# Intrasubunit and Intersubunit Steroid Binding Sites Independently and Additively Mediate $\alpha 1\beta 2\gamma 2L$ GABA<sub>A</sub> Receptor Potentiation by the Endogenous Neurosteroid Allopregnanolone<sup>SI</sup>

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## ABSTRACT

Prior work employing functional analysis, photolabeling, and Xray crystallography have identified three distinct binding sites for potentiating steroids in the heteromeric GABA<sub>A</sub> receptor. The sites are located in the membrane-spanning domains of the receptor at the  $\beta$ - $\alpha$  subunit interface (site I) and within the  $\alpha$ (site II) and  $\beta$  subunits (site III). Here, we have investigated the effects of mutations to these sites on potentiation of the rat  $\alpha 1\beta 2\gamma 2L$  GABA<sub>A</sub> receptor by the endogenous neurosteroid allopregnanolone ( $3\alpha 5\alpha P$ ). The mutations were introduced alone or in combination to probe the additivity of effects. We show that the effects of amino acid substitutions in sites I and II are energetically additive, indicating independence of the actions of the two steroid binding sites. In site III, none of the mutations tested reduced potentiation by  $3\alpha 5\alpha P$ , nor did a mutation in site III modify the effects of mutations in sites I or II. We infer that the binding sites for  $3\alpha 5\alpha P$  act independently. The independence of steroid action at each site is supported by photolabeling data showing that mutations in either site I or site II selectively change steroid orientation in the mutated site without affecting labeling at the unmutated site. The findings are discussed in the context of linking energetic additivity to empirical changes in receptor function and ligand binding.

## SIGNIFICANCE STATEMENT

Prior work has identified three distinct binding sites for potentiating steroids in the heteromeric  $\gamma$ -aminobutyric acid type A receptor. This study shows that the sites act independently and additively in the presence of the steroid allopregnanolone and provide estimates of energetic contributions made by steroid binding to each site.

# Introduction

The  $\gamma$ -aminobutyric acid type A (GABA<sub>A</sub>) receptor is the major ionotropic transmitter-gated inhibitory ion channel in the brain. It normally responds to synaptically released or ambient GABA, but a large number of endogenous and clinical compounds, including neurosteroids, benzodiazepines, and several intravenous anesthetics can activate the receptor and/or modulate its response to the transmitter. Endogenous

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neurosteroids and their synthetic analogs exhibit a large range of efficacies on the GABA<sub>A</sub> receptor (Park-Chung et al., 1999; Li et al., 2007a). Steroids have an apparent affinity to the receptor in the micromolar range, although the actual  $K_{\rm D}$ s may be several log-orders higher because of accumulation of steroid in the lipid bilayer (Akk et al., 2005; Li et al., 2007b).

Functional analysis, photolabeling, and X-ray crystallography have identified three distinct binding sites for potentiating steroids in the membrane-spanning domains of the heteromeric GABA<sub>A</sub> receptor (Hosie et al., 2006; Laverty et al., 2017; Miller et al., 2017; Ziemba et al., 2018; Chen et al., 2019). The sites are located at the  $\beta$ - $\alpha$  subunit interface (dubbed site I) and within the  $\alpha$  (site II) and  $\beta$  subunits (site III) (Fig. 1). Site I is the canonical binding site located in the cleft between the  $\beta$  subunit "+" and the  $\alpha$  subunit "-" surfaces. The steroid positions with its A-ring oriented toward the

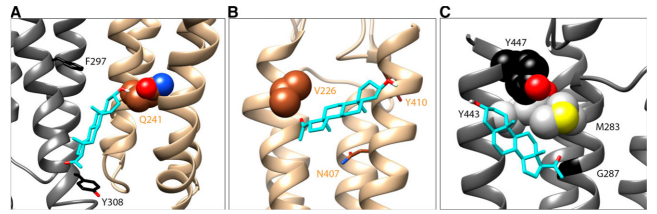
**ABBREVIATIONS:**  $3\alpha5\alpha$ P, 1-[(3R,5S,8R,9S,10S,13S,14S,17S)-3-hydroxy-10,13-dimethyl-2,3,4,5,6,7,8,9,11,12,14,15,16,17-tetradecahydro-1*H*-cyclopenta[a]phenanthren-17-yl]ethanone (allopregnanolone); DDM, *n*-dodecyl- $\beta$ -D-maltoside; DTT, dithiothreitol; ELIC, *Erwinia* ligand-gated ion channel; GABA<sub>A</sub>,  $\gamma$ -aminobutyric acid type A; KK200, ( $3\alpha,5\alpha,17\beta$ )-17-[[4-[3-(trifluoromethyl)-3*H*-diazirin-3-yl]phenyl] methoxy]-androstane- 3- ol; MBP, maltose binding protein; MS, mass spectrometry; PA, probability of being in the active state; PR, probability of being in the resting state; TM, membrane-spanning segment; TMD, transmembrane domain.

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**Fig. 1.** Docking of  $3\alpha 5\alpha P$  in the putative binding sites. The panels show side views of the putative  $\beta$ - $\alpha$  intersubunit binding site (site I; A), the intra- $\alpha$  subunit site (site II; B), and the intra- $\beta$  subunit site (site III; C). The  $\alpha$  subunit is shown in tan, the  $\beta$  subunit in gray.  $3\alpha 5\alpha P$  (cyan) is docked in each site. In site I, the Q241 residue in  $\alpha$ 1TM1 is shown as a sphere. Residues homologous to Y308 (Y309) and F297 (F298) in TM3 are photolabeled by KK200 in ELIC- $\alpha$ 1 (q242L), respectively, and are shown as ball & stick. In site II, the V226 residue in  $\alpha$ TM1 is shown as a sphere. The N407 and Y410 residues in  $\alpha$ TM4 previously identified to contribute to steroid binding are shown as ball & stick. In site IIII, the M283 (light gray;  $\beta$ TM3), Y443 (light gray;  $\beta$ TM4), and Y447 (black;  $\beta$ TM4) are shown as sphere. The G287 residue in  $\beta$ TM3 is identified as black ribbon.

hydrophobic core of the membrane facing the  $\alpha$ (Q241) residue in the first membrane-spanning segment (TM 1), and the Dring near the  $\beta$ (F301) residue in TM3 pointing toward the cytoplasmic side of the membrane. Site II is within a cavity between the extracellular ends of the TM1 and TM4 segments of the  $\alpha$  subunit. The A-ring of the steroid is proximal to the  $\alpha$ (Y415) residue at the extracellular side of the membrane, and the D-ring is positioned near the  $\alpha$ (N407) residue. In TM1, the steroid abuts the  $\alpha$ (V226) residue. Site III is located between the TM3 and TM4 segments of the  $\beta$  subunit. The A-ring of the steroid is positioned toward the extracellular domain near the  $\beta$ (Y443) residue in TM4 and the Dring near  $\beta$ (G287) of TM3.

Mutations introduced to sites I [e.g.,  $\alpha 1(Q241L)$  and  $\alpha Q(241S)$ ] and II [e.g.,  $\alpha 1(V226W)$ ] reduce receptor potentiation by allopregnanolone ( $3\alpha 5\alpha P$ ) and other potentiating steroids thus demonstrating the functionality of these binding sites (Hosie et al., 2006; Akk et al., 2008; Chen et al., 2019). In contrast, no change in potentiation by  $3\alpha 5\alpha P$ or pregnanolone was observed in a recent study probing the effects of amino acid substitutions at a limited set of residues in site III in the  $\alpha 1\beta 3$  GABA<sub>A</sub> receptor (Chen et al., 2019).

The existence of several distinct sites in the relatively compact transmembrane regions of the receptor raises the possibility that the sites interact directly with each other by some form of conformational coupling. Here, we have investigated the effects of mutations to these three steroid binding sites on potentiation of the  $\alpha 1\beta 2\gamma 2L$  GABA<sub>A</sub> receptor by the endogenous steroid  $3\alpha 5\alpha P$ . The mutations were introduced alone or in combination to probe the additivity of effects. We show that the effects of amino acid substitutions in sites I and II are energetically additive, indicating independence of steroid actions at the two binding sites. We also used neurosteroid analog photolabeling to show that mutations in either site I or site II selectively affect steroid orientation and efficacy of labeling only in the mutated site, supporting the absence of allosteric interaction between the sites. In site III, none of the mutations tested reduced potentiation by  $3\alpha 5\alpha P$ . The findings are discussed in the context of linking energetic additivity to empirical changes in receptor function and ligand binding.

## Materials and Methods

Receptor Expression and Electrophysiological Recordings. The rat  $\alpha 1\beta 2\gamma 2L$  GABA<sub>A</sub> receptors were expressed in oocytes from *Xenopus laevis* (African clawed frog). Quarter ovaries were purchased from Xenoocyte (Dexter, MI), and digested at 37°C with shaking at 250 RPM for 15–30 minutes in 2% w/v (mg/mL) collagenase A solution in ND96(96 mM NaCl, 2 mM KCl, 1.8 mM CaCl<sub>2</sub>, 1 mM MgCl<sub>2</sub>, 5 mM HEPES; pH 7.4) with 100 U/ml penicillin and 100 µg/ ml streptomycin. After digestion, the oocytes were rinsed in ND96 and stored in ND96 with supplements (2.5 mM Na pyruvate, 100 U/ ml penicillin, 100 µg/ml streptomycin, 50 µg/ml gentamycin) at 15°C for at least 4 hours before injection.

The cDNAs for rat  $\alpha 1$  (GenBank accession number NM\_183326),  $\beta 2$  (NM\_012957), and  $\gamma 2L$  (NM\_183327) subunits in the pcDNA3 vector were linearized with XbaI (NEB Laboratories, Ipswich, MA). The following mutant clones, generated using QuikChange (Agilent Technologies, Santa Clara, CA), were studied:  $\alpha 1$ (Q241L),  $\alpha 1$ (Q241S),  $\alpha 1$ (V226W),  $\alpha 1$ (Q241L + V226W),  $\alpha 1$ (Q241S + V226W),  $\beta 2$ (M283A),  $\beta 2$ (G287W),  $\beta 2$ (Y443W), and  $\beta 2$ (Y447A). The clones were fully sequenced prior to use. The complementary RNAs were generated from linearized cDNA using Message Machine (Ambion, Austin, TX).

The oocytes were injected with a total of 3.5 ng of complementary RNA per oocyte, in the ratio of 1:1:5 ( $\alpha$ 1: $\beta$ 2: $\gamma$ 2L) to enhance expression and incorporation of the  $\gamma$ 2L subunit. After injection, the oocytes were incubated in ND96 with supplements (2.5 mM Na pyruvate, 100 U/ml penicillin, 100 µg/ml streptomycin, and 50 µg/ml gentamycin) at 15°C for 1 to 2 days prior to conducting electrophysiological recordings.

The electrophysiological recordings were performed using standard two-electrode voltage clamp. The oocytes were clamped at -60 mV. Bath and drug (GABA, steroids, propofol) solutions were gravity-applied from glass syringes with glass luer slips via Teflon tubing to the recording chamber (RC-1Z, Warner Instruments, Hamden, CT) at a flow rate of 6-8 ml/min. Solutions were switched manually using 4-port bulkhead switching valve and medium-pressure 6-port bulkhead valves (IDEX Health and Science, Rohnert Park, CA). All experiments were conducted at room temperature. GABA concentration-response curves were measured by exposing cells expressing GABA<sub>A</sub> receptors to 1, 3, 10, 30, 100, 300, and 1000  $\mu$ M GABA.  $3\alpha5\alpha$ P concentration-response curves were measured by exposing cells to a low concentration of GABA alone, and in the presence of 0.01, 0.03, 0.1, 0.3, 1, 3, and 10  $\mu$ M steroid. The drug applications lasted 20–80 seconds and were aimed at determining the amplitude of the peak response. Successive drug applications were separated by >1-minute washouts in ND96. Each cell was exposed to the full range of GABA or steroid concentrations. Some experiments were conducted by measuring the effect of a steroid on steady-state activity elicited by a low concentration of GABA.

The current responses were amplified with an OC-725C amplifier (Warner Instruments, Hamden, CT), digitized with a Digidata 1200 series digitizer (Molecular Devices), and stored using pClamp (Molecular Devices). The current traces were analyzed using Clampfit (Molecular Devices).

Analysis of Electrophysiological Data. GABA concentrationresponse curves were fitted, separately for each cell, to the Hill equation using raw current amplitudes.  $3\alpha5\alpha P$  concentration-response data were first converted to units of probability of being in the active state (P<sub>A</sub> units) through normalization to the peak response to saturating (1 mM) GABA + 50  $\mu$ M propofol, that was considered to have a peak P<sub>A</sub> indistinguishable from 1 (Shin et al., 2017). The probability of constitutive activity in the absence of any applied agonist was determined by comparing the response amplitudes to 100  $\mu$ M picrotoxin and 1 mM GABA + 50  $\mu$ M propofol. The level of constitutive activity was negligible (< 0.003) for all receptors used and was thus not included in the calculation of P<sub>A</sub>.

The peak current responses expressed in  $P_A$  units were then analyzed using a cyclic two-state concerted transition model (Steinbach and Akk, 2019). The  $3\alpha5\alpha P$  concentration-response curves were fitted, separately for each cell, to the state function:

$$P_{A, \text{peak}} = \frac{1}{1 + L^* \left[\frac{1 + [3\alpha 5\alpha P]/K_{B, 3\alpha 5\alpha P}}{1 + [3\alpha 5\alpha P]/(K_{B, 3\alpha 5\alpha P} - 3\alpha 5\alpha P]}\right]^N}$$
(1)

where L\* expresses the level of background activity in the presence of a low concentration of GABA, and can be calculated as  $(1 - P_{A,GABA})/P_{A,GABA}$ .  $K_{R,3z5zP}$  is the equilibrium dissociation constant for  $3\alpha 5\alpha P$  in the resting receptor,  $c_{3\alpha 5xP}$  is the ratio of the equilibrium dissociation constant for the steroid in the active receptor to  $K_{R,3z5zP}$ , and  $[3\alpha 5\alpha P]$  is the concentration of the steroid.  $N_{3z5zP}$ , the number of binding sites for  $3\alpha 5\alpha P$ , was constrained to 2 in the initial analysis.

Curve fitting was done using Origin 2020 (OriginLab Corp., Northampton, MA) with L\*,  $K_{R,3z5zP}$ , and  $c_{3z5zP}$  as free parameters. Statistical analyses were done using Excel 2016 (Microsoft, Redmond, WA). Steroid-induced potentiation is calculated as ratio of peak responses in the presence and absence of steroid (1 = no effect). The results are reported as mean  $\pm$  S.D. from at least 5 cells for each experiment. All data are included in analysis. The study is exploratory by nature. The findings are reported according to the guidelines detailed in (Michel et al., 2020).

**ELIC**- $\alpha$ **1 Construct Design.** The ELIC- $\alpha$ **1** GABA<sub>A</sub> receptor chimera was constructed by fusing the extracellular domain of ELIC (*Erwinia* ligand-gated ion channel) to the transmembrane domain (TMD) of human  $\alpha$ **1** subunit of the GABA<sub>A</sub> receptor (Kinde et al., 2016; Chen et al., 2018; Sugasawa et al., 2019). The intracellular loop amino acids were substituted by the short linker -GVE-sequence. A decahistidine tag and a maltose binding protein (MBP) were incorporated at the amino-terminal, and a tobacco etch virus protease cleavage site was inserted between MBP and ELIC- $\alpha$ <sub>1</sub> GABA<sub>A</sub> receptor (Chen et al., 2018; Sugasawa et al., 2019).

**Expression and Purification.** Mutagenesis was performed as described previously (Kinde et al., 2016; Chen et al., 2018; Sugasawa et al., 2019) by oligonucleotide-directed mutagenesis using *Phu* polymerase (Thermo Fisher Scientific) verified by sequencing. The ELIC- $\alpha$ 1 GABA<sub>A</sub> receptor was expressed in OverExpress C43(D3) *E. coli*,

derived from Rosetta (DE3) cells under double selection with kanamycin (50 µg/mL) and chloramphenicol (10 µg/mL). A few isolated colonies were used to inoculate 100 mL of LB (Sigma-Aldrich) media with the antibiotics and grown overnight at 37°C in a shaker at 250 rpm. The overnight culture was then diluted 1:100 into  $6 \times 1$  liters of LB media with antibiotics and grown to an optical density of 0.7-0.8. All six liters were harvested (5000 rpm, 15 minutes, 4°C) and resuspended in 2 liters of LB media supplemented with 0.5 M sorbitol. The concentrated cells were equilibrated in a shaker at 15°C and 250 rpm for 1 hour before inducing expression with 0.2 mM isopropyl  $\beta$ -D-1-thiogalactopyranoside. The cells were harvested after 20 hours expression, resuspended in buffer A (20 mM Tris pH 7.5, 100 mM NaCl), complete EDTA-free protease inhibitor cocktail (Roche), and DNase, lysed using an Avestin Emulsiflex C-5 at 15,000 psi, pelleted by ultracentrifugation, and homogenized. The fusion protein was extracted with 1%  $n\text{-}dodecyl\text{-}\beta\text{-}D\text{-}maltoside}$  (DDM). Solubilized protein was purified using amylose resin and eluted using buffer A containing 40 mM maltose and 0.02% DDM. His-MBP-(ELIC- $\alpha_1$  GABA<sub>A</sub> receptor) was digested overnight with tobacco etch virus protease, cleaned up using a reverse nickel on nitrilotriacetic acid (Ni-NTA) purification, and injected onto a Sephadex 200 Increase 10/300 column, which yielded pentameric protein in buffer A + 0.02% DDM.

Photolabeling and Intact Protein MS Analysis. The synthesis of neurosteroid photolabeling reagent  $(3\alpha, 5\alpha, 17\beta)$ -17-[[4-[3-(trifluoromethyl) - 3H- diazirin-3-yl] phenyl] methoxy] -androstane- 3- ol (KK200) has been detailed previously (Cheng et al., 2018). Purified ELIC-a1 GABA<sub>A</sub> receptor was photolabeled and analyzed by intact protein mass spectrometry (MS) as described previously (Budelier et al., 2017; Cheng et al., 2018; Sugasawa et al., 2019). Briefly, 50 µg of detergent-solubilized ELIC-a1 GABAA receptor (0.4 mg/ml in buffer A + DDM) was incubated with 100  $\mu$ M KK200 in a glass tube for 60 minutes at 4°C in the dark. The sample was transferred to a quartz cuvette and then placed in a photoreactor at a distance of 7 cm from the source. The photoreactor uses a 450W Hanovia medium-pressure mercury lamp as the light source. A cold-water jacket cooled the lamp, and the light was filtered through a 1.5 cm thick saturated copper sulfate solution. This filter absorbs all light of wavelength < 315 nm (Katzenellenbogen et al., 1974). Illuminance at the middle of the cuvette was approximately 10 lumens/cm<sup>2</sup> as measured with a Newport model 1918-C optical meter. The samples were irradiated for 5 minutes and continuously cooled to 4°C. For the intact protein MS analysis, the irradiated samples were treated with 250 mM dithiothreitol (DTT) and then precipitated with chloroform/methanol/water. The precipitated protein was washed three times with equal volumes of water and methanol, centrifuged, and the protein pellet was reconstituted in 3 µl of 90% formic acid followed by 100 µl of 4:4:1 chloroform/methanol/water. Reconstituted samples were then analyzed in an Orbitrap Elite mass spectrometer (Thermo Fisher Scientific) by direct injection at 3 µl/min using a Max Ion atmospheric pressure ionization source with a heated electrospray ionization-II probe. Full spectra of photolabeled ELIC-a1 GABAA receptor were acquired on the linear trap quadrupole using spray voltage of 4 kV, capillary temperature of 320°C, and in-source dissociation of 30 V. Deconvolution of intact ELIC-a1 GABAA receptor spectra was performed using UniDec (Marty et al., 2015).

**Tryptic Middle-Down MS Analysis.** Photolabeled ELIC- $\alpha$ 1 GABA<sub>A</sub> receptor was analyzed by middle-down MS as detailed in previous reports (Budelier et al., 2017; Cheng et al., 2018; Chen et al., 2019; Sugasawa et al., 2019, 2020). 20 µg (0.4 mg/ml) of photolabeled ELIC- $\alpha$ 1 GABA<sub>A</sub> receptor was buffer-exchanged to 50 mM triethylammonium bicarbonate buffer with 0.02% DDM using Micro Bio-Spin 6 columns (Bio-Rad). The buffer-exchanged sample was reduced with 5 mM tris(2-carboxyethyl)phosphine for 1 hour, alkylated with 5 mM N-ethylmaleimide for 1 hour, and quenched with 5 mM DTT for 15 minutes. Samples were then digested with 8 µg of trypsin for 7 days at 4°C; extended digestion at low temperature was necessary to obtain maximal recovery of TMD peptides. Next, formic

# 22 Germann et al.

acid was added to 1%, followed directly by liquid chromatography mass spectrometry analysis on an Orbitrap Elite mass spectrometer. 20-µl samples were injected into a home-packed PLRP-S (Agilent) column (10 cm  $\times$  75 µm, 300 Å), separated with a 145-minute gradient from 10% to 90% acetonitrile, and introduced to the mass spectrometer at 800 nl/min with a nanospray source. MS acquisition was set as an MS1 Orbitrap scan (resolution of 60,000) followed by top 20 MS2 Orbitrap scans (resolution of 15,000) using data-dependent acquisition, and exclusion of singly charged precursors.

Fragmentation was performed using high-energy dissociation with normalized energy of 35%. Analysis of data sets was performed using Xcalibur (Thermo Fisher Scientific) to manually search for TM1, TM2, TM3, or TM4 tryptic peptides with or without neurosteroid photolabeling modifications. Photolabeling efficiency was estimated by generating extracted chromatograms of unlabeled and labeled peptides, determining the area under the curve, and calculating the abundance of labeled peptide/(unlabeled + labeled peptide). MS2 spectra of photolabeled TMD peptides were analyzed by manual assignment of fragment ions with and without photolabeling modification. Fragment ions were accepted based on the presence of a monoisotopic mass within 20 ppm mass accuracy. In addition to manual analysis, PEAKS (Bioinformatics Solutions Inc.) data base searches were performed for data sets of photolabeled ELIC- $\alpha 1$ GABA<sub>A</sub> receptor. Search parameters were set for a precursor mass accuracy of 20 ppm, fragment ion accuracy of 0.1 Da, up to 3 missed cleavages on either end of the peptide, false discovery rate of 0.1%, and variable modifications of methionine oxidation, cysteine alkylation with N-ethylmaleimide and DTT, and neurosteroid analog photolabeling reagent on any amino acid.

**Simulations of Function and Muscimol Binding.** Simulations of peak current responses were carried out using a modified state function in which the effects of agonist X were mediated by two classes of sites:

$$P_{A, \text{peak}} = \frac{1}{1 + L^* \left[\frac{1 + |X| / K_{B, X, 1}}{1 + |X| / (c_{X, 1} K_{B, X, 1})}\right]^{N_1} \left[\frac{1 + |X| / K_{B, X, 2}}{1 + |X| / (c_{X, 2} K_{B, X, 2})}\right]^{N_2}}$$
(2)

where  $K_{R,X,1}$  and  $K_{R,X,2}$  are the equilibrium dissociation constants of X for the two classes of binding sites in the resting receptor, and  $c_{X,1}$  and  $c_{X,2}$  are the measures of efficacy.  $N_1$  and  $N_2$  give the number of binding sites per receptor for X within each class of binding sites. L\* was constrained to 1000 to mimic direct activation by X or to 19 to mimic X-induced potentiation of background activity with  $P_{A,peak}$  of 0.05.

Muscimol binding in the presence of an allosteric agonist X was simulated as described previously (Akk et al., 2020):

$$Y_{M} = \frac{QL^{*}\alpha_{M}(1+\alpha_{M})^{N_{M}-1} + Q_{c_{M}}^{(\underline{\alpha}_{M})}(1+(\frac{\alpha_{M}}{c_{M}}))^{N_{M}-1} + (\frac{\alpha_{M}}{d_{M}c_{M}})(1+(\frac{\alpha_{M}}{d_{M}c_{M}}))^{N_{M}-1}}{QL^{*}(1+\alpha_{M})^{N_{M}} + Q(1+(\frac{\alpha_{M}}{c_{M}}))^{N_{M}} + (1+(\frac{\alpha_{M}}{d_{M}c_{M}}))^{N_{M}}}$$
(3)

where  $Y_M$  is the fraction of orthosteric binding sites occupied by muscimol, Q is the ratio of receptors in the active state relative to desensitized state when neither muscimol nor X is bound [assumed 0.29; (Germann et al., 2019)],  $\alpha_X = [muscimol]/K_{R,muscimol}$ , N<sub>M</sub> is the number of muscimol binding sites [assumed 2; (Chang et al., 1996)],  $c_M$  is the ratio of the equilibrium dissociation constant for muscimol in the active receptor (K<sub>A,muscimol</sub>) to K<sub>R,muscimol</sub>, and  $d_M$  is the ratio of the equilibrium dissociation constant for muscimol in the desensitized receptor to K<sub>A,muscimol</sub> [assumed to be 1; (Germann et al., 2019)]. In simulations, the muscimol concentration was selected to generate 10% equilibrium occupancy of muscimol binding sites in the absence of X.  $\mathrm{L}^*$  for eq. 3 was calculated as:

$$\mathbf{L}^{*} = \mathbf{L} \left[ \frac{1 + [X]/K_{\text{R},X,1}}{1 + [X]/(c_{X,1}K_{\text{R},X,1})} \right]^{N_{1}} \left[ \frac{1 + [X]/K_{\text{R},X,2}}{1 + [X]/(c_{X,2}K_{\text{R},X,2})} \right]^{N_{2}} \quad (4)$$

where L is the ratio of resting to active receptors in the absence of agonist [assumed 8000; (Shin et al., 2017)]. Other terms are as described above. In essence, the presence of the allosteric agonist X enhances background activity by reducing the value of L, which then increases muscimol occupancy of high-affinity states (active and desensitized).

**Docking Studies of 3\alpha 5\alpha P.** Site I ( $\beta$ - $\alpha$  interface) was modeled using the cryoEM structure of the human  $\alpha 1\beta 3\gamma 2$  GABA receptor [PDB ID:6I53, (Laverty et al., 2019)]. The site is defined by TM1 of the  $\alpha 1$  subunit and TM3 of the  $\beta 2$  subunit. In this region, there are only two sequence differences between the human  $\beta$ 3 and rat  $\beta$ 2, L294M and F301L. These mutations were made in the modeling program chimera to the structure. Homology models of the rat  $\alpha 1$  and  $\beta 2$  subunits were built using a multitemplate approach using the Modeler program. The structures used as templates were the human  $\alpha 1\beta 3\gamma 2$  GABA receptor [PDB ID:6I53; and 6HUO, (Masiulis et al., 2019)] and the human  $\alpha 1\beta 1\gamma 2$  GABA receptor [PDB ID: 6DW0, (Phulera et al., 2018)]. The best scoring of 100 produced models of each subunit was then used for docking. The docking sites were defined by the photolabeled residues (Sugasawa et al., 2020).  $3\alpha 5\alpha P$  was then docked into each site with the program Autodock Vina using a box size of  $15 \times 15 \times 15$  Å, an energy range of 3 kcal, exhaustiveness of 25, and a num mode of 20.

**Drugs.** The stock solution of GABA was made in ND96 at 500 mM and stored in aliquots at  $-20^{\circ}$ C. The steroids  $3\alpha 5\alpha P$  and alfaxalone were dissolved in DMSO at 20 mM and 10 mM, respectively, and stored at room temperature. Final dilutions were made on the day of experiment.

## Results

Activation of Wild-Type and Mutant Receptors by GABA. The following mutations to the three putative binding sites for potentiating steroids were tested:  $\alpha 1(Q241L)$  and  $\alpha 1(Q241S)$  in site I ( $\beta$ - $\alpha$  interface),  $\alpha 1(V226W)$  in site II (intra- $\alpha$  subunit site), and  $\beta 2(M283A)$ ,  $\beta 2(G287W)$ ,  $\beta 2(Y443W)$ , and  $\beta 2(Y447A)$  in site III (intra- $\beta$  subunit site). We commenced by verifying receptor expression and function by measuring activation of the mutant receptors by GABA. Cells expressing  $\alpha 1\beta 2\gamma 2L$  wild-type or mutant receptors were exposed to a range of concentrations of GABA, and the peak current amplitudes were determined and analyzed using the Hill equation. The EC<sub>50</sub>s and Hill coefficients of the fits are given in Table 1.

#### TABLE 1

Activation of  $\alpha1\beta2\gamma2L$  wild-type and mutant receptors by GABAThe table gives the  $EC_{50}s$  and Hill coefficients (mean  $\pm$  S.D. from 5–8 cells per receptor) for wild-type and mutant receptors.

Receptor	GABA $EC_{50}$	n <sub>Hill</sub>
	$\mu M$	
$\alpha 1\beta 2\gamma 2L$ wild-type	$16.3 \pm 7.4$	$1.32 \pm 0.02$
$\alpha 1(Q241L)\beta 2\gamma 2L$	$444 \pm 199$	$0.76 \pm 0.02$
$\alpha 1(Q241S)\beta 2\gamma 2L$	$23.7 \pm 11.9$	$1.37 \pm 0.13$
$\alpha 1 (V226W) \beta 2 \gamma 2L$	$150 \pm 72$	$0.80 \pm 0.05$
$\alpha 1(Q241L+V226W)\beta 2\gamma 2L$	$309 \pm 241$	$0.78 \pm 0.06$
$\alpha 1(Q241S+V226W)\beta 2\gamma 2L$	$237 \pm 129$	$0.82 \pm 0.06$
$\alpha 1\beta 2(M283A)\gamma 2L$	$22.5 \pm 12.2$	$1.13 \pm 0.13$
$\alpha 1\beta 2(G287W)\gamma 2L$	$51.4 \pm 17.2$	$1.25 \pm 0.08$
$\alpha 1\beta 2(Y443W)\gamma 2L$	$77.1 \pm 44.1$	$0.81 \pm 0.15$
$\alpha 1\beta 2(Y447A)\gamma 2L$	$28.8 \pm 18.0$	$1.24 \pm 0.15$

Overall, the mutations shifted the GABA concentrationresponse relationships to higher transmitter concentrations. The largest effects were seen in receptors containing the  $\alpha 1(Q241L)$  mutation. Mutations to site III in the  $\beta$  subunit had relatively minor effects on receptor activation by GABA. The observations are in agreement with previous studies where these mutations have been used (Hosie et al., 2006; Akk et al., 2008; Bracamontes et al., 2011; Chen et al., 2019).

Analysis of Receptor Activation by GABA +  $3\alpha5\alpha P$ . Receptor potentiation by  $3\alpha5\alpha P$  was tested by coapplying  $0.01 - 10 \ \mu M \ 3\alpha5\alpha P$  with a low concentration of GABA and measuring steroid-induced enhancement of the peak response. The concentration of GABA was selected to generate a response that was 2%-10% of the peak response to 1 mM GABA + 50  $\mu$ M propofol (P<sub>A</sub> = 0.02 - 0.1). Sample current traces and the steroid concentration-response curves are given in Fig. 2. Table 2 gives the low- and highconcentration asymptotes (P<sub>A,low</sub> and P<sub>A,high</sub>) and the steroid EC<sub>50</sub>s in the wild-type and mutant receptors. The mutations had generally small effects on the steroid EC<sub>50</sub>.

We used a resting-active cyclic model to quantitatively analyze the  $3\alpha5\alpha P$  concentration-response relationships (Forman, 2012; Steinbach and Akk, 2019). The model is based on the classic Monod-Wyman-Changeux model, originally used to describe enzyme function and modulation (Monod et al., 1965). The analysis provides estimates of the affinity of the steroid to the resting receptor ( $K_{R,3z5zP}$ ) and its gating efficacy ( $c_{3z5zP}$ ).

The results are summarized in Table 2. The effects of mutations on calculated affinity of the resting receptor to  $3\alpha5\alpha P$  were generally unremarkable. The  $\alpha1(Q241L)$  and  $\alpha1(V226W)$  mutations to sites I (the intersubunit binding site) and II (intra- $\alpha$  site), respectively, reduced  $3\alpha5\alpha P$  gating efficacy. None of the tested mutations to site III affected efficacy of the steroid.

The reduced but still considerable  $(2.3 \pm 0.6$ -fold at 1  $\mu$ M; n = 6 cells; 1 = no effect) potentiating effect of  $3\alpha 5\alpha P$  in the  $\alpha 1(Q241L)\beta 2\gamma 2L$  receptor was surprising given that some previous studies had indicated that the mutation essentially abolishes modulation by classic potentiating steroids (Hosie et al., 2006; Akk et al., 2008; Bracamontes et al., 2011), but see (Bracamontes and Steinbach, 2009). To attempt to exclude any time-dependent effects associated with long recordings and/or errors rising from comparing responses to drug applications separated by several minute-long washes, we tested the effect of 3a5aP on GABA-elicited steady-state current in a continuous recording where no washout between GABA and GABA + steroid was employed. A sample recording showing the potentiating effect of  $3\alpha 5\alpha P$  in this experimental protocol is given in Fig. 3A. In six cells,  $1 \mu M 3\alpha 5\alpha P$ potentiated the steady-state current elicited by 50  $\mu$ M GABA  $(P_{A,peak} = 0.09 \pm 0.04)$  by 1.4  $\pm$  0.3-fold.

During the initial, long GABA application, the receptors equilibrate between resting, active, and desensitized states. The extent of observed potentiation in this experimental protocol is therefore not directly comparable to potentiation observed when peak currents are compared (Fig. 2). Even so, the findings support the notion that the  $\alpha 1(Q241L)\beta 2\gamma 2L$  receptor retains considerable sensitivity to the steroid  $3\alpha 5\alpha P$ .

For further confirmation of sensitivity to potentiating steroids, we tested the ability of the synthetic steroid alfaxalone to potentiate the  $\alpha 1(Q241L)\beta 2\gamma 2L$  receptor. In five cells, 3  $\mu$ M alfaxalone potentiated the peak response to 20  $\mu$ M GABA ( $P_{A,peak} = 0.03 \pm 0.01$ ) by 2.1  $\pm$  0.3-fold. A sample trace is given in Fig. 3B. For comparison, in the wild-type  $\alpha 1\beta 2\gamma 2L$  receptor, 3  $\mu$ M alfaxalone potentiates the response to low GABA by 13-fold (Shin et al., 2017).

**Energetic Contributions of the Steroid Binding Sites.** The parameter expressing agonist efficacy (c) and the postulated number of binding sites (N) for the agonist under consideration are connected through the empirical maximal probability of being in the active state as:

$$P_{A, \text{peak}, \text{max}} = \frac{1}{1 + L^*(c_{\text{effect}})^{N_{\text{effect}}})}$$
(5)

where L\* is determined from the baseline activity as described for eq. 1, and  $c_{\text{effect}}$  and  $N_{\text{effect}}$  are the effective values from the total number of sites available. In the present case, in which  $3\alpha 5\alpha P$  binds to distinct sites, this is:

$$\mathbf{c}_{effect}^{N_{effect}} = \mathbf{c}_{I}^{N_{I}} \mathbf{c}_{II}^{N_{II}} \mathbf{c}_{III}^{N_{III}} = (\mathbf{c}_{I} \mathbf{c}_{II} \mathbf{c}_{III})^{2}.$$
 (6)

Since there are two copies of each site in the receptor, the number of steroid binding sites was constrained to 2 and the product of the three values was estimated.  $\Delta G_{A,binding}$  expresses the total free energy change to stabilize the active state when all steroid binding sites are occupied:

$$\Delta G_{A, binding} = -2RT \ln(c_I c_{II} c_{III}). \tag{7}$$

The values provided in Table 2 indicate that the binding of  $3\alpha 5\alpha P$  to the  $\alpha 1\beta 2\gamma 2L$  receptor provides a total free energy change of -1.9 kcal/mol to stabilize the active state. The energetic contribution is less, i.e., the  $\Delta G$  is less negative in receptors containing a mutation to site I or site II.

Model-independent analysis provides similar information about  $\Delta\Delta G$ , but without any insights into affinity of the interaction. The energetic difference between the active and the resting states of the receptor is proportional to the ratio of the frequencies of the states:

$$\Delta G_A = -RT \ln(P_A/P_R) = -RT \ln(P_A/(1-P_A)), \qquad (8)$$

where in a two-state system such as this,  $P_R = (1 - P_A)$ . At a saturating concentration of  $3\alpha 5\alpha P$ , the total free energy change can be obtained from  $(P_{A,max}/(1 - P_{A,max}))$ . To obtain the free energy provided by binding to all the sites, the base-line free energy difference must be subtracted:

$$\begin{split} \Delta \Delta G_{A,\,binding} &= (-RTln(P_{A,\,max}/(1-P_{A,\,max}))) \\ &\quad - (-RTln(P_{A,\,0}/(1-P_{A,\,0}))) \end{split} \tag{9}$$

where  $P_{A,0}$  is the probability of being active in the absence of the steroid (but in the same baseline condition, for example the presence of a low concentration of GABA). As indicated by the values in Table 2, there is good agreement between the two estimates ( $R^2 = 0.97$ ).

The effects of the mutations on  $\Delta\Delta G_{A,\text{binding}}$  suggest that binding to site I contributes the greater portion of the stabilization energy provided by  $3\alpha 5\alpha P$  (about -1.3 kcal/mol out of -2.0 kcal/mol), whereas occupation of site II contributes about -0.5 kcal/mol, and occupation of site III does not make a meaningful contribution.

Additivity of the Functional Effects of Mutations to Individual Steroid Binding Sites. We employed double mutant cycle analysis to determine pairwise additivity of the

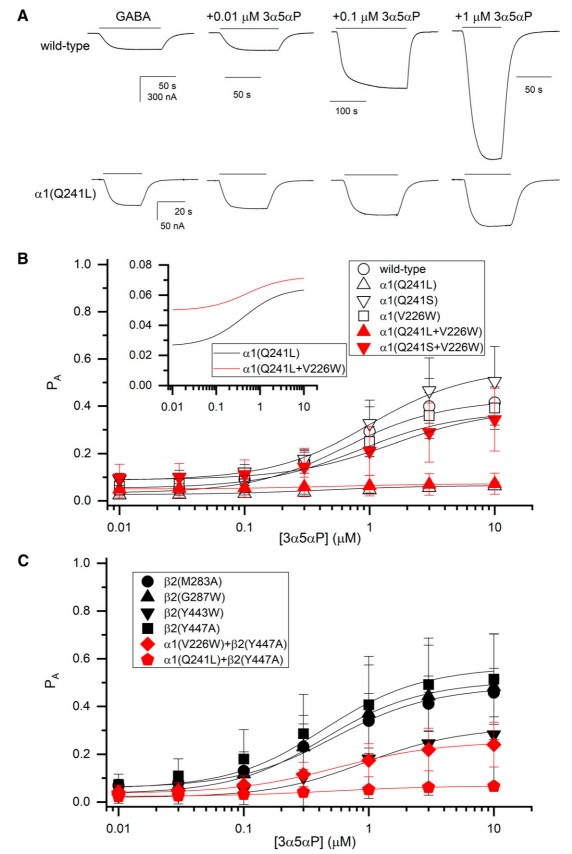


Fig. 2. Modulation of GABA-activated wild-type and mutant  $\alpha 1\beta 2\gamma 2L$  receptors by  $3\alpha 5\alpha P$ . The panels show sample current traces (A) and steroid concentration-response curves for wild-type  $\alpha 1\beta 2\gamma 2L$  and mutant receptors with mutations to steroid sites I and II (B) or site III (C). In (B) and (C), the data points show mean  $\pm$  SD from 5–7 cells. The curves are calculated using the mean fitted parameters provided in Table 2. The inset in (B) gives the concentration-response curves for receptors containing the  $\alpha 1(Q241L)$  and  $\alpha 1(Q241L+V226W)$  mutations at higher resolution.

 TABLE 2
 Potentiation of \$21\\mathcal{L}^2\2L\$ wild-type and mutant receptors by \$352P

eq. 9. The data were also analyzed to provide estimated affinity of  $3.25_{\text{SLP}}$  for the resting receptor ( $\mathrm{K}_{\mathrm{R},3.5,2.P}$ ) and the ratio of the affinities for the active and resting states ( $_{3.35,2.P}$ ) by fitting the concentration-response curves to eq. 1. The values of  $\Delta G$  were calculated from the values of  $c_{3.35,2.P}$  as N<sub>3.55,2.P</sub> as N<sub>3.55,2.P</sub> states ( $a_{3.55,2.P}$ ) by fitting the of changes in  $c_{3.55,2.P}$  was determined using ANOVA with Dunnett's correction (\*P < 0.05; \*\*\*P < 0.001). potentiation curves (mean  $\pm$  S.D. from 5–7 cells per receptor) and the calculated  $\Delta\Delta G$  from the ratios  $P_A/P_R$  using The table gives the low- and high-concentration asymptotes and EC<sub>50</sub>s of the

Receptor	Sites Mutated	${ m P}_{ m A, low}$	$EC_{50}$	$\mathrm{P}_{\mathrm{A,high}}$	$\Delta \Delta G$	$\mathrm{K}_{\mathrm{R},3a5a\mathrm{P}}$	$c_{3\alpha 5 lpha \mathrm{P}}$	ΔG
			$M_{ m H}$		kcal / mol	$M_{ m HI}$		kcal / mol
$\alpha 1\beta 2\gamma 2L$ wild-type	None	$0.03 \pm 0.02$	$0.72 \pm 0.41$	$0.45 \pm 0.08$	$-2.01 \pm 0.29$	$0.58 \pm 0.20$	$0.21 \pm 0.05$	$-1.85 \pm 0.26$
$\alpha 1(Q241L)\beta 2\gamma 2L$	$\beta$ - $\alpha$	$0.02 \pm 0.004$	$0.58 \pm 0.38$	$0.07 \pm 0.02$	$-0.69 \pm 0.16$	$0.35 \pm 0.13$	$0.62 \pm 0.06^{***}$	$-0.57 \pm 0.12$
$lpha 1(0241S)eta 2\gamma 2L$	$\beta$ - $\alpha$	$0.10 \pm 0.03$	$0.89 \pm 0.15$	$0.52 \pm 0.15$	$-1.42 \pm 0.26$	$1.16 \pm 0.40$	$0.27 \pm 0.07$	$-1.57 \pm 0.32$
$\alpha 1 (V226W) \beta 2 \gamma 2L$	$\inf_{\alpha \in \alpha} x$	$0.06 \pm 0.02$	$0.91 \pm 0.32$	$0.41 \pm 0.07$	$-1.46 \pm 0.15$	$0.72 \pm 0.21$	$0.33 \pm 0.03^*$	$-1.32 \pm 0.12$
$lpha 1(\mathrm{Q241L}+\mathrm{V226W})eta 2\gamma 2\mathrm{L}$	$\beta$ - $\alpha$ , intra- $\alpha$	$0.05 \pm 0.04$	$0.68 \pm 0.45$	$0.08 \pm 0.05$	$-0.32 \pm 0.09$	$0.60 \pm 0.40$	$0.76 \pm 0.11^{***}$	$-0.33 \pm 0.18$
$\alpha 1(Q241S+V226W)\beta 2\gamma 2L$	$\beta$ - $\alpha$ , intra- $\alpha$	$0.08 \pm 0.06$	$1.56 \pm 0.82$	$0.34 \pm 0.16$	$-1.15 \pm 0.12$	$1.45 \pm 0.43$	$0.39 \pm 0.04^{*}$	$-1.12 \pm 0.12$
$\alpha 1\beta 2(M283A)\gamma 2L$	$intra-\beta$	$0.06 \pm 0.03$	$0.46 \pm 0.25$	$0.43 \pm 0.13$	$-1.48 \pm 0.16$	$0.53 \pm 0.31$	$0.25 \pm 0.03$	$-1.62 \pm 0.14$
$\alpha 1\beta 2(G287W)y 2L$	$\inf_{\beta \in \mathcal{B}} \beta$	$0.04 \pm 0.04$	$0.45 \pm 0.13$	$0.46 \pm 0.18$	$-2.02 \pm 0.26$	$0.44 \pm 0.14$	$0.18 \pm 0.02$	$-2.04 \pm 0.11$
$lpha 1eta 2(\mathbf{Y}443\mathbf{W})\gamma \mathbf{2L}$	$intra-\beta$	$0.02 \pm 0.03$	$0.89 \pm 0.32$	$0.29 \pm 0.25$	$-1.98 \pm 0.39$	$0.71 \pm 0.26$	$0.20 \pm 0.02$	$-1.90 \pm 0.14$
$lpha 1eta 2(\mathbf{Y}\mathbf{447A})_{\gamma}\mathbf{2L}$	$intra-\beta$	$0.06 \pm 0.04$	$0.53 \pm 0.46$	$0.54 \pm 0.19$	$-1.86 \pm 0.28$	$0.50 \pm 0.36$	$0.21 \pm 0.07$	$-1.89 \pm 0.43$
$\alpha 1(Q241L)\beta 2(Y447A)\gamma 2L$	$\beta$ - $\alpha$ , intra- $\beta$	$0.02 \pm 0.01$	$0.49 \pm 0.14$	$0.07 \pm 0.01$	$-0.80 \pm 0.29$	$0.39 \pm 0.11$	$0.57 \pm 0.10^{***}$	$-0.68 \pm 0.23$
$\alpha 1 (V226W) \beta 2 (Y447A) \gamma 2L$	intra- $\alpha$ , intra- $\beta$	$0.04 \pm 0.03$	$0.65 \pm 0.28$	$0.24 \pm 0.07$	$-1.20 \pm 0.24$	$0.44 \pm 0.15$	$0.33 \pm 0.06^{*}$	$-1.32 \pm 0.19$

Potentiation of the  $\alpha 1\beta 2\gamma 2L$  GABA<sub>A</sub>R by Allopregnanolone 25

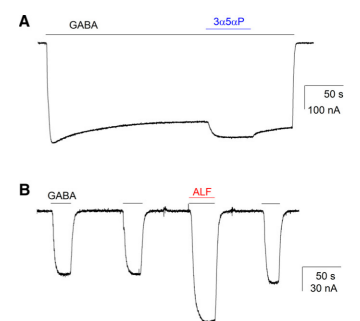


Fig. 3. Potentiation of the  $\alpha 1(Q241L)\beta 2\gamma 2L$  receptor by  $3\alpha 5\alpha P$  and alfaxalone. (A) shows the effect of 1  $\mu$ M  $3\alpha 5\alpha P$  on steady-state current elicited by 50  $\mu$ M GABA. The peak P<sub>A</sub> of the response to GABA was 0.063. (B) shows the effect of 3  $\mu$ M alfaxalone (ALF) on peak responses to 20  $\mu$ M GABA. The mean peak P<sub>A</sub> of the responses to GABA before exposure to the steroid was 0.037.

effects of mutations to the individual steroid binding sites. In this approach, the free energy change in the double mutant is compared with the sum of changes in free energy in receptors containing a single mutation (Hidalgo and MacKinnon, 1995; Horovitz, 1996; Akk, 2001). A large difference in free energy change (typically > 0.5 kcal/mol) is interpreted as the mutated sites being directly or indirectly coupled (LiCata and Ackers, 1995; Schreiber and Fersht, 1995; Horovitz, 1996; Gleitsman et al., 2009).

Figure 4 shows pairwise mutant cycles for sites I and II, I and III, and II and III. For each mutant pair, we calculated the coupling energy,  $\Delta\Delta G$ , as  $\Delta G(\text{wild-type}) + \Delta G(\text{double}$ mutant) –  $\Delta G$  (mutant #1) –  $\Delta G(\text{mutant #2})$  using the values given in Table 2. The calculated  $\Delta\Delta Gs$  with 95% confidence intervals were 0.29 [-0.004 to 0.58] kcal/mol for  $\alpha 1(\text{Q241L} + \text{V226W})$ , 0.09 [-0.28 to 0.46] kcal/mol for  $\alpha 1(\text{Q241L} + \text{V226W})$ , 0.07 [-0.41 to 0.55] kcal/mol for  $\alpha 1(\text{Q241L}) + \beta 2(\text{Y447A})$ , and –0.04 [-0.51 to 0.43] kcal/mol for  $\alpha 1(\text{V226W}) + \beta 2(\text{Y447A})$ . Inclusion of 0 value in the confidence interval was interpreted to indicate a lack of significant coupling energy, i.e., additivity of the effects of mutations. We caution, however, that the relatively wide 95% confidence intervals, at least in part because of experimental imprecision, make it harder to demonstrate small deviation from additivity.

To estimate sensitivity of our approach, we determined the difference in  $\Delta G$  between a hypothetical double mutant and sum of two single mutations that would result in no overlap between their 95% confidence intervals. The mean S.E.M. of  $\Delta G$  in all tested receptors was 0.09, and the propagated S.E.M. for combined two mutations is 0.13. The 95% confidence interval can then be calculated for the double mutant as mean  $\pm 1.96 \times 0.09$  and that for the sum of two single mutants as mean  $\pm 1.96 \times 0.13$ . This calculation gives 0.43 kcal/mol as the minimal  $\Delta\Delta G$  between the sum of effects in

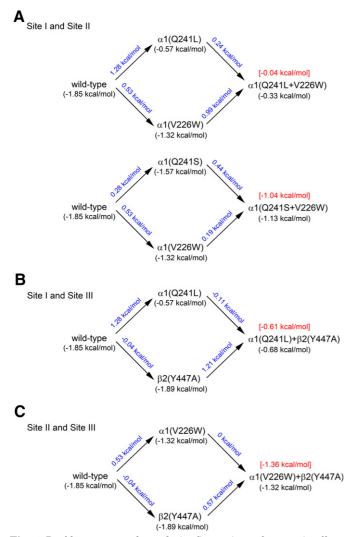


Fig. 4. Double mutant cycle analysis. Comparison of energetic effects of mutations introduced alone or in combination to sites I and II (A), sites I and III (B), and sites II and III (C). The  $\Delta G$  in parentheses indicate the stabilization energy provided by  $3\alpha 5\alpha P$  in each receptor.  $\Delta\Delta G$  values showing the effects of mutations are given in blue. The  $\Delta G$  in brackets (red) for the double mutants gives the expected stabilization energy, calculated based on ideal additivity of the effects of single mutations. The closeness of the measured and calculated  $\Delta G$  values for the double mutants indicates that the mutations act independently and additively.

two single mutants and the effect in double mutant, for us to observe a statistically significant (P < 0.05) effect.

Mutations to Sites I and II Do Not Cause Reciprocal Changes in Photolabeling. Amino acid substitutions leading to changes in local protein structure have been shown to modify the orientation in which a steroid binds in its binding site, which is the likely direct cause of reduced function (Sugasawa et al., 2019). Here, we considered whether mutations in sites I and II differently modify steroid binding in the proximal versus distal site. We reasoned that an independent effect of a mutation in site I may be expected to modify photolabeling in site I but not in site II, and vice versa. Indeed, it has been shown that in the ELIC- $\alpha$ 1 chimeric homomeric receptor, the Q242L or Q242W mutations (homologous to Q241L and Q241W in the rat  $\alpha$ 1 subunit) in site I reorient the steroid analog and change the photolabeled residue from Y309 to F298 in site I, but do not modify the labeled residue (N408 in the fourth membrane-spanning domain) or its photolabeling efficiency (1.7%–1.9%) in site II (Sugasawa et al., 2019). Since a saturating concentration of KK200 (100  $\mu M$ ) was used, the efficiency of labeling does not necessarily depend on the equilibrium occupancy (that is, the affinity of the photolabel).

To confirm the reciprocal relationship, i.e., that amino acid substitution in site II does not affect photolabeling in site I, we compared photolabeling of wild-type and V227W mutant (homologous to V226W in rat  $\alpha$ 1) ELIC- $\alpha$ 1 receptors with the steroid analog KK200. In the wild-type receptor, KK200 labeled N408 in site II and Y309 in site I. In the V227W mutant, KK200 labeling in site II was shifted to F400 at ~2  $\alpha$ -helical turns toward the cytosol (Supplemental Fig. 1), and the labeling efficiency in site II was significantly increased. Neither the residue labeled in site I (Y309) nor the labeling efficiency were affected (Table 3).

Relating Energetic Additivity to Changes in Concentration-Response Measurements. Energetic additivity of mutations can be demonstrated by approaches such as mutant cycle analysis. However, we sought to gain insight into how energetic additivity manifests directly in some commonly measured activation parameters, for example, simple additive or multiplicative relationships among maximal open probabilities for single or multiple mutations. To do this, we simulated direct activation and potentiation concentrationresponse curves for cases mimicking selective modification of one or more binding sites.

In the first case, we modeled activation of a receptor with two binding sites for agonist X. Both binding sites had a  $K_{R,X}$ (equilibrium dissociation constant for X in the resting receptor) of 10 units, and a *c* (ratio of the equilibrium dissociation constant for X in the active receptor to  $K_{R,X}$ ) of 0.01. The value of L was held at 1000 (background  $P_{\rm A}$   ${\sim}0.001).$  From the relationship  $P_{A,peak,max} = 1/(1 + Lc^N) = 1/(1 + Lc_1c_2)$ , we calculate that the maximal peak open probability in this receptor (designated wild-type) is 0.91. We next postulated that introduction of a loss-of-function mutation to a single binding site increases the value of c of that site to 0.9. With either  $c_1$  or  $c_2$  thus modified, the  $P_{A,peak,max}$  in the receptor containing a mutation to one of the binding sites in presence of saturating X is 0.10 (a reduction of  $\sim$ 9-fold). Lastly, we combined mutations to both sites, i.e.,  $c_1 = c_2 = 0.9$ . The  $P_{A,peak,max}$  is 0.0012 (a reduction of  ${\sim}740\mbox{-fold}).$  Thus, the observed effect in the double mutant is much larger, in that the effects are neither additive or multiplicative, than what might be intuitively expected by independent actions of the single mutations. The simulated concentration-response curves are shown in Fig. 5A.

The extent of mismatch depends on the starting values of c. For an agonist-receptor pair with  $c_1 = c_2 = 0.1$ , the  $P_{A,peak,-max}$  is 0.091. With a single mutation increasing c to 0.9 in that site, the  $P_{A,peak,max}$  is reduced to 0.011 (an ~8-fold reduction), and with both sites mutated ( $c_1 = c_2 = 0.9$ )  $P_{A,peak,max}$  is 0.0012 (an ~74-fold reduction). The predictions are illustrated in Fig. 5B.

The dependence on the value of c is illustrated in Fig. 5C, which plots the fold-reduction in the value of  $P_{A,peak,max}$  as a function of the initial value for c for both sites. As can be seen, the effect of the double mutation approaches the effect

## TABLE 3

#### Photolabeling efficiency

The table provides photolabeling efficiency in % (total incorporation) of 100  $\mu$ M KK200 in intact protein and middle-down MS analyses for ELIC- $\alpha$ 1 GABA<sub>A</sub> receptor (n = 2, biologic replicates). The labeling efficiency in TM3 corresponds to labeling at site I and the labeling efficiency in TM4 corresponds to labeling at site II. Photolabeled residues are shown in parentheses. The data for wild-type and Q242L were reported previously (Sugasawa et al., 2019). The Q242 position in ELIC- $\alpha$ 1 is homologous to Q241, and V227 is homologous to V226 in the rat  $\alpha$ 1 subunit.

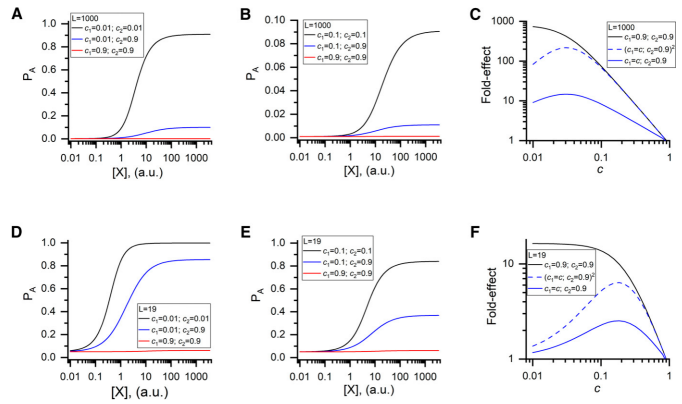
		Middle-Down MS	
ELIC- $\alpha 1$ GABA <sub>A</sub> R	Intact Protein MS	TM3 - Site I	TM4 - Site II
Wild-type	11.3, 10.7	8.0, 7.6 (Y309)	1.8, 1.6 (N408)
Q242L	11.8, 10.2	6.1, 5.5 (F298)	1.9, 1.5 (N408)
V227W	19.9, 15.4	8.5, 5.3 (Y309)	6.7, 4.2 (F400)

squared of a single mutation as the initial value of c approaches 1 (that is, as agonist X becomes a weaker agonist).

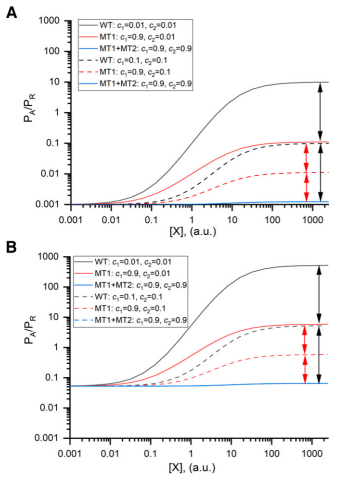
In the second case, we postulated a reduced value of L (L = 19) that generates a background  $P_A$  of 0.05, and examined the effect of X (Fig. 5, D–F). This situation models the potentiation by X of a response to a background agonist that produces a response of 5% of maximal receptor activation. Alternatively, background activity may be elicited by a gain-of-function mutation for which the direct-activating effect of X is measured. The binding of X to two sites, each with  $K_{R,X}$  of 10 units and  $c_X$  of 0.01, increased the  $P_A$  to 0.9981 (~20-fold potentiation of background activity). Introduction of a loss-of-function mutation to one binding site, increasing *c* of

that site to 0.9, reduced  $P_A$  to 0.854 (~17-fold potentiation of background activity; ~1.2-fold reduction from wild-type). When both sites for X are mutated ( $c_1 = c_2 = 0.9$ ), the predicted  $P_A$  is 0.06 (~1.2-fold potentiation; ~15-fold reduction). So, in this model, mutation of one of two independent sites with identical properties has a relatively small effect, whereas modification of both sites virtually eliminates the ability of X to potentiate the receptor.

Again, the apparent mismatch depends on the beginning values of c. With  $c_1 = c_2 = 0.1$ , the application of X increases the  $P_A$  of the response to 0.84 (17-fold potentiation). With  $c_1 = 0.1$  and  $c_2 = 0.9$ , X increases the  $P_A$  to 0.37 (7.4-fold potentiation), and with  $c_1 = c_2 = 0.9$ , to 0.06 (1.2-fold potentiation).



**Fig. 5.** Summary of effects of modifications to one or both binding sites on receptor activation. The graphs show modeled peak open probability  $(P_A)$  versus agonist (X) concentration. Modeling was done using  $K_{R,X}$  of 10 arbitrary units (a.u.) and  $N_X$  of 2. The graphs compare the effects of loss-of-function modifications to one or both agonist binding sites. (A–C) describe a model with L of 1000 mimicking direct activation by X, and (D–F) describe a model with L of 19 mimicking X-mediated potentiation of receptor activity elicited by another agonist. (A, B, D, and E) show the effects of changing the value of c in one (blue lines) or both binding sites (red line) from 0.01 to 0.9. The precise maximal  $P_A$  values for each condition are given in text. (C and F) describe the reduction in maximal  $P_A$  (fold-effect) in receptors containing one (blue solid line) or both (black line) modified binding sites. The value of modified c was held at 0.9. The value of unmodified, i.e., wild-type c is given by the abscissa. The blue dashed line gives a squared fold-effect observed when one binding site is modified.



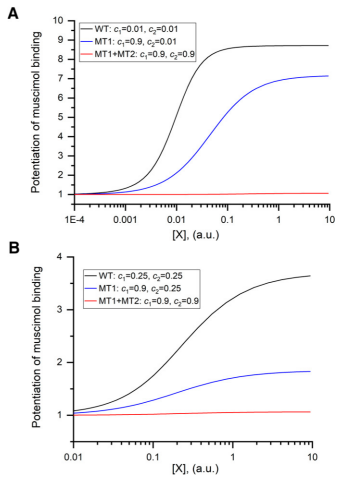
**Fig. 6.** Summary of effects of modifications to one or both binding sites on receptor  $P_A/P_R$ . The graphs show calculated values of  $P_A/P_R$  for the same conditions as were used in Figure 5. (A) shows a logarithmic plot for values when L = 1000, with values for  $c_1$  and  $c_2$  indicated. The low concentration asymptote is 1/L (0.001) for all the curves, whereas the high-concentration asymptote is  $1/(Lc_1c_2)$ . The arrows show the multiplicative relationship between the maximal values (90-fold when the nonmutated c = 0.01 and 9-fold when the nonmutated c = 0.1). (B) shows similar calculations for L=19. Note that the curves in (B) are shifted upwards by the ratio 1000/19 but are otherwise identical. MT, mutant; WT, wild-type.

Comparison of the curves in Fig. 5C and 5F indicates that the extent of the discrepancy depends on both the receptor baseline opening equilibrium and the affinity ratio for the unmutated receptor.F

An alternative experimental parameter is the ratio  $P_A/P_R$ as introduced earlier. Figure 6 shows simulations of this ratio for the same conditions used in the case of  $P_{A,peak,max}$ . Note that the curves for the two values of L are shifted vertically by the multiplicative factor (1/L). The relative maximal values for no, one, and two mutations is  $[1/(c_{wt}c_{mut})]$ , that is 90fold when  $c_{WT} = 0.01$  or 9-fold when  $c_{WT} = 0.1$ . Accordingly, for this experimental parameter, energetic additivity results in multiplicative changes in effect that are clearly seen in logarithmic plots.

Relating Energetic Additivity to Changes in the Binding of Orthosteric Agonist. Changes in equilibrium binding of a radiolabeled orthosteric agonist of the GABA<sub>A</sub> receptor, such as [<sup>3</sup>H]muscimol, in the presence of allosteric modulators have provided insight into the mechanisms and sites of action of allosteric drugs (Harrison and Simmonds, 1984; Peters et al., 1988; Sugasawa et al., 2020). Here, we modeled the effect of allosteric agonist X on binding of muscimol, using a three-state resting-active-desensitized model (Akk et al., 2020). The concentration of muscimol was selected to generate equilibrium occupancy of 10% in the absence of X. We postulated two classes of binding sites for X, with one copy of each in the receptor. The effect of a loss-of-function mutation introduced to one or both binding sites for X was then simulated by changing the value of c of one or both sites from 0.01 to 0.9.

The results of simulations are shown in Fig. 7. Exposure to allosteric agonist X is predicted to enhance muscimol binding because it increases the prevalence of the high-affinity active and desensitized states, and so increases receptor occupancy in the presence of a fixed concentration of orthosteric agonist. The maximal potentiating effect in the receptor with c of both binding sites of 0.01 (a wild-type receptor) is 8.7-fold. In



**Fig. 7.** Summary of effects of modifications to one or both binding sites for allosteric agonist on binding of orthosteric agonist. The simulations show potentiation of muscimol binding by allosteric agonist X. The simulations were based on  $\alpha 1\beta 2\gamma 2L$  GABA<sub>A</sub> receptor with L = 8000 and Q = 0.29 (Shin et al., 2017; Germann et al., 2019). The properties (K<sub>R,muscimol</sub> = 0.5 µM, c<sub>muscimol</sub> = 0.01, N<sub>muscimol</sub> = 2 binding sites) and concentration (0.039 µM) of muscimol were selected so as to generate equilibrium occupancy of 10% of binding sites. The allosteric agonist X was assigned K<sub>R,X</sub> = 0.5 units and N<sub>X</sub> = 2 binding sites, with  $c_1 = c_2 =$ 0.01 (A) or  $c_1 = c_2 = 0.25$  (B). Introduction of a loss-of-function mutation to one or both binding sites was simulated by changing the value of *c* in that binding site to 0.9.

the receptor with one site for X mutated ( $c_1 = 0.9, c_2 = 0.01$ ), X enhances muscimol binding by 7.2-fold, and with both sites mutated, ( $c_1 = c_2 = 0.9$ ) by 1.1-fold (Fig. 7A). Thus, introduction of a single mutation leads to a smaller effect than what might be reasonably expected assuming additive effects. The apparent discrepancy depends on the value of c. With  $c_X$  of 0.25 (i.e., mimicking weaker efficacy of X), muscimol binding is enhanced by 3.7-fold in the wild-type receptor and by 1.8-fold in the receptor containing one mutated site for X (Fig. 7B).

# Discussion

We had two goals in this study. The first was to determine whether the previously identified binding sites for potentiating steroids act independently. An alternative hypothesis of allosteric interaction between sites I ( $\beta$ - $\alpha$  interface) and II  $(intra-\alpha)$  was proposed by Chen et al., 2019, who had observed that mutations to either site reduced potentiation by  $3\alpha 5\alpha P$  more than what appeared intuitively additive. Here, we conclude that the binding sites for potentiating steroids contribute independently and additively to channel activation. Using a more extensive set of mutated residues, we confirm a previous finding (Chen et al., 2019) that the intra- $\beta$ subunit binding site does not contribute to receptor potentiation by  $3\alpha 5\alpha P$ . Our second goal was to test whether energetic additivity between independent binding sites predicts qualitatively interpretable changes in concentration-response curves. From simulations made using the Monod-Wyman-Changeux activation model, we show that additive energetic changes at independently acting binding sites can lead to nonlinear summation of the effects. Many commonly used experimental values, such as the probability of being open or the  $EC_{50}$  for an effect, do not depend in a simple fashion on the energetics of steroid-receptor interaction. Further, the kinetic parameters depend exponentially on the free energies involved. This makes it difficult to intuitively deduce additivity when the plotted experimental values depend nonlinearly on free energy change.

We used a mutational approach to determine independence of effects (Horovitz, 1996). The three previously identified binding sites for potentiating steroids were mutated individually or in combination, and the effects of mutations on channel gating by  $3\alpha 5\alpha P$  quantified. We reasoned that energetic additivity of the effects of mutations in individual binding sites is an indication of independently acting sites, whereas, in the case of a specific allosteric link between sites, a mutation that alters the structure of site I and/or its interaction with the steroid may be expected to modify steroid interaction with site II. The electrophysiological data indicate that the effects of mutations to sites I ( $\beta$ - $\alpha$  interface), II (intra- $\alpha$ ), and III (intra- $\beta$ ) are energetically additive.

Both additivity and nonadditivity between allosteric binding sites in the GABA<sub>A</sub> receptor have been reported previously. By individually mutating the two etomidate binding sites in the  $\beta$ - $\alpha$  interface in the concatemeric  $\alpha 1\beta 2\gamma 2$  receptor, Guitchounts et al., 2012 reported indistinguishable and additive effects of the  $\alpha 1(M236W)$  mutation on potentiation by etomidate. Similarly, the  $\beta 2(Y143W)$  and  $\beta 2(M286W)$  mutations to the two individual  $\beta$  subunits were shown to weaken receptor activation by propofol approximately equally and additively (Shin et al., 2018). In contrast, Szabo et al., 2019 showed nonadditive effects of TM2-15' and TM3-36' mutations at the  $\alpha$ - $\beta$ ,  $\beta$ - $\alpha$ , and  $\gamma$ - $\beta$  interfaces on anesthetic modulation of GABA currents. We do not have an explanation for this difference in observations.

The independent, additive effects of  $3\alpha 5\alpha P$  binding to sites I and II are supported by photolabeling data, which indicate that mutations to sites I and II modify photolabeling in the proximal site but not in the distal site. We previously showed that the  $\alpha 1(Q242L)$  mutation in the ELIC- $\alpha 1$  chimeric receptor modifies the residue photolabeled by a neurosteroid analog in site I, but has no effect on either photolabeling efficiency or the labeled residue in site II (Sugasawa et al., 2019). We now show that the V227W mutation in site II changes the photolabeled residue in site II, but does not affect the photolabeling efficiency or location of the labeled residue in site I. The V227W mutation increases the efficiency of photolabeling in site II (Table 3). Although this increased labeling efficiency could result from either an increase in ligand affinity for mutated site II or an increase in the photochemical efficiency of labeling, it is inconsistent with decreased steroid occupancy of the mutated site. Collectively, the photolabeling data indicate that mutations in site I and II produce their effects by changing ligand orientation in the site, thereby reducing steroid efficacy rather than by decreasing steroid occupancy. This is consistent with electrophysiological data showing that mutations in site I and II substantially reduce  $3\alpha 5\alpha P$  efficacy while producing only minor changes in steroid potency (Table 2).

Steroid-mediated potentiation was observed in the  $\alpha 1(Q241L)\beta 2\gamma 2L$  receptor. Some previous work, including our own, has reported complete elimination of potentiation of GABA-elicited currents by potentiating steroids after this mutation (Hosie et al., 2006; Akk et al., 2008; Bracamontes et al., 2011). Other prior work has shown small but measurable and statistically significant potentiation (Bracamontes and Steinbach, 2009). The ability of a steroid to potentiate the  $\alpha 1(Q241L)\beta 2\gamma 2L$  receptor is critical to the conclusions of the present study, as it would not be feasible to measure the additivity of effects of mutations if modification of one site fully eliminated the response. Potentiation was observed in two experimental protocols, including one in which the effect of the steroid was measured on steady-state response to low GABA, i.e., without a preceding wash (Fig. 3). Qualitatively, this experiment most strongly demonstrates that the  $\alpha 1(Q241L)$ -containing receptor retains sensitivity to  $3\alpha 5\alpha P$ .

To gain insight into relative contributions made by sites I and II to channel potentiation by  $3\alpha 5\alpha P$ , we compared the effects of mutations made to each site. In the wild-type receptor,  $3\alpha 5\alpha P$  provides -1.9 to -2.0 kcal/mol of free energy change. In  $\alpha 1(Q241L)\beta 2\gamma 2L$ ,  $3\alpha 5\alpha P$  provides -0.6 to -0.7kcal/mol. Therefore, if the  $\alpha 1(Q241L)$  mutation fully disables steroid actions through site I, steroid action at site II contributes only -0.7 kcal/mol. This value is comparable to the loss of contribution made by the steroid in the receptor with the  $\alpha 1(V226W)$  mutation [ $\Delta \Delta G_{Site II} = \Delta G_{wild-type} - \Delta G_{\alpha 1(V226W)}$ = -0.5 to -0.6 kcal/mol]. If we assume that the  $\alpha 1(V226W)$ mutation fully abolishes steroid effect through site II, then the remainder (-1.5 kcal/mol) can be assigned to energetic contributions through site I. This value is close to the loss of free energy change observed after the introduction of the  $\alpha 1(Q241L)$  mutation (-1.3 kcal/mol). Thus, we propose that

site I contributes more than 2-fold more free energy change than site II. We note that this estimate applies to the steroid  $3\alpha 5\alpha P$ ; other ligands to sites I and II may differ in their energetic contributions.

One goal of this study was to gain insight into how energetic additivity of combined mutations manifests in commonly measured parameters, including magnitude of effect. In other words, if mutation #1 reduces function (activation or potentiation) by a-fold, mutation #2 by b-fold, and the combination of the two mutations by c-fold, what relationship between a, b, and c indicates independence and additivity versus allosteric interaction. Intuitively, we may expect that  $a \times b = c$ . For example, if mutation #1 reduces channel open probability by 3-fold, and mutation #2 reduces channel open probability by 2-fold, then it may be intuitively expected that the current is reduced by 6-fold in the double mutant if the mutations act independently.

Modeling of current responses using the MWC-based two-state model indicated that energetic additivity predicts responses for combinations of mutations that can be markedly different from the product of effect sizes of individual mutations. The results of simulations are presented in Fig. 5, C and F, where the distance between the blue dashed and black lines indicates the apparent mismatch between the effect predicted by the combination of mutations and the effect predicted by doubling the effect of a single mutation. The lines indicate that the mismatch is most prominent at low values of c, i.e., when the agonist being studied has high efficacy. In contrast, the experimental value P<sub>A</sub>/P<sub>R</sub> results in graphic display of data that qualitatively demonstrates energetic additivity (Fig. 6). The same logic applies to actions of allosteric agonists on enhancement of [<sup>3</sup>H]muscimol binding. 3a5aP enhances the binding of the orthosteric agonist muscimol to the GABA<sub>A</sub> receptor by increasing the occupancy of high-affinity active and desensitized states (Akk et al., 2020). Mutations that reduce the ability of the steroid to bind or gate the receptor reduce the effect on muscimol binding (Sugasawa et al., 2020), whereas our simulations (Fig. 7) indicate that the sum of effects of independent mutations does not need to be algebraically additive.

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#### **Authorship Contributions**

Participated in research design: Evers, Steinbach, Akk. Conducted experiments: Germann, Pierce, Tateiwa, Sugasawa.

Contributed new reagents or analytical tools: Reichert.

Performed data analysis: Germann, Pierce, Tateiwa, Sugasawa, Evers, Steinbach, Akk.

Wrote or contributed to the writing of the manuscript: Germann, Pierce, Tateiwa, Sugasawa, Reichert, Evers, Steinbach, Akk.

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## Potentiation of the $\alpha 1\beta 2\gamma 2L$ GABA<sub>A</sub>R by Allopregnanolone **31**

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