Measurements of the Timescale and Conformational Space of AMPA Receptor Desensitization

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ABSTRACT Ionotropic glutamate receptors are ligand-gated ion channels that mediate excitatory synaptic transmission in the central nervous system. Desensitization of the α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid subtype after glutamate binding appears critical for brain function and involves rearrangement of the ligand binding domains (LBDs). Recently, several full-length structures of ionotropic glutamate receptors in putative desensitized states were published. These structures indicate movements of the LBDs that might be trapped by cysteine cross-links and metal bridges. We found that cysteine mutants at the interface between subunits A and C and lateral zinc bridges (between subunits C and D or A and B) can trap freely desensitizing receptors in a spectrum of states with different stabilities. Consistent with a close approach of subunits during desensitization processes, the introduction of bulky amino acids at the A-C interface produced a receptor with slow recovery from desensitization. Further, in wild-type GluA2 receptors, we detected the population of a stable desensitized state with a lifetime around 1 s. Using mutations that progressively stabilize deep desensitized states (E713T and Y768R), we were able to selectively protect receptors from cross-links at both the diagonal and lateral interfaces. Ultrafast perfusion enabled us to perform chemical modification in less than 10 ms, reporting movements associated to desensitization on this timescale within LBD dimers in resting receptors. These observations suggest that small disruptions of quaternary structure are sufficient for fast desensitization and that substantial rearrangements likely correspond to stable desensitized states that are adopted relatively slowly on a timescale much longer than physiological receptor activation.

INTRODUCTION

Glutamate receptor ion channels mediate most of the fast excitatory synaptic transmission in the vertebrate central nervous system (1). Glutamate binding initiates the opening of an integral ion pore, permitting cations to flow into the postsynaptic cell. The α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) receptor subtype desensitizes rapidly and profoundly in response to the sustained presence of glutamate for more than ~25 ms (2,3). The number of receptors available to respond to glutamate is consequently reduced during a phase of recovery from desensitization, which in turn can determine the amplitude of postsynaptic responses (4,5). The timescale of recovery from desensitization, being for AMPA receptors in the order of tens to hundreds of milliseconds, is pertinent during high-frequency release of glutamate (above 10 Hz). Steady-state desensitization may offer protection during pathological glutamate insults that lead to brain damage (6) and during development (7). Finally, desensitized-like conformations might...
be important during biogenesis, trafficking, or in general, any cellular situation in which activation would be problematic. A connection between surface expression and desensitization has been shown for AMPA and kainate receptors (8,9). These observations provide motivation for understanding the molecular basis of AMPA receptor desensitization.

AMPA receptors assemble from four subunits, each comprising an extracellular amino-terminal domain, a ligand binding domain (LBD) that is connected to the ion channel forming transmembrane domain and a C-terminal domain. The LBD is formed from an upper D1 and a lower D2 lobe. Upon the binding of glutamate, the LBD closes. This motion provokes the separation of the D2 domains, leading to the opening of the receptor gate (10). After activation, the receptor transits to desensitized states in ~10 ms, at least in part because of dissociation of the dimer interface formed by the D1 domains (11).

Recent structures of the full-length AMPA receptor in putative desensitized states suggest that a wide conformational space is sampled. Initial cryo-electron microscopy (cryo-EM) structures of the GluA2 receptor in desensitizing conditions showed a set of three-dimensional classes in which the extracellular domains were progressively spread out (12). A 5-fluorowillardiine-bound structure of GluA2 (13) also showed a large rearrangement of the amino-terminal domain and LBDs. Structures of the related GluK2 kainate receptor show that the LBDs adopt a fourfold symmetric arrangement (14) with individual subunits rotating by more than 120° from their active state dimer positions (Fig. 1; (15)). More recent desensitized state structures of GluA2 in the presence of the accessory proteins GSGL1L or TARP γ-2 revealed compact desensitized arrangements, with the LBD dimers losing their internal twofold rotational symmetry (16,17).

We previously used cysteine and metal-bridge cross-linking to identify compact arrangements of the LBD tetramer associated with the activation of the AMPA receptor. These include the “closed angle” conformation (18), multiple compact forms for LBDs fully bound to glutamate (19), and the partial agonist 5-fluorowillardiine (20). The latter assembly featured a parallel shift of the individual dimers. In this previous work, we largely used cyclothiazide (CTZ) to prevent desensitized arrangements of the LBD layer. More recently, we used bifunctional cysteine crosslinkers to measure the extent to which the LBD tetramer opens up in both active and desensitized states (21).

In this study, we revisited our earlier observation that desensitized receptors can be cross-linked very stably between A and C subunits by the A665C disulfide bond (18). Using a fast perfusion system, we used several strategies on mutant and wild-type (WT) receptors to count conformational states attained during desensitization. Distinct from our previous work in which we trapped active states, here, we employed conditions to enrich desensitized states and also examined the inactivation of apo receptors. Comparing potential desensitized states obtained in crystallographic and cryo-EM to the geometric constraints imposed by disulfide bonds and metal bridges suggests that compact desensitized arrangements can best account for desensitization on the physiological timescale.

MATERIALS AND METHODS

Electrophysiology

All mutants were generated on the rat GluA2flip background by overlap PCR and confirmed by double-stranded DNA sequencing. For consistency with previous reports, the numbering of mutated amino acids assumes a 21-residue signal peptide for GluA2. WT and mutant AMPA receptors were expressed transiently in HEK-293 cells for outside-out patch recording. All patches were voltage clamped between ~30 and ~60 mV. Currents were filtered at 1–10 kHz (~3 dB cutoff, 8-pole Bessel) and recorded using AxoGraph X (AxoGraph Scientific) via an InstruTECH ITC-18 interface (HEKA Elektronik) at a 20-kHz sampling rate.

The external solution in all experiments contained 150 mM NaCl, 0.1 mM MgCl2, 0.1 mM CaCl2, and 5 mM HEPES, titrated to pH 7.3 with NaOH, to which we added drugs, agonists, redox agents, zinc, and ion chelators. CTZ stock solution was prepared in DMSO and added at 100 mM to the external solution. Drugs were obtained from Tocris Bioscience (Bristol, UK), Ascent Scientific (Bristol, UK), or Sigma-Aldrich (St. Louis, MO).

Trapping protocols and chemical modification

To measure the state dependence of cysteine trapping in the desensitized state, we determined the baseline activation by 10 mM glutamate in the presence of 5 mM dithiothreitol (DTT), followed by the application of Cu:Phen (10 μM; prepared as described in (22)) and 100 μM Glu for a range of time intervals. To examine resting state trapping, we applied 10 μM Cu:Phen without agonist, and for some experiments, we added 100 μM CTZ. Each trapping exposure was delivered from the third barrel of the perfusion tool. For all trapping experiments, we quantified the relief of trapping by determining the fraction activated by sequential applications of 10 mM glutamate in 5 mM DTT immediately after trapping, as previously described (18). The envelope of the peak current responses after application of Cu:Phen were fit with a single exponential. By back extrapolating to the end of the Cu:Phen application, we were able to estimate the proportion of receptors that were trapped (22). The amplitude of the fit function was the trapped fraction of receptors, and we subtracted this fraction from 1 to get the active fraction (AF). For metal bridging experiments, Zn2+ was added (10 μM) to the external solution. To achieve zinc-free conditions, we added 10 μM EDTA, a potent Zn2+ chelator (Kd Zn2+ = 10^-16.4 M), to the external solution. We used the same analysis to determine the AF. We applied drugs to outside patches via perfusion tools made from custom manufactured glass tubing with four parallel barrels (VitroCom, Mountain Lakes, NJ) as described in (18). The glass was pulled to a final width of 200 μm, and the tip of the tool was etched in hydrofluoric acid and mounted in a piezo electric lever and controlled via a 100 V amplifier. The command voltage was filtered at 100 Hz to reduce vibration of the tool. When we measured the junction potential, the typical 10–90% rise time was 300 μs. For the fast oxidizing experiments, we determined the time that the patch spent in the Cu:Phen condition (third barrel) by measuring open tip currents, calibrating a voltage ramp protocol with different slopes to vary the dwell time in the third barrel from 5 to 30 ms (Fig. 7 A). We measured concentration-response curves for WT and the mutant A665W. We obtained the half-maximal effective concentration (EC50) from fits to the Hill equation.
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The timing and amplitude of peak currents and rise times of all peaks were measured in AxoGraph. Recordings from 23 patches from different cells were used for further analysis, except for the 5-s conditioning pulses. Because the duration of these measurements was long and the rundown of the current was often substantial, only measurements from four patches could be completed before the patch was lost and had sufficiently good quality for the whole set of records, namely 10–90% rise times of the glutamate response <500 μs and a stable baseline with fluctuations less than 10% of the peak current.

The response after a long (5 s) conditioning pulse was corrected for the slow rundown of current amplitudes caused by either accumulation of receptors into electrically isolated parts of the patch (23) or accumulation of receptors into nonfunctional states. A linear function was fitted to the currents and times of the response to the conditioning (first) pulse from each episode to extrapolate the expected maximum response at the time of each test (second) pulse. For each interval, the normalized responses were averaged, and the SD was used for fitting in IGOR Pro (WaveMetrics).

We and others have previously fitted recovery from desensitization with a Hodgkin-Huxley-type recovery curve (24):

$$f(t) = y_0 + (y_{\text{max}} - y_0) \cdot (1 - \exp(-kt))^n,$$

(2)

where $n$ is the Hill coefficient, $I_{\text{max}}$ is the maximum response, and $[A]$ stands for the agonist concentration. Recovery from desensitization for the A665W mutant was measured with a 400-ms conditioning pulse.

**Recovery from desensitization**

To examine the recovery from desensitization for WT GluA2, patches containing hundreds of receptors were conditioned with applications of 10 mM glutamate for 50, 200, and 800 ms and 5 s. A test glutamate pulse was delivered at 12 different time points between 2 and 790 ms after the conditioning pulse. The protocols with different durations of the first pulse were randomly initiated for each patch and repeated if the patch was stable enough.

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where \( k \) is the rate of recovery, \( m \) is the slope, \( y_{\text{max}} \) and \( y_0 \) are the maximum and minimum, respectively, and \( t \) is the interval between pulses. We also did this here for the data in Fig. 3 A. However, for the recovery after conditioning pulses of longer durations (Fig. 5), this function was insufficient because it could not describe the intermediate phases of recovery. These data could only be well fit by a function that was the sum of two Hodgkin-Huxley terms (with rates \( k_1 \) and \( k_2 \), slopes \( m_1 \) and \( m_2 \)):

\[
f(t) = y_0 + a_1 \cdot (1 - \exp(-k_1 t))^{m_1} + (y_{\text{max}} - a_1 - y_0) \cdot (1 - \exp(-k_2 t))^{m_2}.
\]

(3)

To establish how unique the description by this two-component H.-H. function was, we tried several other different fit functions (three component H.-H. function, sigmoid, Hill) and varied the fit parameters (see Figs. S1 and S2; Tables S1 and S2). Time constants where quoted are reciprocals of rate constants.

**Statistical analysis**

All \( p \)-values were determined by a two-sample unpaired Student’s \( t \)-test. The spread of the data where indicated is the SD of the mean.

Structural analysis and figure preparation was done in PyMOL (version 2.0; Schrödinger).

**RESULTS**

**An interdimer interface forms in a desensitized state**

To place intersubunit bridges in the context of the conformational changes that drive desensitization, we first compared the LBD layers of the apo state structure of GluA2 (Protein Data Bank, PDB: 4U2P (13)) to three candidate-desensitized state structures. We modeled the A665C mutation into these structures and measured the putative distances between the sulfur atoms of the cysteines. In the resting ligand-free structure, the SG-SG distance is 9 Å, partly because of the open angle adopted by the dimers but mainly as a consequence of the distance of the A and C subunits from the central axis of the receptor and ion pore. The structure of the GluA2 receptor in complex with the high-affinity full agonist L-quisqualate and the accessory subunit GSG1L (GluA2-TARP γ-2 Quis structure (PDB: 5VOV (17))) presents LBD dimers. The closest approach of modeled SG-SG distance remains 9 Å (Fig. 1). For comparison, in the active state structure PDB: 5WEO (25), the distance between cysteine sulfur atoms modeled at residue 665 is 11 Å. The GluA2-TARP γ-2 Quis (PDB: 5VOV) shows similar rotations and translations within each of its LBD dimers. Placing the A665C mutations in subunits A and C of this structure allows the SG groups to be only 5 Å distant, within striking distance of forming a disulfide bridge, suggesting the TARP-bound desensitized state is quite compact; this concept was discussed at length in our previous work (21). A yet more dramatic change can be observed in the homologous GluK2 structure in the desensitized state (2S,4R-4-methylglutamate PDB: 5KUF (14)), which shows a rotation of the subunits D and B of 125°, producing pseudo four-fold symmetric arrangement of the LBDs. Similar large-scale disruptions of the LBD layer were observed in single particle analyses of GluA2 without auxiliary subunits (12–14). As a consequence of this movement, the residue T670 (equivalent in GluK2 to A665 in GluA2) is buried between the new interfaces formed between adjacent subunits (Fig. 1). For the same four candidate structures, we measured the distances between the residues that form the site of the T1 lateral Zn\(^{2+}\) bridge (19) built from introduced histidines (D668H T672H K761H; Fig. 1 C). The intersubunit distances are too great to predict zinc bridging for any of the candidate structures. The closest approach was for GluA2-TARP γ-2 Quis (PDB: 5VOV), in which the CA-CA distances were 12–13 Å (for K761 to D668 or T672). Overall, no candidate-desensitized arrangement would support a Zn\(^{2+}\) bridge if histidine residues were placed at these positions.

We reasoned that if we attempted to cross-link the diagonal A-C interface during desensitization, we could determine at which point in the desensitization reaction (either early or late) that these two subunits come together. Likewise, we expected that the lateral interface should not be readily accessible to desensitized receptors, unless a spectrum of different desensitized states are sampled. We used well-characterized cysteine substitutions at three positions in the FG-loop (I664, A665, and V666) (18,20,21). Each mutant was tested for its cross-linking potential in 100 μM glutamate. This concentration was based on the concentration dependence of desensitization of the WT receptor, which reaches full desensitization at 100 μM (22), with minimal activation. We used three barrels of a quadruple barrel fast perfusion system that enabled the application of 100 μM glutamate in the presence of 10 μM Cu:Phen with <10 ms resolution. We observed a dramatic reduction of the current activated by glutamate 10 mM after exposure to oxidizing conditions for the mutants I664C, A665C, and V666C (Fig. 2 A). The untrapped, AF was measured for different intervals of exposure to oxidizing conditions. After ~30 s of oxidizing conditions, the reduction of the AF reached a plateau for I664C, A665C, and V666C (to 31 ± 4, 29 ± 4, and 22 ± 3%, respectively; Fig. 2 B). The maximum extent of inhibition for the three mutants was similar, but this similarity does not imply that the underlying mechanism is the same. Factors that influence the extent of trapping include side-chain angles, state stability, steric hindrance, and the geometry of the subunits. We do note that trapping around the loop between F and G helices is position dependent, with much a reduced trapping extent at position 662, for example (18). The kinetics of recovery after trapping indicate how stable this interface is in the desensitized state. The time constants of recovery after trapping for I664C, A665C, and V666C were 1.4 ± 0.2, 3.3 ± 0.6, and 1.2 ± 0.4 s, respectively, after 10 s in Cu:Phen (Fig. 2 C and
see Discussion for details of the interpretation of these time constants). Strikingly, with the longer exposure to Cu:Phen, the three mutants I664C, A665C, and V666C each showed increased stabilization of the trapped state. We fitted the increase in the recovery time constant against exposure time with a single exponential to obtain the asymptotic maximum recovery time constants of 43 ± 13, 38 ± 19, and 43 ± 5 s for I664C, A665C, and V666C, respectively (Fig. 2C). These results indicate that the A-C interface can be trapped in at least two desensitized states. We cannot discern from these data whether the states are accessed in parallel or series. One state recovers rapidly, but with long exposures, there is a progressive adoption of a state or set of trapped states that are considerably more stable.

**A point mutant at the A-C interface slows recovery from desensitization**

According to previous reports, the introduction of a cysteine mutation at positions 664–666 and WT GluA2. Dotted red line is the exponential fit to recovery of the current in 10 mM glutamate and DTT, following trapping in the presence of 10 μM Cu:Phen and 100 μM glutamate. Open circles with an arrow indicate the presumptive back-extrapolated response immediately after trapping. (B) The AF of receptors after oxidation in the desensitized state plotted against the trapping interval (continuous lines are exponential fits). The probabilities of no difference in the AF after 10-s trapping in 10 μM Cu:Phen were 0.001, 0.0005, and 0.00003 for I664C, A665C, and V666C, respectively (compared to WT A2, Student’s t-test, n = 3–4 patches per point). The arrows indicate the relevant intervals for the traces in (A). (C) The time constant of recovery after trapping is plotted against the trapping interval for I664C, A665C, and V666C, showing a positive correlation (continuous lines are exponential fits; n = 3–4 patches per point). To see this figure in color, go online.

**Adoption of deep desensitized states protect against cross-linking**

We reasoned that if the stable disulfide trapping we detected were unique to the desensitized state, then promoting desensitization should promote trapping and/or slow recovery. However, additional desensitized states might exist that are not readily disulfide linked by cysteines at the A-C interface. Such states would perhaps resemble the four-fold symmetry of the GluK2 structure or, more generally, would stably hinder the approach of cysteines at the otherwise proximal A-C interface because of a substantial conformational change. We tested the formation of intersubunit cross-links in a 20 ms, as previously reported (27). In contrast, we observed a dramatic delay of desensitization recovery of more than sevenfold for the mutant A665W (τrec = 155 ± 5 ms; Fig. 3A). The activation of the mutant and rate of entry to desensitization was indistinguishable from WT GluA2. We constructed a dose response curve for glutamate and observed little difference in the apparent affinity for glutamate (A665W EC50 = 510 ± 130 μM compared to WT EC50 = 330 ± 100 μM, with a p-value of no difference 0.19) (Fig. 3B). We therefore ruled out the possibility that the change in recovery was due to an increase in affinity for glutamate in the A665W mutant. This observation further supports the idea that this intersubunit interface forms during the entry to or exit from desensitization.
putative deep desensitized state using a mutant that strongly stabilizes the desensitized state (E713T and Y768R), with a recovery time constant of about 1 s (27). When we introduced a cysteine in position A665 in the single mutants E713T and Y768R, we observed less profound trapping than for the A665C mutant alone, with a reduction of the AF of 46% for A665C and E713T and 42% for A665C and Y768R after 100 s of application of Cu:Phen (Fig. 4, A and B). Both mutants showed slower recovery after trapping for A665C and E713T of 10.7 ± 1.7 and 9.7 ± 1.6 s for A665C and Y768R (Fig. 4 C). Again, longer exposures to Cu:Phen drove adoption of a yet more stable trapped arrangement. Fitting a single exponential to the recovery time constants versus the time in oxidizing conditions, we determined the asymptotic limiting time constants of a recovery profile in two distinct ways (Fig. 5). (A) Currents evoked by 10 mM glutamate from WT GluA2 (left panel) and the mutant A665W (right panel). The two-pulse protocol had a conditioning pulse of 400 ms, followed by a second pulse at increasing intervals (responses are overlaid). Peak currents (red open circles) were fit with Hodgkin-Huxley functions (with slope fixed to 2, see Materials and Methods). The time constants of recovery from desensitization were 20 ± 2 and 155 ± 5 ms for GluA2 WT and A665W, respectively (probability of no difference = 0.01; Student’s t-test; n = 4). (B) Mean of recovery from desensitization for GluA2 WT (black) and A665W (orange). For each interval, the peak of the second pulse is plotted as the AF (relative to the first peak) and fit with a Hodgkin-Huxley equation (with slope fixed to 2, see Materials and Methods). The time constants of recovery from desensitization were 20 ± 2 and 155 ± 5 ms for GluA2 WT and A665W, respectively (probability of no difference = 0.01; Student’s t-test; n = 4). (C) Concentration-response curves for WT GluA2 (blue circles; EC_{50} = 330 ± 100 μM) and GluA2 A665W (yellow circles; EC_{50} = 510 ± 130 μM). The probability of no difference between the EC_{50} values was 0.19 (n = 3 cells). To see this figure in color, go online.

**Long exposures to glutamate promote entry to stable desensitized states**

From these results, we predicted that the progressively greater stability of trapped receptors after long exposures to desensitized conditions should derive from the selective adoption of more stable desensitized states. However, the WT homomeric GluA2 receptor is known for its rapid and complete recovery from desensitization (see for example Fig. 3 A). To resolve this paradox, we tested if the rate of recovery from desensitization was sensitive to the duration of the conditioning pulse in two-pulse recovery experiments. In each record, a patch containing hundreds of WT GluA2 receptors was first conditioned with an application of glutamate (10 mM) for 50, 200, and 800 ms and 5 s. A second glutamate pulse was delivered at varying times after the conditioning pulse (2–790 ms; Fig. 5). For each patch, we made a series of recordings with the same conditioning pulse but different intervals and then repeated the protocol with a different conditioning pulse length. After short conditioning pulses, we observed prototypical fast recovery from desensitization for GluA2. The recovery after 50- and 200-ms conditioning pulses could be quite well described with a single H.-H.-type function with a time constant of 20 ms (with slope between 2.5 and 3; Fig. S1 B). However, in the same patch, a 5-s conditioning pulse of glutamate slowed the recovery profile in two distinct ways (Fig. 5 E). First, the fastest component, with a time constant of 14 ms, was only about 70% of the total amplitude (Table S1). Second, a smaller (~5%) but very slow (~1 s) component meant that recovery at the end of our protocol (700-ms interval) was always incomplete. Comparison of the recovery after 800-ms and 5-s conditioning pulses revealed that both had a small intermediate component (~20%) with a time constant of about 50 ms (Table S1).

We also observed a slow component of recovery when using 10- or 30-s conditioning pulses, but it was hard to quantify because the recovery protocols using such long
conditioning pulses necessarily lasted for 5–10 min, over which time even stable patches ran down and gave spurious responses. Taken together, these observations emphasize that WT GluA2 receptors can adopt a range of desensitized states with different stabilities, including very stable desensitized states.

**Lateral shifts occur during desensitization**

The GluA2-TARP γ2 Quis (PDB: 5VOV) structure (17) shows a compact packing of the lateral interface of the subunits A, B, C, and D of the LBDs, whereas in the GluA2-2xGSG1LQuis (PDB: 5VHZ) structure (16), this interface is clearly absent (Fig. 1C). To analyze if this interface occurs in early desensitized states that can be adopted over millisecond timescales, we used a metal ion trapping approach. Previously, we engineered a pair of histidine mutants T1 (D668H, T672H, and K761H) and HH (D668H and K765H), which can coordinate Zn$^{2+}$ between subunits A, B, C, and D at intermediate and high concentrations of glutamate, with CTZ present to block desensitization (19). Using these mutants, we detected the formation of the lateral interfaces in the desensitized state applying 100 µM glutamate in the presence of 10 µM Zn$^{2+}$. The mutants T1 and HH showed a reduction of the AF after 1 s of the application of Zn$^{2+}$ in the desensitized state, with a reduction of the AF of 33% for T1 and 23% for HH plateauing after 100 s of Zn$^{2+}$ application (Fig. 6A), with time constants of recovery after trapping for T1 and HH of 1 ± 0.08 s, and 0.6 ± 0.08 s, respectively, after 10 s in zinc (Fig. 6B). The asymptotic time constants of recovery in the limit of long Zn$^{2+}$ exposures were 4 ± 0.6 and 2 ± 0.8 s (for T1 and HH, respectively; Fig. 6C).

We hypothesized that the conformation trapped by the T1 lateral bridge is absent in the deep desensitized states promoted by the ET/YR mutation. To test this hypothesis, we inserted the triple histidine mutant T1 (D668H, T672H, and K761H) in the mutant (E713T and Y768R) background and tested its sensitivity to zinc in desensitized states. As for the A665C mutant, we did not detect the formation of T1 lateral interfaces in the presence of the ET/YR background (upper left) and the GluA2 ET/YR background (upper right) shows no modification. (B) The AF of receptors after oxidation in the desensitized state (continuous lines are exponential fits) is plotted against the trapping interval. The A665C trapping profile (light blue dashed line) and the fit to the ET/YR background (black dashed line) are indicated. The probability of no difference between the AF after 10 s of application of oxidizing conditions was 0.012 and 0.0008 for A665C and Y768R and A665C and E713T (versus A665C and ET/YR, Student’s t-test, n = 3–4 patches per point). Arrows indicate intervals for the traces in (A). (C) Time constants of recovery after trapping plotted against the trapping interval for A665C, Y768R, A665C, and E713T (continuous lines are exponential fits, n = 3–4 patches per point). Arrows indicate intervals for the traces in (A).
Resting state desensitization is rapid and reversible

The differences between the NW-bound, putative desensitized structure (PDB: 4U4F) (28), the apo structure (13), and the resting-like state bound by the bulky competitive antagonist MPQX (29,30) are subtle, with little to no change in the distance between subunits A and C at A665C between 7 and 9 Å. All these structures have preserved intradimer active D1-D1 interfaces, and previous work showed that unbound and singly bound receptors can undergo desensitized transitions (22,31). This raises the question as to whether resting receptors can be trapped at the lateral interface as they desensitize. To investigate this point, we studied the formation of an interdimer cross-link between the residues A665C in the absence of the ligand. Initial experiments suggested that cross-linking was effectively instantaneous, using our regular protocols. To make the briefest application of Cu:Phen possible, we programmed a ramp stimulation for the perfusion tool, as illustrated in Fig. 7A. With this protocol, we could apply oxidizing conditions for less than 5 ms. Exposing the mutant A665C to oxidizing conditions (10 μM Cu:Phen) in the absence of any agonist, we observed trapping of the A665C mutant that developed over ~10 ms, reducing the fraction of active receptors by ~20% (Fig. 7B). This trapping was comfortably the fastest that we have observed in the AMPA receptor (18–20). To investigate whether this trapping requires breaking of the active dimer interface, we constrained the interface by exposing patches to 100 μM CTZ. There was a ~1000-fold delay in the formation of the diagonal disulfide between subunits A and C in resting state in the presence of CTZ, with a reduction of the AF of 20% only after a 100-s application of Cu:Phen (Fig. 7, C and D). The time constant of recovery after trapping in resting conditions was 380 ± 150 ms, but for resting + CTZ, the recovery was much faster (30 ± 5 ms) (after 10 s in Cu:Phen). These results show that the receptor transits between active (D1 intact) and desensitized-like (D1 broken) states on a more rapid timescale at rest than previously thought (22).

DISCUSSION

The idea that the interaction of the neurotransmitters with receptors encompasses more than a simple binding interaction followed by activation was a major step in receptor theory (32). It is now known that receptors in the brain have multiple active and inactive “desensitized” states. For example, the acetylcholine receptor presents at least four desensitized states (33), whereas for the BK potassium channel, which has multiple Ca2+ binding sites, desensitized configurations could represent up to 120 different states (34). Previous work has emphasized that native AMPA receptors, likely in complex with auxiliary subunits, have multiple desensitized states (2,35) as do TARP-associated
receptors expressed recombinantly (36). For GluA2, auxiliary subunits γ-4 and γ-8 and CKAMP44 slow recovery from desensitization, whereas TARPs γ-2 and γ-3 have little effect (37–39). Multiple components in the recovery of AMPA receptors expressed in cell lines are detectable but less pronounced (31,40). However, the GluA2 (Q) homomer that we worked on here, and for which the majority of structural studies were completed, was until now widely reported to have a rapid and monotonic recovery from desensitization (see Fig. 3) (27,41,42).

We mapped inactive conformations of GluA2 over timescales from milliseconds to minutes with metal bridges and disulfide bonds that trap transient intersubunit interactions. This approach facilitated the detection of three distinct classes of desensitized states with glutamate and a fast, inactive state at rest (Fig. 6 A). Recovery from trapping was done in zero glutamate and reducing or chelating conditions; these measurements are closest to recovery from desensitization, and the time constants obtained provide estimates of the stability of the cross-linked desensitized state and transitions out of it. In the same experiments, we measured the time constants for the onset of the trapped states in steady 100 μM glutamate. Although in these conditions receptors are predominantly desensitized, these estimates involve receptors cycling through the desensitized and possibly other states.

We observed a positive correlation between the time of application of oxidizing conditions and the time constant of recovery after trapping for disulfide bonds at the A-C subunit interface and lateral zinc bridges. The simplest explanation is that prolonged exposure drives entry to at least two desensitized states, and the least frequently accessed are the most stable (see Fig. 8 A). The recovery from trapping in these desensitized states was dramatically slower than for the same mutants trapped in active states (18,19). Most surprisingly, by favoring slow recovery by introducing mutations in the D2 lobe of the LBDs, we could access a further, conformationally distinct, stable desensitized state that was immune to cross-linking at the A-C interface and that may or may not be physiological. It is tempting to consider these three distinct states as progressively profound conformational changes in the LBD layer, with more stable
desensitized states corresponding to those seen in some structural biology experiments. In these experiments, ligand exposures are for technical reasons in minutes or hours. In Fig. 8 B, we outline a scheme to link the states identified from their cross-linking behavior to possible LBD arrangements. The conditions for cross-linking were specifically chosen to highly enrich desensitized conditions at the expense of resting and open receptors. However, we cannot be certain that our cross-links identify multiple desensitized states; they might instead act preferentially on transitions either into or out of desensitization. However, the mutants with more stable desensitized states slow down recovery from trapping, without slowing the adoption of deeply trapped states (Fig. 4)—the opposite of what would be expected if transitions alone were responsible for trapping. The detection of multiple time constants in the recovery of GluA2 WT, as predicted from the multiple states detected in our cross-linking data, provide good evidence against only transitions being involved in trapping.

Although cross-links do not define geometry uniquely, we note that they do report a minimum level of complexity in the conformational and dynamic space of GluA2. We identified three recovery time constants for WT GluA2 (~20, 100, and 1 s; from Fig. 5) that may correspond to the “fast,” “deep,” and “protected” classes (Fig. 8 B). Assuming this relation would imply that disulfides and zinc bridges trap these states ~100 times slower than the native states are entered (at the concentrations of Cu:Phen and Zn that we used) and extend the lifetime of the trapped states by ~100-fold. A back of the envelope calculation applying this logic to the observed time constants for the resting state trapping (entry ~10 ms, lifetime ~400 ms) would give the true resting state D1 dimer desensitization with a time constant for an entry of ~100 µs and a lifetime of ~4 ms. Providing some support for these estimates is the separate observation that trapping with bifunctional methanethiosulfonate (MTS) reagents can be accelerated 50-fold by simply increasing the reagent concentration (21). In this and other studies, we typically trap receptors in gentle (slow) conditions to reduce nonspecific cross-linking.

Even though prolonged oxidation can lead to promoting trapping by disulfides in stable inactive states, key weaknesses of this line of approach are that the presence of the bridges themselves could drive nonphysiological conformations, and the trapping bridges necessarily contribute to the lifetime of the trapped states. To address these points, we exposed WT GluA2 receptors to long applications of glutamate in two-pulse protocols and could detect slow components of recovery when we used conditioning pulses of 800 ms or longer. About one third of the population recovered either with an intermediate recovery rate ~4 times slower than the majority or on a timescale longer than the pulse protocol (>1 s). With the brief conditioning pulses that we and other investigators have routinely used (Fig. 3; (27,40,41)), these slow components are either very small or absent. GluA1 recovery could be fit by multiple single exponential functions (40). GluA2 has a steeper recovery profile than GluA1 and needs Hodgkin-Huxley type

FIGURE 7 Rapid resting state desensitization. (A) Using the four-barrel fast perfusion system and switching from barrel 2 to 4, we applied oxidizing conditions for intervals as brief as 5 ms, with a voltage ramp command to the piezo lever (right panel). The open tip junction current shows that the pipette spends between 5 and 30 ms in the third barrel outflow (blue). Dashed lines indicate switch times for the 20-ms exposure. (B) Patch-clamp experiments showing Cu:Phen (10 µM) exposures of 5 ms (left panel) and 20 ms (right panel) to the mutant A665C in the resting state. Arrows indicate reduction of the current after modification. (C) Patch-clamp experiments show applications of Cu:Phen 10 µM for 40 ms (left panel) and 3 s (right panel) to the mutant A665C in resting state with CTZ (100 µM). The inset shows the current profile during recovery; the arrow indicates reduction of the current after trapping. (D) The AF of receptors after oxidization in the desensitized state (continuous lines, exponential fit) is plotted against the trapping interval. Arrows indicate intervals for the traces in (B and C). The probability of no difference between the AF after trapping in resting state without and with CTZ for A665C after 10 s of application of oxidizing conditions was 0.003 (n = 3–6 patches per point). To see this figure in color, go online.
functions, usually fixed to have a slope exponent of 2. Intriguingly, a good description of the early phase of recovery required slope exponents >2 (see Figs. 5 and S1; Table S1). Fixing the slope to 4 offered a marginal improvement in the goodness of fit compared to a slope of 3. This observation is consistent with three or more independent particles being involved in the first recovery phase (24). A more qualitative observation is that, for conditioning pulses of 5 s or longer, we always observed rundown of the response. Although there are multiple explanations of rundown (see Materials and Methods), one source could be the irreversible accumulation of receptors into nonfunctional states. Quantitative measures of receptor conformation during such experiments (for example, from spectroscopy) may be able to provide information in this regard.

The desensitized state structure stabilized by GSG1L does not support the formation of the disulfide bond between A665C residues in subunits A and C (Fig. 1). This observation reinforces the idea that there are multiple desensitized states with common attributes but that some aspects of LBD geometry might be unimportant. Desensitized states in the AMPA receptor may correspond to any number of configurations in which the braced, active dimer arrangements are absent. Dissociation of a single active dimer is enough to desensitize the receptor (31). However, the overall configuration of the four LBDs might otherwise be compact. Auxiliary proteins with very different geometries (for example, TARPs and Shisa variants) seem to have distinct effects on the lifetime of the AMPA receptor desensitized state (43–45), and this could be because they stabilize different LBD arrangements in desensitized states. We cannot exclude the possibility that conformational changes could in principle bring the cysteines from subunits B and D (found on the outer flanks of the receptor in resting and activated states) into potential cross-linking positions. The rotations needed would be even more extreme than those reported for the kainate receptor. Likewise, for a zinc bridge to form diagonally (for example, between subunits B and D), an unprecedented lateral shift of the two LBD dimers would be needed. The requirement to avoid all simpler cross-linking geometries to reach such extreme states was previously discussed (21). Although experiments in heteromeric receptors (for example, GluA1:A2) could in principle provide more insight to subunit interactions, in our hands, cysteine cross-links bias heteromer assembly, and we could not so far generate zinc bridges at the diagonal A-C subunit interface.

Structures of GluA2 in the apo ligand free (13) and the MPQX-bound state (29,30) do not support a contact between subunits A and C in the resting state. Yet inactive states could be trapped by the A665C disulfide bond at “rest” within 10 ms. Therefore, these measurements place an upper limit of the timescale of latent rearrangements of the dimer interface. The S729C mutant forms a precedent for these observations (22), but for that mutant, resting state trapping was ~700-fold slower. Using CTZ, to stabilize the
D1-D1 interface, we could massively slow trapping in the resting state, suggesting that D1 dissociation is rapid and regular (46). It is likely that reformation of the interface at rest is at least as fast, otherwise the majority of receptors should simply desensitize upon binding glutamate. It is likely that the apo state is mobile and that the stability of the D1 dimer interface varies between flip and flop isoforms, impacting AMPA receptor kinetics (47). A second, less likely, effect of CTZ would be to reduce the conformational dynamics of the lower lobe of the LBD (where A665C is located) in the resting state. In the presence of partial agonists, twisting motions and other degrees of freedom have been reported (48–50), and these may be affected by CTZ binding. Even though the mutations used in this study (like the ET/YR mutant) were outside the D1 dimer interface, they could have their own effect on resting state dynamics. But any knock-on effect to alter cross-linking should be quite limited because the inefficiency of our cross-linking meant, on average, many cycles of receptor desensitization must occur before trapping, diluting any influence of resting state dynamics.

Our results are consistent with a previous report that shows differences in the rates of entry and recovery from desensitization in A665C receptors between oxidizing and reducing conditions (26). The introduction of tryptophan in position A665 produces a slow rate of recovery from desensitization of more than sevenfold. Therefore, a movement involving a close contact between subunits A and C at the loop between helices F and G of the LBDs appears necessary for fast recovery from desensitization.

Although the GluA2 homomeric receptor has been taken as the best structural model for synaptic receptors since the first structure was solved in 2009 (51), recent work and common sense suggests limitations in this regard. Whether or not receptors actually desensitize at most synapses remains controversial. Many synaptic receptors are likely triheteromers that include TARPs (52) and probably have richer desensitization behavior. These same limitations clearly apply to our work as well, which by its nature exploits a series of structures of the homomeric receptor.

In summary, our experiments provide insight into the conformations and kinetics of AMPA receptor desensitization. Particularly, this work suggests a hierarchy of AMPA receptor desensitized states. It seems likely that the more dispersed conformations of the extracellular domains detected in some structural biology experiments correspond to slowly attained states that may occur during brain injury or by receptors during biogenesis outside of fast excitatory synapses. However, compact desensitized arrangements of the LBD layer, probably like those stabilized by auxiliary proteins observed in cryo-EM experiments, are rapidly attained by AMPA receptors within milliseconds and on a timescale relevant for desensitization in the brain.

SUPPORTING MATERIAL

Supporting Material can be found online at https://doi.org/10.1016/j.bpj.2020.05.029.

AUTHOR CONTRIBUTIONS

All authors performed experiments, analyzed data, and wrote the article.

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