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Key words: NMR solution structure, conotoxin, Parkinson’s disease, nicotinic acetylcholine receptor

List of abbreviations: NMR; nuclear magnetic resonance, nAChR; nicotinic acetylcholine receptor, AChBP; acetylcholine binding protein, DA; dopamine, CTx; conotoxin, DQF-COSY; double quantum filtered-correlation spectroscopy, NOESY; nuclear Overhauser effect spectroscopy, TOCSY; total correlation spectroscopy, HSQC; heteronuclear single quantum coherence spectroscopy, LBD; ligand binding domain, MPTP; 1-Methyl-4-phenyl-1,2,3,6-tetrahydropyridine

ABSTRACT

Parkinson’s disease is a debilitating movement disorder characterized by altered levels of α6β2* nAChRs localized on presynaptic striatal catecholaminergic neurons. α-Conotoxin MII (α-CTx MII) is a highly useful ligand to probe α6β2 nAChRs structure and function, but it does not discriminate among closely related α6 nAChR subtypes. Modification of the α-CTx MII primary sequence led to the identification of α-CTx MII[E11A], an analog with 500-5300 fold discrimination between α6* subtypes found in both human and non-human primates. α-CTx MII[E11A] binds most strongly (femtomolar dissociation constant) to the high affinity α6* nAChR, a subtype that is selectively lost in Parkinson’s disease. Here we present the three-dimensional solution structure for α-CTx MII[E11A] as determined by two-dimensional 1H NMR spectroscopy to 0.13 +/- 0.09 Å backbone and 0.45 +/- 0.08 Å heavy atom root mean square deviation from mean structure. Structural comparisons suggest that the increased hydrophobic area of α-CTx MII[E11A] relative to other members of the α-CTx family may be responsible for its exceptionally high affinity for α6α4β2* nAChR as well as discrimination between α6β2 and α3β2 containing nAChRs. This finding may enable the rational design of novel peptide analogs that demonstrate enhanced specificity for α6* nAChR subunit interfaces and provide a means to better understand nAChR structural determinants that modulate brain dopamine levels and the pathophysiology of Parkinson’s disease.

INTRODUCTION

Nicotinic acetylcholine receptors (nAChRs) are a diverse family of ligand-gated cation channels found primarily in neuronal and muscle cells in humans. Structurally, each nAChR exists as either a homopentamer of α subunits or a heteropentamer of α and β proteins. nAChRs can be formed by any grouping of α2-α6 subunits together with β2-β4 subunits; leading to a large assortment of different receptor subtype combinations. The α6 containing nAChRs are...
primarily expressed at the presynaptic endings of catecholaminergic neurons in the brain, where their activity modulates the release of dopamine (DA) and norepinephrine.\textsuperscript{4,8} 

\(\alpha\)-CTx MII, isolated from the venom of \textit{Conus magus}, is a potent antagonist of neuronal nAChRs.\textsuperscript{9} \(\alpha\)-CTx MII is a 16 amino acid peptide with the sequence GCCSNPVCHLEHSNLC containing disulfide bonds connecting Cys\textsuperscript{8} to Cys\textsuperscript{8} and Cys\textsuperscript{3} to Cys\textsuperscript{16}. \(\alpha\)-CTx MII binds to the ligand binding domain (LBD) at the interface of the \(\alpha_2\) and \(\beta_2\) subunits of nAChRs and inhibits the receptor binding to acetylcholine (ACh) with an IC\textsubscript{50} of 0.5 nM.\textsuperscript{9,10} At the time of its discovery in 1997, it was thought that the \(\alpha_2\beta_2\) subunit combination in nAChRs of presynaptic neurons in the brain was responsible for regulation of dopamine release into the synapse and transmission of cholinergic responses. More recently, it has been determined that nicotinic drugs acting on the closely related \(\alpha_2\beta_2\) nAChRs can alter dopamine release by DA neurons in the mid-brain and striatum.\textsuperscript{11} The reporting of \(\alpha\)-CTx MII selectivity for \(\alpha\) and \(\beta\) subunits has led to the discovery of an additional five native \(\alpha\)-CTxs that bind both \(\alpha_2\beta_2\) and \(\alpha_2\beta_2\) nAChRs with variable potency and selectivity. \(\alpha\)-CTx MII and \(\alpha\)-CTx GIC bind to and inhibit \(\alpha_2\beta_2\) and \(\alpha_6\beta_2\) with approximately equal preference,\textsuperscript{9,12} while \(\alpha\)-CTx OmlA shows \textasciitilde 20 fold preference for \(\alpha_2\beta_2\) over \(\alpha_2\beta_2\).\textsuperscript{13} \(\alpha\)-CTx BulA and \(\alpha\)-CTx PIA show a \textasciitilde 20 fold and \textasciitilde 75 fold respective preference for \(\alpha_2\beta_2\) over \(\alpha_2\beta_2\).\textsuperscript{14,15} Each of these \(\alpha\)-CTxs is an ACh antagonist with an IC\textsubscript{50} of \textasciitilde 0.5-1 nM (with the exception of OmlA, IC\textsubscript{50} \textasciitilde 11 nM). The single-site mutant, \(\alpha\)-CTx MII[E11A], is a highly selective ligand for a \(\alpha_2\alpha_2\beta_2\#\) subtype of \(\alpha_6\#\) nAChRs. In monkeys, \(\alpha\)-CTx MII[E11A] binds putative \(\alpha_6\alpha_6\beta_2\#\) nAChRs with an affinity of 9.7 fM and \(\alpha_6\#(\alpha_2-\alpha_6))\beta_2\#\) nAChRs with an affinity of 51 pM. Thus, \(\alpha\)-CTx MII[E11A] discriminates between these two closely related subtypes by 5300-fold.\textsuperscript{16}

Treatment with the dopaminergic neurotoxin 1-Methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP) creates irreversible damage that is widely used to mimic Parkinson’s disease. MPTP treatment of rodents and monkeys leads to selective loss of the nigrostriatal DA fibers whose action (DA release) is modulated by dopamine release into the synapse and transmission of cholinergic responses. Importantly, MPTP treatment of rodents and monkeys leads to selective loss of the \(\alpha_0\alpha_0\beta_2\#\) nAChRs in the central nervous system as identified by a decrease in bound \(\alpha\)-CTx MII[E11A] in brain slice studies.\textsuperscript{16,17} Importantly, \(\alpha_0\alpha_0\beta_2\#\) nAChRs are also selectively lost in human Parkinson’s disease.\textsuperscript{18} Furthermore, striatal DA-ergic fiber cells have been shown to be heterogeneous based on nAChR subtype expression, containing two populations of DA-ergic fibers: low action potential threshold fibers versus high action potential threshold fibers for the release of DA. \(\alpha\)-CTx MII[E11A] binds to the low threshold fibers whose action (DA release) is modulated by \(\alpha_6\#\) nAChRs.\textsuperscript{16}

Here we present the solution structure of \(\alpha\)-CTx MII[E11A] and a structural comparison of \(\alpha\)-CTx MII[E11A] to other \(\alpha\)-CTxs that bind neuronal nAChRs. A structure to activity hypothesis is proposed based on primary sequence comparisons, mutagenesis data, and three dimensional (3D) structure features that are thought to contribute to \(\alpha\)-CTx binding potency and specificity.

**MATERIALS AND METHODS**

\textit{NMR Spectroscopy}. An NMR sample of \(\alpha\)-CTx MII[E11A] was initially prepared at a concentration of 4.2 mM in 90\% H\textsubscript{2}O and 10\% D\textsubscript{2}O (\%/\%). Properly folded synthetic peptide was generously donated by the laboratory of Dr. J. Michael McIntosh, University of Utah. High resolution 1D \(^1\)H, DQF-COSY, TOCSY and NOESY spectra were acquired at 11.75 T on a Varian Inova spectrometer at a temperature of 15°C. Time course deuterium exchange experiments were performed on a freeze dried sample that was re-suspended in deuterium oxide followed by data reacquisition in 100\% D\textsubscript{2}O. Subsequent high resolution 1D \(^1\)H, DQF-COSY, TOCSY, NOESY, \(^1\)H/\(^1\)C HSQC, and \(^1\)H/\(^1\)N HSQC NMR spectra were performed on 0.8 mM \(\alpha\)-CTx MII[E11A] in 95\% H\textsubscript{2}O and 5\% D\textsubscript{2}O at a spectrometer field strength of 14.1 T in a Bruker Avance III spectrometer utilizing a TCI cryogenically cooled probe. Two-dimensional \(^1\)H NMR experiments and interpretation of spectra were based on established methods.\textsuperscript{19,20} Proton DQF-COSY,\textsuperscript{21} NOESY,\textsuperscript{22} and TOCSY\textsuperscript{23} spectra were acquired with the transmitter set to 4.76 ppm and a spectral window of 4500 Hz. Spectra in H\textsubscript{2}O were acquired with the transmitter set at 4.8 ppm and a spectral window of 6500 Hz. Transmitter presaturation was applied at the solvent frequency to suppress the H\textsubscript{2}O resonance. A series of NOESY spectra were acquired with mixing times of 150, 250, and 350 ms. TOCSY spectra were acquired with mixing times of 80 ms. Slowly exchanging NH protons were detected by acquiring a series of 1D spectra of the fully protonated peptide immediately following dissolution in 100\% D\textsubscript{2}O. Spectra were processed using Bruker TopSpin (v. 2.1), MestReC, or Varian VNMR software.

\textit{Restraint Set Generation}. The \(^3\)J\textsubscript{N-H} values for Cys\textsuperscript{8}, Ser\textsuperscript{4}, Leu\textsuperscript{10}, Ala\textsuperscript{11}, His\textsuperscript{12} and Cys\textsuperscript{16} were extracted from a high resolution 1D \(^1\)H NMR spectrum. Backbone \textit{phi} dihedral angle restraints were set to \(-120^\circ +/- 40^\circ\) for \(^3\)J\textsubscript{N-H} greater than 8.0 Hz and to \(-65^\circ +/- 25^\circ\) for \(^3\)J\textsubscript{N-H} less than 5 Hz. Inter- and intraprotein distance ranges were derived from

\textsuperscript{2} Turner,Eidemiller,Martin,Narver,Marshall,Zemp,Cornell,McIntosh,McDougall in \textit{Bioorganic & Medicinal Chemistry} (2009)
NOESY spectra recorded at 15°C. A visual analysis of peak intensity was used to assign NOE intensity as strong, medium, or weak. As a distance reference, the intensity of the geminal protons was considered strong and the distance constraint was set to 1.8 Å. Restraints were set to 1.8-2.7, 1.8-3.3, and 1.8-5.0 Å for strong, medium, and weak intensity signals in the NOESY spectra. Only cross peaks that indicated reasonable correlation between the distance constraint was set to 1.8 Å. Restraints were set to 1.8-2.7, 1.8-3.3, and 1.8-5.0 Å.

Structure Calculations. One thousand initial structures generated from random starting conformations were input into the CYANA (V. 2.1) software. The distance geometry and gradient minimization calculations were performed to find the conformations which best modeled the 91 upper and 17 lower distance limits, 6 phi angle, 4 hydrogen bonds, and 2 disulfide bonds consistent with the NMR data and the molecular backbone, respectively. A pseudo atom correction of 0.5 Å was added to the upper bounds of restraints for methyl and methylene groups. The final set of 20 structures demonstrated an overall root-mean-square deviation among backbone atoms of 0.13 +/- 0.09 Å.

RESULTS
Assignment of Resonances. The complete 1H resonance assignment for α-CTx MII[E11A] was achieved using the method of Wüthrich. A combination of DQF-COSY, NOESY, and TOCSY spectra acquired at 15°C in 95% H2O and 5% D2O were used to eliminate any ambiguities in assignment. 1H NMR spectra were observed with good chemical shift dispersion in the amide region ranging from 9.18 to 7.45 ppm. Fourteen of the 16 amino acid spin systems for α-CTx MII[E11A] were identified in the “fingerprint” region of a 80 ms mixing time TOCSY spectrum. Pro lacks a resonance in the “fingerprint” region of the spectrum and was identified later. Gly was never assigned in the TOCSY spectrum. All spin systems with the exception of Gly and Pro were confirmed in the “fingerprint” region of a DQF-COSY spectrum and assigned to Cys, Cys, Ser, Asn, Val, Cys, His, Leu, Ala, His, Ser, Asn, Leu and Cys. Verification of α- and β-proton resonances and the location of the Pro spin system were extracted from TOCSY and DQF-COSY spectra acquired in 100% D2O. A complete and thorough mapping of all amino acid spin systems was accomplished in this manner.

The assignment of spin systems to amino acids in the primary sequence of the peptide went as follows. The methyl region between 0.79 and 1.06 ppm in the TOCSY spectrum acquired in 95% H2O provided information needed to identify Val, Leu and Leu. Diagnostic resonances were also seen for His, His and Pro. The imidizole ring protons of His and His were found in the TOCSY and 1H/13C HSQC spectra between 6.99 and 8.71 ppm. The spin system assigned to Pro originated at 4.26 ppm in the TOCSY spectrum. The δ-protons at 4.07 and 3.93 ppm were helpful for assigning resonances for Pro because its α-proton at 4.26 ppm was within 0.01 ppm of the α-protons of His and Ala. A complete trace of the Pro spin system was the only way to distinguish the amino acid from those with overlapping spin systems. The remaining amino acids, Cys, Cys, Ser, Asn, Cys, His, Ser, Ser, Asn, and Cys all share the same AMX pattern and were sequentially assigned from the 250 ms mixing time NOESY spectrum by way of a “NOESY walk”.

Sequential Assignments. The NOESY data were sufficient to observe proton-proton NOE’s between the amide proton of one residue and the α-proton of proceeding residue for Cys, Ser, Asn, Cys, His, Leu, Ala, His, Ser, Ser, Asn, Leu and Cys. These residues were assigned based on sequential information combined with Hβ to HN+1 cross peaks from the 250 ms NOESY spectrum.

Off-diagonal resonances were observed for the amide and α-protons of Cys at 8.89 ppm and 4.52 ppm to the β-protons of Cys at 3.24 and 2.84 ppm. There was a weak NOE interaction between the amide proton of Cys at 8.89 ppm and the β-proton of Cys at 3.29 ppm that helped to identify Cys and verify the assignment of Cys. Likewise, a weak NOE was observed between the amide proton of Cys at 8.69 and the β-protons of Cys at 2.63 and 2.23 ppm. A summary of restraints entered into CYANA and the results of the molecular dynamics calculations are provided in Table 1.

-Table 1-

DISCUSSION
Structural description of α-CTx MII[E11A]: α-CTx MII[E11A] folds into a highly compact globular structure consisting of a central amphipathic α-helix between residues 6-13 that is flanked by β-turns at the N- and C-termini. Deuterium exchange studies were used to identify four hydrogen bonds that stabilize the α-helix of α-CTx MII[E11A]. This α-helix is held in place by i to i+4 interactions between the carbonyl oxygen atoms of Pro, Val, and...
Cys\textsuperscript{8}, and His\textsuperscript{9}, and the amide protons on Leu\textsuperscript{10}, Ala\textsuperscript{11}, His\textsuperscript{12}, and Ser\textsuperscript{13}, respectively. At the N-terminus, residues 2-5 are involved in a type II \( \beta \)-turn, stabilized by an \( i \) to \( i+3 \) hydrogen bond between the amide proton of Asn\textsuperscript{5} and the carbonyl oxygen of Cys\textsuperscript{2}. The C-terminal region exhibits a type III \( \beta \)-turn between Ser\textsuperscript{13} and Cys\textsuperscript{16}; this turn may be stabilized by an \( i \) to \( i+4 \) hydrogen bond between the amide proton of Cys\textsuperscript{16} and the carbonyl oxygen of Ser\textsuperscript{13}, although the amide proton of Cys\textsuperscript{16} was not observed to persist for more than 20 minutes following dissolution in D\( _2 \)O. The exterior surface of the \( \alpha \)-helix has a hydrophobic surface created by Pro\textsuperscript{6}, Val\textsuperscript{7}, Leu\textsuperscript{10}, and Ala\textsuperscript{11} on one side, and a hydrophilic face consisting of Cys\textsuperscript{8}, His\textsuperscript{9}, and Ser\textsuperscript{13} on the opposite side of the helix. Since the \( \alpha \)-helix has been determined to be significant for peptide activity, the potential for the cis isomer of the proline (Pro\textsuperscript{6}) that initiates the helix was explored. The presence of two strong intensity NOE cross peaks in the 350 ms mixing time NOESY spectrum is indicative of the proximity of the Asn\textsuperscript{5} \( \alpha \)H to Pro\textsuperscript{6} \( \delta \)H's, and is consistent with a structure containing the trans isomer of the Pro\textsuperscript{6} residue. Additional inspection of the 80 ms TOCSY spectrum indicated a single spin system for Pro\textsuperscript{6} originating at 4.26 ppm; again consistent with the trans isomer and the absence of the cis isomer in the overall structure. Surprisingly, the hydrophobic residues of the \( \alpha \)-helix are more exposed to the solvent than the charged/hydrophilic residues, with the exception of His\textsuperscript{12}. This residue, which has been shown to be very significant for biological activity, is oriented toward the surface of the molecule forming its own prominent electropositive surface feature.\textsuperscript{28}

**Structural Comparison between \( \alpha \)-CTx MII[E11A] and wildtype \( \alpha \)-CTx MII:** As expected, the structure of \( \alpha \)-CTx MII[E11A] is very similar to that of native \( \alpha \)-CTx MII (Figure 1). However, there are desirable differences in the electrostatic topography of the peptide that may explain the observed enhancement in binding selectivity for the \( \alpha \beta_2 \) subunit interface as compared to \( \alpha \beta_3 \). As reported by both Shon \textit{et al.} and Hill \textit{et al.}, the \( \alpha \)-helix of \( \alpha \)-CTx MII originates at Pro\textsuperscript{6} and terminates at His\textsuperscript{12}, while in \( \alpha \)-CTx MII[E11A] the \( \alpha \)-helix is observed to extend to Ser\textsuperscript{13}.\textsuperscript{29,30} The increased length of the helix may be attributed to the single point mutation at Glu\textsuperscript{11}. This observation is consistent with the increase in \( \alpha \)-helical content as measured by circular dichroism (CD) reported by Everhart \textit{et al.}\textsuperscript{31} According to their analysis, \( \alpha \)-CTx MII consists of 31.2\%\pm0.5\% \( \alpha \)-helix versus 44.9\%\pm1.3\% for \( \alpha \)-CTx MII[E11A]. The change in \( \alpha \)-helix does not result in a dramatic change in the overall three dimensional (3D) structure, and a change in the backbone helix itself would not be expected to be the ultimate reason for the observed changes in specificity between \( \alpha \)-CTx MII and the [E11A] analog.

**Figure 1**

There appear to be differences in the nature of the C-terminal \( \beta \)-loop between Ser\textsuperscript{13} and Cys\textsuperscript{16} of \( \alpha \)-CTx MII and \( \alpha \)-CTx MII[E11A]. The C-terminal turn of \( \alpha \)-CTx MII is described as a distorted \( \beta \text{_{10}} \)-helix, whereas the C-terminal portion of \( \alpha \)-CTx MII[E11A] more closely resembles a type III \( \beta \)-turn.\textsuperscript{30} There are also differences in the orientation of several amino acid side chains. This is probably due to the relatively few NOE constraints observed in this region of the peptide, and does not significantly affect the results of this study. The helical portion of \( \alpha \text{h7} \) conotoxins (named according to the number of amino acids in each disulfide loop) is generally accepted to be functionally critical in binding.\textsuperscript{12,15}

In \( \alpha \)-CTx MII there is a hydrophobic patch consisting of residues Pro\textsuperscript{6}, Val\textsuperscript{7}, and Leu\textsuperscript{10} on the exposed face of the \( \alpha \)-helix. The size of this hydrophobic area is larger in \( \alpha \)-CTx MII[E11A] with the addition of the nonpolar Ala\textsuperscript{11} side chain. The significance of the hydrophobic regions of \( \alpha \)-CTx MII toward binding selectivity was first demonstrated by consecutive substitution of each peptide amino acid with alanine.\textsuperscript{28} In that study the [V7A] and [L10A] mutations showed a relative shift in binding towards \( \alpha \beta_2 \) versus \( \alpha \beta_3 \) nAChRs. In \( \alpha \)-CTx MII[V7A] and \( \alpha \)-CTx MII[L10A] the hydrophobicity of the amphipathic \( \alpha \)-helix was effectively decreased by substitution of the isopropyl or isobutyl side chains, respectively, for a methyl substituent. Conversely, alanine substitution at His\textsuperscript{9} and Asn\textsuperscript{14} increased the hydrophobic character of the \( \alpha \)-helix, which sharpened selectivity of the peptide mutants for \( \alpha \beta_2 \) versus \( \alpha \beta_3 \) nAChRs. The ratios of [H9A] and [N14A] IC\textsubscript{50} for \( \alpha \beta_2 \) to \( \alpha \beta_3 \) were 74.7 and 24.2, respectively, indicating a significant enhancement of binding preference for the \( \alpha \beta_3 \) subunit interface. To put this in perspective, \( \alpha \)-CTx MII[E11A] has an IC\textsubscript{50} \( \alpha \beta_2 \) to IC\textsubscript{50} \( \alpha \beta_3 \) ratio of 54.5, yet a 500-fold discrimination index for human and 5300-fold for monkey \( \alpha \beta_2 \) versus \( \alpha \beta_3 \) nAChRs.\textsuperscript{16}

\( \alpha \)-CTx MII[E11A] has the same secondary structure as the native peptide \textit{albeit} with an extension of the \( \alpha \)-helix, and a distinct difference in electrostatic topography surrounding His\textsuperscript{12}. In previous studies, His\textsuperscript{12} and Asn\textsuperscript{9} were found to be important determinants of \( \alpha \)-CTx MII potency, as was the protonation state of His\textsuperscript{12,28,31} The difference in selectivity of \( \alpha \)-CTx MII and \( \alpha \)-CTx MII[E11A] may be attributed to a change in the position, orientation, or pK\(_a\)
of His$^{12}$. The distance between the Asn$^5$ and His$^{12}$ side chains appears to be greater in α-CTx MIIE11A than α-CTx MII, but caution must be taken in structural evaluation based on side chain atom position, as the orientation of these atoms is dynamic.

**Structural Comparison between α-CTx MIIE11A and other select α-CTxs:** The NMR solution structures for α-CTxs and the X-ray crystal structures that are currently available have shown that despite differences in primary sequence there is a highly conserved secondary structural motif. It appears that α-CTx peptide binding specificity and potency is dependent on the amino acid side chains which the backbone serves to present to the receptor on the conserv ed scaffold. Table 2 shows the sequences for the α-CTxs that demonstrate binding to αβ[3] and αβ[5] containing nAChR. The recently reported novel α-CTx TxIA was also included due to its similarity to other α-CTxs and because of its binding affinity to an acetylcholine binding protein from *Aplysia californica* (Ac-AChBP). The Ac-AChBP serves as a model for nAChRs because it has highly conserved LBD residues, and biochemical, pharmacological, and functional attributes consistent with indigenous nAChRs in the human body.32,33 Co-crystallization of α-CTx TxIA and Ac-AChBP demonstrate that Arg$^{5}$, Asn$^{11}$, and Asn$^{12}$ are key side chain residues on the peptide ligand that stabilize binding to the receptor by electrostatic and hydrogen bonding interactions.32 As well, disulfide stacking between Cys$^{2}$-Cys$^{8}$ of the peptide ligand and adjacent disulfides in the receptor appear to stabilize binding in the AChBP receptor model.32

-Table 2-

Previous conotoxins have been identified that bind both αβ[3] and αβ[5] nAChRs; these include α-CTxs MII, PIA, GIC, BuIA and OmlA. Interestingly, α-CTxs MIIE11A, BuIA and PIA show an increased size of the hydrophobic face of the peptide as compared to α-CTxs MII, OmlA and GIC. The size of the hydrophobic region of the amphipathic α-helix is hypothesized to be important for selective binding of peptide to αβ[3] versus αβ[5] containing nAChRs. Surface representations for these toxins and for the newly determined α-CTx MIIE11A are shown in Figure 2. As mentioned previously, His$^{12}$ and Asn$^{5}$ were found to be important determinants of α-CTx MII potency.31 The electronic and steric character of a histidine is similar to that of an asparag ine at neutral pH, and it is worth noting that histidine and asparagine are interchangeably found in α-CTx (Table 2, Figure 2). Either histidine or asparagine is found in homologous locations to the His$^{12}$ and Asn$^{5}$ positions of α-CTx MIIE11A in each of the α-CTxs displayed in Figure 2, with the exception of BuIA. α-CTx BuIA has an amidated C-terminus which occupies the same site as His$^{12}$ in [E11A], which may serve as an asparagine side chain homolog for binding. Figure 2 also shows that each of the specified α-CTxs has these residues positioned with the same orientation to create a similar electrostatic surface for receptor interaction. The positive and negative surface charges associated with the peptides in addition to the two disulfide bonds (shown in yellow, Figure 2) provide variation to account for the observed binding differences from one peptide to another.

-Figure 2-

The crystal structures currently available to serve as binding models for the α-CTxs, contain ImI, PnIA, and Tx1A bound in the ligand binding domain of the AChBP from *Lymnea stagnalis*, *Bondinus subtilis*, or *Aplysia californica*. These X-ray crystal structures of the AChBP with bound α-CTxs ImI, PnIA, and Tx1A all indicate that the Cys$^2$ to Cys$^8$ cystine bridge of the peptide is involved via disulfide bond stacking with two adjacent cysteine residues in the receptor.32,35 Figure 2 shows that in each of these conotoxins, the Cys$^2$ to Cys$^8$ disulfide bridge lies directly between either an asparagine or a histidine in the same position as Asn$^5$ and His$^{12}$ in α-CTx MII. In addition, these toxins show a prominent and contiguous hydrophobic region of varying size on the right side of the “front view” and the left side of the “back view” of each structure corresponding to hydrophobic residues in the α-helical portion of these peptides. This hydrophobic region is consistent with the observation that hydrophobic interaction is a predominant factor for ligand-binding to nAChRs.36-38 It appears that the hydrophilic residues at positions 5 and 12 (either Asn or His) in combination with the α-helix and disulfide bonds that dictate the peptide scaffold are significant contributors to binding specificity.

Another interesting observation from this study is that the hydrophilic locus occupied by a serine following the second cysteine in the primary sequence of each toxin is ubiquitous. α-CTx MII, OmlA and GIC have relatively large hydrophilic domains that are much larger than those in the other toxins, while the reduced size of this hydrophilic domain to those found in α-CTxs PIA, BuIA and MIIE11A may be responsible for reduced selectivity toward binding to the αβ[5] subunit interface. The back side of the toxins shows a common hydrophilic region on the...
left formed by three hydrophilic (polar or negatively charged) residues (Figure 2). Thus, the presence of such common hydrophobic and hydrophilic domains, although different in their exact sizes, is likely to represent the structural basis for binding selectivity towards $\alpha_3\beta_2$ or $\alpha_6\beta_2$ nAChR subtypes. These structural comparisons of $\alpha$-CTxs suggest that binding affinity may be caused by the hydrophobic region of the peptide while subunit binding selectivity may be attributed to the hydrophilic patches.

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REFERENCES


Figure Captions

1. A structure comparison of α-CTxs MII (top row) and E11A (bottom row). From left to right, an overlay of the 20 backbone atoms for the final 20 structures (for MII, \textit{Shon et al.} structure) with the N- and C-termini labeled, a tube structure with Pro\textsubscript{6} labeled as the point of initiation of the α-helix and residues in positions 11 and 12 labeled, and lastly, the surface of each peptide colored by atom charge where red is negative, blue is positive, and white is neutral for the peptides at a pH of 3.

2. Surface comparison of α-CTxs PIA, MII[E11A], BuIA, MII, GIC, OmIA from top to bottom, respectively. Blue regions represent electropositive surfaces, red regions are electronegative surfaces, and white/faintly colored regions are hydrophobic/nonpolar surfaces. Sulfur atoms of cystine residues are yellow. Left panel: front view of peptide surface. Right panel: backside view of peptide surface.

Table Captions

1. Restraints and structural statistics for the final 20 structures generated by the CYANA software based on NMR derived constraints for α-CTx MII[E11A].

2. Primary sequence comparison of α-CTxs that bind to both α\textsubscript{3}β\textsubscript{2} and α\textsubscript{6}β\textsubscript{2} subtype nAChRs. The scaffold disulfide bond motif common to these conotoxins is shown for PIA. Included are the IC\textsubscript{50} values for α\textsubscript{3}β\textsubscript{2} and α\textsubscript{6}β\textsubscript{2} subtype nAChR and the PDB code for each conotoxin.
<table>
<thead>
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<th></th>
<th>MII</th>
<th></th>
<th>E11A</th>
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<td><img src="image" alt="P6 E11 H12" /></td>
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<td><img src="image" alt="H12" /></td>
<td><img src="image" alt="A11" /></td>
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Figure 1
Figure 2
Table 1

<table>
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<tr>
<th>Parameter</th>
<th>Value</th>
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<tbody>
<tr>
<td>Number of NOE distance constraints</td>
<td>92</td>
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<tr>
<td>Number of intra-residue NOE constraints</td>
<td>16</td>
</tr>
<tr>
<td>Number of sequential NOE constraints</td>
<td>27</td>
</tr>
<tr>
<td>Number of medium range NOE constraints</td>
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<tr>
<td>Number of long range NOE constraints</td>
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<tr>
<td>Dihedral angle restraints</td>
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<tr>
<td>Hydrogen bond constraints</td>
<td>4</td>
</tr>
<tr>
<td>Ramachandran plot regions (%)</td>
<td></td>
</tr>
<tr>
<td>Most favored region</td>
<td>51.9</td>
</tr>
<tr>
<td>Additionally allowed region</td>
<td>40.4</td>
</tr>
<tr>
<td>Generously allowed region</td>
<td>7.7</td>
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<tr>
<td>Disallowed region</td>
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<tr>
<td>RMSDs from the average structure (Å)</td>
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</tr>
<tr>
<td>Backbone atoms</td>
<td>0.13 ± 0.09</td>
</tr>
<tr>
<td>Heavy atoms</td>
<td>0.42 ± 0.08</td>
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Table 2

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<th>IC₅₀ ααβ₂ nM</th>
<th>Ratio IC₅₀ α₁β₂ to ααβ₂</th>
<th>Reference</th>
<th>PDB</th>
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