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Membrane Transport, Structure, Function, and Biogenesis:
β2 Subunit Contribution to 4/7 α-Conotoxin Binding to the Nicotinic Acetylcholine Receptor

Sébastien Dutertre, Annette Nicke and Richard J. Lewis
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The structures of acetylcholine-binding protein (AChBP) and nicotinic acetylcholine receptor (nAChR) homology models have been used to interpret data from mutagenesis experiments at the nAChR. However, little is known about AChBP-derived structures as predictive tools. Molecular surface analysis of nAChR models has revealed a conserved cleft as the likely binding site for the 4/7 α-conotoxins. Here, we used an α3β2 model to identify β2 subunit residues in this cleft and investigated their influence on the binding of α-conotoxins MII, PnIA, and GID to the α3β2 nAChR by two-electrode voltage clamp analysis. Although a β2-L119Q mutation strongly reduced the affinity of all three α-conotoxins, β2-F117A, β2-V109A, and β2-V109G mutations selectively enhanced the binding of MII and GID. An increased activity of α-conotoxins GID and MII was also observed when the β2-F117A mutant was combined with the α4 instead of the α3 subunit. Investigation of A10L-PnIA indicated that high affinity binding to β2-F117A, β2-V109A, and β2-V109G mutants was conferred by amino acids with a long side chain in position 10 (PnIA numbering). Docking simulations of 4/7 α-conotoxin binding to the α3β2 model supported a direct interaction between mutated nAChR residues and α-conotoxin residues 6, 7, and 10. Taken together, these data provide evidence that the β subunit contributes to α-conotoxin binding and selectivity and demonstrate that a small cleft leading to the agonist binding site is targeted by α-conotoxins to block the nAChR.

Neuronal nicotinic acetylcholine receptors (nAChRs) comprise a large family of ion channels formed by the heteropentameric assembly of homologous subunits. α-Conotoxins, small disulfide-rich peptides isolated from the venom of predatory cone snails, potently and selectively block nAChRs. α3β2, α3β4, and α4β2 nAChR subtypes are already providing useful insights into ligand-receptor interactions at the molecular level. Our previous docking simulations of α-conotoxins ImI and PnIB on an α3 nAChR homology model revealed that these two peptides target a small cleft forming one of two possible entrances to the ACh binding site. We could also identify an equivalent cleft on an α3β2 model as the likely region of the α3β2 nAChR targeted by these peptides. Upon examination of this model, it was apparent that the β2 subunit contributed more than the α subunit to the formation of this cleft, suggesting an important role of β-residues for binding and/or selectivity. In this study, we mutated three residues in the β2 subunit residues located in this cleft to characterize the contribution of the β subunit to α-conotoxin binding. Each of these mutants influenced the binding of α-conotoxins MII, GID, and PnIA. Additionally, molecular surface analysis revealed striking shape complementarity between the highly conserved N-terminal half of the 4/7 α-conotoxins structure and the cleft. Finally, docking experiments showed that both MII and PnIA bind deep into the α3β2 nAChR cleft, explaining at the molecular level the experimental results obtained from mutagenesis studies. Based on these results, we propose that the conserved cleft above the β9β10 hairpin is the binding site for 4/7 α-conotoxins active at α7, α3β2, and α4β2 nAChRs and that the

β2 Subunit Contribution to 4/7 α-Conotoxin Binding to the Nicotinic Acetylcholine Receptor*

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† The abbreviations used are: nAChR, nicotinic acetylcholine receptor; ACh, acetylcholine; AChBP, acetylcholine-binding protein.

The 4/7 (4 residues in loop 1 and 7 in loop 2) α-conotoxins are of particular interest given their ability to discriminate between diverse neuronal α-β nAChR subunit combinations. For example, MII was found to be selective for the α3 and especially the α6 subunit containing nAChRs (3, 4) and aided the identification of the nAChR subunit composition in monkey striatum implicated in Parkinson disease (5, 6). The 4/7 α-conotoxin PnIA preferentially blocks α3β2 but also α7 rat nAChRs (7, 8), whereas α-GID potently blocks α7 and α3β2 nAChRs but also acts at α4β2 nAChR at higher concentrations (9). MII, PnIA, and GID share an identical backbone structure common to all 4/7 α-conotoxins investigated (9–11). Therefore, their specific selectivities are thought to arise from their different amino acid side chains. As a consequence, understanding the molecular determinants of their interaction with the receptor may help in the design of optimized pharmacological tools for the nAChRs with novel or improved selectivities. However, the lack of detailed information on the three-dimensional structure of the α-conotoxin binding site has hampered such projects.

The acetylcholine-binding protein (AChBP) crystal structure solved by Sixma and colleagues (12) recently provided a template to create homology models of the ligand binding domain of receptor members of the Cys-loop ligand-gated ion channel family. Homology models of the ligand binding domain of α7, α3β2, α3β4, and α4β2 nAChR subtypes are already providing useful insights into ligand-receptor interactions at the molecular level (13–16). Our previous docking simulations of α-conotoxins ImI and PnIB on an α7 nAChR homology model revealed that these two peptides target a small cleft forming one of two possible entrances to the ACh binding site (17). We could also identify an equivalent cleft on an α3β2 model as the likely region of the α3β2 nAChR targeted by these peptides. Upon examination of this model, it was apparent that the β2 subunit contributed more than the α subunit to the formation of this cleft, suggesting an important role of β-residues for binding and/or selectivity. In this study, we mutated three residues in the β2 subunit residues located in this cleft to characterize the contribution of the β subunit to α-conotoxin binding. Each of these mutants influenced the binding of α-conotoxins MII, GID, and PnIA. Additionally, molecular surface analysis revealed striking shape complementarity between the highly conserved N-terminal half of the 4/7 α-conotoxins structure and the cleft. Finally, docking experiments showed that both MII and PnIA bind deep into the α3β2 nAChR cleft, explaining at the molecular level the experimental results obtained from mutagenesis studies. Based on these results, we propose that the conserved cleft above the β9β10 hairpin is the binding site for 4/7 α-conotoxins active at α7, α3β2, and α4β2 nAChRs and that the...
differences in the cleft residues provide the determinants directing α-conotoxin selectivity.

EXPERIMENTAL PROCEDURES

Materials—cDNAs encoding neuronal nAChRs were provided by J. Patrick (Baylor College of Medicine, Houston, TX) and subcloned into the oocyte expression vector pNKS2 (18). MII, PnIA, and GID were gifts from J. T. Blanchfield (School of Pharmacy, The University of Queensland, Australia), G. Hopping and M. Loughnan (Institute for Molecular Bioscience, The University of Queensland, Australia), respectively, and were synthesized as described previously (9, 19, 20).

Homology Modeling—The FASTA format of the α3β2 and α9β2 rat sequences were retrieved from the ligand-gated ion channel database (pasteur.fr/recherche/banques/LGIC/). Their extracellular ligand-binding domains were aligned with the AChBP sequence as described previously (21).

A homology model of the α3β2 nAChR subtype was built on a Silicon Graphics Octane R12000 work station using the MODELLER program (22). The AChBP structure (1I9B) was loaded in the INSIGHT II (Accelrys, San Diego, CA) environment and used as a template. Three models were built with a high “optimize level.” The two options “optimize loop models” and “loop optimize level” were set to 3 and high, respectively. User disulfide was selected to assign the disulfide bonds as in AChBP except for the missing vicinal disulfide in the α9 subunit. The model with the lowest root mean square deviation compared with AChBP was refined further and used for these studies. Steepest descent energy minimizations were applied using the AMBER force field and DISCOVER program implemented in Insight II.

Mutagenesis—Mutagenesis of the β2 nAChR subunit cDNA was achieved using the QuikChange™ site-directed mutagenesis kit (Stratagene) following the manufacturer’s instructions. Primers used to generate the mutants were from Proligo (Lismore, Australia). All mutations were confirmed by cDNA sequencing.

Electrophysiological Recordings—cRNA was synthesized from linearized plasmids with SP6 RNA polymerase using the mMessageMachine kit (Ambion, Austin, TX). Xenopus laevis frogs were purchased from Nasco International (Fort Atkinson, WI) or Firma Kahler (Hamburg, Germany). X. laevis oocytes were prepared as described previously (6) and injected with 50-nl aliquots of cRNA (0.5 μg/μl).

Two-electrode voltage clamp recordings were performed in oocytes 1–10 days after cRNA injection at a holding potential of −70 mV. Pipettes were pulled from borosilicate glass and filled with 3 M KCl. Resistances were below 1 megohm. Membrane currents were recorded using a Turbo Tec-10CX or a Turbo Tec 05X Amplifier (npi electronic, Tamm, Germany) filtered at 200 Hz and digitized at 400 Hz. Version


### Table I

<table>
<thead>
<tr>
<th>α-Conotoxins</th>
<th>IC₅₀ values and 95% confidence interval for α-conotoxins on wild type and mutant nAChR combinations</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>α-Conotoxins</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>MII</td>
<td>α3β2</td>
</tr>
<tr>
<td></td>
<td>α3β2F117A</td>
</tr>
<tr>
<td></td>
<td>α3β2V109A</td>
</tr>
<tr>
<td></td>
<td>α3β2V109G</td>
</tr>
<tr>
<td></td>
<td>α3β2L119Q</td>
</tr>
<tr>
<td>GID</td>
<td>α3β2</td>
</tr>
<tr>
<td></td>
<td>α3β2F117A</td>
</tr>
<tr>
<td></td>
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<td>α3β2L119Q</td>
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<td></td>
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</tbody>
</table>

* ND, not determined.  
* The 95% confidence interval values are shown in parentheses.

### Table II

<table>
<thead>
<tr>
<th>α-Conotoxins</th>
<th>kᵣ⁻¹ min⁻¹</th>
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<tr>
<td>MII</td>
<td>α3β2</td>
<td>0.07</td>
<td>(0.04–0.1)</td>
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<tr>
<td></td>
<td>α3β2F117A</td>
<td>ND*</td>
<td>(8.9 ± 0.7% recovery in 20 min)</td>
</tr>
<tr>
<td></td>
<td>α3β2V109A</td>
<td>ND</td>
<td>(14.1 ± 2.5% recovery in 20 min)</td>
</tr>
<tr>
<td></td>
<td>α3β2V109G</td>
<td>ND</td>
<td>(15.2 ± 3.0% recovery in 20 min)</td>
</tr>
<tr>
<td></td>
<td>α3β2V109G</td>
<td>0.03</td>
<td>(0.01–0.05)</td>
</tr>
<tr>
<td>GID</td>
<td>α3β2</td>
<td>0.19</td>
<td>(0.15–0.22)</td>
</tr>
<tr>
<td></td>
<td>α3β2F117A</td>
<td>ND</td>
<td>(15.2 ± 3.0% recovery in 20 min)</td>
</tr>
<tr>
<td></td>
<td>α3β2V109A</td>
<td>0.09</td>
<td>(0.06–0.12)</td>
</tr>
<tr>
<td></td>
<td>α3β2V109G</td>
<td>0.03</td>
<td>(0.01–0.05)</td>
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<tr>
<td>PnIA</td>
<td>α3β2</td>
<td>ND</td>
<td>(100% recovery in 2 min)</td>
</tr>
<tr>
<td></td>
<td>α3β2F117A</td>
<td>ND</td>
<td>(100% recovery in 4 min)</td>
</tr>
<tr>
<td></td>
<td>α3β2V109A</td>
<td>ND</td>
<td>(100% recovery in 2 min)</td>
</tr>
<tr>
<td>[A10L]PnIA</td>
<td>α3β2</td>
<td>ND</td>
<td>(100% recovery in 2 min)</td>
</tr>
</tbody>
</table>

* ND, not detectable.  
* The 95% confidence interval values are shown by numbers in parentheses.

8.53 Pulse software (HERA Elektronik, Lambrecht, Germany) or CellWorks software were used. The perfusion medium was automatically switched between ND96 with or without agonist (100 µM ACh) using a custom-made magnetic valve system. A fast and reproducible solution exchange (<300 ms) for agonist application was achieved using a 50-µl funnel-shaped oocyte chamber combined with a fast solution flow (~150 µls) fed through a custom-made manifold mounted immediately above the oocyte. ACh pulses were applied for 2 s at 4-min intervals. After each application, the cell was superfused for 1 min with agonist-free solution, and the flow was then stopped for 3 min. Peptide was mixed from a 10-fold stock into the static bath when responses to three consecutive agonist applications differed by less than 10%. Addition of toxin directly to the recording chamber conserved material and avoided adhesion of the toxin to tubing surfaces. To obtain estimates of potency, dose-response curves were fit to the data by the equation % response = 100/(1 + ([toxin]/IC₅₀)^µH) using Prism software (GraphPad version 3.0 for Macintosh, San Diego, CA). To obtain estimates of toxin dissociation rates, agonist responses were measured at 2-min intervals under constant superfusion, and after stabilization of responses, oocytes were incubated with toxin in a static bath for 1 min. To obtain estimates of the toxin association rate, oocytes were continuously superfused with solution containing the indicated toxin concentration and ACh pulses were applied in 1- or 2-min intervals. Association and dissociation curves were fit to the data by the equations % response = (100 – plateau) x e^(-K x time) + plateau and % response = span (1 – e^(-K x time)) + plateau, respectively.

**Docking Experiments**—Docking simulations of α-conotoxins were carried out as described in Dutertre et al. (17). Briefly, MII and PnIA structures were retrieved from the protein data bank (Protein Data Bank codes 1MI1 and 1PEN, respectively) and docked onto an α3β2 homology model using the program GOLD, version 1.2 (Genetic Optimization for Ligand Docking, Cambridge Crystallographic Data Centre, Cambridge, UK). One NMR/crystal structure was chosen for each α-conotoxin, as GOLD treats ligands as flexible molecules with side chain orientations optimized during calculations. As conotoxins are competitive antagonists, the active site radius was set at 20 Å from
Trp-147 to ensure the analysis focused on residues around the ACh binding site. From the 100 docked structures obtained for each conotoxin, the selection of the final docked structure was based on a low scoring function determined in GOLD (23).

RESULTS AND DISCUSSION

Homology Modeling Identifies Val-109, Phe-117, and Leu-119 in the β2 nAChR Subunit as Likely Residues Interacting with α-Conotoxins MII and PnIA—We previously identified two cavities at opposite sides of the β9/10 hairpin (C-loop of an α subunit) on the surface of the nAChR, one large and easily accessible and one small and narrow, from which the ACh binding site could be reached (24). To differentiate between these two cavities and avoid confusion, we propose the use of “cavity” to refer to the larger one (below the β9/10 hairpin, close to the cell membrane), and “cleft” to refer to the smaller one (above the β9/10 hairpin). The cavity has recently been confirmed to represent the binding site used by the large snake neurotoxins to block the nAChR (24), whereas the cleft represents the likely region targeted by α-conotoxins (17, 21). Docking simulations with ImI and PnIB structures on an α3 subunit model revealed overlapping but different binding sites for α-conotoxins and snake toxins (17). Both α-conotoxins block the receptor by targeting a small cleft above the β9/10 hairpin that is also blocked by the small peptide antagonist toxin waglerin (25) and the non-peptidic antagonist d-tubocurarine (26). Concerning the α3β2 nAChR, only one residue (β2-Thr-57) weakly affecting 4/7 conotoxin binding has been identified in the β2 subunit compared with four on the α3 subunit (27, 28). Interestingly, β2-Thr-57 is located above the β9/10 hairpin, forming part of the cleft in α3β2 together with three other residues, Val-109, Phe-117, and Leu-119. A potentially important role for additional residues in the β subunit is suggested from binding kinetic studies where exchange of the β2 subunit by the β4 subunit (in combination with α3 or α6 subunits) strongly reduced the off-rate constants of PIA (29). Likewise, BuIA showed much slower dissociation if α2, α3, or α6 subunits are combined with β4 instead of β2 (30). In an attempt to identify additional determinants in the β subunit that contribute to the binding and selectivity of 4/7 α-conotoxins, we mu-
tated Leu-119, which appears deep in the cleft, as well as Phe-117 and Val-109, which form part of the wall of the cleft (Fig. 1).

Residue Leu-119 Is Generally Important for α-Conotoxin Binding—A leucine in position β2-119 lies at the bottom of the cleft and is conserved in neuronal β4 and α7 nAChR subunits as well as in the muscle δ, γ, and ε nAChR subunits. Thus, a leucine is present at this location in all non-α subunits (and the −− face of the α7 nAChR) for which highly active α-conotoxins have been identified. Indeed, no potent α-conotoxin has been found for the α9 and α10 subunits, which have a negatively charged aspartic acid at this position. In combination with the docking studies, this suggests that the hydrophobic environment created by β2-Leu-119 and the corresponding leucine residues in other subunits plays an important role in the high affinity interaction with α-conotoxins. To test the hypothesis that Leu-119 is generally important for α-conotoxin binding, the activity of three different α-conotoxins that bind with high affinity to the α3β2 nAChRs was tested on the hydrophilic α3β2-L119Q mutant. The α3β2/αβ2-selective α-conotoxin MII and as well as the α3β2/αβ2-selective α-conotoxins PnIA and GID had a strongly reduced activity (Fig. 2, A–C), with 40-fold (PnIA), 165-fold (GID), and 300-fold (MII) higher IC50 values at this mutant α3β2 receptor (Table 1). This is the strongest effect described thus far for a non-α subunit mutant affecting α-conotoxin binding. Based on its position at the bottom of the cleft and the fact that the activity of all three conotoxins was reduced by at least 40-fold upon introduction of a similar length hydrophilic residue (Gln), Leu-119 may play an important stabilizing role for the interaction with α-conotoxins, allowing them to bind deep into the nAChR cleft.

In addition to the four cysteines, which are involved in two disulfide bonds and are mostly buried within the peptide core, only serine and proline residues are common to all three α-conotoxins. Because mutation of Ser-4 in MII did not affect potency (∼1), we suspect that Leu-119 might interact with the α-conotoxin backbone or the conserved proline found in 4/7 α-conotoxins (third residue of loop 1). However, this possibility is difficult to test by double mutant cycle analysis because alteration of the conserved cysteine or proline residues in the conotoxin alters the peptide backbone conformation.

Alanine Exchange of Val-109 and Phe-17 Specifically Enhances Binding of MII and GID—In our model of the β2 subunit, residues Val-109 and Phe-117 lie adjacent to each other on two neighboring antiparallel β-strands and above the Leu-119 residue, potentially forming the walls of the α-conotoxin binding cleft (Fig. 1). By mutating the residues into the smaller alanine, we expected to weaken the interaction with α-conotoxins. Unexpectedly, both the F117A and V109A mutations produced IC50 values for MII, GID, and PnIA little different from those at the wild type receptor, suggesting that Phe-117 and Val-109 do not have close interactions with the α-conotoxins.
molecular surface complementarities for α-conotoxins and the α3β2 cleft. Molecules are shown at the same scale, allowing direct size comparison and shape complementarity between α-conotoxins and the nAChR cleft. α-Conotoxins PnIA and MII are shown with their N-terminal sequences facing the reader. The contour of the cleft was flipped 180° to allow a direct comparison with the α-conotoxin shapes. This figure was produced using MOLCAD in Sybyl7.0 (Tripos software, St. Louis, MO).

FIG. 7. Molecular surface complementarities for α-conotoxins and the α3β2 cleft. Molecules are shown at the same scale, allowing direct size comparison and shape complementarity between α-conotoxins and the nAChR cleft. α-Conotoxins PnIA and MII are shown with their N-terminal sequences facing the reader. The contour of the cleft was flipped 180° to allow a direct comparison with the α-conotoxin shapes. This figure was produced using MOLCAD in Sybyl7.0 (Tripos software, St. Louis, MO).

FIG. 8. Docking of PnIA (A) and MII α-conotoxins (B) onto the α3β2 nAChR. Each of these conotoxins penetrated deep into the cleft with its conserved N-terminal hydrophobic patch, which includes the conserved Pro-6. Toxin residues make contacts with both the α3 and β2 subunits. The C-terminal part of both conotoxins protrudes beyond the cleft.
interactions with the Phe-117 and Val-109 mutants. To further test this hypothesis, we investigated the activity of the A10L-PnIA analogue on both /H92513/H92522 and /H92514/H92522 mutants. As predicted, replacing Ala-10 by a longer leucine residue enhanced PnIA affinity for /H92522 receptors containing the Phe-117 or Val-109 mutations (Figs. 2D and 3D; Tables I and II). Indeed, a markedly reduced off-rate constant and an up to 23-fold lower IC50 value were observed for A10L-PnIA at the /H92522 mutants (Table I: 3.99 nM on /H92522-F117A; 3 nM on /H92522-V109A; 2.4 nM on /H92522-V109G) compared with the wild type /H92513/H92522 receptor (55 nM). Moreover, this A10L modification converted PnIA into a potent antagonist at the /H92514/H92522-V109G receptor (Fig. 5B).

Residue 10 in PnIA has previously been shown to be an important determinant of selectivity between /H92513/H92522 and /H92517/H92527 receptors (8). A comparison of a range of conotoxins with sequence similarity in the second loop showed that the length of the side chain in position 10 correlated with an /H92517 versus /H92513/H92522 selectivity. This correlation was also found to be valid for GID and PnIA but not for MII, suggesting that MII has an altered binding mode. However, the similar influence of /H92522 mutants on GID, PnIA, and MII affinity indicate that these /H92513-conotoxins orientate similarly within the cleft formed by the /H92513/H92522 interface and that the critical position 10 side chain likely faces the /H92522 subunit in all three /H92513-conotoxins.

Shape and Size Complementarity between 4/7 /H92513-conotoxins and the Cleft in nAChRs—A high affinity complex requires either a large contact area between receptor and ligand surfaces with multiple points of interaction or shape complementarity. 

**FIG. 9.** α-Conotoxin and β2 subunit pharmacophores. PnIA (A) and MII (B) conotoxins are presented in their docked orientation in the cleft (C). The minimal pharmacophore shown is consistent with mutagenesis results and previous α-conotoxin alanine scan data. Distances (Å) between residues are measured from the last carbon of the side chain and reported on the vectors in the right-hand panels. Also shown are the equivalent residues for the β3 (green), β4 (blue), and α7 (red) subunits. Note that Phe-117 is positioned under Glu-59.


tility that favors the formation of a smaller number of specific interactions (e.g., a ligand bound in a protein cleft). Given their relatively small size, the high affinity complex observed for α-conotoxins suggests that a tight fit between the α-conotoxin and a cleft on the nAChR surface is likely to occur. The α7 nAChR cleft appears to be ~7 Å deep, 13.5 Å long, and 9.5 Å wide (6.6 Å at the narrowest point). These dimensions are almost identical to those found for the α3β2 nAChR (7.5 Å deep, 14.5 Å long, and 9.8 Å wide (6.8 Å at the narrowest)), and are mostly conserved in shape. For comparison, we measured the dimensions of the N-terminal hydrophobic patch of α-conotoxins. In MII it is 12 Å long (Gly-to-Val-7) and 7 Å wide, and in PnA it is 11.2 Å long (Gly-1 to Pro-7) and 6.5 Å wide, whereas GID has a flexible tail that complicates the measurements (a tail-truncated analogue has similar dimension to PnIA and MII, and the tail could be accommodated by the flexible loop F of nAChR).

These comparisons reveal a striking size and shape complementarity between 4/7 α-conotoxins and the nAChR cleft, strongly suggesting a lock-and-key interaction (Fig. 7).

Docking Models of PnIA and MII Interactions with α3β2 nAChR—To verify that 4/7 α-conotoxins can interact with the nAChR cleft via their conserved hydrophobic patch (residues 6, 7, and 10), docking of PnIA and MII structures onto an α3β2 nAChR model were simulated using the program GOLD. Analysis of the results from docking simulations confirmed both hypotheses: the small cleft is the binding site for 4/7 α-conotoxins, and the α-conotoxins bind to the α3β2 nAChR by presenting their conserved N-terminal shape (Fig. 7). Indeed, although an active site radius set to 20 Å would have allowed ligands to dock at other locations around the C-loop, all structures were found docked in the cleft above the β7/10 hairpin.

The final orientation of PnIA and MII docked into the receptor cleft presented in Fig. 8 was chosen from the cluster of lowest docking results (Fig. 9). PnIA has a very hydrophobic N-terminal patch, comprising Leu-5, Pro-6, Pro-7, Ala-9, and Ala-10 (Fig. 9A). MII also possesses a similar hydrophobic core (Pro-6, Val-7, and Leu-10) that we take to represent a minimum antagonist pharmacophore (Fig. 9B). When compared with the β2 subunit pharmacophore (Fig. 9C), the distances found in conotoxins are compatible with the direct interaction between the conserved proline in position 6 and β2-Leu-119 that we observed in our docking model. We propose that Pro-6 in 4/7 conotoxin acts as an anchor and stabilizes the conotoxin-receptor complex. Additional hydrophobic contacts such as Pro-7 (PnIA) or Val-7 (MII) and Ala-10 (PnIA) or Leu-10 (MII) with β2-Val-109 and β2-Phe-117 strengthen the interaction, whereas secondary structures such as His-9 (MII) and β2-Glu-57 determine the selectivity. A comparison of residues in the equivalent position of the receptor pharmacophore in other β subunits and (−) α7 reveals important differences likely to influence α-conotoxin selectivity (Fig. 9C). For example, the polar Thr-119 residue present in β3 may explain the absence of α-conotoxins acting at nAChRs containing this subunit.

Conclusions—We have demonstrated the predictiveness of nAChR homology models based on the AChBP structure and characterized a small cleft as the common binding site for 4/7 α-conotoxins. Point mutations in this cleft revealed a major effect of β2-Leu-119 on α-conotoxin affinity and important influences of β2-Phe-117 and β2-Val-109 on α-conotoxin binding kinetics. Furthermore, we could demonstrate a specific interaction of residue 10 in α-conotoxins with the β2 subunit. Docking simulations of PnIA and MII at α3β2 confirmed these results by showing a direct interaction between β2 and α-conotoxin residues. These studies indicate that α-conotoxins block the nAChRs by a lock-and-key interaction, where a primary hydrophobic interaction between β2-Leu-119 and the conserved Pro of 4/7 α-conotoxins "locks" the ligand in its cleft, and secondary complementary interactions contribute to ligand selectivity. Based on homology models of nAChRs, the identification of ligand-accessible residues in the cleft (Leu-119, Val-109, and Phe-117) and the determination of a minimal antagonist pharmacophore, a rational approach to the design of subtype-selective nAChR modulators can now be pursued.

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