Mutational Analysis and Modeling of Negative Allosteric Modulator Binding Sites in AMPA Receptors

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ABSTRACT

The \(\alpha\)-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptors (AMPA\left(\text{AMPARs}\right)\) are a subclass of the ionotropic glutamate receptor superfamily, which functions as glutamate-gated cation channels to mediate the majority of excitatory neurotransmission in the central nervous system. AMPARs are therapeutic targets in a range of brain disorders associated with abnormal glutamate hyperactivity. Multiple classes of AMPAR inhibitors have been developed during the past decades, including competitive antagonists, ion channel blockers, and negative allosteric modulators (NAMs). At present, the NAM is the only class of AMPAR ligands that have been developed into safe and useful drugs in humans in the form of perampanel (Fycompa), which was recently approved for treatment of epilepsy. Compared with the detailed understanding of other AMPAR ligand classes, surprisingly little information has been available regarding the molecular mechanism of perampanel and other classes of NAMs at AMPARs; including the location and structure of NAM binding pockets in the receptor complex. However, structures of the AMPAR GluA2 in complex with NAMs were recently reported that unambiguously identified the NAM binding sites. In parallel with this work, our aim with the present study was to identify specific residues involved in the formation of the NAM binding site for three prototypical AMPAR NAMs. Hence, we have performed a mutational analysis of the AMPAR region that links the four extracellular ligand-binding domains to the central ion channel in the transmembrane domain region. Furthermore, we perform computational ligand docking of the NAMs into structural models of the homomeric GluA2 receptor and optimize side chain conformations around the NAMs to model how NAMs bind in this specific site. The new insights provide potentially valuable input for structure-based drug design of new NAMs.

SIGNIFICANCE STATEMENT

The \(\alpha\)-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) receptors are glutamate-gated ion channels that mediate the majority of excitatory neurotransmission in the brain. Negative allosteric modulators of AMPA receptors are considered to have significant therapeutic potential in diseases linked to glutamate hyperactivity. The present work employs mutational analysis and molecular modeling of the binding site for prototypical NAMs to provide new molecular insight into how NAMs interact with the AMPA receptor, which is of potential use for future design of new types of NAMs.

Introduction

The \(\alpha\)-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptors (AMPA\left(\text{AMPARs}\right)\) mediate the majority of excitatory postsynaptic currents in the central nervous system (Traynelis et al., 2010) by utilizing energy from the binding of glutamate (Glu) to drive ion channel opening. In addition to a central role in basal excitatory neurotransmission, AMPARs are involved in synaptic plasticity mechanisms such as long-term potentiation and depression that underlie learning and memory (Kessels and Malinow, 2009). AMPARs assemble as tetramers from combinations of GluA1-GluA4 subunits as well as a range of auxiliary subunits of different classes that can modulate receptor function (Jackson and Nicoll, 2011; Haering et al., 2014), generating a range of subtypes with overall similar core function but with differences in biophysical properties, pharmacology, and regulatory mechanisms (Cokić and Stein, 2008; Jackson and Nicoll, 2011). Structures are currently available of homomeric and heteromeric AMPARs in resting (Sobolevsky et al., 2009; Zhao et al., 2019; Herguedas et al., 2019), active (Chen et al., 2017; Twomey et al., 2017a), and

ABBREVIATIONS: Ala, alanine; AMPAR, \(\alpha\)-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptor; ATD, amino-terminal domain; blac, \(\beta\)-lactamase; \(\text{Ca}^{2+}\), intracellular calcium; CP (CP-465,022), 3-(2-chloro-phenyl)-2-[2-(6-diethylaminomethyl-pyridin-2-yl)-vinyl]-6-fluoro-3H-quinazolin-4-one; CTZ, cyclothiazide; DMEM, Dulbecco’s modified Eagle’s medium; Glu, glutamate; Gly, glycine; GYKI (GYKI-53,655), 1-(4-Aminophenyl)-3-methylcarbamyl-4-methyl-3,4-dihydro-7,8-methylenedioxy-5H-2,3-benzodiazepine; HB, hydrogen bond; HKE, human embryonic kidney; LBD, ligand-binding domain; NAM, negative allosteric modulator; PAM, positive allosteric modulator; PDB, Protein Data Bank; PMP, perampanel; TEVC, two-electrode voltage clamp; TMD, transmembrane domain; WT, wild type.
dissected (Chen et al., 2017; Twomey et al., 2017b) states. These have advanced the molecular understanding of AMPARs by providing insight into the structural mechanisms underlying receptor function and pharmacology (Kumar and Mayer, 2013; Karakas et al., 2015; Zhu and Gouaux, 2017; Twomey and Sobolevsky, 2018).

Abnormal AMPAR activity is involved in neurologic diseases, and compounds that can modify AMPAR signaling have for decades been pursued as potential therapeutics in treatment of Alzheimer’s disease, Parkinson’s disease, depression, and epilepsy (Zarate and Manji, 2008; Traynelis et al., 2010; Citraro et al., 2014; Di Bonaventura et al., 2017). AMPARs contain multiple binding sites where small-molecule ligands can modulate receptor function positively or negatively via allosteric effects on core channel function (Traynelis et al., 2010; Karakas et al., 2015). This includes negative allosteric modulators (NAMs), which bind outside the orthosteric binding site or the ion channel and inhibit the receptor by noncompetitive mechanisms. Several chemical classes of AMPAR NAMs are known and include 2,3-benzodiazepines (Sólyom and Tarnawa, 2002), quinazolinones (Lazzaro et al., 2002), and pyridones (Zwart et al., 2014). Among these, the pyridone-based perampanel (PMP) under the trade name Fycompa was recently approved as a first-in-class AMPAR drug for treatment of epilepsy (Hanada, 2014), thereby demonstrating the potential of negative allosteric modulation of AMPARs in treatment of neurologic diseases linked to glutamatergic hyperfunction (Di Bonaventura et al., 2017). However, the molecular mechanisms underlying NAM inhibition of AMPARs are surprisingly poorly understood for all NAM classes; including the location and structure of the NAM binding pockets and how ligand binding can inhibit channel gating.

The AMPAR structure is highly modular and contains an extracellular amino-terminal domain (ATD) and a ligand-binding domain (LBD), a transmembrane domain (TMD) with the ion channel, and a cytoplasmic carboxyl-terminal domain (Sobolevsky et al., 2009; Kumar and Mayer, 2013). Channel gating involves allosteric coupling between LBD motions and the TMD via short flexible linkers. Binding studies have indicated that NAMs of the 2,3-benzodiazepine, quinazolinone, and pyridone classes share the same binding site (Menniti et al., 2000; Sólyom and Tarnawa, 2002; Micale et al., 2008; Mittapalli and Roberts, 2014), which is distinct from the well-characterized binding sites for positive allosteric modulators (PAMs) at the LBD interfaces (Sun et al., 2002). Early work suggested that NAM binding sites were located in the LBD-TMD linker region and identified residues in the linkers connecting the S1-M1 and S2-M4 regions to affect NAM inhibitory potency (Balannik et al., 2005; Coki and Stein, 2008). Currently, structures of homomeric GluA2 in complex with NAMs are available that unambiguously identify each subunit containing a NAM binding pocket formed between the upper M3 transmembrane helix and the S1-M1 and S2-M4 linkers (Yelshanskaya et al., 2016). Although these structures contain electron densities for the bound NAM ligands, the resolution of approximately 4 Å does not allow direct identification of NAM-protein interactions (Twomey and Sobolevsky, 2018).

We have conducted a mutational analysis of the LBD-TMD linker regions in the GluA2 subunit to further study the role of LBD-TMD linker residues for the function of three prototypical NAMs in the form of PMP, 1-(4-Aminophenyl)-3-methylcarbamyl-4-methyl-3,4-dihydro-7,8-methylenedioxy-5H,2,3-benzodiazepine [GYKI-53,655 (GYKI)] (Moncada et al., 1991), and quinazolinone 3-(2-chloro-phenyl)-2-(2-(6-diethylaminomethyl-pyridin-2-yl)-vinyl)-6-fluoro-3H-quinazolin-4-one [CP-465,022 (CP)] (Menniti et al., 2000). Our initial aim was to use the resulting data set to specify the location of NAM binding pockets in the linker region and employ computational ligand docking of NAMs into GluA2 structures to create binding mode models. We used the new GluA2/NAM structures as templates to create binding models that include all side chains. In combination with our mutational analysis, the models provide molecular insight into ligand orientations and specific protein-ligand contacts for the investigated NAMs.

Materials and Methods

Materials. All chemicals were obtained from Sigma-Aldrich (St. Louis, MO) unless otherwise stated. Dulbecco’s modified Eagle’s medium (DMEM), FBS, trypsin, and penicillin-streptomycin were obtained from Invitrogen (Carlsbad, CA). DNA-modifying enzymes were obtained from New England Biolabs (Ipswich, MA), except PfuUltra II Fusion HS DNA Polymerase (Agilent, Carlsbad, CA) and Rapid T4 Ligase (Roche Molecular Systems, Pleasanton, CA). All tissue cell culture plasticware were obtained from Sarstedt (Numbrecht, Germany) unless otherwise stated. 6-Cyano-7-nitroquinoxaline-2,3-dione, cyclothiazide (CTZ), and CP-465,022 hydrochloride were obtained from HelloBio (Bristol, UK). GYKI-53,655 or GYKI-53,655 hydrochloride was obtained from Axon MedChem (Groningen, Netherlands) or Tocris Bioscience (Bristol, UK). Perampanel was obtained from Toronto Research Chemicals (Toronto, Ontario, Canada) or Apexmol (Beijing, China). The calcium dyes Fluo-2 or Fluo-8 AM were obtained from TEFlabs (Austin, TX) or AAT Bioquest (Sunnyvale, CA).

Molecular Biology. The plasmid vectors pRK5 (BD PharMingen, San Diego), pGEM-HE (Liman et al., 1992) and pXOOF containing cDNA for the unedited flip isoform of the rat GluA2 gene were used for expression in mammalian cells or generation of mRNA for microinjection in Xenopus laevis oocytes. The pXOOF vector is a modified version of the dual-purpose mammalian and Xenopus expression vector pXOON (Jespersen et al., 2002), in which cDNA encoding enhanced GFP is removed. Specifically, we created pXOOF by polymerase chain reaction amplification of the entire pXOON sequence enhanced GFP encoding cassette, followed by digestion with the restriction enzymes NheI-HF and SbfI and religation using T4 DNA Ligase (Roche, Mannheim, Germany) to generate pXOOF. The pXOOF plasmid sequence was verified by DNA sequencing. Generation of GluA2 point mutants was performed by site-directed mutagenesis using the QuickChange Mutagenesis Kit (Stratagene, La Jolla, CA). The mutations were verified by DNA sequencing of the entire GluA2 gene (GATC Biotech, Constance, Germany). For analysis of cell-surface expression of wild-type (WT) and mutant GluA2 receptors, cDNA encoding β-lactamase (blac) was inserted in the GluA2 cDNA in between the segments encoding the N-terminal signal sequence and the ATD. Specifically, using In-Fusion cloning (Mountain View, CA), a polymerase chain reaction–amplified DNA fragment encoding blac flanked by two short amino acid linkers (GGSOS and GSGS) was inserted in-frame into an XbaI site introduced by site-directed mutagenesis of codons 26 and 27 in the GluA2 sequence to create blac-tagged GluA2. For use as a template for in vitro transcription of complementary RNA, pGEM-HE and pXOOF plasmid constructs were linearized downstream from the 3′-untranslated region of the Xenopus β-globin sequence using the NheI-HF restriction enzyme, and purified by ethanol precipitation.
with Pellet Paint NF Co-Precipitant (Merck, Billerica, MA). Complementary RNA transcription was performed using the AmpIICap-Max T7 High Yield Message Maker Kit (CellScript, Madison, WI) and RNA was column purified using the NucleoSpin RNA Clean-up XS Kit (Macherey-Nagel, Düren, Germany).

**Mammalian Cell Culturing and Expression.** Human embryonic kidney (HEK) 293T cells (American Type Culture Collection, Manassas, VA) were cultured in growth medium (DMEM supplemented with 10% v/v FBS, 100 U/ml penicillin, and 100 μg/ml streptomycin) at 37°C in a humidified 5% CO₂ environment. For expression of WT and mutant GluA2 in HEK293T cells, TransIT-LT1 DNA Transfection Reagent (Mirus, Madison, WI) was used as described previously (Sørensen et al., 2014). Briefly, HEK293T cells in suspension (1 × 10⁶ cells/ml) were prepared and injected with 1–25 ng mRNA, as described previously (Poulsen et al., 2013). The care and use of *Xenopus laevis* were in strict adherence to a protocol approved by the Danish Veterinary and Food Administration (license 2014-15-0201-00031). Injected oocytes were incubated at 18°C in standard Barth’s solution containing 88 mM NaCl, 1 mM KCl, 0.4 mM CaCl₂, 2.4 mM NaHCO₃, 0.33 mM Ca(NO₃)₂, 0.82 mM MgSO₄, and 5 mM Tris (pH 7.4), supplemented with 50 μg/ml gentamicin and used for two-electrode voltage-clamp electrophysiology (TEVC) measurements 2–7 days after injection.

**Intracellular Ca²⁺ Imaging Assay.** For determination of concentration-response curves for glutamate activation and NAM inhibition of WT and mutant GluA2 receptors, imaging of changes in the concentration of intracellular Ca²⁺ in transfected HEK293T cells was performed using a FlexStation I Plate Reader (Molecular Devices, Sunnyvale, CA) to measure changes in the fluorescence of a calcium-sensitive fluorophore dye (dissolved in plain DMEM and incubated for 30 minutes at 37°C). The dye fluorescence in response to the addition of agonist solution were repeated on three independent days. Preincubation with NAMs in some instances produced concentration-dependent inhibition of the baseline measurement before Glu addition, which resulted in an apparent increase in response amplitude with increasing NAM concentration. This effect likely arises from activation by small amounts of contaminating Glu produced by the cells that were not removed by thorough washing. The observation might indicate an effect of the desensitization state of the receptor on NAM potency.

**Determination of Cell-Surface Expression.** For determination of mutational effects on cell-surface expression of GluA2 receptors, mutants were created in a GluA2 subunit construct tagged with a bacterial β-lactamase. Relative levels of surface-expressed β-lactam-tagged GluA2 (i.e., β-lacGluA2) can be accurately quantified in living cells by measuring the conversion rate of the membrane-impermeable substrate nitrocefin by simple absorption spectroscopy (Lam et al., 2013). For analysis in HEK293T cells, transfected cells were cultured in poly-D-lysine–coated clear-bottom 96-well plates for 2 days. Cells were washed twice in PBS supplemented with CaCl₂ and MgCl₂, followed by addition of nitrocefin to a final concentration of 50 μM in a total volume of 100 μl per well. Immediately following nitrocefin addition, plates were placed in a microplate reader (Safire2; Tecan, Maennedorf, Switzerland), and well absorbance at 486 nm of the nitrocefin conversion product was recorded every minute for 60 minutes at ~30°C. For each well, the absorbance at 486 nm was plotted as a function of time using GraphPad Prism version 6.01 (GraphPad Software, San Diego, CA) and the rate of nitrocefin conversion per minute was determined by linear regression analysis of the slope of the curve in the linear range.

**TEVC Electrophysiology.** For TEVC recordings, glass micropipettes (0.69 mm i.d./1.2 mm o.d.; Harvard apparatus, Holliston, MA) were pulled on a Sutter P-1000 to a tip resistance of 0.5–2.0 MΩ and filled with 3 M KCl. Oocytes were clamped using a TEVC amplifier (OC-725C; Warner Instruments, Hamden, CT) and continuously perfused with frog Ringer’s solution containing 115 mM NaCl, 2 mM KCl, 5 mM HEPES, and 1.8 mM BaCl₂ (pH 7.6 adjusted with NaOH), via the force of gravity at flow rates of 5–10 ml/min into a vertical oocyte flow chamber with a volume of 0.3 ml (Joshi et al., 2004). Compounds dissolved in frog Ringer’s solution were added by bath application. This perfusion system yields a solution exchange time of 100–500 milliseconds. Concentration-response data were generally recorded at a holding potential of −30 or −60 mV, and each compound solution was applied for 30 or 45 seconds to ensure that steady-state currents were obtained. NAM concentration-response curves were obtained by adding increasing concentrations of NAM in the continuous presence of 300 μM glutamate after evoking a full agonist response by adding 300 μM glutamate alone. All experiments were performed at room temperature (20–23°C). Data acquisition was accomplished using an analog-digital converter (CED 1401 Plus; Cambridge Electronic Design, Cambridge, UK) interfaced with a PC running WinWCP software (available from Strathclyde Electrophysiology Software, University of Strathclyde, Glasgow, UK). Concentration-response and concentration-inhibition measurements were performed by measuring agonist-evoked current during stepwise application of increasing concentrations of agonist or NAM (as illustrated in Fig. 6).

**Patch-Clamp Electrophysiology (HEK Cells).** For electrophysiological experiments involving rapid ligand-application measurements, outside-out membrane patches were excised from transiently transfected HEK293 cells using thin-wall glass micro-pipettes (TW150F-4; World Precision Instruments), heat polished to 3–10 MΩ tip resistance filled with internal solution containing 140 mM KCl, 10 mM HEPES, and 1 mM 1,2-bis(o-aminophenoxy) ethane-N₂,N₂',N₂'-tetraacetic acid (BAPTA) (pH 7.20, adjusted with KOH) and osmolality adjusted to 285 mosm by adding 10 μM D-Glucose. Pipettes had a tip resistance of 3–10 MΩ. External recording solution for all experiments was comprised of 150 mM NaCl, 3 mM KCl, 10 mM HEPES, 0.01 mM EDTA, and 1 mM CaCl₂ (pH 7.40 with NaOH (290 mosm)). Currents were recorded at a holding...
potential of 60 mV with an Axopatch 200B amplifier (Molecular Devices), low-pass filtered at 8 kHz using a 8-pole Bessel filter (Frequency Devices, Ottawa, IL), and digitized at 40 kHz using a Digidata 1440A (Molecular Devices). Rapid application and removal of ligand solutions at excised membrane patches were performed using a piezo (Burleigh Instruments, Fishers, NY) bimorph-driven double-barreled theta tube perfusion system. The time course of solution exchange across the laminar flow interface was estimated by liquid junction potential measurements of 0.2–0.4 milliseconds (10%–90% rise time) for a 10-fold difference in ionic strength. Current kinetics results were analyzed using ChannelsLab version 2 (SynaptoSoft, Decatur, GA).

Data and Statistical Analyses. For construction of concentration-response curves from intracellular calcium ([Ca\(^{2+}\)]\(_{i}\)) imaging experiments, peak fluorescence response (corresponding to the difference between the maximum and baseline 538 nm emission) from quadruplicate wells representing identical NAM or agonist concentrations were averaged using SoftMax Pro version 5.4 software (Molecular Devices). For each individual plate experiment, the average responses were normalized to the average corresponding maximal agonist response or the agonist response in absence of NAM. Composite concentration-response curves were generated by plotting normalized data from multiple experiments as a function of NAM or agonist concentration using GraphPad Prism software. Concentration-response relationships were fitted as composite curves using GraphPad Prism version 6.01 (GraphPad Software) to the Hill equation:

\[
\text{response} = \frac{\text{bottom} + \text{top} - \text{bottom}}{1 + 10^{(\text{EC}_{50} - X) / nH}}
\]

where bottom is the fitted minimum response; top is the fitted maximum response, \(nH\) is the Hill slope; \(X\) is the concentration; and \(\text{EC}_{50}\) is the IC\(_{50}\) (half-maximally effective concentration of inhibitor) or EC\(_{50}\) (half-maximally effective concentration of agonist), respectively. For the construction of concentration-response curves from electrophysiological data, agonist-evoked current responses from individual oocytes were determined from TEVC traces using ClampFit version 10.3 (Molecular Devices) and normalized to the current response obtained in the absence of NAM (for IC\(_{50}\) determinations) or by maximal agonist concentration (for EC\(_{50}\) determinations). Concentration-response curves were constructed as composites of normalized responses from three to six oocytes and analyzed as described for the [Ca\(^{2+}\)]\(_{i}\) assay. Unless otherwise noted, results are expressed as mean ± S.D. Statistical analysis by pairwise and multiple comparisons was performed using Student’s t test or ANOVA with multiple comparisons procedures as indicated.

Molecular Modeling. The crystallographic structures of the GluA2 receptor used as input models for ligand docking included the apo state structure [Protein Data Bank (PDB) 5L1B], the GYKI-bound structure (PDB 5L1H), the CP-bound structure (PDB 5L1E), and the PMP-bound structure (PDB 5L1F). All structures were determined by Yelshanskaya et al. (2016) and obtained from the Brookhaven Protein Data Bank (Berman et al., 2000). All four structures have several missing residues and residues with missing side chain atoms. For the initial docking calculations, missing atoms in locations that were distant from the NAM binding sites (located at the extracellular side of the TMD) were not modeled; this included residues located in the ATD-LBD linker region, the M1-M2 linker region, and the C-terminal region. Within and around the NAM binding pockets, positions with missing atoms are located in the S1-M1 linker region, the M3 transmembrane helix, the M3-S2 linker, and the S2-M4 linker. For these regions the missing atoms were added using the WHAT-IF webserver interface (Vriend, 1990), following the method described by China et al. (1995). The ligand molecules were cocrystallized with the GluA2 receptor in PDB files 5L1H, 5L1E, and 5L1F were imported into the Maestro software (Schrodinger Release 2013-2; Maestro, version 9.5; Schrödinger, LLC, New York), where hydrogen atoms were added. The ligands were either left in the crystal conformation or subjected to short energy minimization using the clean-up geometry function; giving rise to two conformations for each imported structure. The original ligand structures (with hydrogen atoms added) and the equivalent energy-minimized structures were used as input for docking calculations (i.e., in total, eight structures per ligand from the four different subunits; each in either original or optimized conformation). Additionally, for each ligand two conformations were generated by constructing the ligand from scratch in Maestro, meaning that 10 input ligand structures were used for docking of each of the ligands CP, PMP, and GYKI. It should be noted that PMP, CP, and GYKI have low solubility in water, suggesting that they are neutral at physiologic pH; therefore, they were docked in their unprotonated states. All ligand dockings into the GluA2 receptor structures were performed using the Autodock Vina docking software (Trott and Olson, 2010), utilizing a search box that covered all four binding pockets. Twenty poses were saved in each docking run during which the ligands were treated flexibly, while the protein remained rigid. In general, the poses showed variations in the position of the ligand due to the relatively large search box. However, the binding pocket was identified for all three ligands when docked into their parent protein structure and assumed orientations similar to those proposed by the existing GluA2/NAM structures (Yelshanskaya et al., 2016). Based on the assumption that these ligand orientations are overall correct, we constructed up to 100 models for each compound (98 for GluA2/GYKI and 99 for GluA2/PMP and GluA2/CP), i.e., generating approximately 400 binding sites since the NAM-GluA2 stoichiometry (at least in the crystal structures) is 4:1. In this alternative approach, different side chain conformations were sampled around the ligand-binding sites using modeling software MODEL-ELLER (Sali and Blundell, 1993) to build models of the GluA2/NAM complex for which all missing atoms and residues were added. Furthermore, all residue positions within 12 Å of the ligands were optimized. MODELLELLER can accommodate the ligands in terms of space; however, without parameters for the ligands, the interactions with the protein atoms of the binding site are not included in the optimization. Thus, to account for protein-ligand interactions and to optimize the ligand conformations, all of the generated models were subjected to energy minimization in GROMACS 5.0 (Van Der Spoel et al., 2005) using the AMBER99-ILDN (Lindorff-Larsen et al., 2010) force field for the protein and ligand parameters from the general AMBER force field (Wang et al., 2004) generated using Antechamber in AmberTools14 (Salomon-Ferrer et al., 2013). Autodock (Sousa da Silva and Franks, 2012) was used to generate GROMACS topologies from the AMBER ones. Minimizations were performed in vacuum until the maximum force on a single atom was less than 100 kJ/mol per nanometer or machine precision was reached. Analysis and image generation were performed using the Maestro software along with VMD software (Humphrey et al., 1996) and PyMOL software.

## Results

**Glycine-Scanning Mutagenesis of the S1-M1 and S2-M4 Linker Regions.** Previous mutational analysis had identified the LBD-TMD linker segments as overall candidate regions to form the NAM binding sites in the AMPAR complex (Balannik et al., 2005). Specifically, mutation of residues in the S1-M1 and S2-M4 segments can perturb NAM inhibition, and these residues may thus be involved in forming NAM binding pockets as well as interacting directly with NAMs (Fig. 1). However, the NAM binding site likely involves additional residues in the linker segments as well as upper M1 and M4 segments. To expand the knowledge of the contribution of the side chains of individual residues in the S1-M1 (including the pre-M1 helix), upper M1, S2-M4, and upper M4 segments for NAM binding, we initially performed glycine (Gly) scanning mutagenesis of all amino acid positions

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**References:**

between K511 and M527 in the S1-M1 and upper M1 segments and E782 and Y797 in the S2-M4 and upper M4 segments (Fig. 3). We selected the flip isoformal of the GluA2 subunit in the unedited form as the model AMPAR subunit for the mutational study based on the robust expression and well-characterized functional properties of homomeric GluA2 receptors. In addition, several X-ray crystallographic and cryo-electron microscopy structures exist for homomeric GluA2 receptors (Sobolevsky et al., 2009; Yelshanskaya et al., 2014; Twomey et al., 2016, 2017b), which provide an excellent framework for interpreting mutagenesis results in a structural context and can serve as target structures for ligand docking. Notably, the target LBD-TMD linker regions are conserved between subunits GluA1-4. Thus, if the NAM binding sites are harbored in the linker region, homo- or heteromeric AMPARs formed of any subunit combination may potentially be equivalent with regard to the structure of the NAM binding pocket. To allow determination of mutational effects on surface expression, we used a version of GluA2 tagged in the N-terminal of the ATD with the reporter enzyme blac (blac-GluA2; see Materials and Methods) as the parent construct for all mutants. To ensure that fusion of blac to the ATD does not change the function of the GluA2 receptor, we compared current responses from WT GluA2 and blac-GluA2 in excised outside-out membrane patches from HEK293 cells to the application of a maximally effective concentration (10 mM) of glutamate by using a rapid perfusion system with a solution exchange time <0.4 milliseconds (see Materials and Methods) (Supplemental Fig. 1); the results showed that the rates of receptor activation, desensitization, and deactivation were similar with and without blac fused to the ATD (Supplemental Fig. 1). Furthermore, we determined the IC50 values for NAM inhibition of WT GluA2 and blac-GluA2 currents in Xenopus oocytes (see Materials and Methods) and did not observe any significant effect of blac on NAM pharmacology (Supplemental Fig. 1; Tables 1 and 2).

For initial characterization of the mutant library, we expressed mutants individually in HEK293 cells and used an intracellular calcium concentration imaging assay to measure concentration-dependent Glu-evoked increases in [Ca2+]i (see Materials and Methods) and assessed the impact of Gly substitutions on basal receptor function by determining the EC50 and maximum response amplitude (Emax) for Glu (Fig. 2). Also, we measured blac activity to determine mutational effects of the ability of the mutant GluA2 subunits to correctly fold, assemble, and traffic to the cell surface (Supplemental Fig. 2; Supplemental Table 1). The results of the Gly scanning mutagenesis are summarized in Fig. 3 and Table 1. For the 16 positions in the S1-M1 and upper M1 segments, Gly substitution (which removes the WT residue side chain) appeared to disrupt receptor function at five positions (L518, P520, L521, I525, and W526) since these mutants, when expressed in HEK293 cells, did not produce clearly detectable Ca2+ signals in the [Ca2+]i imaging assay in response to Glu concentrations up to 1 mM (for a representative example, see Fig. 2A). However, expression of all five mutants generated extracellular blac activity in HEK293 cells that were within the range of the activity observed for expression of WT blac-GluA2 receptors (Supplemental Table 1), indicating that these Gly substitutions maintain the ability of the GluA2 subunit to fold and traffic to the cell surface. These results suggest that Gly substitutions at these positions may alter or completely disrupt receptor function. The Gly mutants for the remaining 11 positions displayed robust Glu-evoked Ca2+ signals that allowed determination of Glu EC50 values. For the majority of functional mutants, the Glu EC50 value was identical to or within the 3-fold range of the EC50 value for WT blac-GluA2 (Fig. 3; Table 1). For the 15 Gly substitutions in the S2-M4 and upper M4 segments, four mutants (L787G, V792G, F796G, and Y797G) were nonfunctional with intact cell surface expression (Supplemental Fig. 2; Supplemental Table 1), whereas Gly substitution at the remaining 11 positions yielded mutants with robust Glu-evoked responses and EC50 values that were unchanged or within the 3-fold range of WT blac-GluA2 (Fig. 3B). Thus, in total, we generated 31 Gly mutants, of which 22 were functional and could be subjected to characterization of potential changes in GluA2 NAM pharmacology.

Effect of Glycine Substitutions on Inhibitory Potency of Perampanel, CP, and GYKI. We next used the [Ca2+]i imaging assay to generate full concentration-inhibition relationships for PMP, CP, and GYKI (see Materials and Methods) to determine the IC50 value for each NAM across all functional Gly mutants (Table 2). Representative examples of concentration-response curves are shown in Fig 4. Fig 5 summarizes the relative changes in NAM potency produced by the Gly substitutions compared with WT blac-GluA2. Determination of specific IC50 values was not possible at some mutants that displayed decreased sensitivity to NAMs due to limitations in NAM solubility.

For the S1-M1 segment, we generally observed similar patterns of mutational effects for the three NAMs (Fig. 5). Specifically, removal of residue side chains at K511 and V514...
produce loss-of-potency effects for PMP, CP, and GYKI, whereas removal at F517 produces gain-of-potency effects for all three NAMs (albeit only 3-fold for CP at F517G) (Fig. 5; Table 2). The similar effect patterns suggest that the S1-M1 linker is involved in a NAM binding or inhibitory mechanism, and furthermore corroborate the previous suggestions that PMP, CP, and GYKI have overlapping binding sites (Menniti et al., 2000; Hanada et al., 2011). For GYKI, additional changes in potency occur by removal of the F515 and D519 side chains (leading to gain of potency), and the S516 side chain (leading to loss of potency). These results indicate that GYKI might interact with additional residues in the S1-M1 region compared with PMP and CP. For the Gly mutations in the upper M1 segment (A522G to M527G), no substantial
potency changes (here defined as more than a 5-fold change in the IC$_{50}$ value) were observed for any NAM (Fig. 5; Table 2); indicating that the upper M1 transmembrane domain may not be involved in NAM binding. However, notably, only four Gly mutants within the M1 region were characterized since the I525G and W526G mutants were not functional in the [Ca$^{2+}$], assay and thus not amenable to pharmacological characterization.

For Gly mutants of the seven positions in the S2-M4 segment, A786G displayed a gain-of-potency effect for PMP and GYKI, whereas the remaining Gly substitutions did not substantially change the potency of any of the NAMs (Fig. 5). In the upper M4 segment, no substantial changes in NAM potencies were seen upon Gly substitutions from L787 to V795 (Fig. 5). The exception to this pattern is the effect of the N791G mutation on CP potency, for which removal of the asparagine side chain decreases the IC$_{50}$ value of CP by more than 10-fold, whereas PMP and GYKI potency does not substantially change.

In summary, we assessed the effects of Gly substitutions on NAM potency for 11 positions in the S1-M1/upper M1 segments and 11 positions in the S2-M4/upper M4 segments. Among these positions, six Gly substitutions in the S1-M1 segment displayed more than a 5-fold change in IC$_{50}$, whereas no effects were observed for the upper segment of the M1 transmembrane helix. Furthermore, one substitution in the S2-M4 region and one substitution in the upper M4 segment displayed substantial effects. Thus, overall, these results of the Gly scan implicate eight residues in potential formation of the NAM binding site in GluA2.

**Mutational Analysis of Key NAM Binding Pocket Residues.** To assess the role of side chains at the initial positions identified in the Gly scan (Fig. 5), we made additional mutants of potential key positions to introduce side chains with
different physicochemical properties (e.g., change the bulk size and/or hydrogen bonding properties of the side chain). Positions included mainly those where Gly substitution induced more than a 5-fold change in NAM potencies or was previously reported to change NAM activity (Balannik et al., 2005) (Fig. 5). At this point in the mutational analysis, X-ray crystallography structures were reported of homomeric GluA2 in complex with the same NAMs included in the present study (PMP, GYKI, and CP) (Yelshanskaya et al., 2016), showing that four nearly identical NAM binding sites are located within the LBD-TMD linker region of the tetramer. Although the electron densities around the identified NAM binding pockets in all of the GluA2/NAM structures were insufficient to resolve several protein residue side chains, the structures provide unambiguous identification of the segments that form the binding pocket. Specifically, each NAM binding pocket is formed almost exclusively within each subunit by segments in the subunit S1-M1 linker (from K509 to P520, including the pre-M1 helix) and the extracellular facing regions of the M4 helix (from L787 to V795) and M3 helix (from Y616 to R628). The identification of these parts of the S1-M1 and upper M4 segments was in good agreement with the results from the Gly scan in that all of the eight positions identified to change NAM potency upon Gly substitution are contained herein (Fig. 5). Also, the GluA2/NAM structures suggest that several residues within the upper M3 segment are in proximity to form direct interactions with the NAMs (Y616, L620, and F623, and S615 from the neighboring subunit). We included mutations of three of these residues to probe their potential involvement for direct interactions with the NAMs.

In total, we made additional mutations at 16 positions (Fig. 7) and determined effects on NAM potency using TEVC electrophysiology, which more directly measures AMPAR activation. Specifically, we expressed mutant and WT GluA2 receptors in Xenopus oocytes and determined the NAM IC₅₀ values for inhibition of steady-state agonist-evoked currents (see Materials and Methods). The standard recording protocols are illustrated in Fig. 6. In general, the IC₅₀ values obtained from the TEVC experiments were lower than those obtained from the [Ca²⁺]ᵢ experiments (Table 2). For example, PMP and GYKI displayed IC₅₀ values for WT GluA2 that were 5- and 10-fold lower than the corresponding values obtained in the [Ca²⁺]ᵢ experiments (Table 2). Additionally, relative changes in NAM potencies for mutants investigated by TEVC electrophysiology were generally smaller than the changes observed in the [Ca²⁺]ᵢ assay. It should be noted that during agonist application in the [Ca²⁺]ᵢ assay, desensitization is inhibited by coapplication of CTZ to produce robust Ca²⁺ responses (see Materials and Methods), whereas desensitization is not blocked during TEVC recordings. The difference in CTZ coapplication with NAM may underlie these differences since previous work has shown that CTZ, which binds in the LBD dimer interfaces to stabilize in a nondesensitizing conformation (Sun et al., 2002), allosterically influences the distinct NAM binding site to decrease apparent inhibitory potency (Johansen et al., 1995).

We created additional mutants at eight positions within the pre-M1 helix, the flanking S1-M1 linker segments, and the upper M1 and determined NAM potency (Fig. 7). At three positions in the pre-M1 helix (V514, F515, and F517), Gly substitutions indicated potential roles for NAM potency (Fig. 5). Since Gly has no side chain, Gly mutation is excellent for disrupting all potential ligand-protein interactions with the residue side chain. However, Gly also introduces flexibility to the protein backbone that might perturb local secondary structures or exert allosteric effects on the adjacent binding
pocket structure. Therefore, we mutated V514, F515, and F517 to alanine (Ala) to reduce their hydrophobic side chains, but maintain backbone structural constraints. Also, for the L518 position, where Gly substitution destroyed measurable activity, we created the functional L518A mutant to probe the effect of shortening the hydrophobic side chain. In general, we considered performing Ala substitutions at all positions where Gly substitution produced nonfunctional receptors; in particular, for residues with large hydrophobic side chains (such as, for example, W526 in the M1 region and F796 and Y797 in the M4 region) to create potentially functional mutants that would enable characterization of the impact of side chain reduction on NAM potency. However, since GluA2/NAM structures were available at this point in the mutational analysis, we excluded residues that appeared to be outside the NAM binding pockets from further analysis under the assumption that further substitution of such residues was outside the primary aim of this part of the mutational analysis, which focused on probing potential direct ligand contacts. In agreement with previous observations (Yelshanskaya et al., 2016), Ala substitutions at F515, F517, and L518 generally produced no or minor changes in NAM potency. For F515 and F517, these results indicate that the observed changes by Gly substitutions may involve allosteric effects. In contrast, the V514A mutation produced reductions in potency for all NAMs that were similar to those produced by the V514G mutation (Fig. 7). To further explore V514, we mutated this residue to aspartate (Asp; introducing a negatively charged side chain) and tryptophan (Trp; increasing side chain size). Interestingly, the pattern of effects of these mutations differed among the NAMs, with GYKI potency being reduced at V514D and V514W, whereas PMP and CP were not affected or displayed smaller increases in IC50 values of up to 5-fold (Fig. 7). In the GluA2/NAM structures, V514 appears to be located outside the binding pocket (Fig. 7). Thus, the effect pattern of large decreases by Gly and Ala substitutions and less effect of Asp and Trp substitutions may not result from perturbation of direct interactions between the NAMs and the V514 side chain, but rather reflect a role of V514 in shaping the NAM binding site.

At position D519 in the pre-M1 region and A522, Y523, and E524 in the upper M1 region (Fig. 7), we made further mutations based on previous reports of combinations of mutations at these positions in GluA3 and GluK2 to change NAM potency (Balannik et al., 2005). D519N displayed decreased potency for CP, which is in agreement with a suggested role of the γ-carboxylate in direct interaction with CP (Yelshanskaya et al., 2016). No other single mutations at A522, Y523, and E524 produced substantial effects on NAM potency (Fig. 7; Table 2). These results are in contrast to the previously suggested roles of these residues in interaction with GYKI (Balannik et al., 2005), but in general agreement with their positions outside the NAM binding pockets in the GluA2/NAM structures (Fig. 7).

We created additional mutants at five positions within the S2-M4 linker and upper M4 region (Fig. 7). The mutations were observed to produce similar effect patterns on PMP and...
GYKI potency in the \([\text{Ca}^{2+}]\), assay (Fig. 7; Table 2), which both were sensitive to mutations at N791 and insensitive to mutations at V792, A793, and G794 (Fig. 7). In contrast, CP displayed sensitivity toward mutation at all positions (Fig. 7; Table 2), despite the side chains of positions 792–794 being located outside the NAM binding pocket. In the TEVC experiments, the effects on NAM potencies by mutations at V792, A793, and G794 were generally less than 3-fold, indicating that neither of these residues forms key interactions with the NAMs (Table 2). These results indicate that blocking desensitization affects the role of S615 in NAM inhibition. F623A displayed substantial loss of potency for PMP and CP, and no change in CP potency, similar to the results obtained in the \([\text{Ca}^{2+}]\) assay and the previously reported effect of the S615A mutation on PMP and GYKI potency (Yelshanskaya et al., 2016). These results indicate that blocking desensitization affects the role of S615 in NAM inhibition. F623A displayed substantial loss of potency for PMP and CP in both the \([\text{Ca}^{2+}]\), and TEVC assays, supporting a role of the side chain phenyl group in direct ligand interactions as previously suggested (Yelshanskaya et al., 2016).

**Generation of NAM Binding Models in GluA2.** The GluA2/NAM crystal structures have inadequate resolution in the NAM binding pockets for assignment of side chain conformations for many residues around the bound NAMs; therefore, they provide limited information on specific ligand-protein interactions. To address this issue, Yelshanskaya et al. (2016) modeled several side chains based on conformations observed in a higher resolution structure of GluA2 obtained in complex with the competitive antagonist ZK200775 (PDB code 3KG2) (Sobolevsky et al., 2009), and then used docking calculations to create models of NAM binding modes. However, the GluA2/ZK200775 structure also has several side chains missing in the NAM binding pocket; therefore, these were not modeled in the docking calculations (M. Kurnikova, personal communication). The missing side chains close to the NAM binding pocket include K508, K509, K511, E627, R628, E782, K783, and T784. Thus, the conformations of several side chains that might form critical interactions with PMP, CP, and GYKI remain unknown. Therefore, we repeated the docking with the inclusion of the missing side chain atoms in the binding sites using the GluA2/NAM structures as a template (see Materials and Methods).
Methods. Initially, we employed a docking procedure with a flexible ligand docked into a rigid NAM binding pocket. To make the search as unbiased as possible, the search box used for the docking calculations was made large enough to include all four binding sites as well as the majority of the TMDs and the nearest part of the LBD. Of the 200 generated poses for each ligand (10 input conformations/poses per docking), at least 25% showed the ligand positioned in the binding site (27% for PMP, 35% for GYKI, and 79% for PMP). The remaining poses were located either in the pore, between transmembrane helices, or in between the TMDs and LBDs. The ligand poses in the binding sites showed considerable variability, possibly because only very few hydrogen bonds (HBs) were formed. However, we confirmed that poses similar to the binding modes observed in the crystal structures by Yelshanskaya et al. (2016) could be identified for each of the three ligands. The predicted binding energies for poses docked in the binding sites for all three ligands were similar, ranging from $-27.4$ to $-10.5$ kcal/mol; comparable to the results presented previously (Yelshanskaya et al., 2016). Thus, when docked into the GluA2/NAM structures with all side chains around the binding site added, all three NAM ligands can find binding modes very similar to those suggested by Yelshanskaya et al. (2016) (Supplemental Fig. 4). On this basis, we proceeded with the ligand orientations that were proposed from the GluA2/NAM structures to create three sets of binding models based on sampling flexible side chain orientations followed by energy minimizations to generate a total of 400 binding site models for each ligand (see Materials and Methods). Each model set represents a range of possible conformations of all binding pocket side chains and their potential ligand-protein interactions. Figure 8 illustrates the observed variability in side chain orientation for all residues within 5 Å of the NAMs (26 residues in total). For all three NAMs, we found the majority of proposed protein-ligand interactions to be primarily hydrophobic. Specifically, among the 26 residues surrounding the NAMs, 14 were purely hydrophobic, and four were mostly hydrophobic with a single polar group (K509, K511, Y616, and W526). Notably, the positions of the hydrophobic side chains showed only very little variability among models. In contrast, the side chain of K511, which was not resolved in the GluA2/NAM X-ray crystal structures (Yelshanskaya et al., 2016), displayed a wide range of conformations with the K511 side chain pointing toward the NAM molecules in some models, and more toward the lipid bilayer region in others. Figure 9 shows a representative model for each NAM, representing frequently observed side chain orientations among the 400 models. In these representative models, the phenyl ring of PMP is stacked between F623 and P520 (Figs. 5A and 5B).
8A and 9A); similar to the proposal by Yelshanskaya et al. (2016). F517 and V795 are lining the tip of the PMP pyridine ring, and the benzo-nitrile is lined by P512 and the carbon chain of K511. Similar to the X-ray crystal structures (Yelshanskaya et al., 2016), the methylene-dioxy group of GYKI is stacked against F623 (Figs. 8C and 9C). The amino-phenyl group of GYKI is surrounded by a hydrophobic pocket consisting of F517, P520, L620, L624, and L787, and the 5-methyl group is lined by P512 and the carbon chain of K511. The quinazolinone moiety of CP (surrounded by F517, P520, I611, L620, L787, V792, and V795) and the pyridine moiety are orientated parallel to F623 with its amino-substituent lined by the carbon chain of K511 and the cyano-phenyl pointing toward L787 and L624 (Figs. 8B and 9B).

In terms of potential hydrogen bond interactions, the NAM molecules may form HBs to both side chain or backbone atoms of the protein. Overall, 10 residues were observed within 5 Å distances of the NAM ligands that have side chains containing HB donor or acceptor moieties (Figs. 8 and 9). Bulk analysis of the orientations of these side chains in the model sets showed five of these (S510 and K511 in the S1-M1 linker, S516 in the pre-M1 helix, S615 and Y616 in the upper M3 helix, and N791 in the upper M4 helix) to frequently be oriented to potentially form HB interactions with the ligand. It should be noted that the initial side chain orientations for the optimized models were generated using MODELLER (Sali and Blundell, 1993), which ensures that there is space for the ligands but does not take into account the chemical nature of the ligand. The models were subsequently energy minimized to optimize the protein-ligand interactions, using proper parameters for the ligands. However, minimization will only locate a nearby local minimum; therefore, if a side chain is not reasonably close to the ligand initially, it will not form a HB during minimization. Thus, the method may not allow for the identification of all possible HBs, and the frequency of a given HB is not necessarily a measure of the importance of that HB for the binding of the ligand. However, the method allowed us to identify several possible HBs that were not indicated in the crystal structures or through docking. With this in mind, the five residues (K511, S516, S615, Y616, and N791) can be considered as potential key HB interaction points for NAM binding (Supplemental Figs. 5–7). Of these, K511 has not previously been suggested as an interaction point since the side chain was not resolved or modeled in the GluA2/NAM X-ray crystal structures (Yelshanskaya et al., 2016). For PMP and GYKI, three side chains are implicated in potential HB interaction with the ligand (K511, S516, and N791 for PMP and S510, S615, Y616, and N791 for GYKI), whereas only N791 is suggested for CP (Fig. 9). The side chain of N791 is thus suggested to participate in HB interactions with all three NAMs, similar to the proposal by Yelshanskaya et al. (2016).

For PMP, the side chain most frequently observed in an orientation to form potential HBs was N791, through the interaction of the δ-amide group of N791 with either the pyridine nitrogen or the nitrile of PMP (45% of our models in total) (Supplemental Fig. 5). The γ-hydroxyl group of the S516 side chain was within the distance for HB formation with the nitrile of PMP as suggested by Yelshanskaya et al. (2016) in 28% of the models. The ω-ammonium group of the K511 side chain was less frequently (14% of the models) within the distance for interaction with the nitrile and/or the carbonyl...
groups of PMP. Potential HB interactions between PMP and the side chains of S615 or Y616 as suggested by Yelshanskaya et al. (2016) were generally not observed in our models. For GYKI, N791 was also frequently observed in an orientation to form potential HB interactions, through interaction of the d-amide group of N791 with the amide oxygen and/or the benzodiazepine nitrogen of GYKI (26% and 33% of the models, respectively) (Supplemental Fig. 7). Similar to the X-ray crystal structures (Yelshanskaya et al., 2016), the g- and h-hydroxyl groups of S615 and Y616 were within the distance for hydrogen bonding with the amine of GYKI in 72% and 75% of the models, respectively. In contrast to suggestions by Yelshanskaya et al. (2016), the γ- and η-hydroxyl groups of S615 and Y616 were within the distance for hydrogen bonding with the amine of GYKI in 72% and 75% of the models, respectively. In contrast to suggestions by Yelshanskaya et al. (2016), the γ-hydroxyl group of the S516 and S788 side chains was infrequently or not at all within the distance to form HB interactions with GYKI in our models. For CP, the δ-amide group of N791 was within the HB distance of the carbonyl of CP in 52% of the models (Supplemental Fig. 6). Backbone-mediated HBs were not frequently observed in the models. Specifically, the analysis only suggests the backbone amide hydrogen of K511 to be oriented for potential HB interaction with the amide oxygen of GYKI (in 35% of the models) and the nitrile of PMP (in 24% of the models) (Supplemental Figs. 5–7). In general, in combination with the GluA2/NAM structures, our models fit well with most of the key experimental observations from the mutational analysis and provide an additional framework for understanding molecular details of AMPAR interactions with NAMs of three different chemical classes.

**Discussion**

The molecular pharmacology of AMPARs involves multiple binding pockets targeted by many chemical classes of exogenous ligands acting as synthetic agonists, channel blockers, competitive antagonists, PAMs, and NAMs. The location, structure, and molecular properties of binding pockets for competitive antagonists (Sobolevsky et al., 2009), agonists (Dürr et al., 2014; Yelshanskaya et al., 2014; Twomey et al., 2017a; Zhu and Gouaux, 2017), and positive modulators (Dürr et al., 2014; Yelshanskaya et al., 2014; Twomey et al., 2017a; Zhu and
Gouaux, 2017) are now defined at the atomistic level in the context of near full-length tetrameric AMPAR structures; leading to detailed models for how these ligands influence AMPAR dynamics to modulate function. In contrast, progress in NAM molecular pharmacology has been limited and essential aspects of NAM binding and functions have remained unknown. This is surprising since NAMs at present are the only class of AMPAR ligands from which a compound has emerged to become a therapeutic drug in humans. In the present study, our first aim was to use mutational analysis to pinpoint specific residues within the LBD-TMD linker regions as candidates for forming direct protein-ligand interactions. Our second aim was to use the mutational data to construct NAM binding models using existing GluA2 structures as docking templates. During work on both aims, X-ray crystallography structures of homomeric GluA2 in complex with the same prototypical NAMs selected for our study were reported (Yelshanskaya et al., 2016); thereby unambiguously identifying the number, location, and overall structure of the NAM binding pockets. However, the resolution of the X-ray structures was not sufficient for unambiguous identification of specific ligand-protein interactions involved in NAM binding. Thus, the data from our mutational analysis in combination with the recent GluA2/NAM structures form an improved basis for the construction of detailed binding mode models, which can be used for detailed analysis of ligand-protein interactions and serve as a framework for beginning to understand all molecular aspects of NAM function at AMPARs.

Overall, our mutational data fit well with the GluA2/NAM structures since most of the mutations changing NAM potencies were located in or near the NAM binding pockets, and these residues may thus potentially be critical for shaping the pocket or involved in direct ligand interactions. Some of these mutations concerned residues that were not resolved in the crystal structures. For example, we found that K511 in the S1-M1 linker is important for the potency of all three NAMs (Fig. 5; Table 2). Mutational effects at K511 could not be interpreted from the existing GluA2/NAM structures since the side chain of K511 is not resolved or modeled (Yelshanskaya et al., 2016). Our results from the side chain optimizations...
suggest that K511 can adopt a position to form HBs with the NAM ligands, either by interaction with the side chain ω-ammonium group or the backbone hydrogen (Figs. 8 and 9). Also, the aliphatic carbons of the K511 side chain may line up against hydrophobic moieties of the NAMs, and thus form hydrophobic interactions. Interestingly, in other GluA2 structures for which the side chain of K511 has been resolved, several different conformations and orientations of the K511 side chain were observed. These included structures in which the side chain is orientated such that the primary amine points directly toward the empty NAM binding pocket (e.g., PDB codes 4U4F and 4U4G) (Yelshanskaya et al., 2014). In other structures, the K511 side chain points toward the membrane region; such as observed in a recently determined structure of a heteromeric GluA1/2 receptor in complex with transmembrane AMPAR regulatory protein y8 (PDB code 6QKC) (Herguedas et al., 2019). Interestingly, in this structure, the K511 side chain is pointing directly toward the extracellular region of y8 and a membrane lipid molecule. Hence, the variation in side chain conformations observed in the modeling appears convincing. Because we only tested the mutation of K511 to Gly, and Gly introduces backbone flexibility, it is plausible that the mutation may have affected hydrogen bonding capacities of both the backbone and side chain as well as hydrophobic interactions of the aliphatic carbons. Thus, our models and mutational data together point toward K511 as a novel interaction point for the NAMs. However, there are also examples of mutations of residues that we found exerted substantial effects on NAM potencies that cannot be readily explained by direct interactions in the current structures or our models. For example, V514 in the S1-M1 linker region is in the periphery of the binding pocket, and its side chain is pointing away from the NAM ligands (Fig. 7). Yet, several mutations of this residue produced substantial changes in potency for all three NAMs (Figs. 5 and 7). V514 is located in the pre-M1 helix of the linker, and this residue potentially has a critical role for the stability or position of this structural element. The pre-M1 helix is part of the NAM pocket, and several residues in the pre-M1 helix make direct contacts to the NAMs. Thus, the results for V514 are likely an example of how point mutations can affect ligand potencies via indirect allosteric effects on ligand binding pockets.

Key interactions that were previously suggested from the GluA2/NAM crystal structures and mutational analysis by Yelshanskaya et al. (2016) are supported by our mutational data and models. These include hydrogen bonding interactions involving N791, and hydrophobic and π-stacking interactions involving F520 and F623 (Figs. 8 and 9). In contrast, the idea of key HB interactions of the side chains of S788 and Y616 with all of the tested NAMs is not supported by our mutational data or modeling. Specifically, in our modeling we only observed an HB interaction between Y616 and GYKI. However, the Y616F mutation did not change GYKI potency; therefore, the presence of this interaction was not supported (Fig. 7). S615 has also been suggested to form HBs to the NAM ligands, and mutation to Ala was previously reported to cause an order of magnitude loss of potency for GYKI and PMP (Yelshanskaya et al., 2016). When desensitization was blocked by CTZ in the [Ca2+]i assay and TEVC experiments [similar to the patch-clamp electrophysiology experiments by Yelshanskaya et al. (2016)], we observed similar loss-of-potency effects by the S615A mutation, while CP potency remained unchanged (Fig. 7; Supplemental Fig. 3). However, in the TEVC experiments without CTZ, S615A did not change potency for PMP and GYKI, but increased potency of CP by an order of magnitude. These results suggest that the role of S615 in NAM binding is state dependent, e.g., that block of desensitization by CTZ causes a different conformation of S615 compared with the receptor population in our TEVC experiments without CTZ in which the vast majority of receptors are desensitized. Interestingly, the S615A mutant has been previously reported to display slower rates of deactivation and desensitization and an increased fraction of steady-state to peak current (Yelshanskaya et al., 2017). Several other mutants display differential effects on NAM potencies between assays, albeit not to the same extent as the S615A mutant. Possibly, other residues in the NAM binding region play state-dependent roles in NAM inhibition due to state-dependent variations in their structural conformations or mutational effects on receptor desensitization as reported for several residues in the region (Yelshanskaya et al., 2017). This aspect of NAM molecular pharmacology is of potential interest since AMPAR activation and desensitization are thought to involve transitions between multiple open and desensitized states (Robert et al., 2001; Jin et al., 2003; Robert and Howe, 2003). Also, the receptor activation mechanism may involve multiple preactivated states (Durr et al., 2014). The dependence of NAM binding on different receptor conformations underlying this potential multitude of functional states and which state transitions are affected by NAM binding are at present unknown. Concerning this, it is interesting to note early observations (Johansen et al., 1995) that NAM potency appears to be affected by PAM ligands that now are known to bind in the LBD dimer interfaces and stabilize nondesensitized conformations of the LBD layer. Interestingly, our results indicate that the interactions of specific AMPAR residues with the NAMs can be altered by the presence of PAMs. Future modeling studies, potentially involving molecular dynamics simulations, aiming to compare the configuration of the NAM binding site in AMPAR structures representing different functional states seem warranted. In addition, inclusion of a lipid bilayer in the models may likely provide additional insight into how membrane lipids influence the NAM binding pocket. Such studies may reveal potentially important effects of the presence of different types of lipids on side chain orientation of residues such as V514, which in our models appears to orientate toward the membrane. Furthermore, an important open question to answer will be the role of cooperability of the four NAM sites per receptor complex; including the number of sites needed to be occupied to achieve inhibition of channel gating. Moreover, auxiliary subunits also influence AMPAR states and transitions; adding a layer of complexity to understanding the structural basis of AMPAR function (Dawe et al., 2016; Greger et al., 2017). In this relation, it is interesting to note that novel classes of NAMs have recently been reported that modulate AMPARs depending on the presence of certain members of the transmembrane AMPAR regulatory protein family (Gardinier et al., 2016; Maher et al., 2016; Azumaya et al., 2017).

In summary, we have performed mutational analysis and modeling of the NAM binding pockets and ligand binding modes in the GluA2 AMPAR, which in combination with recent X-ray crystal structures published by others (Yelshanskaya et al., 2016) constitute a step toward achieving a full molecular understanding of this important class of AMPAR inhibitors.