Structural and Functional

Analysis of Two New Positive Allosteric Modulators of GluA2 Desensitization and Deactivation


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Abbreviations: AMPA (alpha-amino-3-hydroxy-5-methyl-4-isoxazole propionic acid), CMPDA (a phenyl-1,4-bis-alkylsulfonamide), CMPDB (a phenyl-1,4-bis-carboxythiophene), deact (deactivation), des (desensitization), CTZ (cyclothiazide), CX614 (pyrrolidino-1,3-oxazino benzo-1,4-dioxan-10-one), LBC (ligand-binding core), PDB (Protein Data Bank)
ABSTRACT

At the dimer interface of the extracellular ligand-binding domain of AMPA receptors a hydrophilic pocket is formed that is known to interact with two classes of positive allosteric modulators, represented by cyclothiazide and the ampakine CX614. Here we present structural and functional data on two new positive allosteric modulators of AMPA receptors, CMPDA and CMPDB. Crystallographic data show that these compounds bind within the modulator-binding pocket, and that substituents of each compound overlap with distinct moieties of cyclothiazide and CX614. The goals of the present study were to determine 1) the degree of modulation by CMPDA and CMPDB of AMPA receptor deactivation and desensitization; 2) whether these compounds are splice isoform-selective; and, 3) whether predictions of mechanism of action could be inferred by comparing molecular interactions between the ligand binding domain and each compound to those of cyclothiazide and CX614. CMPDB was found to be more isoform-selective than would be predicted from initial binding assays. Importantly, these new compounds are both more potent and more efficacious, and also may be more clinically relevant, than the previously described AMPA receptor modulators.
INTRODUCTION

Alpha-amino-3-hydroxy-5-methylisoxazole-4-propionic acid (AMPA) receptors mediate the initial peak of excitatory postsynaptic potentials and are critical for the strengthening and weakening of synapses that underlies the cellular basis of learning and memory. Positive allosteric modulators of AMPA receptors act by enhancing normal AMPA receptor activity to mimic long-term potentiation (Staubli et al., 1994a) and prolong open-channel time by slowing or preventing channel closure (Arai et al., 1996a; Partin et al., 1996; Suppiramaniam et al., 2001; Vyckicky et al., 1991; Yamada and Rothman, 1992). Slowing the termination of a glutamate-evoked response enhances excitatory postsynaptic potentials by increasing ion flux through AMPA receptor channels.

Initial studies indicated that potentiating AMPA receptors could alleviate cognitive deficits (Arai et al., 1996b; Staubli et al., 1994b). More recently developed compounds such as the benzamides improve normal cognitive function, as well as cognitive function impaired by aging or schizophrenia (Goff et al., 2001; Ingvar et al., 1997). However, in animal models or human studies several of these compounds either were not efficacious, or exhibited adverse side-effects, demonstrating the challenge to discover compounds which are both effective and tolerated (Ward et al., 2010a). Reduced glutamatergic signaling is involved in Alzheimer’s disease, schizophrenia, attention deficit hyperactivity disorder, narcolepsy, autism and Parkinson’s disease. Understanding the molecular mechanism of positive modulators that enhance glutamatergic signaling will advance the development of more efficacious treatments for these disorders (Black, 2005; Lynch, 2002; O’Neill et al., 2004).

Understanding the molecular mechanism of allosteric modulation is contingent upon understanding the fine details of AMPA receptor gating, and in particular, the decay of the AMPA receptor-mediated synaptic response, which may be terminated by two different, agonist-dependent
mechanisms. When the exposure to glutamate is brief, AMPA receptor channels will close and
agonist may dissociate (deactivation), leaving receptors in an activatable state. Alternatively,
prolonged exposure to glutamate causes channels to desensitize. In this case, receptors occupy a
conformational state in which the bound agonist reduces the probability of channel opening. The
channel must rearrange and recover before it can be activated again (Armstrong et al., 2006;
Horning and Mayer, 2004; Robert et al., 2005; Sun et al., 2002; Weston et al., 2006; Zhang et al.,
2008).

AMPA receptors are heterotetrameric membrane proteins comprised of subunits GluA1-4 (Traynelis
et al., 2010). Receptor functional heterogeneity is expanded through RNA splicing and RNA editing,
and also through differential assembly with auxiliary accessory transmembrane proteins and
cytoplasmic proteins that interact with the carboxyl terminus (Tomita, 2010). Each subunit is
composed of four domains: an amino-terminal domain (ATD), a ligand-binding core or domain (LBC
or LBD) to which both agonists and allosteric modulators bind, the membrane spanning domains
(M1, M3, and M4) and a re-entrant pore loop (M2), and the cytoplasmic domain. Recently, it has
been shown that the quaternary structure of the membrane spanning domain has 4-fold symmetry.
In contrast, ATD and LBC domains of the channel have two 2-fold axes of symmetry such that a
dimer of dimers is created (Sobolevsky et al., 2009).

The LBC is composed of an upper lobe, domain 1, and a lower lobe, domain 2, which are brought
together when agonist binds within the cleft formed between the domains (this structure may also
be referred to as a “clamshell”). Beta-strand hinges (clamshell hinges) that connect domains 1 and
2 impart stability upon the closed-cleft conformation. Once agonist is bound, the receptor may open
and then desensitize, entailing a structural re-arrangement of the interface between the paired
LBCs of different subunits (Furukawa et al., 2005; Horning and Mayer, 2004; Jin et al., 2005; Quirk
et al., 2004; Sun et al., 2002). Alternatively, upon brief exposure to glutamate, the receptor may
deactivate by allowing the LBC cleft to re-open and glutamate to dissociate. Although time constants representing decays due to deactivation and desensitization can be determined experimentally, the structural relationship between these two processes is unclear.

Two classes of cognition-enhancing modulators for which there is structural data, the benzothiadiazides, (e.g., cyclothiazide, CTZ), and the pyrrolidinones (e.g., ampakine CX614) are thought to favor the open channel configuration by primarily slowing the onset of desensitization or rate of deactivation, respectively. CTZ predominantly modulates desensitization by preventing rearrangement at the dimer interface, thereby blocking entry into the desensitized state, with only a weak effect on deactivation of flip isoforms (Mitchell and Fleck, 2007; Partin et al., 1996; Sun et al., 2002). In contrast, CX614 slows desensitization with a profound effect on deactivation (Arai et al., 2000; Arai et al., 2002; Jin et al., 2005) (Nagarajan et al., 2001).

Recent high throughput screening assays have led to the development of a number of new positive allosteric modulators, including biarylsulfonamides (Miu et al., 2001; Ornstein et al., 2000), the series of amino indane sulfonamide derivatives (Ward and Harries, 2010; Ward et al., 2010b; Ward et al., 2010c), as well as compounds based on the structure of LY404187 that then underwent “Structure-Based Drug Design” (Jamieson et al., 2010a; Jamieson et al., 2010b). Although excellent structural, and in some cases preclinical, information has been published on these compounds, less is understood about their biophysical mechanisms of action.

Here we describe structural and functional analyses of two new potent allosteric modulators of AMPA receptors, CMPDA and CMPDB, and compare their actions on GluA2 to CTZ and CX614.
MATERIALS AND METHODS

X-ray Crystallography. Crystals of the iGluA2 flop extracellular domain were prepared using minor modifications of the methods previously described (Armstrong et al., 1998). The polypeptide sequence used for crystallization corresponds with that described as HS1S2H by Chen et al. (Chen et al., 1998), and was refolded and crystallized as described by Armstrong and Gouaux (Armstrong and Gouaux, 2000), and Chen et al. in the presence of glutamate by the hanging drop vapor diffusion method, using a precipitant solution containing 13-18% PEG 8000, 0.1 M Zn(OAc)$_2$, and 0.1 M sodium cacodylate at pH 6.5. Crystals were transferred into precipitant solutions containing 13-18% PEG8000, 0.1 M Zn acetate, 0.1 M sodium cacodylate pH 6.5, 3 mM of CMPDA or CMPDB and 3% DMSO. Crystals were incubated for 24 hours and then flash frozen in liquid nitrogen using cryoprotectant solution containing 30% PEG8000, 20% glycerol, 0.1 M Zn acetate and 0.1 M sodium cacodylate pH 6.5. Diffraction data were collected on the Advanced Photon Source beamlines 17ID & 31ID and were integrated, merged and scaled using the software package, HKL2000 (HKL Research, Inc., Charlottesville, VA). Data scaling statistics are given in Table 1. Structures were solved using rigid body refinement of starting models derived originally from the “1LBC” PDB coordinates for CTZ structure in complex with the GluA2 LBC (Sun et al., 2002) against the diffraction data. Structures of CMPDA and CMPDB were fit to clear difference density maps using Afitt (Openeye, Inc., Santa Fe, NM) software. The models were iteratively refined using CNX (Accelyrs, Inc., San Diego, CA) and Refmac (Murshudov et al., 1999) software, with manual model building carried out using the program Coot (Emsley and Cowtan, 2004). Model refinement statistics are given in Table 1. Figures were generated from protein data bank (PDB) files using Pymol (DeLano, 2002). Molecular coordinates for CMPDA and B in complex with the GluA2 LBC have been deposited in the Protein Data Bank as 3RN8 and 3RNN, respectively.
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**Molecular Biology.** Plasmids encoding cDNAs for the flip (i) and flop (o) variants of rat wild-type GluA2 were gifts of Dr. Peter Seeburg (University of Heidelberg, Germany). The “WT GluA2” DNA plasmid contains a substitution of one residue within the pore, GluA2 R<sub>607</sub>Q (QuikChange II XL Site-Directed Mutagenesis Kit, Stratagene; La Jolla, CA), which recapitulates RNA editing of the pore, to facilitate electrophysiological studies. The receptors made from this mutant form homomers more efficiently, have greater conductances, and have an inwardly rectifying current-voltage relationship (Hume et al., 1991; Verdoorn et al., 1991).

**HEK293 Cell Culture.** Human embryonic kidney 293 fibroblasts (HEK 293 cells; ATCC CRL 1573 from American Type Culture Collection, Rockville, MD) were cultured as described previously (Cotton and Partin, 2000). Cells were cultured in DMEM supplemented with 10% fetal bovine serum (Gemini Bio-Products, Inc, Calabasas, CA), penicillin/streptomycin (100 units/ml each) and 1% GlutaMax-1 (both from Gibco, Grand Island, NY). Cells were transiently transfected using FuGene 6 reagent (Roche Diagnostic Corp., Indianapolis, IN) with GluA2 (GluA2i, flip or GluA2o, flop) cDNA and enhanced yellow fluorescent protein (EYFP) cDNA (1 and 0.2 μg/35 mm dish, respectively). For some experiments, 10μM NBQX was added after 18-24 hrs.

**HEK293 Electrophysiology.** Currents were recorded 24-72 hours after transfection as described previously (Cotton and Partin, 2000). Outside-out membrane patches from transfected HEK 293 cells were held under voltage-clamp at a holding potential of -60 mV using an Axopatch 200B amplifier (Molecular Devices, Sunnyvale, CA). Synapse software (version 3.6d) (Synergy Research, Inc., Bromma, Sweden) was used on a PowerPC Macintosh computer for trace analysis and data acquisition through an ITC-16 interface (HEKA, Bellmore, NY) which included control and timing of perfusion tubing movement by a piezoelectric device (Burleigh Instruments, Fishers, NY). Responses were filtered at 5 kHz, digitized at 10-500 μsec/point. Electrodes of 2-5 MΩ were filled
with (in mM): 135 CsCl, 10 CsF, 10 HEPES, 5 Cs-BAPTA (Invitrogen, Carlsbad, CA), 1 MgCl₂, 0.5 CaCl₂, pH 7.2. 2 mM Na₂-ATP was added to the internal pipette solution immediately before recording each day. During recordings, cells were perfused continuously with extracellular control solution containing (in mM): 20 sucrose, 145 NaCl, 5.4 KCl, 5 HEPES, 1 MgCl₂, 1.8 CaCl₂ and 0.01 mg/ml phenol red, pH 7.3. Currents were evoked with test solutions containing 10 mM glutamate. Control and test solutions were perfused through quartz tubing (Sutter Instrument Company, Novato, CA). The patch pipette tip was positioned in the control solution stream near the interface between the control and glutamate containing streams. When modulators were used, they were added to both test and control solutions in the following concentrations (in µM): 100 CTZ, 100 CX614, 10 CMPDA or CMPDB. Modulator stock solutions (10 mM) were dissolved in DMSO before dilution in extracellular solutions; final DMSO concentrations were 0.3-1%. When applying glutamate in the absence of modulator, after having perfused modulator through the tubing for a previous patch, we noticed that the steady-state current responses were not completely desensitized. This indicated that the modulators were sticking to the perfusion tubing and slowly leaching into the perfusion stream. We alleviated this problem by changing the tubing after using the modulator. Consequently, this prevented us from collecting control and modulator data on the same patches. For this reason, and because of channel rundown, we do not report on differences in current amplitudes comparing control glutamate responses with those in modulator. Continuous solution flow was driven by a syringe pump (KD Scientific, New Hope, PA) at a rate of 0.2 ml/min; patches were pre-incubated in modulator for at least 2 minutes prior to switching into glutamate. No change in holding current was observed when the patches were initially brought into control solution containing modulators indicating that the new modulators do not have intrinsic agonist properties. A piezoelectric device was used to drive rapid solution exchanges of 1 ms or 500 ms to measure channel deactivation or desensitization kinetics, respectively. Solution exchange rates were determined at the end of each experiment by measuring open-tip junction currents; for this purpose the salts in the control solutions (no agonist) were diluted by 10%. Most of the decays of responses
in the presence of modulator were best fit by the sum of two exponents. Data fit with a two-
exponential function with both fast and slow components were combined into a weighted average
time constant of exponential decay based on their respective amplitude contributions (\(\%A_{Fast} \times \tau_{Fast} + \%A_{Slow} \times \tau_{Slow}\)) where \(A\) = the amplitude of each time constant, \(\tau\). Data reported as mean ± SEM
were compared using ANOVA with Dunnett’s test for multiple comparisons within each receptor
type to L-glutamate “control”. Current traces and graphs were plotted using KaleidaGraph 3.6
(Synergy Software, Reading, PA).

**Calcium Flux Assay.** The ability of CMPDA and CMPDB to potentiate glutamate-evoked influx of
calcium into HEK293 cells stably expressing human GluA2 (R607Q; Quirk and Nisenbaum, 2003)
was evaluated using Fluorescent Imaging Plate Reader (FLIPR) technology (Molecular Devices,
Sunnyvale, CA). Confluent monolayers of stably transfected cells were prepared by delivering cells
into 96-well plates at a density of approximately 60,000 cells/100 microliter well and incubating
plates at 37°C in 95% O₂/5% CO₂ overnight. The following day, the tissue culture medium in the
wells was discarded, and 50 μl of fluo3-AM dye (8 μM; Molecular Probes Inc., Eugene, Oregon; Cat
#F-1241) in HBSS buffer (Biowhittaker; Cat #12-614Q plus 3.7 mM CaCl₂ and 20 mM HEPES) was
added to each well. The plates were then incubated for 60 minutes at 25°C in the dark. After the
incubation period, the fluo3 dye was removed and 50 μl of the HBSS buffer was added to each well
and the plate was placed into the FLIPR.

Test compounds were dissolved in 100% DMSO to yield 10 mM stock solutions. The stock
solutions were diluted in buffer and tested in duplicate 10-point curves at concentrations of 0.0003-
10.0 μM. The final DMSO concentration in the assay was 1.2%. Fluorometric readings were taken
from each well following addition of the compound alone (1\textsuperscript{st} addition) and after addition of
glutamate (100 μM) (2\textsuperscript{nd} addition). Additional control wells contained glutamate (100 microM) alone
and glutamate plus a maximal concentration of cyclothiazide (100 microM) used as a positive control compound. Compound effects were determined by subtracting the background fluorescence reading of each concentration of compound alone from the fluorescence reading obtained after addition of each concentration of compound plus glutamate. These responses were then normalized to a signal window determined by the response to 100 microM cyclothiazide plus 100 microM glutamate (maximum response) and 100 microM glutamate alone (minimum response) and plotted. The data were then evaluated using non-linear curve analyses and the effective concentration yielding a 50% maximal response (EC\textsubscript{50}) value was determined by fitting the data with a four-parameter logistic equation.

Mathematical Simulations of Currents. Simulations of currents under voltage clamp to an AMPA receptor model were performed using code originally written by John Clements (Benveniste et al., 1990), revised and converted by M.B. to an Igor Pro XOP. Receptor state occupancies at each time point were determined numerically by calculating the change in each state occupancy resulting from transitions into and out of each state according to first order reaction rate kinetics as detailed in Benveniste et al. (Benveniste et al., 1990). This was done iteratively at least 20 times per time point.

The model used in this study (Figure 8A) is based on the activation and desensitization models for homomeric AMPA receptors of Robert and Howe, in which three open states of different conductance are possible, based on the binding of two, three or four molecules of agonist (Robert et al., 2005; Robert and Howe, 2003; Rosenmund et al., 1998). This model has an added transition after agonist binding which represents the conformational change between the open and closed clamshell (cleft) states. The closed clamshell state, precedes both the transition to the open state and the desensitized state. In addition, transition to desensitized states was also possible from the open state (Figure 8A). Starting values for rate constants were taken from published values for GluA2 (Robert et al., 2005) and adjusted in the new model so as to yield similar values to the
published peak and steady state EC$_{50}$ values and the time constants for the onset and recovery of desensitization. A full listing of the transitions for each model is presented in the Supplemental Material. The model was optimized to reflect experimental values for the flip isoform of GluR2. Flip and flop isoform-specific deactivation and desensitization kinetics have been simulated by others using a strategy of altering the rate of channel closing ($k_{cl}$ or $\alpha$) (Pei et al., 2007a; Pei et al., 2007b). Although we tried to replicate that finding with the present model, we were unable to model the faster flop kinetics by simply increasing the rate of channel closing. With the present model, flip and flop differences were largely replicated by simply changing the rate of onset of desensitization by 10-fold. This apparent discrepancy exemplifies the model-dependency of kinetic analyses.

Simulations and fitting and related analyses were run on a Macintosh MacBook Pro computer utilizing Igor Pro 6.1 (Wavemetrics Inc., Lake Oswego, OR).

Gibbs Free Energy Calculations – The changes in stability resulting from the manipulation of rate constants for five basic receptor states (unbound; agonist bound, clamshell open; agonist bound, clamshell closed; open channel; and, desensitized channel) were compared by converting forward and reverse reaction rates to Gibbs free energy of activation, $\Delta G^\ddagger$, according to the following equation:

$$\Delta G^\ddagger = RT \ln \left( \frac{k_B T}{h k} \right)$$

where $R$ is the universal gas constant, $T$ is the temperature (293°K), $k_B$ is the Boltzmann constant, $h$ is the Planck constant and $k$ is either the forward or reverse rate constant (Eyring, 1935; Wilkinson., 1997). Changes in energy states are relative, with the unbound receptor state assumed to have a $\Delta G^0 = 0$ kcal/mol. In Figure 9, the agonist bound closed clamshell state is represented twice on successive rows, in order to illustrate transitions from that state to either the open state or desensitized state.
All chemicals were purchased from Sigma-Aldrich (St. Louis, MO) unless otherwise indicated.
RESULTS

CMPDA and CMPDB bind between two subunits at the clamshell hinges

Two positive allosteric modulators of AMPA receptors, phenyl-1,4-bis-alkylsulfonamide (CMPDA) and a phenyl-1,4-bis-carboxythiophene (CMPDB) were identified through a calcium influx screening assay of GluA2i and GluA2o receptors. Both compounds are derivatives of a biarylpropylsulfonamide series (Shepherd et al., 2002; Vandergriff et al., 2001) but are composed of a single central aryl ring with symmetrical R-groups. The alkylsulfonamide R-groups of CMPDA and the nitrile/thiophene R-groups of CMPDB extend above the plane of the aryl ring, while only the carboxy groups of CMPDB extend below the plane of the aryl ring (Figure 1). The ability of CMPDA and CMPDB to allosterically modulate GluA2i and GluA2o receptors stably transfected into HEK 293 cells was assessed by measuring the glutamate-evoked influx of calcium using FLIPR technology. Potency estimates for the two compounds were determined by measuring the change in fluorescence each well following addition of the compound alone at concentrations of 0.0003-10.0 microM and after addition of glutamate (100 microM). The effects of CMPDA and CMPDB were determined by subtracting the background fluorescence reading of each concentration of compound alone from the fluorescence reading obtained after addition of each concentration of compound plus glutamate. For each compound, the effects of each concentration were normalized to the response to 100 microM of cyclothiazide plus 100 microM glutamate and plotted. Evaluation of the concentration-response plots revealed that the two compounds potently enhanced glutamate-induced calcium influx into HEK293 cells transfected with human GluA2i and GluA2o receptors (Supplemental Figure 1). The potency for CMPDA at GluA2i receptors was EC$_{50}$ = 45.4 ± 4.2 nM (mean ± sem; n=4) and GluA2o receptors was EC$_{50}$ = 63.4 ± 5.6 nM (n=4). The potency for CMPDB at GluA2i receptors was EC$_{50}$ = 122.5 ± 12.9 nM (n=2) and GluA2o receptors was EC$_{50}$ = 470.5 ± 3.7 nM (n=2). Results showed that CMPDA was nearly equipotent at modulating the two
isoforms of GluA2 receptors; whereas, CMPDB displayed a modest preference for the flip splice variant.

High-resolution co-crystal structures of the isolated GluA2o LBC (HS1S2H) with L-glutamate and either CMPDA or CMPDB were solved (Table 1). Like the ampakine CX614 (Jin et al., 2005), a single molecule of CMPDA or CMPDB binds at the dimer interface, within a cleft situated near the interdomain hinges. The 2-fold symmetry of each compound is coincident with the 2-fold axes that relate subunits within the extracellular domain dimer (Figure 2).

N1 and N2 of the alkylsulfonamide groups of CMPDA form hydrogen bonds with the main chain oxygens on residue Pro 494 of both protomers. O2 and O4 form hydrogen bonds with the nitrogens on Gly 731 for both protomers (Figure 3A & B). Residues Pro 494 and Gly 731 lie within the two interdomain β-strands of the AMPA receptor, making up the clamshell hinges (Hinge 1, residues 494-498; Hinge 2, residues 729-733). The two carboxy groups of CMPDB also form hydrogen bonds with the interdomain hinges through the main chain nitrogen of residue Ser 497 in Protomer B and through water-mediated hydrogen bonds with the main chain oxygen of Phe 495 of Protomer B, main chain oxygen and nitrogen of Lys 730 and main and side chains of Ser 729 of Protomer A and B (Figure 3C - E).

**CMPDA and CMPDB bind to the GluA2o ligand binding core in a space that overlaps the CTZ and CX614 Binding Sites**

Previous crystallographic studies have revealed that while CTZ and CX614 (and its parent compound, aniracetam) bind within the same region, at the dimer interface between two protomers, and interact with some of the same amino acid residues, they bind with different stoichiometries. CTZ binds with a modulator to protomer stoichiometry of 2:2, whereas CX614 (and aniracetam) has
a 1:2 binding stoichiometry (Jin et al., 2003; Sun et al., 2002). Results from the current studies show that CMPDA and CMPDB also bind with a 1:2 stoichiometry similar to CX614. However, superposition of the crystal structures of the LBC bound to all four compounds clearly demonstrates that the set of residues that interact with CMPDA and CMPDB partially overlaps with those of both CTZ and CX614. The central phenyl rings of CMPDA and CMPDB are oriented similarly to the central aromatic ring of CX614, while the alkylsulfonamide and carboxythiophene groups partially overlap the space occupied by CTZ (Figure 4).

CTZ and CX614 also have distinct effects on the deactivation and desensitization of AMPA receptors that are splice isoform-specific. For example, CTZ is more potent and efficacious in blocking desensitization of flip versus flop isoforms of AMPA receptors, but only weakly modulates deactivation of flip splice variants with little effect on flop-expressing receptors. In contrast to CTZ, CX614 is more efficacious at attenuating desensitization of GluA2o receptors but also significantly slows deactivation in both GluA2i and GluA2o receptors (Arai et al., 2000; Jin et al., 2005; Mitchell and Fleck, 2007). Because the binding domains of CMPDA and CMPDB within the LBC are shared with both CTZ and CX614, we initially investigated whether the physical binding interactions could be used to predict the effects of CMPDA and CMPDB on the gating of GluA2i and GluA2o receptors.

A detailed analysis of the crystallographic structures of the four compounds (CTZ, CX614, CMPDA and CMPDB) is necessary to assess the extent to which all four compounds interact with the same sets of residues but through differing types of interactions (Figure 5). For example, Pro 494 forms direct hydrogen bonds with CMPDA and CTZ, water-mediated bonds with CX614, and has a hydrophobic interaction with CMPDB; Ser 497 forms water-mediated hydrogen bonds with CMPDB, water-mediated hydrogen bonds with CX614, and has a hydrophobic interaction with CMPDA. Ser
729 forms a direct hydrogen bond with CTZ, water-mediated bonds with CMPDB and CX614, and has a hydrophobic interaction with CMPDA.

Most of the residues that interact with all four compounds reside in the hinge β-strands that connect the upper lobe (Domain 1) to the lower lobe (Domain 2) of the ligand-binding clamshell. The interaction between CMPDA or CMPDB and the hinges might suggest that they act to stabilize the closed-clamshell conformation of the protein, as proposed for CX614, resulting in the slowing of receptor deactivation. Stabilization of the clamshell closed conformation would enhance agonist-binding. This has been demonstrated for CX614, which enhances AMPA and fluorowillardiine binding, and is in contrast to the effects of CTZ, which reduces agonist-binding (Kessler and Arai, 2006). However, because CMPDA and CMPDB also have moieties that overlap with CTZ, modulation of deactivation might be superseded by effects on dimer interface stability. In addition, the crystal structure is static, representing a single snapshot of how CMPDA and CMPDB can interact with the ligand binding domain. It is difficult to predict the extent to which the efficacy of compound modulation may differ depending on direct interactions with a residue versus making indirect, water-mediated contacts that may be perturbed by competing water molecules. We therefore assessed the functional effects of CMPDA and CMPDB on the deactivation and desensitization processes of GluA2 flip and flop receptors and compared these effects with those of CTZ and CX614.

**Functional analysis of CMPDA and CMPDB reveals potent modulation of deactivation**

To assess the effects of the modulators on channel activity, we measured responses evoked by ultrafast perfusion of glutamate to outside-out membrane patches from HEK293 cells expressing recombinant rat GluA2 AMPA receptors. Modulator effects on deactivation and desensitization were assessed using short (1 ms) and long (500 ms) pulses of glutamate (10 mM), respectively in the
presence and absence of concentrations of each of the four modulators that were predicted to be at “saturating” doses, based either on electrophysiological data (CX614 and CTZ) or FLIPR data (CMPDA and CMPDB). Both the flip (i) and flop (o) isoforms of rat GluA2 were tested, in order to determine whether the modulators showed isoform selectivity.

Consistent with previous reports (Arai et al., 2000) (Jin et al., 2005) (Mitchell and Fleck, 2007), CX614 preferentially slowed deactivation of GluA2o receptors, whereas CTZ had no effect on deactivation of either GluA2i or GluA2o receptors (Figure 6, Table 2). Similar to CX614, CMPDA slowed the rate of deactivation of GluA2o receptors approximately 2-fold, but had no effect on GluA2i receptor deactivation. In contrast, CMPDB attenuated the rate of deactivation of both GluA2i and GluA2o receptors, although to differing degrees. The time constants of deactivation for GluA2i and GluA2o receptors were approximately 7- and 5-fold slower in the presence of CMPDB (Figure 6, Table 2). Collectively, these results indicate that CMPDA was modestly flop-selective and CMPDB was generally more flip-selective in attenuating receptor deactivation.

As we have reported previously (Kessler et al., 2000; Partin et al., 1995; Quirk and Nisenbaum, 2003), CTZ eliminated desensitization of GluA2i receptors and significantly slowed the rate of desensitization of GluA2o receptors (Figure 7, Table 2). CX614 had similar effects on the extent of desensitization of GluA2i and GluA2o receptors such that the degree of desensitization (steady-state-to-peak) was approximately 65%. Results from the present study also showed that CMPDA robustly modulated receptor desensitization of both the flip and flop isoform of GluA2 receptors, virtually blocking the macroscopic onset of desensitization similar to the effects of CTZ on GluA2i receptors (Figure 7, Table 2). CMPDB was as effective as CX614 on flip receptors (although it was not as effective as CTZ in blocking desensitization), but markedly less effective on flop receptors. The waveform of the decay was unusual in that onset of desensitization in the presence of CMPDB was similar to control, but the remaining large steady-state current for GluA2o reflect that the
equilibrium desensitization differed. CMPDA and CX614 were non-selective for flip and flop isoforms, whereas CMPDB and CTZ exhibited a greater block of desensitization for the flip isoform.

**Mathematical modeling of allosteric modulation of AMPA receptor gating.**

Computer simulations of AMPA receptor gating have been used successfully to facilitate interpretation of complex electrophysiological data (Benveniste et al., 1990) (Clements and Westbrook, 1991; Edmonds et al., 1995; Kessler et al., 1996; Mitchell and Fleck, 2007; Partin et al., 1996; Raman and Trussell, 1992; Vyklicky et al., 1991), including a widely accepted model that was recently developed and refined in a series of papers from the Howe laboratory (Robert et al., 2005; Robert and Howe, 2003; Zhang et al., 2008).

For the present studies, we were interested in understanding whether we could distinguish different mechanisms of modulation of receptor deactivation and desensitization. According to the Howe model, the macroscopic rate of deactivation would largely be determined by agonist affinity and/or by channel closing (α), whereas the macroscopic rate of desensitization would be governed by the onset and recovery from desensitization (δ, γ). Because crystal structures indicate that modulators of AMPA receptor desensitization may affect binding cleft stability (Jin et al., 2005), we modified the Howe model to explicitly account for agonist binding (k_f) and dissociation (k_-f), as well as clamshell closing (CC) and opening (CO) (Figure 8a). This model allows for sequential, independent agonist binding and cleft closing, which then leads to either channel opening or channel desensitization. We adjusted the parameters such that analysis of simulated currents yielded EC_{50} values at peak and steady state that were close to those of experimental data for GluA2 (Robert et al., 2005). For example, we increased the rate of β two-fold, and the rate of γ three-fold. These adjustments also
yielded time constants for the onset of and recovery from desensitization consistent with ours and other experimental data (Figures 8b, 8c), further validating the model.

Utilizing simulations, we next wished to discern if the effects of AMPA receptor modulators could be predicted by changes in isolated rate constants in our model. The manipulation of a single rate constant can influence the stability of receptor states in addition to the manipulated transition. For this reason, we have also included Gibbs free energy diagrams in Figure 9. Since the binding of modulator is not explicitly modeled, the predicted results represent what would happen in the presence of a saturating concentration of modulator. The limits of solubility or availability of a modulator is a persistent technical issue for the current study; subsaturating concentrations result in partial effects on deactivation or desensitization. Thus, we are only sure of modulator potency when desensitization is totally blocked. Slowing the transition from the closed clamshell conformation to the open clamshell conformation by 100-fold (“Slow CO”, Figure 9), produced a 2.4-fold slowing of in $\tau_{\text{Deact}}$, but also produced a slower low amplitude decay, in response to a simulated 1 ms pulse of 10 mM glutamate (Figure 9, top row). The rate of onset of desensitization could be examined by simulating a 500 ms pulse of 10 mM glutamate (Figure 9, bottom row). This same manipulation of the CO rate constant also yielded a decrease in equilibrium desensitization, a phenomenon that was not observed with any of the modulators tested (Figure 7).

Slowing the channel closing rate $\alpha$ by 20-fold, caused an 4.4-fold decrease in $\tau_{\text{Deact}}$ and a 68% increase in the steady state current indicating partial block of desensitization without changing the rate of onset of desensitization (“Slow $\alpha$”, Figure 9). These results are reminiscent of the effect of CMPDB on the flip isoform. Decreasing the rate of entry into the desensitized state $\delta$ by 100-fold had almost no effect on the decay of current after a 1 ms glutamate pulse, but almost completely blocked desensitization (“Slow $\delta$”, Figure 9). CTZ can block desensitization of the flip isoform.
(Figure 7) while only weakly modulating deactivation (Figure 6). These simulations suggest that this may occur by destabilizing the desensitized conformation.

Manipulation of a single rate constant could not reproduce experimental results for CMPDA on either AMPA receptor isoform. Thus, we proceeded with the simultaneous manipulation of two rate constants. Decreasing $\alpha$ by 20-fold and $\delta$ by 100-fold produced a 4.6-fold slowing of $\tau_{\text{Deact}}$ and complete block of desensitization ("Slow $\alpha$ & $\delta$, Figure 9). However, similar results could be obtained by reducing the CO transition by 10-fold and the $\delta$ transition by 100-fold ("Slow CO & $\delta$, Figure 9).

**DISCUSSION**

We describe here two new positive allosteric modulators of AMPA receptors: the bis-alkylsulfonamide CMPDA and the bis-carboxythiophene CMPDB. Analysis of co-crystals of each compound with L-glutamate and the GluA2 flop ligand binding core reveal that both compounds have overlapping binding sites with CTZ and CX614, within the previously described modulator binding pocket located at the inter-dimer interface and the clamshell hinges. Our studies suggest that this pocket is a powerful target for drug discovery of compounds that modulate AMPA receptors, as well as being a possible site of action for endogenous metabolites (Prescott et al., 2006).

An initial concern about developing modulators of AMPA receptors was the perceived need to increase their affinity, as well as improve their subunit and splice isoform selectivity. In this regard, CMPDA and CMPDB provide important new insight. CMPDA and CMPDB effectively block desensitization at lower concentrations than some of the previously described modulators (Figures
6 and 7), indicating that the site is amenable to the development of new, therapeutically-relevant compounds.

Recently, it was proposed that the allosteric modulatory site can be delineated into five overlapping subsites at the dimer interface between subunits (Ptak et al., 2009). The central subsite, A, lies parallel to the axis of symmetry between the protomers, and is occupied by the planar rings of aniracetam, CX614 and the biaryl compounds. Hinge residues (connecting Domain 1 and Domain 2 of one protomer), particularly Pro 497, Ser 497, and Ser 729 form hydrogen bonds with these compounds, and through these interactions are thought to stabilize closed clamshell conformations (Jin et al., 2003). Subsite B, or B' on the opposing protomer, is an exposed, hydrophilic pocket formed by residues Tyr 424, Phe 495, Ser 497, Lys 763 and Ser 729; while subsite C (C') is a deep, hydrophobic pocket lined by residues Ile 481, Lys 493, and Leu 751. CMPD A fully blocks desensitization in both flip and flop isoforms and binds directly to subsites A and C. Residues that form subsite C play an important role in dimer interface stability, and may be critical for dimer interactions that permit receptor desensitization (Horning and Mayer, 2004; Sun et al., 2002). CMPD B, which binds to subsites A and C but also has extensive interactions in subsite B, shows surprising isoform selectivity for blocking desensitization that is not predicted from its binding characteristics (Figure 7). This may suggest that modulator binding in subsite B can influence isoform selectivity through direct interaction or by destabilizing interactions through subsite C. Consistent with this, CTZ preferentially blocks desensitization of the flip isoform through the interaction of its N4 substituent with the Ser/Asn site on helix J (Ptak et al., 2009; Sun et al., 2002). Although Ser/Asn 750 lies between the B and C subsites, the asparagine limits access to the C subsite. The occupancy of CMPD A and B may also be compared to that of some of the newly-described positive allosteric modulators, such as compound 7a that was characterized in Ward et al., 2010c. Similar to CMPD A, 7a has a phenyl moiety that occupies site A and a trifluoromethyl group that interacts with the hydrophobic pocket of subsite C, and therefore would be predicted to
impede desensitization as its related compound, 9a, was shown to do (Ward et al., 2010c).

Additional insight has been garnered from the “hybridization” studies of Jamieson et al. (Jamieson et al., 2010a; Jamieson et al., 2010b), which demonstrate the chemical synthesis of a molecule that hybridizes structural features from LY404187 and compound 1 derived from a high throughput screen. Although the biophysical attributes of the hybridized structure have not yet been published, functional screens indicate that a degree of flexibility between the moieties that occupy the central subsite and the two pockets (A and C), is critical to potent modulation (Jamieson et al., 2010b).

Co-crystals of the flip and flop interface of the GluA2 LBC with agonist are not markedly different (Ahmed et al., 2009), suggesting that the modulator bound crystal structure may not be the dominant conformer for one or both isoforms. To pursue this, we performed a site-directed mutational analysis of residues in GluA2i and GluA2o that were predicted to contribute to the binding site based upon their proximity to the modulator in the crystal structure. Mutations of residues found closest to each of the different modulators did, in fact, have the greatest impact on the ability of the modulator to block receptor desensitization (manuscript in preparation).

The functional implications of the selectivity of CMPDB may be significant. Regarding the selectivity, one of the concerns about using AMPA receptor positive modulators clinically is our insufficient understanding of how the balance of inhibition and excitation in neuronal circuitry is changed by enhancing AMPA receptor activity at excitatory synapses, which belong to either excitatory or inhibitory postsynaptic neurons. For instance, the CA1 region of the hippocampus contains pyramidal neurons that are excitatory but also receive input from inhibitory interneurons which are themselves activated by excitatory input. Effects of ampakines CX516 and CX546 (which are slightly flop-selective) on excitation have been shown to outweigh their effects on inhibition, while CTZ seems to have a similar effect on both pyramidal cells and interneurons (Xia et al., 2005). Importantly, principal neurons express AMPA receptors composed predominantly of flip
isoforms, whereas interneurons express AMPA receptors with a substantial flop component (Geiger et al., 1995). Thus, CMPDB would be predicted to enhance the excitatory component of principal neurons, while not effectively modulating the AMPA receptors on inhibitory cells, whereas CMPDA would affect both types of cell equally. This isoform selectivity, together with the potent modulation by CMPDB of deactivation rather than desensitization, makes it an ideal candidate for further drug discovery efforts. Future studies in an *in vitro* brain slice preparation using CMPDB could provide important new information relevant to the development of the AMPA receptor complex for therapeutic benefit.

**Insight into the molecular mechanisms of allosteric modulation using modeling**

*Figures 6* and *7* clearly show differential effects of modulators on deactivation and desensitization which can also be isoform-specific. However, without employing single channel electrophysiology techniques, we can only measure *macroscopic* rates of modulation. Our interpretation of the macroscopic data is confounded by the fact that changes in desensitization rates can influence the macroscopic rate of deactivation, and changes in deactivation rates can affect macroscopic rates of desensitization. Furthermore, the structural determinants that alter deactivation (the cleft and clamshell hinges) spatially overlap with the structural determinants of desensitization (the dimer interface).

The goal of kinetic modeling is to test different mechanisms of action. In many cases, this also provides a way to utilize macroscopic data to support or refute mechanisms with microscopic rates and transitions, in the absence and presence of drugs. The unique features of our kinetic model over the well-established Howe model (Robert et al., 2005; Robert and Howe, 2003) are that it explicitly incorporates clamshell opening and closing, as well as allows for ligand-binding transitions between open states (we could think of no valid reason why channel opening prevents binding access to other independent subunits). The simulated data presented suggest that these alterations
did not invalidate the Howe model, but rather allowed us to specifically investigate how opening and closing of the clamshell influenced deactivation and desensitization kinetics. Although these four modulators all bind to the interface between LBC protomers, could we reproduce their effects on desensitization and deactivation by manipulating the same transitions for each drug? In fact, interpretation of the modeling data suggests that the drugs are mechanistically quite different depending upon how they interact with each of the five subsites. To a first approximation, CTZ modulation can be simulated by slowing $\delta$, consistent with its full occupancy of subsite C, residues of which play an important role in hydrogen and salt bridge interactions which are broken to allow receptor desensitization (Horning and Mayer, 2004). CMPDB can be simulated by slowing $\alpha$, consistent with its occupancy of subsites A and B, through which CMPDB interacts with hinge domain sidechains and backbone moieties. CMPDA can be simulated by slowing $\alpha$ and $\delta$ or CO and $\delta$.

A second goal was to model explicitly whether modulators of deactivation act primarily on the rate of channel closing ($\alpha$) (Partin et al., 1996; Suppiramaniam et al., 2001; Vylicky et al., 1991) or clamshell re-opening (CO) (Jin et al., 2005). Unfortunately, the model did not distinguish between direct effects on clamshell stability through interactions with the hinges (slowing of CO) or allosteric effects on channel gating through dimer interactions (slowing of $\alpha$). We might expect that slowing CO and $\delta$ might be a more likely mechanism for modulator action because the slowing CO and $\delta$ are associated through their common closed clamshell state, whereas affecting $\alpha$ and $\delta$ are more likely two independent processes. Yet, aniracetam has been shown to increase single channel open times in native AMPA receptors in cultured neurons, indicating that two separate effects of modulator are possible (Vylicky et al., 1991). Modulation by CX-614 has also been simulated by slowing $\alpha$ using the original Howe model (Mitchell and Fleck, 2007; Robert et al., 2005; Robert and Howe, 2003). Aniracetam and CX614 bind primarily within the A subsite of the allosteric modulatory
site at the dimer interface and have a strong effect on deactivation kinetics but a weaker effect on
desensitization (Figures 6 and 7). This evidence, together with the modeling data (Figure 9),
suggests that the hinge region coinciding with subsite A may modulate deactivation by slowing
\( \alpha \). Another way to look at this issue is to ask if stabilizing the closed clamshell conformation could
defully explain efficacious block of desensitization and a slowing of deactivation. However, our
modeling data suggest that only destabilization of the desensitized state can easily produce non-
desensitizing currents (Figure 9) and that stabilizing the agonist bound closed cleft state by slowing
CO is not sufficient to reproduce our findings with the various modulators (Figure 7).

Future experiments directed at understanding the molecular mechanism of existing allosteric
modulators, as well as the discovery and development of new modulators, will also have to take into
account the important contributions that accessory subunits make towards determining the efficacy
and potency of modulation (Tomita et al., 2006). In general, TARPs such as stargazin shift
modulator efficacy dose response curves leftward, significantly increasing the activity of the AMPA
receptors they associate with, and thereby enhancing the postsynaptic responses elicited.

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**Authorship Contribution**

Participated in research design: Timm, Benveniste, Weeks, Nisenbaum & Partin
Conducted experiments: Timm, Benveniste, Weeks & Partin
Contributed new reagents or analytic tools: Benveniste & Nisenbaum
Performed data analysis: Timm, Benveniste, Weeks, Nisenbaum & Partin
MOL #70243

Wrote or contributed to the writing of the manuscript: Timm, Benveniste, Weeks, Nisenbaum & Partin

Other: Benveniste & Partin. Acquired funding for the research.
References


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Footnotes

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**Figure Legends**

**Figure 1.** Chemical structures (top) and omit electron density maps (bottom) of two positive allosteric modulators of AMPA receptors, **a**, Bis-Alkylsulfonamide 506091 (CMPDA) and **b**, Bis-Carboxythiophene 2152080 (CMPDB). Omit density map for CMPDA (A) was calculated using |F₀| - |Fₑ| coefficients, view is shown perpendicular to the two-fold axis. A simulated omit density map for CMPDB is shown (B).

**Figure 2.** CMPDA and CMPDB co-crystals with GluA2o HS1S2H and L-glutamate. **a**, One molecule of CMPDA (magenta CPK) binds within the dimer interface at the clamshell hinges. View looking down the two-fold axis. Glutamate (GLU) is shown as red CPK. HS1S2H is shown in cartoon representation. **b**, View of CMPDA perpendicular to the two-fold axis, HS1S2H is shown in surface representation. **c**, One molecule of CMPDB (pink CPK) binds within the dimer interface at the clamshell hinges. View looking down the two-fold axis. Glutamate is shown as red CPK. HS1S2H is shown in cartoon representation. **d**, View of CMPDB perpendicular to the two-fold axis, HS1S2H is shown in surface representation. D1 and D2, Domains 1 and 2, respectively. Coordinates have been submitted to the Protein Data Bank as 3RN8 and 3RNN, respectively.

**Figure 3.** Although both modulators bind at the dimer interface and hinges, they make different contacts with the receptor. **a**, Top view of CMPDA (magenta stick) looking down the two-fold axis of symmetry showing the relationship of CMPDA to the hinge regions of HS1S2H **b**, Side view of CMPDA rotated 90° about the x-axis from view A **c**, Top view of CMPDB (pink stick) showing the relationship of CMPDB to the flip/flop and hinge regions of S1SJ2 **d**, Side view of CMPDB , rotated 90° about the x-axis from view c, omitting the flip/flop regions of Protomers A and B for clarity. **e**, Side view of CMPDB rotated 90° about the y-axis from view d, omitting the hinges of Protomer B for clarity. Residues within 3.2Å of CMPDA or CMPDB are shown in ball-and-stick representation with...
CPK colors and carbons colored according to protomer (green, A and blue, B). Yellow residues indicate sites of point mutations. Water molecules are shown as cyan spheres. Calculated hydrogen bonds are shown as black dashed lines. Yellow dashed lines represent Asn 754-Ser 729 hydrogen bond.

**Figure 4.** CMPDA (magenta stick, a and b) and CMPDB (pink stick, c and d) share common groups with both classes of potentiators, cyclothiazide (CTZ, green stick, left panel) and ampakine, CX614 (slate blue stick, right panel). Aside from carbons, colors are shown in CPK. Top (upper) and side (lower) views of each overlay are provided. e, Alignment of CMPDA and CMPDB. f, Top view of same alignment in panel E rotated 90º about the x-axis illustrating the position of both compounds at the Hinges and flip/flop regions. Protomer A (green) and Protomer B (blue) shown in ribbon format. The coordinates for CMPDA and CMPDB in complex with the HS1S2H LBC have been submitted. The coordinates use for CTZ and CX614 were 1LBC and 2AL4, respectively.

**Figure 5.** Pharmacophores of CTZ (a), CX614 (b), CMPDA (c) and CMPDB (d) show different profiles of modulator binding. Water molecules and amino acid residues within 3.2 Å of each compound are defined by red semi-circles (amino acids) or circles (water), between 3.3 and 4.9 Å (green) and greater than 5.0 Å from modulator (blue). Calculated hydrogen bonds are illustrated by black dashed arrows.

**Figure 6.** CMPDA and CMPDB modulate deactivation of GluA2 flip and flop receptors. Representative traces for homomeric WT GluA2i (left panel) and GluA2o (right panel) receptors heterologously expressed in HEK293 cells exposed to 1 ms of glutamate alone or glutamate plus each of 4 modulators (to measure channel deactivation). The inverted trace (outward current) above the flip control trace is a representative open-tip junction potential, which reflects the rapidity of solution exchange. Fits to the sum of two exponentials are shown in red (see Table 2). Bar plot
shows mean ± sem time constant of deactivation for flip (black bars) or flop (grey bar) isoforms of rat GluA2 receptors as described in Materials and Methods.

**Figure 7.** CMPDA and CMPDB modulate desensitization of GluA2 flip and flop receptors. Representative traces for homomeric WT GluA2i (left panel) and GluA2o (right panel) receptors heterologously expressed in HEK293 cells exposed to 500 ms of glutamate alone or glutamate plus each of 4 modulators (desensitization protocol). The inverted trace above the flip control trace is a representative open-tip junction potential indicating when glutamate was applied. Fits to the sum of two exponentials are shown in red and represent the onset of desensitization kinetics (see Table 2). Note that the calibration bar represents 200 ms for all panels. Also, three panels show an expanded trace for which the calibration bar represents 5 ms. Bar plot shows mean ± sem of the steady state divided by peak current amplitudes (ss/pk) for flip (black bars) or flop (grey bars) GluA2 receptors.

**Figure 8.** Simulations of AMPA receptor gating. a, State diagram describing a model for AMPA receptor gating in which the closed clamshell (CC) and open clamshell (CO) are explicitly modeled. Blue circles represent closed states, green circles represent open states and red circles represent desensitized states. Sequential glutamate binding is represented by transitions from left to right (κf and κ-i) and can occur in closed open and desensitized states. Conformational transitions in the clamshell are represented by vertical transitions. Two clamshell closures are required prior to the transition to the open state (β and α transitions). A complete description of the states and rates of transitions is reported in the Supplemental data (Table S1). b, Simulated responses to 1 and 500 ms pulses of 10 mM glutamate. A single exponential fit of the decays results in a τ_deact=0.7 ms and τ_des=6.2 ms, similar to GluA2i experimental data. c, Simulated dose response for peak (filled circles) and steady-state (open squares) currents in response to 1-10,000 µM pulses of glutamate (left); simulated recovery from desensitization in a paired pulse paradigm (right). Fits to the simulated
data yield EC$_{50}^{\text{peak}}$ = 292 µM, EC$_{50}^{\text{ss}}$ = 139 µM, and a $\tau_{\text{recovery}}$ = 59 ms. These fits are close approximations to experimental data presented in this paper and to data that has been previously reported (Zhang et al., 2008).

**Figure 9.** Changes in rate constants in the model predict changes in the kinetics of deactivation and desensitization observed by modulators. Simulations of GluA2 currents under voltage-clamp in response to 10 mM glutamate for either a 1 ms pulse to observe deactivation kinetics (top row), or a 500 ms pulse to observe desensitization kinetics (bottom row). Simulated GluA2 currents under control conditions (no modulator) are represented by gray traces. Potential effects of modulator (black traces) are segregated by column and have been simulated by slowing either the CO, $\alpha$ or $\delta$ transitions in isolation or in combination. The middle row shows how changes in the rate constants influence the stability of various receptor states, as measured by estimated changes in Gibb’s free energy ($\Delta G$). The troughs from left to right represent: the unbound receptor, the agonist bound receptor in the open clamshell state, the agonist bound receptor in the closed clamshell state and the open channel. The lower middle row repeats the trough found above it and shows the transition to the agonist bound desensitized state (states not shown are identical to those shown above). Results indicate that CTZ can be simulated by slowing $\delta$, CMPDB can be simulated by slowing $\alpha$; and, CMPDA can be simulated by slowing $\alpha$ and $\delta$ or CO and $\delta$. 
Table 1. Scaling and Refinement Statistics

<table>
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<tr>
<th></th>
<th>CMPDA</th>
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<td>Dmax (Å)</td>
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<td>1.7</td>
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<tr>
<td>Rsym</td>
<td>0.067 (0.288)</td>
<td>0.055 (0.355)</td>
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<td>I/SigI</td>
<td>22.3 (7.0)</td>
<td>25.8(4.4)</td>
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<tr>
<td>Completeness (%)</td>
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<td>98.5 (95.1)</td>
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<tr>
<td>Observations Unique (Total)</td>
<td>90,630 (735,725)</td>
<td>96,319 (596,873)</td>
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<td>R (Rfree)</td>
<td>0.216 (0.247)</td>
<td>0.226 (0.2404)</td>
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<td>rmsd bonds (Å)</td>
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<tr>
<td>rmsd angles (°)</td>
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Rsym = \sum |I| / \sum I, where I is the integrated intensity of a given reflection. The refinement residual, 

R = \sum |F_{obs} - F_{calc}| / \sum F_{obs}. Rmsd, Root mean square deviation from ideal values. Values for highest resolution bin are shown in parenthesis. Rfree, calculated from 4546 observation test set for CMPDA and 4820 observation test set excluded for CMPDB.
Table 2. Summary of mean modulation of WT GluA2 flip and flop.

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<th></th>
<th>GluA2i</th>
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<td></td>
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<td><strong>10 mM Glu</strong></td>
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<tr>
<td>Weighted $\tau_{des}$ (ms)</td>
<td>6.7 ± 0.4</td>
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<td>49.0 ± 20.8</td>
<td>2.3 ± 0.2</td>
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<td>$A_{fast \ tau_{des}}$ (%)</td>
<td>51.8 ± 5.4</td>
<td>56.6 ± 4.5</td>
<td>57.6 ± 4.8</td>
<td>92.1 ± 2.6</td>
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<tr>
<td>% Des (1-ss/pk)</td>
<td>96.3 ± 0.8</td>
<td>97.1 ± 0.7</td>
<td><strong>34.4 ± 10.9</strong></td>
<td><strong>81.5 ± 3.6</strong></td>
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<td>Weighted $\tau_{deact}$ (ms)</td>
<td>0.8 ± 0.1</td>
<td>0.7 ± 0.07</td>
<td>5.6 ± 1.2***</td>
<td>3.7 ± 1.5</td>
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<td>$A_{fast \ tau_{deact}}$ (%)</td>
<td>55.4 ± 6.3</td>
<td>62.0 ± 6.2</td>
<td>43.9 ± 10.2</td>
<td>72.0 ± 13.9</td>
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<td><strong>10 µM CMPD A</strong></td>
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<td>Weighted $\tau_{des}$ (ms)</td>
<td>Non-Des(5)</td>
<td>Non-Des(12)</td>
<td><strong>34.4 ± 10.9</strong></td>
<td><strong>81.5 ± 3.6</strong></td>
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<td>$A_{fast \ tau_{des}}$ (%)</td>
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<td>Non-Des</td>
<td>43.9 ± 10.2</td>
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<tr>
<td>% Des</td>
<td>8.0 ± 2.9**</td>
<td>6.1 ± 1.1**</td>
<td>11.9 ± 2.5f</td>
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<td>$A_{fast \ tau_{deact}}$ (%)</td>
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<td><strong>100 µM CX614</strong></td>
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<tr>
<td>Weighted $\tau_{des}$ (ms)</td>
<td>7.25 ± 0.5*</td>
<td>121 ± 24.3**</td>
<td>4.3 ± 2.3***</td>
<td>67.3 ± 5.1***</td>
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<td>$A_{fast \ tau_{des}}$ (%)</td>
<td>76.6 ± 3.4*</td>
<td>31.1 ± 5.2**</td>
<td>1.3 ± 0.3(8)</td>
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<tr>
<td>% Des</td>
<td>34.4 ± 5.5**</td>
<td>33.4 ± 2.1**</td>
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<td>59.1 ± 9.6</td>
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<td>Weighted $\tau_{deact}$ (ms)</td>
<td>1.9 ± 0.7</td>
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<td>$A_{fast \ tau_{deact}}$ (%)</td>
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Summary of electrophysiology data for homomeric GluA2 WT receptors heterologously expressed in HEK293 cells exposed to glutamate alone or glutamate and each of 4 modulators. Data reported as mean ± SEM compared using ANOVA with Dunnett’s test for multiple comparisons within each
receptor type to L-glutamate “control”, (***, p<0.0001; **, p<0.001; *, p<0.05). Data were fit with a two-exponential function and both fast (F) and slow (S) components were combined into a weighted average based on their respective amplitude contributions (%A_{Fast} \times \tau_{Fast} + %A_{Slow} \times \tau_{Slow})", where A= the amplitude of each time constant. Non-des, cells that did not desensitize in response to modulator. Comparisons between, a-c; or, within, d-f. Number of cells is given in parenthesis. #, for WTGlup2i exposed to CX614, the initial phase of decay only was fit with a single exponential function, due to “resensitization” as seen by the increasing inward current during the application of glutamate.
Figure 1

(a) Bis-Alkylsulfonamide CMPDA

(b) Bis-Carboxythiophene CMPDB
Figure 3
Figure 4
Figure 5
Figure 6
**Figure 7**

The figure illustrates the effects of different compounds on the desensitization of Flip and Flop channels. The top panels show the time course of desensitization with and without the addition of CMPDA, CX614, CMPDB, and CTZ, respectively. The control (no added compound) is shown on the left, and the experimental conditions are on the right.

- **Flip**
  - Control: Initial rapid desensitization followed by a slower phase.
  - With CMPDA, there is a significant reduction in desensitization.
  - With CX614, desensitization is moderately reduced.
  - With CMPDB, desensitization is slightly reduced.
  - With CTZ, desensitization is significantly increased.

- **Flop**
  - Control: Similar to Flip, initial rapid desensitization followed by a slower phase.
  - With CMPDA, desensitization is reduced.
  - With CX614, desensitization is moderately reduced.
  - With CMPDB, desensitization is slightly reduced.
  - With CTZ, desensitization is significantly increased.

The bar graph at the bottom shows the quantitative comparison of desensitization efficiencies for each condition. The x-axis represents the conditions (Control, CMPDA, CX614, CMPDB, CTZ), and the y-axis represents the desensitization (ss/pk) values. The bars indicate the average desensitization with error bars for variability.
Figure 8
Figure 9