Identifying Key Amino Acid Residues That Affect \( \alpha \)-Conotoxin AuIB Inhibition of \( \alpha \beta \delta \) Nicotinic Acetylcholine Receptors*

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**Background:** \( \alpha \)-Conotoxin AuIB interacts with \( \alpha \beta \delta \) nAChRs and GABA\(_{\beta}\) receptors, but structural determinants of these interactions are unknown.

**Results:** Using alanine scanning mutagenesis and molecular dynamics, we identified residues crucial for AuIB\( \alpha \beta \delta \) nAChR interaction.

**Conclusion:** We identified the key residues that mediate AuIB\( \alpha \beta \delta \) nAChR interaction.

**Significance:** Ability to direct \( \alpha \)-conotoxin binding to nAChRs or GABA\(_{\beta}\) receptors will improve analgesic conopeptides.

\( \alpha \)-Conotoxin AuIB is a selective \( \alpha \beta \delta \) nicotinic acetylcholine receptor (nAChR) subtype inhibitor. Its analgesic properties are believed to result from it activating GABA\(_{\beta}\) receptors and subsequently inhibiting Ca\(_{\delta}\) voltage-gated calcium channels. The structural determinants that mediate diverging AuIB activity at these targets are unknown. We performed alanine scanning mutagenesis of AuIB and \( \alpha \beta \delta \) nAChR, homology modeling, and molecular dynamics simulations to identify the structural determinants of the AuIB\( \alpha \beta \delta \) nAChR interaction. Two alanine-substituted AuIB analogues, [P6A]AuIB and [F9A]AuIB, did not inhibit the \( \alpha \beta \delta \) nAChR. NMR and CD spectroscopy studies demonstrated that [F9A]AuIB retains its native globular structure, so its activity loss is probably due to loss of specific toxin-receptor residue pairwise contacts. Compared with AuIB, the concentration-response curve for inhibition of \( \alpha \beta \delta \) by [F9A]AuIB shifted rightward more than 10-fold, and its subtype selectivity profile changed. Homology modeling and molecular dynamics simulations suggest that Phe-9 of AuIB interacts with a two-residue binding pocket on the \( \beta \) nAChR subunit. This hypothesis was confirmed by site-directed mutagenesis of the \( \beta 4\)–Trp-59 and \( \beta 4\)–Lys-61 residues of loop D, which form a putative binding pocket. AuIB analogues with Phe-9 substitutions corroborated the finding of a binding pocket on the \( \beta 4\) subunit and gave further insight into how AuIB Phe-9 interacts with the \( \beta 4\) subunit. In summary, we identified critical residues that mediate interactions between AuIB and its cognate nAChR subtype. These findings might help improve the design of analgesic conopeptides that selectively “avoid” nAChR receptors while targeting receptors involved with nociception.

Peptides isolated from the venom of cone snails belonging to the genus Conus are valuable pharmacological tools, and some are also promising drug leads (1–5). \( \alpha \)-Conotoxins are a subfamily of these peptides and consist of 12–19 amino acid residues, including four cysteines with a characteristic CC-C-C arrangement (type I cysteine framework) (6, 7). These four cysteines can yield three possible disulfide connectivities: globular (I-III, II-IV), ribbon (I-IV, II-III), and beads (I-II, III-IV). However, naturally occurring \( \alpha \)-conotoxins typically exhibit the “globular” conformation (7). The number of amino acids in each of the two loops between the framework cysteine residues is used to divide \( \alpha \)-conotoxins into subclasses. For example those with four amino acids in loop 1 and six in loop 2 are referred to as 4/6-\( \alpha \)-conotoxins.

Nicotinic acetylcholine receptors (nAChRs) are transmembrane proteins that form cationic ligand-gated channels that mediate fast excitatory cholinergic neurotransmission in the central nervous system (CNS). They also have an important regulatory role in the body, modulating the release of several neurotransmitters. The importance of nAChRs is emphasized by their involvement in various CNS disorders, including Alzheimer disease, schizophrenia, pain, nicotine addiction, and cancer (8–11).

nAChRs belong to the Cys-loop family of ligand-gated ion channels that is characterized by a pentameric composition and subunit arrangement in the receptor (12, 13). Seventeen nAChR subunits have been identified (12). Six of them are muscle nAChR subtypes (\( \alpha 1\beta 1\delta\gamma \) or \( \alpha 1\beta 1\epsilon\gamma \)) found exclusively at neuromuscular junctions, whereas the rest (\( \alpha 2\)-\( \alpha 10\), \( \beta 2\)-\( \beta 4\)) can combine in numerous homomeric (having only \( \alpha \) subunits) or heteromeric (having \( \alpha \) and \( \beta \) subunits) neuronal nAChR subunits.

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6 The abbreviations used are: nAChR, nicotinic acetylcholine (ACh) receptor; TOCSY, total correlation spectroscopy; AChBP, acetylcholine binding protein; MD, molecular dynamics; NAL, 3-(2-naphthyl)-alanine; TrpD, tryptophan of loop D.
types (8). Heteromeric receptor subtypes exhibit two distinct subunit stoichiometries of α:β ratios (2:3 or 3:2) and display different pharmacological properties (11, 14). The diversity of neuronal heteromeric nAChR subtypes in vivo increases even further when more than one α/β subunit is included within the same pentamer (e.g. α3β2β4 or α3α6β2) (11).

AuIB is a 4/6-α-conotoxin isolated from the venom of Conus aulicus and consists of 15 amino acid residues (15). In contrast with most α-conotoxins that inhibit 2–3 nAChR subtypes with similar potency, AuIB only inhibits the α3β4 nAChR subtype and with micromolar potency (15). Two other α-conotoxins, BuIA, an unusual 4/4-α-conotoxin isolated from the venom of Conus bullatus, and the recently described 4/7-α-conotoxin RgIIA from Conus regius, have been found to be considerably more potent inhibitors of the α3β4 subtype, blocking it at low nanomolar concentrations (16, 17). However, unlike AuIB, BuIA and RgIIA are not selective for the α3β4 subtype but potentially block other nAChR subtypes too. Additionally, despite α-conotoxins generally being described as competitive nAChR antagonists (18–20), AuIB is a non-competitive α3β4 antagonist (14).

α-Conotoxins with a different disulfide bond connectivity from the native form typically show losses in biological activity. However, the ribbon disulfide isomer (I-IV, II-III) of AuIB retains its nAChR inhibition with greater potency than that of the native globular (I-III, II-IV) AuIB disulfide isomer in rat parasympathetic ganglion neurons (21). Furthermore, the AuIB ribbon isomer exhibits stoichiometry-dependent blockade of oocyte-expressed α3β4 nAChRs, and unlike globular AuIB, it competitively inhibits the α3β4 nAChR (14).

Interestingly, AuIB and several other α-conotoxins (e.g. RgIA and Vc1.1) exhibit analgesic properties when tested in animal models of pain (22–24). These conotoxins inhibit various nAChR subtypes (α3β4, α9α10, and α7), but all inhibit high voltage-activated calcium channels via G protein-coupled GABAβ receptors with much more potency than their corresponding nAChR target (25–27). Therefore, GABAβ receptor-mediated suppression of N-type calcium channels (Ca2,2.2) has been suggested as the common mechanism of their analgesic action (28).

To design more effective α-conotoxin analogues that could lead to improved analgesic drugs with fewer side effects, it is important to know the structural determinants of inhibition at nAChRs and GABAβ receptors. This would help us focus on analgesic activity and remove the undesired side effects in engineered α-conotoxin analogues. Furthermore, it might help to identify previously overlooked naturally occurring analgesic conotoxins. Here we take the first step toward this goal by characterizing the residues that are critical for α-conotoxin AuIB inhibition of α3β4 nAChRs. We show that the interaction between phenylalanine (Phe) at position 9 of AuIB and β4-Tyr-59–β4-Lys-61 of the loop D on the (−) face of the β4 subunit are essential for this inhibition. A preliminary report of these results in part has been presented in abstract form (29).

**EXPERIMENTAL PROCEDURES**

**Peptide Synthesis**—All peptides were assembled on MBHA-Rink-Amide resin (Novabiochem) by solid-phase peptide synthesis using a Liberty Microwave Peptide Synthesizer with the in situ activation/N,N,N′,N′-tetramethyl-O-(1H-benzotriazol-1-yl)urani um hexafluorophosphate protocol and Fmoc (N-(9-fluorenylmethoxycarbonyl) chemistry (30). Cys-3 and Cys-15 were incorporated into the peptide chain with acetamidomethyl side chain protection during assembly to facilitate regioselective disulfide formation. Peptides were cleaved from the resin using trifluoroacetic acid (TFA) with tri-isopropylsilane and water as scavengers (9:0.5:0.5 (v/v) TFA:triisopropylsilane:water) at 22 °C for 2 h. The TFA was then removed under vacuum, and peptides were precipitated with ether, filtered, dissolved in 50% acetonitrile containing 0.05% TFA, and lyophilized. Crude peptides were purified by reversed-phase high performance liquid chromatography (RP-HPLC) on a C18 column using a gradient of 0–80% B (A, H2O, 0.05% TFA; B, 90% CH3CN, 10% H2O, 0.045% TFA) in 80 min. Analytical RP-HPLC and electrospray mass spectrometry confirmed the purity and molecular mass of synthesized peptides.

The first disulfide bond in each peptide was formed by incubating peptides in 0.1 M NH4HCO3 (pH 8.2, 0.3 mg/ml) overnight at 22 °C, then purifying them by RP-HPLC. The second disulfide bond was formed by incubating peptides with iodine in acidic conditions. Peptides were dissolved in 50% aqueous acetic acid (0.5 mg/ml). To this solution 100 μl of 1 M HCl/mg of peptide was added, then a solution of 0.1 M I2 in 50% acetic acid was slowly added until the solution became pale yellow. The reaction mixture was stirred for 12 h at 22 °C in the dark. The reaction was quenched by adding 1 M ascorbic acid until the mixture became colorless. The peptide was purified by RP-HPLC, and fractions were combined after analytical RP-HPLC confirmed purity. Electrospray mass spectrometry was used to confirm the peptide identity.

**Nuclear Magnetic Resonance (NMR) and Circular Dichroism (CD)**—NMR spectra for all AuIB analogues were recorded on samples dissolved in 90% H2O and 10% D2O at a pH of ~4. Bruker Avance 500 and 600 MHz NMR spectrometers were used to acquire spectra, including 1H, total correlation spectroscopy (TOCSY) and nuclear Overhauser effect spectroscopy (NOESY) data, as described previously (18, 31), and processed using Topspin (Bruker). All spectra were recorded at 290 K, and chemical shifts were referenced to the residual water signal at 4.85 ppm. Processed spectra were analyzed and assigned within the program Sparky (32).

For CD spectroscopy experiments, 70 μM concentrations of each peptide were dissolved in 20 mM sodium phosphate buffer at pH 7. To examine the helical propensity of each isomer, CD data were also obtained for each product after 30% tetrafluoroethene was added to the solution. Spectra were acquired on a Jasco J-810 spectropolarimeter, which was routinely calibrated using 0.6% (w/v) ammonium-D-camphor-10-sulfonate. All experiments were conducted at room temperature (21–23 °C) under a nitrogen atmosphere (15 ml/min). The experimental parameters were set to a scanning speed of 50 nm/min, response time of 1 s, sensitivity range of 100 millidegrees, and a step resolution of 1 nm. Absorbance was measured in the far UV region (185–260 nm) using a 1-mm path length quartz cuvette. Each recording was an accumulation of four scans. To eliminate any possible interference from the solvent, cuvette,
and spectropolarimeter optics, we subtracted CD spectra of the pure solvents from each sample.

Protein Sequence Alignment—The National Institute of Health’s online Constraint-based Multiple Alignment Tool, COBALT (www.ncbi.nlm.nih.gov), was used to align protein sequence, and conservative domains were taken into account.

Residues of nAChR subunits were numbered according to the sequences of the mature proteins that lacked the signal peptide at the start of their sequences.

Site-directed Mutagenesis of α3 and β4 nAChR Subunits—
Constructs in which Trp-59, Lys-61, or both β4 nAChR residues were substituted by alanine ([W59A]β4, [K61A]β4, and [W59A]+[K61A]β4, respectively) were generated using the Geneart Site-directed Mutagenesis System (Invitrogen catalog no. A13282) and the following primers: 5′-cat gac cac cag cgc gta gac tga tca cc-3′ and 5′-ggt cag tcc att cct gtg cca gcc aga tgc tgg tgg tca tg-3′ for [W59A]β4; 5′-cat gac cac cag cgc gct gcc aca gga atg gag tga tca cc-3′ and 5′-ggt cag tcc att cct gtc gca cgc tgc tgg tca tg-3′ for [K61A]β4; 5′-cat gac cac cag cgc gct gcc aca gga atg gag tga tca cc-3′ and 5′-ggt cag tcc att cct gtc gca cgc tgc tgg tca tg-3′ for [W59A]+[K61A]β4. Gln-198 of the α3 subunit was substituted to alanine (Q198A) in the same way using primers 5′-ctg tga gga gat ctc cgc aca gga atg gag tga tca cc-3′ and 5′-ggt cag tcc att cct gtc gca cgc tgc tgg tca tg-3′ for Q198Aα3. The same day using identical procedures to maximize the consistency of concentration and purity between subunits.

RNA Preparation—Plasmid DNAs encoding rat α3, α4, α9, α10, β2, and β4 nAChR subunits and human α7 nAChR subunits were obtained from Dr. J. Patrick, Baylor College of Medicine, Houston TX, Dr. J. Lindstrom University of Pennsylvania, Philadelphia PA, and OriGene Technologies Inc., Rockville MD. The plasmids were linearized with appropriate restriction enzymes, and cRNA was synthesized in vitro using a SP6 or T7 in vitro transcription kit (mMessage mMachine; Ambion, Foster City, CA). RNA for different nAChR subunits to be co-injected into the same oocytes was synthesized in parallel on the same day using identical procedures to maximize the consistency of concentration and purity between subunits.

Oocyte Preparation and Microinjection—Stage V-VI oocytes were obtained from Xenopus laevis, defolliculated with collagenase (Type I, Sigma) at 3 mg/ml, and incubated at 18°C in sterile ND96 solution (96 mM NaCl, 2 mM KCl, 1.8 mM CaCl2, 1 mM MgCl2, and 5 mM HEPES at pH 7.5) supplemented with 50 μg/liter gentamycin (Sigma). Glass pipettes for microinjection were pulled from glass capillaries (3–000–203 GX, Drummond Scientific Co.). The cRNAs were diluted in water to the appropriate concentrations (−0.1 ng/nl), and 5 ng of RNA was injected into each oocyte using a microinjector (Nanjoet, Drummond Scientific Co.). Electrophysiological recordings were carried out 2–7 days after microinjection.

Electrophysiological Recordings and Data Analysis—Two-electrode voltage clamp recordings from oocytes were carried out at room temperature using a GeneClamp 500B amplifier (Molecular Devices Corp., Sunnyvale, CA) at a holding potential of −80 mV. The voltage-recording and current-injecting electrodes were pulled from borosilicate glass (GC150T-15, Harvard Apparatus Ltd.) and had resistances of 0.3–1.5 megohms when filled with 3 M KCl. Oocytes were continuously perfused in a recording chamber with a volume of ~50 μl, with ND96 solution at 2 ml/min, applied by a gravity-fed perfusion system. nAChR-mediated currents were evoked by pipetting 100 μl of acetylcholine (ACH) into the bath when the perfusion was temporarily halted. ACh concentration was 50 μM unless specified otherwise. Oocytes were preincubated with the peptide for ~5 min, then ACh and the peptide were co-applied. Peak ACh-evoked current amplitude was recorded before and after peptide incubation using pClamp 9 software (Molecular Devices). The effects of native AuIB and its peptide analogues on ACh-evoked nAChR-mediated currents were defined as peak current amplitudes relative to the average peak current amplitude of 3–5 control ACh applications, recorded before preincubation with the peptides. Concentration-response curves for AuIB and [F9A]AuIB were fitted by unweighted nonlinear regression to the logistic equation,

\[ E_x = \frac{E_{\text{max}} X^n}{X^n + I_{C50}^{\text{act}}} \]  

where \( E_x \) is the response, \( X \) is the peptide concentration, \( E_{\text{max}} \) is the maximal response, \( n \) is the slope factor (Hill slope), and \( I_{C50}^{\text{act}} \) is the peptide concentration that gives 50% inhibition of the maximal response. All electrophysiological data were pooled (\( n = 3–6 \) for each data point) and represent the arithmetic means ± S.E. of the mean. One-way analysis of variance followed by Bonferroni’s post hoc test was used to compare current amplitudes affected by AuIB analogues with those of the native peptide. Data were statistically analyzed using SigmaPlot Version 11.0 (Systat Software Inc., San Jose, CA) or Prism5 (GraphPad Software Inc., La Jolla, CA).

Molecular Modeling and Docking Simulation—Homology models of the extracellular ligand binding domain of the rat (α3)3(β4)n nAChR bound to AuIB, [F9A]AuIB, or [F9Y]AuIB were generated using the crystallographic coordinates of Aplysia californica acetylcholine-binding protein (ACHBP) co-crystallized with the double mutant α-conotoxin PnIA[A10L,D14K] (Protein Data Bank accession code 2BR8) as a template. This template was chosen to provide a suitable 4/7 α-conotoxin-bound conformation of the receptor for subsequent molecular dynamics (MD) simulations and analyses.

AuIB and mutant peptides were modeled bound to the two α(3+)β4(−) receptor binding sites using the geometry of the PnIA mutant in the AChBP crystal structure as a template. Rat α3 and β4 sequences were obtained from the Swiss-Prot database (codes P04757 and P12392, respectively) and aligned with the template sequence using the ClustalW server. BLOSUM was used as the scoring matrix. With the multiple alignment as input, 10 AuIBα3β4, [F9A]AuIBα3β4, and [F9Y]AuIBα3β4 complex models were generated using Modeler9v6. The top-ranking models were selected and validated using PROCHECK. MD simulations were carried out on the top models of both receptor complexes.

MD Simulations—Before MD simulations, the energies of the complexes were minimized using the steepest descent algorithm and an energy gradient convergence criterion of 0.01 kcal/mol Å. Each receptor complex was placed in icosahedral simulation boxes with edge lengths of 100 × 100 × 100 Å and solvated with 34,000 TIP3P water molecules. To neutralize charge and pro-
vide a salt concentration of ~150 mM, 94 Na\(^+\) and 70 Cl\(^-\) ions were added to the solvent. All simulations were performed using GROMACS Version 4.5 (75) with the CHARMM27 forcefield (with cmap) (76, 77). All subsequent simulations were performed using a constant particle number, pressure, and temperature ensemble. Temperature was maintained at 300 K using the Nose-Hoover temperature coupling algorithm, and pressure was maintained at 1 atm using Berendsen's pressure coupling algorithm. Time steps of 2 fs were used to integrate all simulations. Solvent equilibration simulations of 100 ps lengths were performed. The non-hydrogen atoms of the receptor and peptides were positionally restrained so the solvent and ions could relax from an initially semi-crystalline structure. “Data collection” simulations of both complexes were then conducted, and all atoms of the system were free to undergo dynamics. Each complex was simulated for 100 ns. All analyses were performed on the final 20 ns of the trajectories to reduce bias from initial homology model conformations. Molecular graphics were produced using VMD Version 1.9.2 (78). All analyses were done using a combination of VMD, GROMACS analysis software suite, and in-house scripts.

**RESULTS**

**Alanine Scanning Mutagenesis Identifies Residues in the AuIB Sequence That Are Critical for Its Interaction with the α3β4 nAChR Subtype**—To find residues in the AuIB sequence that contribute most to α3β4 nAChR inhibition, we performed alanine scanning mutagenesis of the peptide. We systematically substituted each of the original residues to alanine, except for the four cysteines essential for maintaining the peptide globular structure and the native alanine at position 10 (see Fig. 1, A and B). All peptides were successfully synthesized using solid-phase peptide synthesis and a regioselective approach to form disulfide bonds to produce the globular disulfide framework. For future reference, we refer to globular AuIB as native AuIB. Each alanine mutant structure was analyzed using NMR and CD spectroscopy. The NMR spectral data for each peptide except [P6A]AuIB were successfully assigned whereby the sequential assignment of the individual spin systems determined from TOCSY spectra.

The [P6A]AuIB mutant exhibited broadened signals and multiple conformations in the TOCSY and NOESY spectra, so could not be assigned. The loss of secondary structure in the [P6A]AuIB mutant was confirmed by CD spectroscopy (Fig. 1C). Pro-6 is the only highly conserved amino acid residue in α-conotoxins apart from the cysteines and is responsible for helix initiation by inducing the \(3_\alpha\) helix turn in the backbone \(33, 34\). Mutation of Pro-6 to alanine has also been shown to disrupt the α-conotoxin Vc1.1 structure \(35\).

Secondary αH chemical shifts represent the difference between an observed αH chemical shift and that of the corresponding residue in a random coil peptide. They are strong indicators of the presence of a secondary structure \(36\). Secondary shift analysis of the chemical shift data indicated that, for native AuIB and analogues \([\text{G1A}]\text{AuIB}, [\text{S4A}]\text{AuIB}, [\text{P7A}]\text{AuIB}, [\text{F9A}]\text{AuIB}, [\text{T11A}]\text{AuIB}, [\text{P13A}]\text{AuIB}\) and \([\text{D14A}]\text{AuIB}\) residues 6–10 have negative αH secondary shifts (Fig. 1D). These negative αH secondary shifts indicate a helical region, which is consistent with the previously reported three-dimensional structure of native AuIB \(21\) and other α-conotoxin structures reported to date \(37\). Interestingly, NMR secondary shift data for [Y5A]AuIB and [N12A]AuIB were more consistent with the ribbon isomer of AuIB \(21\) despite using a regioselective disulfide bond formation strategy to form the native (globular) isomer (Fig. 1D). This was confirmed by CD spectroscopy (Fig. 1C).

The effect of point modifications in the AuIB analogues on α3β4 nAChR-mediated current inhibition was examined by two-electrode voltage clamp recordings in oocytes expressing the α3β4 nAChR subtype. The relative amount of inhibition AuIB alanine-substituted analogues produced was compared with that of non-modified peptides at a fixed concentration (3 \(\mu\)M; ~1 \(\text{IC}_{50}\) AuIB \(14\)) (Fig. 2, A and B). Only three residues in the AuIB sequence significantly reduced inhibition of relative peak ACh-evoked current amplitude when alanine was substituted for them. These were Gly-1 (0.82 \(\pm\) 0.11, \(n = 3; p < 0.05\)), Pro-6 (1.02 \(\pm\) 0.07, \(n = 3; p < 0.001\)), and Phe-9 (1.07 \(\pm\) 0.15, \(n = 3; p < 0.001\)) compared with a relative peak current amplitude of 0.51 \(\pm\) 0.07 \((n = 6)\) obtained in the presence of native AuIB (Fig. 2B). Interestingly, Gly1 replacement by Ala reduced α3β4 nAChR inhibition, although Gly1 does not belong to either loop 1 or loop 2 of AuIB, which are thought to be the primary mediators of α-conotoxin interaction with nAChRs \(6, 7\). Unlike the [G1A]AuIB mutation, which retained minor inhibitory activity, P6A and F9A analogues exhibited a complete loss of inhibitory activity. As the NMR and CD data showed that secondary structure of [P6A]AuIB is irregular (Fig. 1, C and D), we concluded that this peptide’s loss of inhibitory activity was due to disruption in the three-dimensional structure. This is not surprising, as Pro-6 is believed to be responsible for inducing the \(3_\alpha\) helix turn in the backbone of α-conotoxins \(33, 34\). In contrast, [F9A]AuIB NMR and CD spectrum data were consistent with the native AuIB structure (Fig. 1, C and D). This led us to conclude that the F9A substitution is specific and relevant to the peptide-receptor interaction rather than a general disruption of the peptide structure.

**Characterization of [F9A]AuIB Inhibition of the α3β4 nAChR Subtype**—Our next step was to probe the degree to which [F9A]AuIB inhibitory activity is impaired by modifying the Ala to Phe substitution. Because the [F9A]AuIB analog was not active at ~1 \(\text{IC}_{50}\) concentration of the native AuIB (3 \(\mu\)M) (Fig. 2A), we tested this analog at higher concentrations, up to the highest practical concentration available (30 \(\mu\)M). The [F9A]AuIB analog applied at 10 \(\mu\)M mildly inhibited ACh-evoked currents, with the amplitude reduced to 0.77 \(\pm\) 0.04 \((n = 4)\) of normalized control currents (Fig. 3, A and B). This indicates that Phe to Ala substitution at position 9 of AuIB may strongly reduce the affinity of the modified peptide for α3β4 nAChRs but does not disrupt peptide-receptor interaction altogether. However, native AuIB completely blocked the current at this concentration (0.05 \(\pm\) 0.01, \(n = 6)\) (Fig. 3, A and B). A concentration of 30 \(\mu\)M [F9A]AuIB inhibited 55% of current amplitude (0.45 \(\pm\) 0.07, \(n = 3\)). The resulting concentration-response curve demonstrates a rightward shift, indicating
interaction with \( \alpha 3\beta 4 \) nAChRs is markedly reduced (Fig. 3B).

The Selectivity Profile of [F9A]AuIB for nAChR Subtypes—Previously, most \( \alpha \)-conotoxins that act on neuronal nAChRs were shown to have only relative selectivity in the sense that they inhibit two to three nAChR subtypes with similar potency and rarely inhibit just one nAChR subtype (6, 7). To date, AuIB is the only \( \alpha \)-conotoxin shown to selectively inhibit the \( \alpha 3\beta 4 \) nAChR subtype (15). As described above, substituting Phe-9 to Ala in AuIB led to complete loss in activity at \( \alpha 3\beta 4 \). We also assessed the pharmacological profile of [F9A]AuIB at other nAChR subtypes. When tested, [F9A]AuIB (3 \( \mu \)M) exhibited only 16.2% inhibition of relative current amplitude (0.84 ± 0.1, \( n = 3 \)) at \( \alpha 3\beta 2 \) nAChRs, whereas no activity was observed at \( \alpha 4\beta 4 \) nAChRs (\( n = 5 \)) (Fig. 4A). However, we found that 3 \( \mu \)M [F9A]AuIB produced minor potentiation at the homomeric \( \alpha 7 \) nAChR subtype (1.27 ± 0.04, \( n = 3 \)) (Fig. 4A). We next tested higher concentrations of [F9A]AuIB at \( \alpha 7 \) and \( \alpha 4\beta 4 \) nAChR subtypes. Unlike the \( \alpha 3\beta 4 \) nAChR subtype, [F9A]AuIB exhibited minimal activity at these subtypes. The pharmacological profile of [F9A]AuIB at other nAChR subtypes is detailed in the supplementary material.
nAChR subtype, where higher concentrations of [F9A]AuIB significantly inhibited ACh-evoked currents (Fig. 4B, see also Fig. 3B), no significant change in relative current amplitude was observed at α3β4 nAChRs by 10 μM (0.88 ± 0.07, n = 4) or 30 μM (0.85 ± 0.1, n = 4) of [F9A]AuIB. Similarly, at α7 nAChRs neither 10 μM (1.01 ± 0.15, n = 4) nor 30 μM (0.83 ± 0.35, n = 6) [F9A]AuIB produced any potentiation or inhibition of ACh-evoked current amplitude (Fig. 4B).

**Homology Modeling and MD Simulations Suggest Loss of Interactions between AuIB and Key Receptor Residues When Ala Is Substituted for Phe-9**—Having established that position 9 Phe in AuIB is crucial for the peptide inhibiting α3β4, we used atomistic simulations of native AuIB and the [F9A]AuIB mutant bound to α3β4 to provide molecular-level explanations of why the mutation so markedly reduces inhibition. In particular, whereas experimental alanine scanning mutagenesis on AuIB identified peptide residues crucial for its efficacy, homology modeling and MD simulations identified receptor residues that might play important roles in AuIB inhibition of α3β4 nAChR. Representative homology models of AuIB-bound α3β4 are shown in Fig. 5, A and B.

To elucidate which receptor residues probably need contact with the peptide for inhibition to occur, we first calculated the average number of interatomic contacts between native AuIB and the [F9A]AuIB mutant bond to α3β4 to provide molecular-level explanations of why the mutation so markedly reduces inhibition. In particular, whereas experimental alanine scanning mutagenesis on AuIB identified peptide residues crucial for its efficacy, homology modeling and MD simulations identified receptor residues that might play important roles in AuIB inhibition of α3β4 nAChR. Representative homology models of AuIB-bound α3β4 are shown in Fig. 5, A and B.
**α-Conotoxin AuIB Interaction with α3β4 nAChRs**

![Figure 4](image)

**FIGURE 4. nAChR subtype selectivity profile of [F9A]AuIB.** A, bar graph showing relative ACh-evoked current amplitude in the presence of 3 μM [F9A]AuIB tested at different nAChR subtypes. B, comparison of effects of three different [F9A]AuIB concentrations (3, 10, and 30 μM) on ACh-evoked current amplitude at α3β4, α7, and α3β2 nAChR subtypes. Data represent means ± S.E., n = 3–6. * indicates p < 0.05 for relative change of current inhibition at various nAChR subtypes versus α3β4 nAChRs (A) and for relative change of current inhibition at 10 and 30 μM compared with 3 μM peptide concentration (B).

5C shows the α3β4 receptor residues and mutation-induced changes in terms of the number of contacts with the peptide.

At the α3(+) face, loss of peptide contact at several residues (e.g. Tyr-93, Cys-192, and Cys-193) is offset by increased contact at others (e.g. Ile-188, Tyr-190, Glu-195, Tyr-197, Gln-198, and Asp-199). Therefore, on average, peptide contact at the (+) face slightly increased. In contrast, at the β4(−) face, the F9A mutation caused widespread loss of contact across many residues (with notable exceptions at Arg-115). Therefore, removing the bulky phenyl side chain at position 9 makes AuIB detach from the β4(−) face. This slightly increased the peptide’s number of interatomic contacts at the opposing α3(+) face.

Based on this analysis, we propose several specific receptor residues that are especially important for AuIB inhibition of α3β4. First, because we demonstrated that the F9A mutation substantially reduces AuIB inhibitory efficacy, we examined the receptor residues that bind Phe-9, because these are probably crucial for AuIB inhibition of α3β4. The homology model and MD simulation trajectory of wild-type AuIB/α3β4 suggested that Phe-9 makes contact exclusively at the β4(−) face, with its phenyl ring sandwiched between the Trp-59 indole ring and hydrocarbon segment of the Lys-61 side chain (Fig. 6B). As expected, both of these residues lose contact with the peptide when position 9 of AuIB is substituted (Fig. 5C).

The geometry of contacts between AuIB-Phe-9 with β4-Trp-59 and β4-Lys-61 suggests that cation-π and π-π interactions may be important in AuIB inhibition of α3β4. However, we caution that the CHARMM27 force field does not explicitly account for interactions involving π electrons and acknowledge that methods which explicitly model cation-π interactions are needed to produce quantitatively accurate geometries involving charged and aromatic side chains. Nonetheless, the force field we used partially mimicked cation-π interactions and predicted Lys-Phe and Phe-Trp interaction geometries that are qualitatively in agreement with similar interactions observed in the PDB. In particular, our simulation suggests that contacts are made between β4-Lys-61 and the AuIB-Phe-9 phenyl ring primarily through the methylene carbons adjacent to the amine. In the PDB, Lys is known to engage with π systems more commonly via carbon (38, 39). Furthermore, our simulation suggests an offset-stacked interaction between β4-Trp-59 and AuIB-Phe-9. This geometry was identified as a common structural motif in Trp-Phe interactions in the PDB (40).

In addition, at the α3(+) face, there is modest but statistically significant loss of contact between AuIB and both C-loop cysteines when position 9 of AuIB is substituted (Fig. 5C). This is due to reduced contact between the AuIB-Cys2/8 cysteines and α3-Cys192/193. A snapshot from the MD trajectory of the native AuIB (Fig. 6C) illustrates close contact between the Cys2–8 sulfurs of the peptide and the Cys192/193 sulfurs of α3. In contrast, a similar snapshot for the [F9A]AuIB trajectory (Fig. 6D) illustrates loss of close contacts between the peptide and receptor disulfides. This result is especially intriguing, given that direct contact between peptide and C-loop cysteines is proposed to be essential for the peptide competitive antagonism of nAChRs, based on results from dicarba-conotoxin variants (41). Fewer close interatomic contacts between AuIB and α3 C-loop cysteines, indirectly caused by substituting Ala for Phe at position 9 (which lies on the opposite side of the peptide), may contribute to loss of inhibitory efficacy of this mutant. Therefore, we propose that Phe-9 has two roles in the AuIB inhibition of α3β4: 1) direct contact with β4-Trp-59 and Lys-61 and 2) indirect facilitation of close contact between AuIB Cys2/8 and α3-Cys192/193.

**Site-directed Mutagenesis of the β4 nAChR Subunit Verifies the Putative Binding Pocket on the β4 as the Site of Interaction between AuIB Phe-9 and α3β4 nAChRs**—Homology modeling and MD simulations indicated that the two residues on the (−) (complementary) side of the β4 subunit sandwich Phe-9 of AuIB. We further focused on the contribution of these residues to AuIB activity at the α3β4 nAChR. We used site-directed mutagenesis of the β4 nAChR subunit to verify what role β4-Trp-59 and β4-Lys-61 have in forming a binding pocket for AuIB Phe-9. When β4-Trp-59 and β4-Lys-61 were both mutated to alanines ([W59A]+[K61A]β4), the mutant α3β4 receptor produced much smaller ACh-evoked peak current amplitudes than those of wild-type nAChR. The same effect...
FIGURE 5. MD simulation suggests that substituting Ala for Phe at position 9 of AuIB reduces interaction between AuIB and the β4 nAChR subunit. A model shows α3β4 nAChR in the (α3)2(β4)3 stoichiometry in top view (A) and its α3(+)/β4(−) interface in side view (B). AuIB binding to the receptor is shown in red. C, contact difference bar graph representing quantification of results from homology modeling and MD calculations (20 ns). Shown is the number of interatomic contacts in the α3(+)/β4(−) subunit interface (<0.45 nm) between each receptor residue and the bound peptide lost (negative y axis values) or gained (positive y axis values) as a result of substituting Ala-9 for Phe-9. Note that the effect at the β4(−) side is a loss of interaction between [F9A]AuIB and the β4 subunit. D, a contact difference bar graph for [F9Y]AuIB indicates mutation to Tyr results in substantial loss of contacts between the peptide and several aromatic cage residues at the α3 (Tyr-93, Trp-149, Tyr-197) and β4 (Trp-59) subunits. Furthermore, the number of contacts with the α3-Cys192/193 sulfurs is reduced.
a control mutation of a tryptophan further down the sequence from the ACh binding site ([F9W]Aβ4), but no ACh-induced current was elicited at the mutated receptor.

Our model suggested that Tyr-5 of AuIB probably interacts with the backbone of Gln-198 of the α3 subunit (Fig. 6A). A mutation of α3-Gln-198 to alanine ([Q198A]α3) had only a minor, but significant, effect on currents evoked by ACh (50 μM) and inhibited by AuIB (10 μM) (0.25 ± 0.02, n = 5, p = 0.032 for [Q198A] mutant; 0.17 ± 0.02, n = 4 for native AuIB). However, mutation of its putative partner, Tyr-5 of AuIB ([Y5A]AuIB), did not reduce wild-type α3β4 nAChR inhibition at 3 μM (0.33 ± 0.02, n = 5) (see Fig. 2B). Therefore, α3-Gln-198 interaction with AuIB-Tyr-5 is unlikely to determine AuIB activity.

How Side Chain Size and Hydrophobicity at Position 9 of AuIB Affect Inhibition at the α3β4 nAChR—Given the critical interaction between AuIB-Phe-9 and β4-Trp-59 and the proposed importance of cation-π and π–π interactions for the inhibitory activity of AuIB, we assessed how side chain size, aromaticity, and hydrophobicity of the amino acid residue at position 9 in the AuIB sequence affect nAChR inhibition. In particular, we sought to find position 9 substituents that may better fit the Trp-59–Lys-61 pocket on the β4 nAChR subunit than Phe. We probed the inhibitory action of AuIB analogues substituted at position 9 by large aromatic (Tyr, Trp, 3-(2-naphthyl)-L-alanine (NAL)) and small (Gly) side chains on wild-type α3β4 nAChRs. CD analysis confirmed that the globular peptide structure was retained in all of these analogues ([FG9]AuIB, [F9Y]AuIB, [F9W]AuIB, [F9NAL]AuIB) (Fig. 8A).

Substituting Phe-9 to glycine ([FG9]AuIB, 10 μM) dramatically reduced inhibition (0.90 ± 0.04, n = 3; p < 0.001) compared with native AuIB (0.047 ± 0.01, n = 6) (Fig. 8B). This is consistent with our proposed model in which AuIB interaction with the binding pocket on the β4 subunit is crucial. This interaction is presumably absent when Phe-9 is replaced by the side chain-free Gly.

Phe-9 mutation to the slightly larger and less hydrophobic Tyr ([F9Y]) significantly reduced α3β4 nAChR inhibition (0.42 ± 0.07, n = 3; p < 0.001) (Fig. 8B) compared with native AuIB. Homology modeling and MD simulation of the [F9Y]AuIB-α3β4 interaction suggest that mutation to Tyr results in substantial loss of contacts between the peptide and several aromatic cage residues at the α3 (Tyr-93, Trp-149, Tyr-197) and β4 (Trp-59) subunits, as shown by contact difference plot (Fig. 5D). The latter residue is of special interest given its known importance for AuIB inhibition (discussed above). Inspection of the structural model indicates that the aromatic ring of Tyr-9 in [F9Y]AuIB directly contacts the hydrocarbon segment of β4-Lys-61. But unlike the Phe-9 of native AuIB (Fig. 6B), the Tyr-9 side chain does not contact Trp-59 β4. Furthermore, mutation to Tyr results in loss of contact with the α3-Cys-192/193 sulfurs (Fig. 5D). Together with the electrophysiology data, our model of [F9Y]AuIB-α3β4 interaction further supports our hypothesis that position 9 contact with the β4-Tyr-59 and α3 disulfide structural motif is essential for AuIB activity against α3β4.

Another AuIB analog in which Trp was substituted for Phe-9 ([F9W]) substantially reduced inhibition (0.80 ± 0.07, n = 3; p <
presumably due to steric clashes. Indeed, automated blind docking of [F9W]AuIB to /H9251 3/H9252 4 using Autodock (42) with all peptide side chain torsions free to rotate showed that none of the docking solutions involved binding of analog [F9W]AuIB to the canonical C-loop pocket. In contrast, docking of both native and [F9Y]AuIB predicted binding conformations very close to that of the double mutant PnIA-AChBP co-crystal structure (2BR8) (data not shown). An analog with an even larger unnatural amino acid, NAL, to substitute Phe-9 ([F9NAL]AuIB) also showed dramatically reduced inhibition (0.82 ± 0.07, n = 3; p < 0.001) (Fig. 8B). The NAL was attached to the 2-position of the Ala residue. Unlike the Trp residue, which has an N-heterocyclic five ring, NAL has a homocarbocyclic six ring.

Together, these results demonstrate that side chain size, aromaticity, and hydrophobicity at position 9 of AuIB are important for interaction between the peptide and β4 subunit of the α3β4 pentamer. In contrast with the above mentioned analogues with substitutions at position 9, inhibition in a control AuIB analog with a histidine substitution at position 12 (N12H) was not significantly impaired (0.069 ± 0.003, n = 3) (Fig. 8B).

**DISCUSSION**

Using alanine scanning mutagenesis, we identified three residues in the AuIB sequence (Gly-1, Pro-6, and Phe-9) that affect inhibition of α3β4 nAChRs. The Gly to Ala substitution only moderately reduced inhibition. Homology modeling of the AuIB-α3β4 complex suggests that the N terminus NH$_2$ of AuIB forms a salt bridge with the β4-Asp-172 side chain. The G1A mutation introduces a non-polar CH$_3$ side chain that may weaken the favorable interaction between the peptide N ter-

FIGURE 7. The effects of point mutations in the β4 nAChR subunit on AuIB inhibition. A, representative ACh-evoked currents showing native AuIB (10 μM) inhibition of α3β4 nAChRs compared with the β4 subunit double mutant (W59A + K61A) and two single point mutants of β4, W59A and K61A. B, bar graph showing quantification of AuIB (10 μM) inhibition at the β4 nAChR subunit mutants and wild-type α3β4 nAChR. AuIB inhibition is abolished in [W59A]β4 and the [W59A]+[K61A]β4 double mutant, confirming these residues have a key role in AuIB binding. Note the intermediate effect of K61A mutation, suggesting it has an auxiliary role in forming the binding pocket for the hydrophobic interaction with Trp-59, which is key for inhibition. Note that only high ACh concentration elicited measurable currents in the W59A+K61A mutant. Therefore, 2.5 mM ACh was used as an agonist to compare inhibition in the mutants and wild-type receptors. Data represent the means ± S.E., n = 3–6. ** indicates p < 0.005; *** indicates p < 0.001 for relative reduction of current inhibition versus wild-type α3β4 nAChRs.
minus and β4-Asp-172 side chain. Additionally, in a recent study investigating N-terminal post-translational modification of α-conotoxin M1, truncation of the N-terminal residue decreased the affinity for the muscle-type nAChR, a direct pharmacological effect indicating the N-terminal residues are important for bioactivity (43). α-Conotoxin LsIA has also shown reduced potency at α7 and α3β2 nAChRs when N-terminally truncated (44), which corroborates the potential importance of the N-terminal amino acid in α-conotoxin potency and selectivity.

The other two substitutions, P6A and F9A, caused an even greater decrease in peptide activity on α3β4 nAChR than G1A. Our NMR and CD data revealed that the [P6A]AuIB analog structure is disrupted, but that of [F9A]AuIB is not. This is not surprising considering the structural effect Pro has on protein structures in general and α-conotoxin structures in particular (34). Therefore, we reasoned that the ability of Phe-9, but not Pro-6, to reduce inhibition is due to Phe-9 specifically interacting with the α3β4 nAChR. We found that the number of mutations that reduce AuIB inhibition of the α3β4 nAChR is small compared with other α-conotoxins. For instance, alanine substitutions of almost all intercysteine residues in another α-conotoxin, Vc1.1, affected α9α10 nAChR inhibition (35). We also found no alanine mutants that inhibited the α3β4 nAChR significantly more than native AuIB.

As well as substantially reducing α3β4 inhibition, [F9A]AuIB selectivity for nAChR subtypes also shifted. This indicates that Phe-9 has a role in the peptide specific interaction with the
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dihydro-β-erythroidine and bungarotoxin and to agonists ACh and nicotine were explained by the same difference in the sequence (Lys-61 at β4; Thr-59 at β2 numbering) (55, 56). A residue in this position underlies a difference between insect and vertebrate nAChRs in sensitivity to neonicotinoid insecticides (57). A difference in two residues preceding TrpD accounts for a remarkable difference in human versus rat α3β4 nAChR sensitivity to the agonist 5-(trifluoromethyl)-6-(1-methyl-azepan-4-yl)methyl-1H-quinolin-2-one (TMQA) (54).

Using α3β2 receptor chimeras, determinants for MII conotoxin specificity were found on α and β subunits (58). Interestingly, one of the determinants was mapped to loop D of the β2 subunit and identified as Thr-59 (homologous to β4-Lys-61). Therefore, the Lys to Thr substitution in the β2 subunit was one factor that defined β2/β4 selectivity for MII. BuIα conotoxin has a different wash-off rate for α3β2 than α3β4 nAChRs. The same difference in the sequence (Lys/Thr-59) in β4 and β2 was also a determinant for the different wash-off rates for the various nAChR subtypes (59).

Consistent with the studies mentioned above, our modeling and experimental data show that the β4 single-point mutation [K61A]β4 only reduces AuIB inhibition, but [W59A]β4 (TrpD) completely abolishes it. Therefore, TrpD is essential for AuIB to inhibit α3β4 nAChRs. TrpD involvement in nAChR pharmacology is well documented. Mutations of TrpD in AChBP have been shown to affect its conformation as well as alter its potency, desensitization, efficacy, and selectivity for various ligands (60–64). However, its effects depend on the subunit and ligand in question and are more ambiguous than mutations of the principal subunit’s aromatic residues. For instance, the effects of the TrpD mutation on ACh affinity for muscle nAChRs differed between δ and γ subunits (20,000 versus 7,000-fold reduction) (61). TrpD mutation caused opposite shifts in the potency of 4OH-GTS21 agonists in α7 and α4β2 nAChRs (62).

When expressed in Xenopus oocytes and tested by two-voltage clamp recordings, the [W59A]β4 mutation dramatically reduced the ACh-induced current without altering its kinetics. To induce measurable currents, we had to use the saturating ACh concentration of 2.5 mM instead of 50 μM. In a previous study we showed that high ACh concentration has practically no effect on AuIB inhibition of α3β4, as this inhibition is non-competitive (14).

There are examples where the binding/action of pharmacological agents is critically dependent on interaction with TrpD. Apolipoprotein E inhibits α7 nAChRs via hydrophobic interactions with α7 TrpD (Trp55 in α7 numbering) (64). Varenicline, a smoking cessation drug, has been shown to interact with β2-TrpD in the α4β2 nAChR. TrpD substitution to Ala converted varenicline from a partial to full agonist and abolished varenicline-induced desensitization at high concentrations (63).

Nature of AuIB-Phe-9β4-Trp-59 Interaction and Its Implications for AuIB Pharmacology—The sequence alignment of several nAChR subunits (Fig. 9) centered at the AuIB binding pocket shows that only TrpD (Trp-59 in β4 numbering) is absolutely conserved across the different subunits. However, substitutions in the WLK pocket are often homologous. For example,
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instead of β4-Lys at the TrpD +2 position, α9 and α10 subunits have a similar positively charged Arg. There is a single different residue, Thr instead of Lys, between β2 and β4, and this can cause differences in pharmacology. It is unlikely that Lys/Thr substitution underlies AuIB selectivity for the α3β4 subtype, as the [K61A]β4 mutation only reduces inhibition, and AuIB is not active at the α9a10 subtype when Lys is changed to an essentially homologous Arg. Interestingly, loop D of the α3 subunit has the same WLK pocket as the β4 subunit.

AuIB could also bind at the non-canonical binding site (+)/β/(-)α, as suggested for galanthamine and cocaine binding to AChBP (65), anti-helminthic compound morantel binding to α3β2 nAChRs (66), and recently for α-conotoxin Vc1.1 binding to α9a10 nAChRs (67). However, this is unlikely because the docking model does not support binding at a non-canonical site and the [W59A]β4 mutant loses inhibition despite identical WLK pockets being available on the α3 subunit (−) side. Another possibility could be that AuIB anchoring at the WLK pocket is controlled by its interactions with the α subunit.

Our homology and MD-simulated model suggested that Tyr-5 of AuIB interacts with the α3 subunit Gln-198. However, [Q198A]α3 mutation did not affect AuIB inhibition, and mutation of its putative partner, Tyr-5-AuIB, did not reduce inhibition on the wild-type receptor. Mounting evidence suggests that differences in three-dimensional structure across subunits probably underlies distinct bond patterns for the same ligand despite the presence of identical residues, as demonstrated for nicotinic ligands (68, 69). This is certainly possible for larger molecules, such as AuIB.

There is also the question of the mechanism behind AuIB inhibition of nAChRs and how it can act non-competitively despite overlapping with the ACh-binding site. At least two larger peptides, apolipoprotein E and Aβ1–42 amyloid, block nAChRs non-competitively as well. In addition, apolipoprotein E binds to loop D by interacting with TrpD (64, 70). AuIB may be able to act non-competitively because it can bind to α3β4 in the presence of ACh. Because TrpD may be involved in desensitizing or transducing the gating signal (54, 63, 71), AuIB may work by stabilizing the desensitized state or blocking the gating signal. Another possibility is steric hindrance to the movement of loop C, which is thought to be associated with agonist activation of nAChRs (47, 72).

What could be the nature of the AuIB-Phe-9 interaction with TrpD? It is well established that residues around ligand binding sites and those that form the aromatic box in Cys-loop family receptors often make cation–π bonds with their cognate ligands (73). The reverse situation, when aromatic moieties of ligands engage in a cation–π bond formation with positively charged residues of the receptor, is also possible (74). AuIB does not have positively charged residues so cannot make a cation–π bond with β4-Trp-59 (TrpD). A situation in which a positively charged β4-Lys-61 forms a cation–π bond with Phe-9 of AuIB is possible. However, our experiment with single-point mutations of the WLK pocket showed that removing Lys-61 does not abolish inhibition but that Trp-59 mutation does. This suggests that Trp-59 is indispensable for AuIB inhibitory effect and Lys-61 plays an auxiliary role in it.

Our experiments with second-generation AuIB mutants at position 9 demonstrated that size and hydrophobic/aromaticity of the residue in this position are important for the peptide to inhibit α3β4 nAChRs. Although simple hydrophobic interaction of AuIB-Phe-9 and β4-Trp-59 is possible, it is more likely that Phe-9 and Trp-59 interact via π–π stacking due to the deep insertion of Phe-9 in the WLK pocket. Because removing positively charged Lys-61 reduces inhibition, it will likely interact with Phe-9 of AuIB and/or stabilize AuIB-Phe-9 interaction with β4-Trp-59. In conclusion, we identified determinants of AuIB binding/action on the α3β4 nAChR.

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