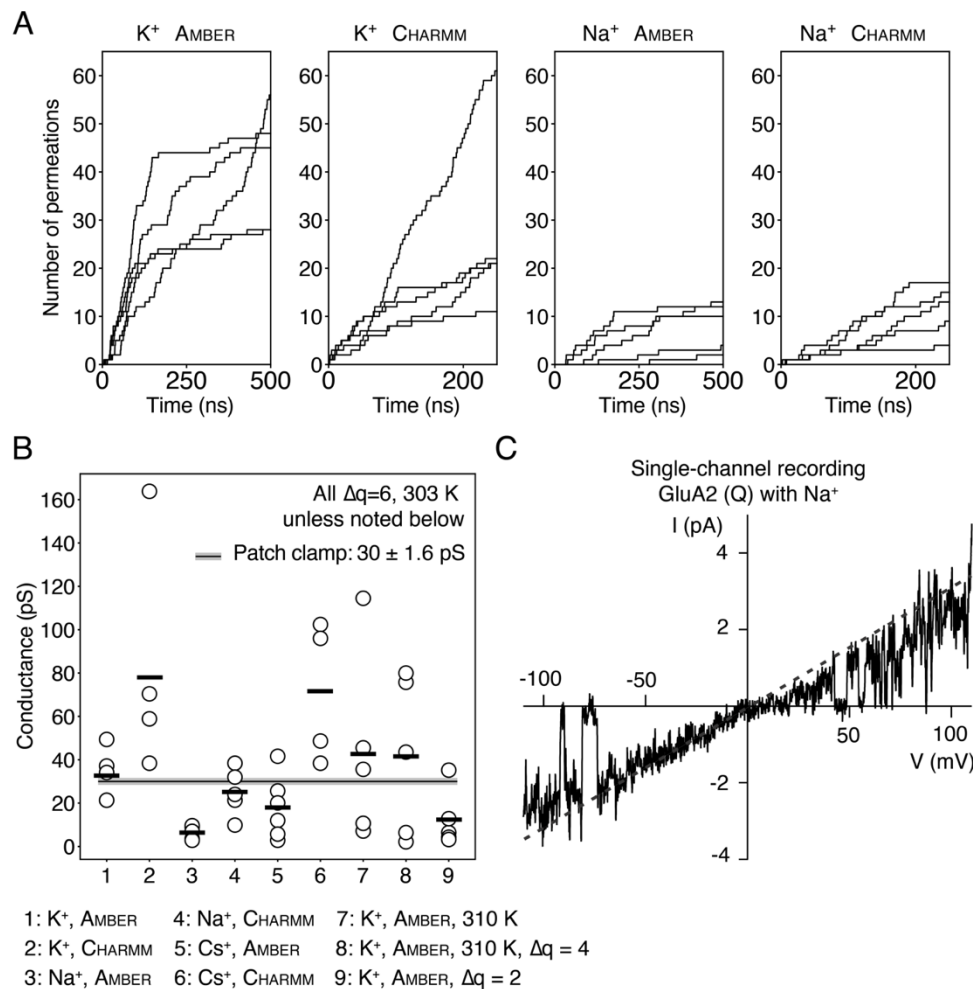


Figure 2. Ion conductances in GluA2 (A) Cumulative ion permeation events in the GluA2 pore for individual trajectories of K⁺ and Na⁺ simulations with the respective force fields. The simulations were performed at 303 K using an ion imbalance of 6e⁻ between two compartments α and β . (B) Conductances derived from the K⁺, Na⁺ and Cs⁺ simulations performed with AMBER99sb and CHARMM36 force fields. Simulations were carried out at different temperature and with different ion imbalance between two compartments α and β . The experimental value derived from single-channel recordings for Na⁺ is displayed as a grey line. (C) Outside-out patch clamp electrophysiology of a single GluA2 (Q) channel in symmetrical Na⁺. The voltage ramp was made in the presence of 10 mM glutamate and 100 μ M CTZ. The dashed line shows the fitted chord conductance, which for this patch was 33 pS.

123 with a very large variability. Higher conductance using CHARMM36 compared to AMBER99sb
 124 was previously reported in simulations of K⁺ selective and non-selective NaK-CNG channels (9).

125 *Binding sites for permeant ions*

126 Selectivity filter architectures differ widely amongst tetrameric ion channels. Whilst K⁺
 127 channels generally have long, straight filters, non-selective cation channels tend to have shorter
 128 filters that display more plastic or eccentric geometries (22, 23). To understand how the
 129 AMPAR selectivity filter supports cation-selective ion conduction, we determined one- and two-
 130 dimensional ion occupancies in the AMPA channel, which were sampled from cumulative

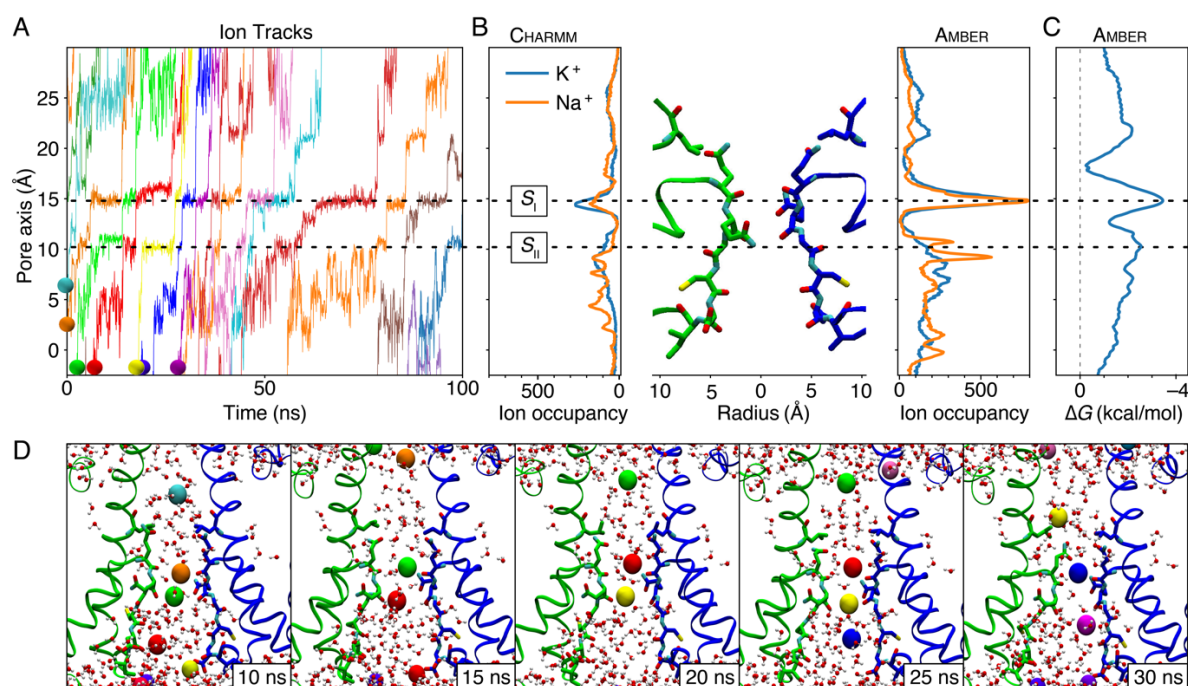


Figure 3. Simulated Na⁺ and K⁺ permeation of GluA2. Simulations were performed at 303 K using an ion imbalance of 6 e⁻ between two compartments α and β . (A) Representative traces of K⁺ passing through the SF of AMPA channel pore. Ions drawn in panel D are indicated with balls. (B) One-dimensional ion occupancy of cations within the SF of the AMPA channel pore. Left panel: occupancy from simulations with the CHARMM36 force field, right panel from AMBER99sb force field. (C) Free-energy profile for K⁺ permeation constructed from AMBER99sb force field. (D) Representative snapshots of a simulation (from Movie S1) showing multiple K⁺ ions passing the selectivity filter during ion conduction. The color code of ions is identical with the ion track plot in panel A.

131 simulations of K⁺ and Na⁺ with the respective force fields (Figure 3B, 4A and S2). For each ion
 132 species, these plots reveal only one principal binding site (which we call S_I) in the SF, which is
 133 located around 15 Å along the pore axis (Figure 3). At this site, the primary coordination of
 134 permeant ions is by backbone carbonyls of Q586 that replace some but not all of waters of
 135 hydration (Figure S3). Secondary, less frequent interactions occur with the mobile side chain
 136 of Q586 and the backbone carbonyls of Q587. In some trajectories using AMBER99sb force field,
 137 solvated Na⁺ ions stay at the S_I longer than 100 ns (Figure 4B), which may explain the slower
 138 permeation. During outward permeation, ions leaving the S_I site frequently pause above the tip
 139 of the SF (25 Å at the pore axis in Figure 4A) and then rapidly enter bulk water as they traverse
 140 the upper reaches of the channel. Before reaching the major binding site S_I, multiple ions can
 141 simultaneously enter the inner mouth of the region, classically expected to be part of the SF
 142 (2, 24), and composed by the residues G588, C589 and D590. K⁺ and Na⁺ paused frequently
 143 but briefly at a secondary site (which we denote S_{II}) which is positioned around 10 Å into the
 144 pore, cytoplasmic to the S_I site (Figure 3B). Compared to the primary site, the S_{II} site was
 145 weakly populated with shorter residence times (Figure 4B), Therefore we conclude that the

146 true SF region of the AMPA is the shortest seen in any tetrameric channel to date, comprising
147 essentially only Q586 and Q587 for accommodating S_I and S_{II} sites.

148 *Cation conduction mechanism and selectivity*

149 During the ion conduction, most of the time we observed multiple ions reside simultaneously
150 in the canonical SF region of $^{586}\text{QQGCDI}^{591}$ (Figure 3D, Movie S1 and S2), with an overall time-
151 average number of 2.4 ± 0.6 and 2.3 ± 0.7 for K^+ and Na^+ respectively in the AMBER simulation
152 runs (Figure S4). Ions followed a loosely coupled knock-on mechanism, where in most cases
153 the exit of an ion to the upper cavity was closely followed by the entry of an ion to the S_I site
154 (Figure 3D, Movie S1 and S2). From the free-energy profiles (Figure 3C), we conclude that the
155 energy barriers bracketing S_I (at around 13 Å and 18 Å along the pore axis) are the major
156 obstacles for ion conduction in the SF. The ion entering the S_I site was normally supplied from
157 the adjacent S_{II} site, presumably aiding permeant ions to overcome the main energy barriers
158 before and after the S_I site (6).

159 To understand better where cation selection occurs, we examined the paths of chloride (Cl^-)
160 ions during our simulations. Consistent with a very short selective region, Cl^- ions occasionally
161 entered the cone-shaped void below the SF and readily approached the QQ-filter from the
162 bundle crossing side. Surprisingly, Cl^- ions found stable binding sites immediately above the
163 QQ-filter (Figure S5) in all simulations. We also observed one Cl^- ion permeation event (Movie
164 S3), providing a crude estimate of the $\text{Cl}^-:\text{K}^+$ permeability ratio at around 1:220. In practical
165 terms, this relative selectivity would correspond to an approximately 15 μV shift in the reversal
166 potential (GHK equation, see Methods), compared to no Cl^- permeation at all. In other words,
167 at this level of selection between cations and chloride, any reversal potential shifts are too small
168 to be measured experimentally. Further context comes from the observation that R-edited
169 homomeric GluA2 channels are anion permeable (25). The observation of infrequent yet stable
170 anion approach to the face of the QQ-filter favoured for anions by the electric field provides a
171 simple explanation as to how this point mutation can switch selectivity simply by changing
172 local electrostatics.

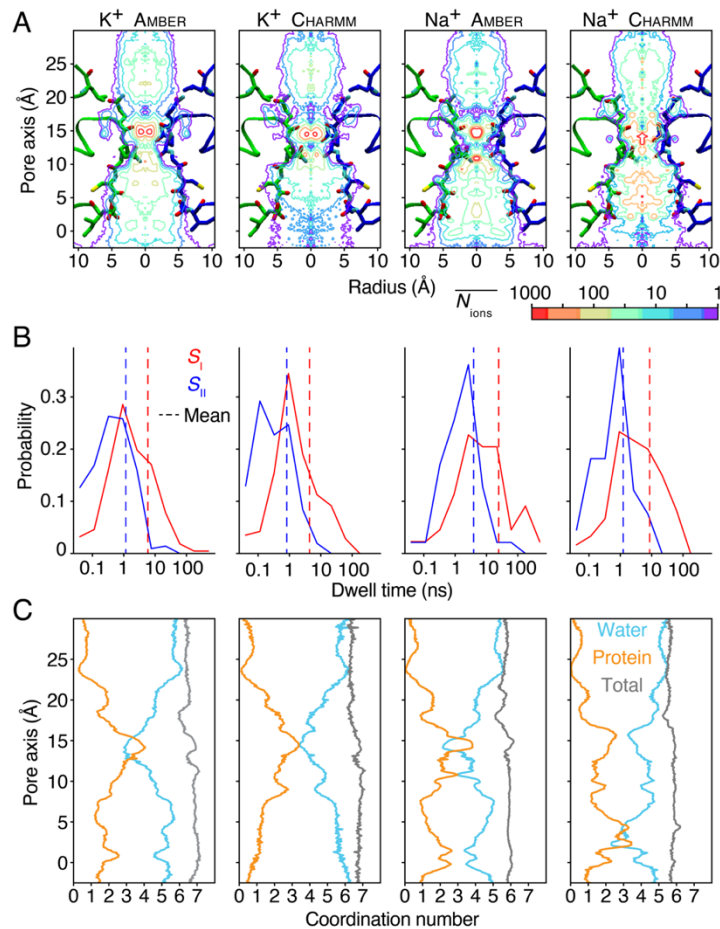


Figure 4. Ion occupancy, dwell time distribution and hydration states. (A) Two-dimensional ion occupancy within the SF resolved radially and in pore axis (z axis) as a contour plot calculated from MD simulations of GluA2 with different ion types (K⁺ and Na⁺) and force fields (AMBER99sb and CHARMM36). The simulations were performed at 303 K using an ion imbalance of 6e⁻ between the two compartments, α and β . The ion occupancy in number of ions per 0.001 Å³ per 500 ns was normalized according volume change along radius. (B) Dwell time histograms for S_I and S_{II} binding sites. Vertical dashed lines are means of the respective dwell times. (C) The number of oxygens within the first hydration shell of the according ion type along the pore axis. The number of coordinating water oxygens in blue, of coordinating protein oxygens in orange and the sum in grey.

173 *Ion hydration states during permeation*

174 To determine the ion hydration states during permeation, we calculated the number of ion-
 175 coordinating oxygens from both water molecules and protein residues within each ion's first
 176 solvation shell. Hydration shells are dynamic; here we defined the waters of hydration using
 177 the radii corresponding to the minimum in the radius of gyration profiles: 3.1 Å for Na⁺, 3.4 Å
 178 for K⁺, and 3.8 Å for Cs⁺ (26, 27). All monovalent ions were principally coordinated by waters
 179 in the lower SF region (Figure 4C), while the contribution of protein oxygens (backbone
 180 oxygens of Q586, backbone and side-chain oxygens of Q586 and Q587, Figure S3) increases
 181 substantially at the major ion binding site and surpasses the water oxygens for K⁺ and Cs⁺. The
 182 average number of ion-coordinating oxygens in the first hydration shell (black line) remained

183 remarkably stable along the pore axis, being 6, 7 and 8 for Na^+ , K^+ and Cs^+ , respectively (Figure
 184 4C). This total number of ion-coordinating oxygens was in very good agreement with the
 185 coordination number for hydrated ions determined experimentally (28). The simulations
 186 indicate a qualitatively similar mechanism for each ion species. Cations are never fully
 187 dehydrated, leading to appreciable water co-permeation. The retention of water is probably a
 188 strong factor in determining the lack of selectivity across the alkali metal series.

189 *Caesium permeation*

190 Classical electrophysiological measurements of reversal potentials for cloned glutamate
 191 receptors show at most minor differences in the permeation of Cs^+ (van der Waals radius 343
 192 pm) from that of Na^+ and K^+ (radii 227 pm and 280 pm respectively) (3). We reasoned that to
 193 achieve similar permeation statistics despite its greater size, Cs^+ should either alter selectivity
 194 filter geometry, occupy distinct binding sites or have radically different hydration state during
 195 permeation. To resolve this point, we simulated Cs^+ permeation in the same conditions as for
 196 K^+ and Na^+ . From both AMBER and CHARMM simulations we observed a moderately higher Cs^+
 197 permeation rate (Table 1 and Figure 2B) compared to Na^+ . However, the geometry at the tip
 198 of the pore loop did not change. The hydration of Cs^+ is similar to K^+ , with protein oxygens
 199 outnumbering the water oxygens at the S_I and S_{II} sites in the QQ selectivity filter. However, in
 200 AMBER simulations the major binding site (S_I) for Cs^+ is axially displaced by about 0.5 Å towards
 201 the bundle crossing compared to those of K^+ and Na^+ (Figure 5B, Figure S2), with a

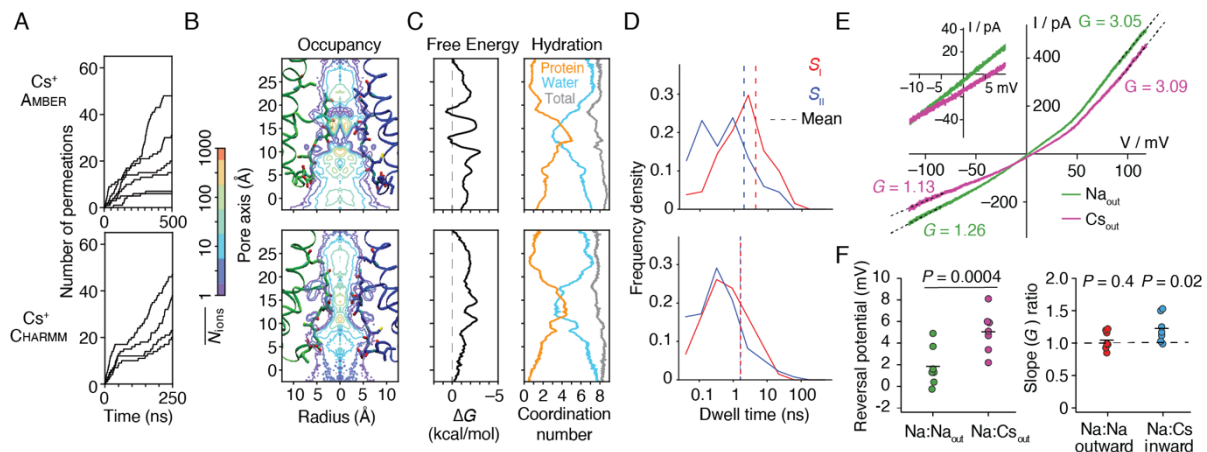


Figure 5. Cs^+ permeation of GluA2 (A) Permeation events for Cs^+ in the AMBER99sb and CHARMM36 force fields. (B) Two-dimensional ion occupancy plots (C) Free energy and hydration profiles for Cs^+ along the pore axis. (D) Dwell time histograms for Cs^+ in the S_I and S_{II} sites, with mean dwell time for each site indicated with a dashed line. (E) Outward rectifying responses to voltage ramps in Na^+ (green) and Cs^+ (purple) measured with patch clamp electrophysiology. The outward rectification (ratio of chord conductances at +100/-100 mV) was 2.7 ± 0.3 for Na^+ and 3.1 ± 0.3 for Cs^+ ($n = 7$ patches). Inset shows the typical shift in reversal potential on exchange of external Cs^+ and Na^+ . The pipette solution contained Na^+ . (F) Reversal potential of AMPAR currents in external NaCl and CsCl solution and conductance ratios of Na^+ versus Cs^+ for inward and outward permeation. Probability of no difference in the slope ratio was versus a ratio of 1 with paired Student's t-test.

202 concomitant longer distance to the S_{II} site. Most critically, in both AMBER and CHARMM
203 simulations, the mean residence time for Cs^+ at the S_I site was much less (~ 1 ns) than for Na^+
204 or K^+ and matched that at the S_{II} site (Figure 4B and Figure 5D). A prediction from its serial
205 binding to two effectively equivalent sites is that Cs^+ may be selected over Na^+ and K^+ , and
206 thus be more permeant.

207 The permeation of cations in GluA2 (Q) may be different to that observed in extensive studies
208 of GluA1(3). We performed patch clamp electrophysiology experiments on unedited GluA2
209 channels expressed in HEK 293 cells to reexamine the reversal potentials and slope
210 conductances for Na^+ and Cs^+ . We readily detected a small but consistent shift in the reversal
211 potential in external Cs^+ of 3.7 ± 0.5 mV ($n = 7$ patches), meaning Cs^+ was more permeant
212 than Na^+ ($P_{Cs} : P_{Na} = 1.16 \pm 0.02$) like in GluA1. The slope conductance for the inward arm of
213 the I-V relation in Cs^+ was similar to that of Na^+ ($82 \pm 6\%$; $n = 7$ patches). These observations
214 are consistent with the AMPAR pore being mildly selective for Cs^+ over Na^+ and Cs^+ retaining
215 a high conductance despite using a subtly different permeation mechanism on the same overall
216 scaffold.

217 *Conformational flexibility in the vicinity of the SF*

218 Examining the RMSF of heavy atoms in residues in the traditional SF ($^{585}MQQGCDI^{591}$) region
219 reveals a clear split in dynamics (Figure 6). The backbone atoms at the pore loop ($^{585}MQQG^{588}$)
220 are comparatively immobile, similar to the M3 backbone adjacent at L610. In the lower filter
221 region, the backbone and side chains of CDI residues are comparatively unrestrained. The side
222 chain of Q586 is more flexible than the Q587, as it points toward the cavity and has thus more
223 freedom for movement. Furthermore, the dynamics of the SF were indistinguishable across the
224 different ionic conditions, with the RMSF profiles of both backbone and side-chain atoms in
225 the Na^+ , K^+ and Cs^+ simulations being highly similar (Figure 6). These observations further
226 underline that the stable structure of the QQ filter is sufficient for selecting cations.

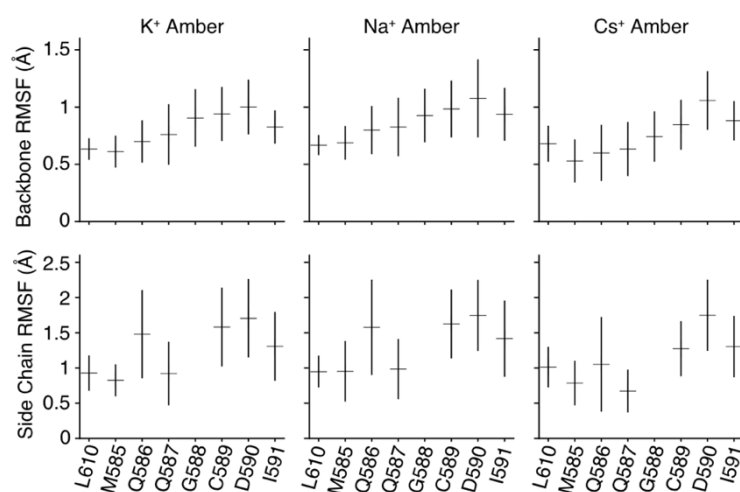


Figure 6. Root mean square fluctuations (RMSFs) of residues surrounding the selectivity filter. RMSFs were calculated separately for backbone and side-chain atoms of selectivity filter residues $^{585}MQQGCDI^{591}$ and L610. The simulations were performed at 303 K using an ion imbalance of $6 e^-$ between the two compartments α and β .

227 Discussion

228 In this study, we simulated permeation of various monovalent cations through AMPA channel
229 pore using the computational electrophysiology approach. Our direct measurements of ion
230 conduction confirm that the Cryo-EM structure reported by Twomey *et al.* (1) is bona-fide open
231 state, with a conductance large enough to be the fully open state. Particularly, the upper bundle
232 crossing in this structure, which we held open mechanically, provides almost no barrier to
233 conduction. Our simulations provide the first answer to how this channel can allow a range of
234 cations to pass with similar rates. The simulated conductance shows a striking agreement with
235 the single-channel recording data of Na⁺ in GluA2, suggesting that to a first approximation,
236 the ion conduction we observe resembles physiological permeation. The simulations suggested
237 that Cs⁺ permeates with a subtly different mechanism to Na⁺ and K⁺, and we could provide
238 evidence for this using patch clamp electrophysiology.

239 Although in principle using computational electrophysiology approach we should simulate both
240 inward and outward ion permeations simultaneously from one computational setup, we only
241 observed outward ion flux (from intracellular to extracellular) through the AMPA channel pore
242 in all simulation setups. All simulated conductance listed in Table 1 and Figure 3B are therefore
243 derived from the outward ion permeations. Similar directional flux has also been observed in
244 previous computational electrophysiology simulations of K⁺ selective channels such as KcsA
245 (29). Although electrophysiology reveals that the channel is outward rectifying (Figure 5) for
246 both Na⁺ and Cs⁺, the magnitude of the effect (about 3 fold) is too small to be consistent with
247 an total absence of inward permeation in our simulations. We hypothesize that inward and
248 outward ion permeations may use different SF conformations. In this interpretation,
249 simulations with a longer time scale than performed in the current study would be necessary
250 to reach the conductive states required for inward ion permeations.

251 For the major physiological ions Na⁺ and K⁺, we observed only one strongly populated ion
252 binding site (S_I) within the SF. This main ion binding site is remarkably congruent with the
253 single ion density observed in the Cryo-EM structure (PDB ID: 5WEO, Figure S6). The centres
254 of the ion densities along the pore axis are within 1 Å. This difference may in part be attributed
255 to the simulations being performed under transmembrane voltages, while the Cryo-EM
256 structure was determined at 0 mV in the absence of ion gradients. In contrast to the major
257 binding site S_I , ion binding to a second on-axis site (S_{II}) in our simulations was rather short-
258 lived and less ordered. This observation markedly contrasts with the four adjacent ion binding
259 sites of canonical K⁺ channels that were determined by X-ray crystallography (15) and MD
260 simulations (29). In several non-selective cation channels such as NaK (30, 31), CNG (14) and

261 HCN channels (13), the SF consists of either two or three consecutive sites. It has been
262 suggested that a single ion binding site within the NMDA receptor SF is the key to the ion non-
263 selectivity of these cation channels (32). Our finding from the AMPA simulations revealing a
264 single major ion binding site with twin energy barriers is clearly in line with this mechanism.
265 In other words, the AMPA receptor channel is minimally selective and has one of the simplest
266 observed selectivity filters comprising only two residues (Q586 and Q587).

267 Moreover, we found that during permeation hydration states of the ions in the SF also
268 significantly differ from the ones in the K^+ -selective and several non-selective cation channels.
269 Previous simulations suggested K^+ pass desolvated through the SF of potassium channels (9),
270 while an entire hydration shell surrounding Na^+ ions was observed whilst they traverse the SF
271 (9, 27) (10). There is no narrow constriction in the AMPA receptor pore to match those in K^+
272 and non-selective NaK channels, with the narrowest portion at Q586 (7.6 Å between CB atoms
273 in 5WEO). Therefore, all monovalent ions remain mostly hydrated in the SF and are only
274 partially dehydrated at Q586. Dehydration of smaller Na^+ ions requires much more energy
275 (33) (34), and consistent with this expectation, the Na^+ ions were more hydrated at the QQ
276 filter compared to K^+ and Cs^+ . The very short SF has a permissive architecture that
277 accommodates the different alkali metals in distinct solvation states.

278 Based on these observations, we propose that the mechanism of ion conduction through the
279 AMPA channel pore is remarkably different from that seen to date in other tetrameric channels.
280 For K^+ (29) and several other non-selective cation channels such as NaK (10) and the NaK-
281 CNG mutant (9), ions strictly follow a direct knock-on mechanism when passing through the
282 SF. In contrast, although multiple cations generally occupied the SF-region in the AMPA
283 receptor pore, only one stable axial binding site could be resolved, resulting in a quasi-
284 uncoupled knock-on for K^+ and Na^+ conduction. All ions retained a degree of hydration and
285 water was co-permeant. Furthermore, quite distinct from the non-selective NaK channel (10),
286 all monovalent ions investigated in this study (K^+ , Na^+ and Cs^+) follow a similar path during
287 permeation, with no requirement for conformational change for physiological permeation
288 rates. The conduction mechanism in the AMPA channel displays some similarities with Na^+
289 conduction in the voltage-gated sodium (NaV) channels, where both loosely coupled knock-on
290 and “pass-by” processes compete during Na^+ permeation (27, 35).

291 Using extensive MD simulations combined with Markov state model analyses, Furini and
292 colleagues suggested that ion-triggered conformational change within the SF determines ion
293 selectivity in bacterial sodium channels (36). Along the same lines, in our previous study on
294 non-selective NaK channels we proposed that K^+ and Na^+ permeation is coupled to distinct SF

295 conformations that are stable in the microsecond time scale and did not readily interconvert
296 (10). In marked contrast to these findings, the short AMPA selectivity filter does not show ion-
297 dependent conformations, and rather, achieves non-selectivity in a quite simple way. The
298 AMPAR provides a minimal selection architecture that is insensitive to ion diameter; the ion
299 hydration shells provide a compensatory variation. However, it is probably not correct to think
300 of the AMPA receptor SF region as a single permissive structure. Instead, the SF of the AMPA
301 receptor is flexible at the nanosecond time scale and allows different ions to permeate using
302 distinct mechanisms but at similar rates. It is also possible that the selectivity filter may be
303 coupled to the upper gate movement or fluctuation as in KcsA (37) and that by holding the
304 upper gate in a single stable conformation, we reduced the SF dynamics. Recent results
305 suggesting that SF perturbations can change the desensitization lifetime are consistent with
306 coupling to regions outside the pore domain (38). On this point, results from CryoEM are
307 inconclusive, with some open and closed structures having similar SF architectures, whereas
308 one open state structure (edited to R586) has marked two fold-symmetry (2). Restraining the
309 linkers was enough to keep the channel stably open, even though we removed flanking
310 Stargazin molecules. Although longer timescales may be required to see any putative relaxation
311 into a TARP-free state, this observation provides further evidence that the modulation of
312 channel gating by Stargazin and other TARPs is principally extracellular (39). Future work will
313 further clarify the relation between conduction properties, receptor conformation and
314 dynamics of the AMPAR pore (40).

315 **Methods**

316 *Computational Electrophysiology*

317 To prepare the Cryo-EM structure of the AMPAR in an open state (PDB ID: 5WEO) (1) for MD
318 simulations, we removed the four copies of the auxiliary protein Stargazin. We truncated the
319 receptor linkers before I504 and G774 and after I633 and A820 and added N-methyl amide
320 (NME) and acetyl (ACE) caps at the newly created C- and N-termini. We used MODELLER (41)
321 to build missing loops (residues 550 to 564 of the intracellular loops of each subunits) as well
322 as missing side chains (sidechains of S818 and R819 of subunits B and D, K505 of subunit D).
323 To hold the truncated structure together and in an open state, we restrained protein at the
324 heavy atoms of all four peptide chain termini with a harmonic potential of 10000 kJ/mol.

325 We performed MD simulations with both CHARMM36 (21) and AMBER99sb (20) force fields. In
326 the AMBER setup, insertion of the AMPAR TMD into a palmitoylcholine phosphatidylcholine
327 (POPC) lipid bilayer was performed with the Gromacs internal embedding function, whereas
328 in the CHARMM setup this process was carried out in CHARMM-GUI (42). The concentration of
329 KCl, NaCl and CsCl was 274 mM in simulations with AMBER, except for one simulation setup
330 of the closed channel with the AMBER force field was 284 mM KCl. For CHARMM simulations,
331 we used a salt concentration of 212 mM. Improved lipid parameters (43) and ion parameters
332 (44) were used in the AMBER simulations. We used the Tip3p water model (45) in all
333 simulations.

334 All the MD simulations were carried out with the GROMACS software package (versions 2016.1
335 and 2019.5) (46). Short-range electrostatic interactions were calculated with a cutoff of 1.7
336 nm, whereas long-range electrostatic interactions were treated by the particle mesh Ewald
337 method (47). The cutoff for van-der-Waals interactions were set to 1.7 nm. The simulations
338 were performed at 303 K or 310 K with an enhanced Berendsen thermostat (GROMACS V-
339 RESCALE thermostat) (48). A surface-tension Berendsen barostat was employed to keep the
340 pressure within the membrane-plane (XY-axis) at 250 bar * nm per lipid surfaces and at one
341 bar in z-axis direction (49). All bonds were constrained with the LINCS algorithm (50).
342 Interatomic forces (van-der-Waals and Coloumb) were calculated with a 1.5 nm cutoff. Long
343 range electrostatic forces were computed with the Particle-Mesh-Ewald method. To decrease
344 computational cost, we employed virtual sites for hydrogens in all AMBER simulations. AMBER
345 simulations were calculated with an integration time step of 4 fs. All CHARMM-based simulations
346 used an integration time step of 2 fs.

347 Before starting the simulation of the AMPAR with an ion gradient, the system was firstly energy-
348 minimized and equilibrated. After the systems energy was minimized to below 1000 kJ/mol
349 with GROMACS “steepest descent” implementation a 10 ns free equilibration without any
350 restraints (except those for the linkers) was performed. For CHARMM-based simulations the
351 recommended energy minimization and equilibration steps from CHARMM-GUI were
352 performed.

353 For the computational electrophysiology study (17), two copies of the equilibrated system of
354 the AMPAR TMD in the lipid bilayer were included in a simulation box of typically $10 \times 10 \times$
355 20 nm (Figure 1D). A charge gradient and therefore transmembrane potential was generated
356 by introducing an ion difference of 2, 4 or 6 ions between the two compartments separated by
357 the two lipid bilayers. Since the instantaneous introduction of a charge gradient into the system
358 can produce large perturbations in the system we omitted the first 20 ns of every simulation.
359 During MD simulations, the number of ions in each chamber was kept constant alchemically
360 by an additional algorithm (17). The resulting membrane potential can be calculated by
361 double-integration of the charge distribution using the Poisson equation as implemented in the
362 GROMACS tool *gmx potential* (51). During the simulations a permeation event was counted
363 when an ion traversed the entire filter region. Individual simulation runs were 500 ns in the
364 AMBER setups and 250 ns in the CHARMM setups (Table 1).

365 The majority of the production runs were conducted at 303 K using a charge imbalance (Δq)
366 of $6 e^-$ (elementary charges) between the two compartments α and β (Figure 1D). Although
367 ion channel recordings are possible at these voltages, they lay well outside the physiological
368 range. We additionally simulated K^+ permeation using Δq of 4 and $2 e^-$ with AMBER99sb,
369 respectively, corresponding to a voltage difference of 275 mV and 100 mV. The simulation at
370 275 mV shows a similar conductance to the one at 450 mV. At 100 mV, the conductance is less
371 but still substantial (Table 1, Figure 2B), indicating that similar conduction mechanisms are at
372 play. The principal consequence of high membrane voltage is merely statistical – it allows us
373 to record more permeation. Furthermore, compared to the simulations conducted at 303 K we
374 noted that simulations at 310 K revealed only slightly larger currents but with much greater
375 variations in the ion permeation rate within a given simulation run (Figure 2B). Finally, to
376 buttress our observations and validate the relevance of the starting structure, we performed
377 equivalent MD simulations of a closed AMPA structure (PDB ID: 5WEM; GluA2-GSG1L-apo-1)
378 with the ATD, LBD and GSG1L removed, and extracellular peptide termini restrained. We
379 observed no ion permeation events over the same time scale as the open structure simulations,
380 and the structure did not spontaneously open on this timescale.

381 To align results over different setups we used the center of mass of the backbone carbons of
382 the classical selectivity filter residues (⁵⁸⁶QQGCD⁵⁹⁰) as a reference point. All trajectories were
383 analysed with GROMACS tools and PYTHON using MDANALYSIS (52). Molecular visualizations
384 were made with VMD (53).

385 *Patch Clamp Electrophysiology*

386 HEK293 cells were plated on glass coverslips in dishes and incubated for 20-44 hr before
387 calcium phosphate transfection with 3 μ g of cDNA. For macroscopic current recordings, the
388 cDNA transfection was done with the Rat GluA2 (Q) pRK5 vector encoding enhanced green
389 fluorescent protein after an IRES. Single channel recordings from outside-out patches were
390 performed 24 hr after transfection. For single channel recordings, we used a plasmid ratio
391 approach to obtain sparse expression(54), whereby the rat GluA2 Q vector was cotransfected
392 with enhanced green fluorescent protein and empty vector (pCDNA3.1+) in a ratio of 1 : 63 :
393 313. Standard extracellular solution contained 150 mM NaCl, 0.1 mM MgCl₂, 0.1 mM CaCl₂, 5
394 mM HEPES, 10 μ M EDTA and was titrated with NaOH to a pH of 7.3. We included EDTA to
395 chelate trace divalent ion contamination. The internal solution contained 115 mM NaCl, 1 mM
396 MgCl₂, 0.5 mM CaCl₂, 10 mM NaF, 5 mM Na₄BAPTA, 10 mM Na₂ATP, 5 mM HEPES, titrated
397 to a pH of 7.3 with NaOH. Pipettes were mounted in an ISO holder (G23 Instruments, London,
398 UK) and had a resistance of 3 M Ω for macroscopic current recordings. For single channel
399 recording, pipettes were fire polished to a resistance of 10-25 M Ω and were coated with
400 Sylgard. The junction potential between the pipette and Na⁺-bath solution ($E_{\text{bath}}-E_{\text{pip}}$,
401 considering Na⁺, Cl⁻ and F⁻ mobilities) was 3.7 mV (55). For reversal potential experiments
402 with Cs, we substituted NaCl in the extracellular solution with CsCl and titrated the pH to 7.3
403 with CsOH. CTZ (Hello Bio) was prepared as a 100 mM stock solution in DMSO and used at
404 100 μ M (giving a final concentration of 0.1% DMSO. EDTA stock solution was prepared in
405 NaOH. Reagents were obtained from Carl Roth GmbH (Karlsruhe, Germany), Sigma Aldrich
406 (Munich, Germany), or Toronto Research Chemicals (Toronto, Canada). Ultra-fast perfusion
407 to outside-out patches was used for drug application. The perfusion tools were made with
408 custom designed four-barrel glass (Vitrocom, Mountain Lakes, USA). Currents were filtered at
409 10 kHz (-3 dB cutoff, 8-pole Bessel) with an Axopatch 200B amplifier (Molecular Devices,
410 Sunnyvale, USA). For analogue-digital conversion an InstruTECH ITC-18 digitizer (HEKA
411 Elektronik Dr. Schulze GmbH, Lambrecht, Germany) was used at 40 kHz sampling rate. Data
412 were recorded and analysed with AxoGraph X (AxoGraph Scientific, Sydney, Australia).

413 For single channel conductance measurements, in each record, we ran a ramp protocol (-120
414 to +120 mV, 1.2 V s⁻¹) both before and during glutamate application (10 mM) to the outside

415 out patch. The leak current recorded during the no-glutamate ramp was subtracted from the
416 current recorded during glutamate application. Stretches of the recording corresponding to one
417 open channel were selected, the open levels were selected, and fit with a linear relation to
418 obtain the chord conductance.

419 For macroscopic measurements of reversal potential and conductance, we alternated washing
420 each patch with Cs⁺ and Na⁺ solution. Slope conductances were fit to the traces over the 30
421 mV ranges at the extreme positive and negative ends of the ramp. For the slope conductance,
422 we compared the conductance ratios (Cs vs Na inward and Na vs Na outward) from the same
423 patch against the null (ratio = 1) using a paired t-test. We assumed that there was a similar
424 junction potential (within 0.5 mV) in both cases, because the junction potential from the
425 pipette to the Cs⁺ solution (-1.7 mV) is cancelled by a second junction potential from Cs⁺
426 solution back to the bath electrode in Na⁺ (4.9 mV) (56). We added this junction potential to
427 the measured mean reversal potential. The flow rate through the local perfusion tool was low
428 (< 200 μL per minute), meaning the overall bath Cs⁺ concentration remained low.

429 We used the Goldman-Hodgkin-Katz equation (57) to calculate permeability ratios based on
430 reversal potentials:

$$431 \quad P_X/P_{Na} = [Na^+]_i/[X^+]_o \cdot e^{\frac{FE_{rev}}{RT}} \quad (1)$$

432 where X was Na⁺ or Cs⁺.

433 Likewise, to calculate the putative shift in E_{rev} due to a minor Chloride permeability, we
434 assumed $P_{Cl:Na}$ of 1:220 and $P_{Cs:Na}$ of 1.16:1, as measured.

$$435 \quad E_{rev} = \frac{RT}{F} \cdot \ln \left(\frac{[Na^+]_o + P_{Cs:Na} \cdot [Cs^+]_o + P_{Cl:Na} \cdot [Cl^-]_i}{[Na^+]_i + P_{Cs:Na} \cdot [Cs^+]_i + P_{Cl:Na} \cdot [Cl^-]_o} \right) \quad (2)$$

436 Statistical analysis and data plotting was done in IGOR Pro (Wavemetrics).

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448 **Author contributions**

449 All authors designed the experiments, performed the experiments and/or analysed data, wrote
450 and edited the manuscript.

Cation	Force Field	Δq (e ⁻)	[C] (mM)	V_m (mV)	T (K)	Replicates	Ion Permeations	γ (pS)
K ⁺	AMBER	6	273.8	440	303	5	220	33 ± 11
K ⁺	AMBER	2	273.8	100	303	5	18	12 ± 12
Na ⁺	AMBER	6	273.8	490	303	5	50	6 ± 3
K ⁺	CHARMM	6	212.1	230	303	5	144	78 ± 41
Na ⁺	CHARMM	6	212.1	325	303	5	63	25 ± 10
Cs ⁺	AMBER	6	273.8	400	303	6	133	18 ± 13
Cs ⁺	CHARMM	6	212.1	280	303	4	121	71 ± 28
K ⁺	AMBER	6	273.8	450	310	5	300	43 ± 39
K ⁺	AMBER	4	273.8	270	310	5	163	42 ± 33
K ⁺ closed	AMBER	6	284	350	303	3	0	0
Na ⁺	<i>Single Channel Recording</i>	-	150	-120 to +120	296	3 patches	-	30 ± 1.6

451 **Table 1. Summary of computational electrophysiology simulations**

452 Simulations of the GluA2 transmembrane domain for individual permeating monovalent
453 cations Na⁺, K⁺ and Cs⁺. In addition to changing cations, we also varied temperature (T),
454 charge imbalance (Δq) and the force field employed. C is the salt concentration for each
455 simulation. Each simulation replicate was 500 ns except for CHARMM runs where the duration
456 was 250 ns. These simulations represent an aggregate time of 20.5 μ s. AMBER refers to
457 AMBER99sb and CHARMM to CHARMM36. For comparison, the conductance of GluA2 (γ) in Na⁺
458 determined from 3 single channel recordings is also included.

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