A new level of conotoxin diversity, a non-native disulfide bond connectivity in α-conotoxin AuIB reduces structural definition but increases biological activity

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Abstract
α-Conotoxin AuIB and a disulfide bond variant of AuIB have been synthesized to determine the role of disulfide bond connectivity on structure and activity. Both of these peptides contain the 15 amino acid sequence GCCSYPPCFATNPDC, with the globular (native) isomer having the disulfide connectivity Cys(2-8 and 3-15) and the ribbon isomer having the disulfide connectivity Cys(2-15 and 3-8). The solution structures of the peptides were determined by NMR spectroscopy, and their ability to block the nicotinic acetylcholine receptors on dissociated neurons of the rat parasympathetic ganglia was examined. The ribbon disulfide isomer, although having a less well defined structure, is surprisingly found to have approximately 10 times greater potency than the native peptide. To our knowledge this is the first demonstration of a non-native disulfide bond isomer of a conotoxin exhibiting greater biological activity than the native isomer.

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A New Level of Conotoxin Diversity, a Non-native Disulfide Bond Connectivity in α-Conotoxin AuIB Reduces Structural Definition but Increases Biological Activity*

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α-Conotoxin AuIB and a disulfide bond variant of AuIB have been synthesized to determine the role of disulfide bond connectivity on structure and activity. Both of these peptides contain the 15 amino acid sequence GCCSYPFCDPNC, with the globular (native) isomer having the disulfide connectivity Cys(2–8 and 3–15) and the ribbon isomer having the disulfide connectivity Cys(2–15 and 3–8). The solution structures of the peptides were determined by NMR spectroscopy, and their ability to block the nicotinic acetylcholine receptors on dissociated neurons of the rat parasympathetic ganglia was examined. The ribbon disulfide isomer, although having a less well defined structure, is surprisingly found to have approximately 10 times greater potency than the native peptide. To our knowledge this is the first demonstration of a non-native disulfide bond isomer of a conotoxin exhibiting greater biological activity than the native isomer.

The main component of the venom of cone snails, carnivorous mollusks from tropical marine environments, is a group of disulfide-rich peptides that act as potent and highly specific antagonists of ion channels and receptors (1–3). These channels and receptors play vital physiological roles, and envenomation results in rapid immobilization of prey. One family of these peptides is the α-conotoxins, which target nicotinic acetylcholine receptors (nAChRs) in skeletal muscle and neurons (4–6). These peptides are just 12–18 amino acids in length and feature four conserved cysteine residues.

As is depicted in Fig. 1, the four cysteine residues may, in principle, be joined to form disulfide bonds in three different ways, resulting in formation of “globular,” “ribbon,” and “beads” disulfide bond isomers. In nature the α-conotoxins are present exclusively in the globular form, raising the following question: “What are the advantages of this linkage pattern compared with the two other possibilities?” Their relative ease of synthesis, the small number of possible disulfide bond isomers, and their structural definition (7–17) make α-conotoxins ideal model peptides for the study of the effects of disulfide bonding on protein structure, which in turn may lead to answers to this question.

The cysteine framework (i.e. the placement and spacing of cysteine residues within the peptide sequence) of the α-conotoxins is highly conserved and correlates with their functional and structural features. The first and second cysteines in α-conotoxins are always adjacent to one another, but the number of residues between the second and third cysteines and between the third and fourth cysteines varies (Fig. 1). Based on the intracysteine loop spacing the toxins are divided into four framework classes, with loop spacings of 3/5, 4/3, 4/6 or 4/7. All of the 3/5 α-conotoxins antagonize muscle nAChRs, whereas the 4X classes, with the exception of EI, are active against the neuronal nAChRs (Table I). The difference in the functions of the classes is reflected in distinct differences between their sequences and three-dimensional structures. Whereas all of the α-conotoxins feature a central helix flanked by turns, the helix is 3½ in 3/5 α-conotoxins and takes an α configuration in members of the 4X classes.

The 3/5 α-conotoxin GI and derivatives thereof have been used recently to examine the cooperative formation of disulfide bonds (18). GI, and related 3/5 α-conotoxins GI2 and MI, have been used in studies concerned with determining the effect of changes to the number or connectivity of disulfide bonds on peptide function (19–21). Gehrmann et al. (22) determined the NMR solution structures of the three possible disulfide bond isomers of GI and found considerable structural variations among them. Whereas the globular form exhibited a single well defined conformation in solution, the ribbon and beads isomers displayed an increasing degree of conformational heterogeneity. The connectivity of the disulfides in the beads isomer so dramatically reduced the stability of the structure that a single superposition of the ensemble of NMR structures was not possible.

Zhang and Snyder (23) have shown that the spacings of the cysteines can affect the preferred disulfide connectivity of α-conotoxins. In their study, they oxidized GI and various mutants of GI, including Pro-5→Ala GI mutants with or without residues Cys-7 and Ala-6 swapped. Oxidation of the GI analogues, in both denaturing and folding solvents, and the sub-

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The atomic coordinates and structure factors (code 1MXN and 1MXP) have been deposited in the Protein Data Bank, Research Collaboratory for Structural Bioinformatics, Rutgers University, New Brunswick, NJ (http://www.rcsb.org/).

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The abbreviations used are: nAChR, nicotinic acetylcholine receptor; NMR, nuclear magnetic resonance; HPLC, high performance liquid chromatography; MeNBz, meryl benzyl; MS, mass spectrometry; DQF-COSY, double quantum filtered correlation spectroscopy; ECOSSY, exclusive correlation spectroscopy; TOCSY, total correlation spectroscopy; NOESY, nuclear Overhauser spectroscopy; NOE, nuclear Overhauser effect; RMSD, root mean square deviation; ACh, acetylcholine; HF, hydrogen fluoride; ppb, parts per billion.
sequent analysis of the resulting isomers by high performance liquid chromatography (HPLC) revealed that shortening the first loop of the framework from three to two non-cysteine residues resulted in a reversal of the preferences for different isomers. Mutation of the proline to an alanine did not affect the preferences. They concluded that an even number of residues in either one is preferred over an odd number of residues for loop sizes of fewer than five residues. This conclusion is consistent with earlier studies in which they examined the formation of a different framework classes because of the significant effects of disulfide bonding on structure in /H9251

In general, studies of disulfide bonding in /alpha/-conotoxins have been confined to the 3/5 class. It was of interest to examine the effects of disulfide bonding on structure in /alpha/-conotoxins from different framework classes because of the significant effects of cysteine spacing on disulfide bonding preferences seen in model systems and the differences in structure and function between the various /alpha/-conotoxin classes. Such studies are given further impetus by the fact that new families of conotoxins with alternative disulfide connectivities are now being found. For example a new family of conotoxins that displays the ribon disulfide connectivity as their native form has been isolated recently (25–27). These molecules, referred to as /gamma/- and /lambda/-conotoxins, are of similar size but exhibit different biological activity, primary sequences, and cysteine spacings to the /alpha/-conotoxins (Table 1).

The observed preference for globular disulfide bond connectivity over the ribbon and beads forms in 3/5 /alpha/-conotoxins is consistent with the rules established by Zhang and Snyder (23). By applying the same rules to the 4/7 /alpha/-conotoxins, it is clear that the globular and ribbon forms would be preferred over the beads isomer. A disulfide bond between adjacent cysteines is energetically unfavorable. The reason for the absolute preference for the globular connectivity over the ribbon connectivity in 4/7 /alpha/-conotoxins is not obvious from analysis of cysteine spacings. As is shown in Table II, small conotoxins with 4/7 spacings may exhibit either globular or ribbon disulfide bond connectivities in their native forms.

In this paper we report the chemical synthesis and NMR solution structures of the globular (native) and ribbon disulfide bond isomers of a 4/6 /alpha/-conotoxin, AuIB (28). We also report the finding that the AuIB ribbon isomer has 10-fold greater activity in dissociated neurons of rat parasympathetic ganglia than native AuIB, and we consider this in light of the structural data. Although some studies have looked at the function of /alpha/-conotoxins with altered numbers or arrangements of disulfide bonds, none of these studies included members of the 4/6 or 4/7 classes (19–21). In those classes the reason for the preference for globular connectivity over ribbon connectivity is unclear. In addition, although activity of some /alpha/-conotoxins has been increased through mutation of specific residues, there have been no previous reports of an increase in biological activity as a result of altering disulfide bond connectivity from native.

Experimental Procedures

Peptide Synthesis—The peptides were assembled on Boc-Phe-OCH2Cl/Butt-Pam resin using standard BOC chemistry. A linker, 4-(α-$\text{L}$-tert-butyloxy carbonyl)amino benzoylethyl) phenoxyacetic acid, was attached to the resin to form the C-terminal amide. The amino acids Boc-Ala, Asn(Xan), Asp(O-$\text{B}$-Hex), Cys(methylbenzyl (Mebzl)), Gly, Phe, Pro, Ser(Bzl), Thr(Bzl), and Tyr(BrZ) were obtained from Auspep (Melbourne, Australia). All reagents were of synthesis or HPLC grade.

AuIB globular and ribbon isomers were synthesized using MeBzl protecting groups on the cysteines. The assembled peptides were cleaved from the resin using liquid hydrogen fluoride (HF)/(p-cresol)/p-toluenesulfonic acid (18:1:1) at 0–2 °C for 2 h. The crude peptides were purified on a Vydac C18 column, with a gradient of 1% B per min (A = 0.1% trifluoroacetic acid in H2O, B = 0.09% trifluoroacetic acid in 90% acetonitrile/H2O). Pure peptide, assayed by analytical C18 HPLC and electrospray mass spectrometry (MS), was prepared. The MeBzl protecting groups were removed in the HF cleavage reaction. The globular and ribbon isomers were oxidized by oxidation of the fully reduced peptide at a concentration of 0.08 mg/ml in 20% isopropanol in 0.1 M NH4HCO3 (pH 8.0). The oxidized peptides were purified by preparative C18 HPLC.

NMR—Samples for NMR experiments contained 2–0.5 mM peptide in either H2O or 90% H2O/10% H2O2 at pH 5.5. The pH values are meter readings at 295 K that were uncorrected for deuterium isotope effects. NMR experiments were carried out on Bruker 500 and 750 MHz NMR spectrometers. These included double quantum-filtered correlation spectroscopy (DQF-COSY) (29), exclusive correlation spectroscopy (EXOSY) (30), total correlation spectroscopy (TOCSY) with a mixing time of 80 ms (31), and nuclear Overhauser effect spectroscopy (NOESY) with mixing times of 150, 300, 350, and 400 ms (32–35). Spectra were routinely acquired at 280 and 290 K. They were internally referenced to sodium 2,2-dimethyl-2-silapentane-5-sulfonate at 290 K after acquisition of all other experiments.

Identification of slowly exchanging amide protons was achieved by acquisition of one-dimensional and TOCSY spectra immediately upon dissolution of fully protonated peptide in H2O. Additional information for the backbone NH resonances was acquired from one-dimensional spectra in 90% H2O, 10% 2H2O recorded at 280, 285, 290, 295, and 300 K for both the globular and ribbon forms and also at 297.5, 302.5 and 305 K for the globular form. This information was used to determine NH temperature coefficients. Additional spectra were acquired using samples of globular AuIB at pH 6.1.6, 4.2, and 6.8.

Spectra were processed on a Silicon Graphics Indigo Workstation using XWIN-NMR (Bruker).

Structure Determination—Nuclear Overhauser effect (NOE) restraints were derived from the intensity of cross-peaks in 150- or 280-ms NOESY spectra in 90% H2O, 10% 2H2O recorded at 280, 285, 290, 295, and 300 K for both the globular and ribbon forms and also at 297.5, 302.5 and 305 K for the globular form. This information was used to determine NH temperature coefficients. Additional spectra were acquired using samples of globular AuIB at pH 6.1.6, 4.2, and 6.8.

Spectra were processed on a Silicon Graphics Indigo Workstation using XWIN-NMR (Bruker).

Structure Determination—Nuclear Overhauser effect (NOE) restraints were derived from the intensity of cross-peaks in 150- or 280-ms NOESY spectra. The restraints were derived from 280 and 290 K experiments for the ribbon and native isomers, respectively. They were classified as either strong (1.8–2.7 Å), medium (1.8–3.5 Å), weak (1.8–5.0 Å), or very weak (1.8–6.0 Å) (34, 35). Pseudo-atom corrections of 1.5 Å for methyl, 1 Å for methylene, and 2.0 Å for tyrosine ring protons were added (36).

JHNH1 coupling constants were measured from DQF-COSY spectra. Backbone dihedral restraints of 65–15° were applied for 3 ≤ $\text{JHNH1}$ ≤ 5.8 Hz and 120–30° for 8 ≥ $\text{JHNH1}$ ≥ 5.9 Hz. $\text{JHNH1}$ and $\text{JHNH2}$ coupling constants were measured from TOCSY spectra of the peptide in H2O. Consideration of the coupling constants and NOE intensities allowed inclusion of χ1 angle restraints and stereo-specific assignment of β carbon protons. χ1 angles restraints of 180 = 30° for a “+” conformation around the Co-Cδ bond (NH-Hδ2 NOE = strong/medium, NH-Hj1 NOE = strong, Hα-Hj2 NOE = strong, Hα-Hj1 NOE = weak, $\text{JHNH2}$ < 5 Hz, and $\text{JHNH1}$ > 10 Hz) and χ1 angles of −60–30° for a “−” conformation around the Co-Cδ bond (NH-Hj2 NOE = strong, NH-Hj1 NOE = weak, Hα-Hj2 NOE = weak, Hα-Hj1 NOE = strong, $\text{JHNH2}$ < 5 Hz, and $\text{JHNH1}$ > 10 Hz) were included as restraints.

Three-dimensional structures were calculated using simulated annealing and energy minimization protocols in X-PLOR (37, 38) as described previously (39). NOE distance and dihedral angle restraints for the globular and ribbon isomers were used in X-PLOR version 3.851 to

**FIG. 1.** α-Conotoxin framework. The number of residues between the second and the third cysteines (referred to as loop 1) and between the third and fourth cysteines (referred to as loop 2) are represented by “m” and “n,” respectively. α-Conotoxin framework classes are defined by m and n, with the 3/5 class, for example, having three residues in loop 1 and five residues in loop 2, exemplified by the α-conotoxin GL. The three theoretically possible disulfide bond isomers are illustrated.
Table II

| Name | Sequence | Receptora | Dibb | m/n | Ref.
|------|----------|-----------|------|-----|------
| GI   | ECC-NPACORHYS-Ca | M(a1)αβγδ | G | 3/5 | α | 53 |
| EI   | DDCCYHPTMKNSPSQIC | M(a1)αβγδ | G | 4/7 | α | 54 |
| PnIA | GCCSLPCAPNNDPYC | N α3 β2 | G | 4/7 | α | 55 |
| Epi  | GCCSDPRCMNNPDPYC | N α3 β2/α3 β4 | G | 4/7 | α | 56 |
| MII  | GCCSNPVCYHEHSLNC | N α3 β2 | G | 4/7 | α | 57 |
| AuIB | GCCSTYPCFATPD-C | N α3 β4 | G | 4/6 | α | 58 |
| lml  | GCCSDPRCAWRC | N α7 | G | 4/3 | α | 59 |
| TlxIX | eCCeDGGw-C-CTAAO | ND | G | 4/0 | ε | 60 |
| TIA  | FNWRCCILPAACRNHKKC | α1Adrenoceptor | G | 4/7 | ρ | 61 |
| MrIA | NGVCCGYKLCOH-C | NET | R | 4/2 | γ/λ | 25-26 |

a M, muscle nicotinic acetylcholine receptor; N, neuronal nicotinic acetylcholine receptor; NET, noradrenaline transporter.
b Dib, disulfide connectivity.
c m and n refer to the number of residues between the second and third cysteine and between the third and fourth cysteine.
d F, family.

Table II

Loop sizes of the three possible disulfide bond isomers fo
4 different framework classes of conotoxin

G, R, and B refer to globular, ribbon, and bead disulfide bond connectivities, respectively. In the abbreviation for the framework given in the boxheads, m/n, m refers to the number of residues between the 2nd and 3rd Cys, and n refers to the number of residues between the 3rd and 4th Cys. The framework (m/n) is written in italic type and roman type for α-conotoxin and other small conotoxins, respectively. Loop spacing is abbreviated in the body of the table as x:y where x is the number of residues between the 1st Cys and the Cys it is disulfide-bonded to, and y is the number of residues between the Cys residues that form the other disulfide bond. The loop spacing found in the native peptides is written in boldface type.

Peptide Synthesis—The amino acid sequence of α-conotoxin AuIB was assembled using rapid t-butyloxy carbonyl amino acid/O-benzotriazol-1-yl-N,N,N′,N′-tetramethyluronium hexafluorophosphate/diisopropylethylamine in situ neutralization chemistry (46). An average coupling yield of 99.80% (assessed using a mixture of 10:1 0.05 M NH4HCO3 and isopropyl alcohol) was achieved. Fully reduced peptide was isolated in high purity after HF cleavage and HPLC purification and characterized using MS. The purified peptide was oxidized using a mixture of 10:1 0.05 M NH4HCO3 and isopropyl alcohol to give two isomers, the native (globular) and ribbon isomers, which were purified by HPLC and characterized using MS. The native AuIB isomer was produced in greater yield and was in the range 280–290 mmol/kg. All chemicals used were of analytical grade. ACh chloride and atropine sulfate were supplied by Sigma.

RESULTS

Peptide Synthesis—The amino acid sequence of α-conotoxin AuIB was assembled using rapid t-butyloxy carbonyl amino acid/O-benzotriazol-1-yl-N,N,N′,N′-tetramethyluronium hexafluorophosphate/diisopropylethylamine in situ neutralization chemistry (46). An average coupling yield of 99.80% (assessed using ninyhdrin assay) was achieved. Fully reduced peptide was isolated in high purity after HF cleavage and HPLC purification and characterized using MS. The purified peptide was oxidized using a mixture of 10:1 0.05 M NH4HCO3 and isopropyl alcohol to give two isomers, the native (globular) and ribbon isomers, which were purified by HPLC and characterized using MS. The native AuIB isomer was produced in greater yield than the ribbon isomer under the selected oxidation conditions. Quantities of highly purified peptides were obtained for the structural and functional studies.
**NMR Data**—High quality spectra were obtained for both isomers. The amide signals were spread over the ranges 8.99 to 7.45 and 8.89 to 7.69 ppm for the native and ribbon isomers, respectively. The reasonable dispersion indicates that they adopt ordered structures, with the slightly greater spread of signals for the native form providing a first indication that it adopts the most well defined structure. Sequential assignments were made based on TOCSY, double quantum filtered correlation spectroscopy (DQF-COSY), and nuclear Overhauser effect spectroscopy (NOESY) spectra. Representative spectra showing the assignment of the native and ribbon forms are shown in Fig. 2. Despite the fact that the samples were chemically pure, as judged by MS and HPLC, extra spin systems were present in spectra of the native isomer, suggesting the presence of a second conformation in solution. Because this conformer was of low relative population (<10%), it was not possible to characterize it fully, and subsequent studies focused on the major isomer. The observation of minor conformers in the spectra of small disulfide-rich peptides is not uncommon and may result from cis/trans-isomerization of Pro residues or pro-R/pro-S disulfide bond configurations. Only peaks corresponding to a single isomer were observed in the spectra of the ribbon isomer.

Chemical shifts were examined to identify which residues in the peptides were involved in secondary structure. The differences between observed and random coil α-proton shifts (referred to as secondary shifts) for the native and ribbon isomers are shown in Fig. 3. From the contiguous sequence of negative secondary shifts from residues Pro-6 to Ala-10, it is clear that the native isomer incorporates a helix over this region. In contrast, the ribbon isomer has a less regular pattern of secondary shifts, indicative of less well defined secondary structure. The fact that the two isomers have quite different structures is exemplified by the large difference in α-proton shifts of Tyr-5 in the two structures, presumably resulting from different relative orientations of the aromatic side chain and subsequently different ring current effects.

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**Fig. 2.** Two-dimensional NMR spectra that summarize the sequential assignments for native (a) and ribbon AuIB (b). The lower panels are parts of NOESY spectra of native or ribbon AuIB in 90% H2O, 10% D2O acquired at 290 and 280 K, respectively. The upper panels are parts of TOCSY spectra for the same samples. In the upper panel the different spin systems are identified. In the lower panel the sequential assignment is shown. The abbreviation i/*H11*1 is used to label the NOE that results from interaction between the α proton of residue i and the backbone amide proton of the following residue.

**Fig. 3.** Secondary shifts (top) and temperature coefficients (bottom) for native and ribbon AuIB. Native AuIB is indicated by the black bar, and ribbon AuIB is indicated by the white bar. In the top graph, the differences between the actual shifts of the α protons and their respective random coil shifts are graphed for each residue. A continuous stretch of residues that exhibit negative deviations from random coil indicates helix. A sequence specific correction factor of 0.29 ppm was deducted from residues preceding Pro. In the bottom graph a change in temperature coefficient from that for random coil of less than −3 ppb/K indicates residues that may be solvent-protected.
The variation in amide proton chemical shifts with temperature provides a measure of exposure to solvent, and hence potentially yields information on hydrogen bonding (47, 48). With the exception of Phe-9 all residues from Ser-4 to Cys-15 of the native isomer of AuIB have temperature coefficients less negative than $-4$ parts per billion (ppb/K, a standard measure of amide proton protection from solvent. This was not the case for the ribbon isomer of AuIB where only two residues, Thr-11 and Asn-12, had temperature coefficients less negative than $-4$ ppb/K. As the amide proton temperature coefficients greatly differ for different amino acid residues in random coil peptides, the temperature coefficients for the native and ribbon isomers of AuIB were further considered relative to random coil temperature coefficients. From Ser-4 to Cys-15 in the native AuIB peptide, the temperature coefficients relative to random coil temperature coefficients are smaller than $-3$ ppb/K (Fig. 3) but greater than $-3$ ppb/K for other residues. When the data are considered in this way, it appears that the amide proton of Phe-9 in the native isomer of AuIB and amide proton of Tyr-5 of the ribbon isomer of AuIB are also solvent-protected.

The conclusions drawn from the temperature coefficient data are strongly supported by measurement of the exchange rates of amide protons with solvent D$_2$O. The backbone amide protons of Tyr-5, Cys-8, Phe-9, Ala-10, Thr-11, Asn-12, and Cys-15 were still detected in NMR spectra recorded 4 h after dissolution of the native isomer in D$_2$O. All backbone amide protons had exchanged in the native isomer after 24 h in D$_2$O. No backbone amide protons were detected for the ribbon isomer 5 min after dissolution of the peptide in D$_2$O. This confirms that the native isomer has a substantially more stabilized structure than the ribbon form.

Fig. 4 summarizes NMR data that were used to characterize fully the secondary structures of the native and ribbon AuIB isomers. These include coupling constants that were used to assign dihedral angle restraints, slow exchanging amides that are involved in H-bonds, and a summary of some pertinent NOE restraints. The strong dNN NOEs relative to dαN NOEs in the middle region of the native isomer, for example, suggest the presence of helical structure. The relative intensity of dNN NOEs to dαN NOEs is quite different in the ribbon isomer and is more indicative of an extended structure. Couplings that were used to calculate χ$_1$ and ϕ angles are also indicated in Fig. 4.

**Structure Calculations**—Analysis of the NMR spectra for the native isomer yielded a total of 184 NOE restraints including 32 medium and 18 long range NOE restraints. Seven angle restraints were included with the ϕ angles of Ser-4, Tyr-5, Thr-11, and Asn-12 restrained to $-120 \pm 30^o$, the ϕ angle of Ala-10 restrained to $-65 \pm 15^o$, and the χ1 angles of Asn-12 and Cys-15 restrained to $180 \pm 30^o$. 140 NOE restraints, including 4 long and 13 medium range NOEs, were identified in the NMR spectra of the ribbon isomer. Three dihedral angles were included, with the ϕ angles for Cys-8 and Ala-10 restrained to $-120 \pm 30^o$ and $-65 \pm 15^o$, respectively, and the χ1 angle for Cys-8 restrained to $-60 \pm 30^o$. These restraints were used to calculate three-dimensional structures for the native and ribbon isomers.

The ensemble of native AuIB structures overlays very well (Fig. 5a) with a pairwise backbone RMSD of 0.36 ± 0.13 Å over residues Cys-2 to Cys-15. Table III summarizes the structural and energetic statistics for the calculated structures. The native isomer features an α-helix from Tyr-5 to Asn-12 (Fig. 5c) which is flanked by turns. In 6 of 20 structures, a 3$_{10}$ helical region is present in the N-terminal region of the peptide from Gly-1 to Ser-4. The disulfide bond joining Cys-2 and Cys-8 is present in a right-handed hook conformation in 17 of the 20 structures, but there is no consensus configuration for the disulfide between Cys-3 and Cys-15. The average χ2, χ3, and χ2’ angles for the right-handed hook Cys-2 to Cys-8 disulfide bonds are $-65, 101, and 141^o$ in 13 structures and $-128, 78,$ and $153^o$ in the other four structures. The average χ2, χ3, and χ2’ angles for the non-right-handed hook Cys-2 to Cys-8 disulfide bonds are $53, -62,$ and $57^o$. The structure is stabilized by the presence of hydrogen bonds between the CO groups of Cys-2, Pro-6, Cys-8, Phe-9, and Asn-12 and the NH groups of...
Tyr-5, Ala-10, Thr-11, Asn-12, and Cys-15, respectively. The CO of Tyr-5 forms a bifurcated hydrogen bond with the NH groups of both Cys-8 and Phe-9.

In contrast with the native form, the individual structures of the ribbon isomer of AuIB do not align well over the whole molecule and have a pairwise backbone RMSD of 2.50 ± 0.73 Å (Fig. 5b). The alignment is better over smaller regions within the structure. For example, residues 2–10 overlay with a pairwise backbone RMSD of 1.08 ± 0.38 Å and residues 11–15 overlay with a pairwise backbone RMSD of 1.20 ± 0.50 Å. The global fold is shown in Fig. 5d. Whereas there is an overall similarity in the shape of the fold relative to the native isomer, the ribbon isomer lacks a defined helical region. The region that is helical in the native form is straightened out, and the backbone bends over residues Tyr-5 to Pro-7 and at Ala-10 and Pro-13 in the ribbon isomer. The disulfide bond joining Cys-3 and Cys-8 in the ribbon isomer is present in either a right- or left-handed spiral conformation in the individual structures within the ensemble. The average χ2, χ3, and χ4' angles for the left-handed spiral Cys-2 to Cys-8 disulfide bonds are −114, −83, and −105°. The average χ2, χ3, and χ4' angles for the right-handed spiral Cys-2 to Cys-8 disulfide bonds are 68, 91, and 135°. There is no consensus conformation for the disulfide bond joining Cys-2 and Cys-15 of the ribbon isomer.

The first loops of the ribbon and native isomers overlay well (Fig. 6), with a mean global backbone RMSD of 0.92 ± 0.00 Å over residues Cys-3 to Cys-8. The second loop of the molecule is “looser” in the ribbon isomer relative to the native. Overlays of the ribbon and native AuIB structures with the structures of related α-conotoxins are also shown in Fig. 6.

Ramachandran plots show the likely ψ and φ angle combinations for residues in proteins and are based on the measurement of these angles in 163 known structures. All of the residues in the native isomer (excluding Pro and Gly) lie in the most favored and additionally allowed regions of the Ramachandran plot. 89.5% of the residues in the ribbon isomer (excluding Pro and Gly) were in the most favored and additionally allowed regions, and 9.5% were in the generously allowed regions and 1% in the disallowed regions. The residues that lie in the disallowed regions were not confined to just one or two residues in the ensemble and were distributed over different residues, from Phe-9 to Asn-12 and Cys-2 and Cys-3. A large proportion of Thr-11 residues lay in the generously allowed regions. This is not surprising as Ramachandran statistics are based on angles seen in structures of native proteins and may not necessarily reflect the angles observed in small constrained peptides, particularly those with non-native folds.

Electrophysiology—Rapid focal application of ACh (500 μM) to dissociated neurons of the rat parasympathetic ganglia held at −70 to −80 mV, in the presence of 100 nM atropine, resulted in a characteristic biphasic inward current comprising an initial transient peak that rapidly decayed to a steady-state current (Fig. 7). The ratio of peak to steady-state current amplitude varied between neurons.

Effects of native and ribbon AuIB on nicotinic ACh responses were investigated in 30 neurons. Native AuIB inhibited the ACh-evoked current in a dose-dependent manner in 12 of 14 neurons and ribbon AuIB in 12 of 16 neurons. In the remaining six neurons, the ACh-evoked current was not inhibited in the presence of >1 μM AuIB. This is most likely due to heterogeneity in the expression of the β4 subunit that has been shown using single cell PCR analysis to be expressed in a proportion (>50%) of neurons (49). The half-maximal inhibitory concentration (IC50) value for native AuIB was 1.2 nM and maximal block ranged from 21 to 62% of the total ACh-induced current. The ribbon AuIB isomer had an IC50 of 0.1 nM and caused a maximal block of 31 to 62% of the ACh-evoked current. The inhibition of ACh-evoked currents by the ribbon AuIB isomer was slowly reversible, whereas <50% recovery was obtained after 30 min of washout.

The finding that the ribbon isomer has an activity 10-fold higher than the native was unexpected and to our knowledge

### Table III

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<thead>
<tr>
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<th>AuIB native</th>
<th>AuIB ribbon</th>
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<tr>
<td>Mean RMSD from experimental restraints</td>
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<tr>
<td>NOE (Å)</td>
<td>0.009 ± 0.001</td>
<td>0.025 ± 0.005</td>
</tr>
<tr>
<td>Dihedral (*)</td>
<td>0.244 ± 0.141</td>
<td>0.180 ± 0.279</td>
</tr>
<tr>
<td>Mean RMSD from Idealized covalent geometry</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bonds (Å)</td>
<td>0.0058 ± 0.0003</td>
<td>0.0687 ± 0.0011</td>
</tr>
<tr>
<td>Angles (*)</td>
<td>2.06 ± 0.04</td>
<td>2.24 ± 0.11</td>
</tr>
<tr>
<td>Impropers (*)</td>
<td>0.14 ± 0.02</td>
<td>0.18 ± 0.04</td>
</tr>
<tr>
<td>Restraint violations</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NOE violations &gt;0.1Å</td>
<td>0</td>
<td>2.65 ± 1.63</td>
</tr>
<tr>
<td>Maximum NOE violation (Å)</td>
<td>0.081</td>
<td>0.197</td>
</tr>
<tr>
<td>Dihedral angle violations &gt;1°</td>
<td>0.05 ± 0.22</td>
<td>0.10 ± 0.31</td>
</tr>
<tr>
<td>Maximum angle violation (°)</td>
<td>1.2</td>
<td>1.8</td>
</tr>
<tr>
<td>Mean energies (kJ mol⁻¹)</td>
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<td></td>
</tr>
<tr>
<td>E_NOE</td>
<td>0.49 ± 0.13</td>
<td>2.76 ± 1.08</td>
</tr>
<tr>
<td>E_dih</td>
<td>0.03 ± 0.03</td>
<td>0.00 ± 0.00</td>
</tr>
<tr>
<td>E_abw</td>
<td>−62.23 ± 1.75</td>
<td>−45.16 ± 2.42</td>
</tr>
<tr>
<td>E_bond</td>
<td>1.41 ± 0.10</td>
<td>87 ± 0.51</td>
</tr>
<tr>
<td>E_propers</td>
<td>0.23 ± 0.06</td>
<td>0.35 ± 0.16</td>
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<tr>
<td>E_angle</td>
<td>22.44 ± 0.76</td>
<td>26.52 ± 2.71</td>
</tr>
<tr>
<td>E_total</td>
<td>−37.64 ± 1.91</td>
<td>−13.66 ± 5.05</td>
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<tr>
<td>Pairwise RMSD</td>
<td></td>
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<tr>
<td>Backbone atoms (N,Ca,C) (Å)</td>
<td>(2–15) 0.36 ± 0.13</td>
<td>(2–10) 1.08 ± 0.38</td>
</tr>
<tr>
<td></td>
<td>(1–15) 0.65 ± 0.25</td>
<td>(11–15) 1.20 ± 0.50</td>
</tr>
<tr>
<td></td>
<td>(1–15) 2.50 ± 0.73</td>
<td>(1–15) 2.47 ± 0.87</td>
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<tr>
<td></td>
<td>(1–15) 3.18 ± 0.90</td>
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<tr>
<td>Heavy atoms (Å)</td>
<td>(2–15) 0.95 ± 0.15</td>
<td>(2–10) 1.69 ± 0.51</td>
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<tr>
<td></td>
<td>(1–15) 1.04 ± 0.19</td>
<td>(11–15) 2.47 ± 0.87</td>
</tr>
<tr>
<td></td>
<td>(1–15) 2.18 ± 0.80</td>
<td></td>
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<tr>
<td>Ramachandran Plot</td>
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<tr>
<td>Most favored regions (%)</td>
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<td>56.0</td>
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<tr>
<td>Additional allowed regions (%)</td>
<td>17.0</td>
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<tr>
<td>Generously allowed regions (%)</td>
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<td>9.5</td>
</tr>
<tr>
<td>Disallowed regions (%)</td>
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<td>1.0</td>
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</tbody>
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represents the first observation of an increase in activity being produced by non-native disulfide connectivity in a conotoxin.

**DISCUSSION**

In the current study, we have synthesized and have determined the three-dimensional solution structures of the ribbon and native forms of α-conotoxin AuIB. Native α-conotoxin AuIB features an α-helical region from Tyr-5 to Asn-12 that is flanked by turns, whereas in the ribbon form the helical nature of AuIB is lost. We have also investigated the potency of the native and ribbon AuIB peptides as antagonists of the nAChRs present in rat parasympathetic neurons and have made the surprising finding that the ribbon isomer is approximately 10 times more potent at this site than native AuIB.

The consensus features of the native AuIB structure described here are similar to those in a recently reported structure (17), with the α-helical region spanning the same residues in both. The two structures are virtually identical in the first loop, but there are some minor changes, over residues 12–14, in the second loop. Comparison of the NOE and angle restraints used in calculation of the two AuIB structures identifies some differences that may explain this structural change. The use of 750 MHz NMR data in our study allowed us to include more medium and long range NOE restraints in the structure calculations than the previous study, which used 600 MHz NMR data. In particular, we included 50 medium and long range NOEs, and the other study included 32 (although they stated 38 in the paper there were 32 in the deposited restraints). Twenty of these NOE restraints were identical, and many of the other restraints although not identical were derived from interactions between the same amino acid residues. The Asn-12 χ1 angle was restrained in our structure but not in the Cho et al. (17) structure. In addition, the Cho et al. (17) structure included NOEs between Asn-12 and Cys-15 and between Cys-8 and Thr-11 that were not observed in our spectra. We included NOEs between Phe-9 and Thr-11, Asp-14, and Cys-15 that were not observed by Cho et al. (17). These restraint differences explain the observed structural differences, but the origin of the restraint differences is not clear as similar experimental conditions were used. The major difference was that they used a peptide concentration of 6 mM whereas we used 2 mM.

AuIB is a member of the 4/6 family of α-conotoxins, and because it is the only member of this class for which structural information is available, it was of interest to compare its structure with members of other framework classes. Fig. 6b shows that the first loop of AuIB overlays well with the representative 4/7 α-conotoxin PnIB. Indeed AuIB features the α-helix that is typical of the 4/7 class. The only significant difference between the structure of AuIB and α-conotoxins from the 4/7 class is that the second loop is tighter in AuIB due to the smaller loop size. The 4/6 α-conotoxin AuIB also overlays well with the 4/3 α-conotoxin ImI over the first loop but exhibits large structural
probably not the factor that prevents PnIA from binding to the position 13 rather than a Pro. Tyr-15 in PnIA is therefore an additional residue in the second loop, Tyr-15, and by having a Ser with less potency than AuIB. They differ from AuIB in having an amide bond as the peptides are crystallized.

The general similarity in the backbone fold of the 4/7 family but the second loop is significantly altered (13).

The reduction in structural homogeneity that results from altering the disulfide connectivity from the globular to ribbon form provides an insight into the role of the disulfide bonds in defining the structure. The trend in backbone atom pairwise RMSDs between the native and ribbon forms of AuIB is similar to that seen for the native and ribbon forms of GI. In both cases the ribbon isomer is less precisely defined (i.e., has lower RMSD values) than the respective native peptide. Increases in RMSD values can result either from a lack of sufficient NOE or dihedral angle restraints or from inherent flexibility. Based on the trends in slow exchange amide protons and temperature coefficients, it appears that the latter is the more likely explanation and that the large RMSD for the ribbon isomer of AuIB reflects a high degree of flexibility in this structure.

In the case of GI, the ribbon connectivity resulted in a structure that, although relatively well defined, was different from that of the native structure. The inactivity of the ribbon isomer of GI was attributed to its non-native structure and to only a moderate increase in structural flexibility over the native form (22). That was presumed to prevent it from adopting the fold required for binding to the muscle nAChR. It is interesting to note that the ribbon isomer of AuIB is a potent antagonist of nAChRs in dissociated neurons of rat parasympathetic ganglia. The ability of the AuIB ribbon isomer to bind to the neuronal nAChR may reflect greater flexibility in this peptide, as compared with the GI ribbon isomer, allowing it to adopt a conformation that is capable of binding. The greater flexibility of the ribbon isomer of AuIB is

Fig. 8. Solvent-accessible surfaces of AuIB (a), PnIA (b), MII (c), and Epl (d). Hydrophobic residues are light green; positive residues are blue; negative residues are red; Cys residues are gold; and all other residues are white. Views on the right-hand side are rotated 180° from the views on the left-hand side.
reflected not only in the higher RMSD than GI but also in the fact that it has no slowly exchanging protons at all, whereas Ala-6 is slowly exchanging in the GI ribbon isomer.

A further understanding of the reasons for the activity of the ribbon isomer would be assisted by mutagenesis studies. There are currently no extensive mutagenesis data available in AuIB analogues, but some insight into residues important for binding can be obtained from a comparison of the structure of AuIB and the structural activity relationship data of related α-conotoxins. Residues in PnIB and ImI that interact with the α7 receptor have been identified (50–52). Whereas AuIB binds to the α3β4 receptor rather than the α7 receptor, it is interesting to note that for both PnIB and ImI two of the important residues for binding are on the first loop and one is in the second loop. All of the important residues are in the central region of the peptides, corresponding to the helical region. In the ribbon isomer AuIB the first loop is essentially unchanged from that in the native isomer, whereas the ribbon isomer of AuIB, it is more active than the native isomer. It appears that once bound, the ribbon isomer is able to have a highly complementary fit with the receptor site.

It is interesting that despite the apparent flexibility of the ribbon isomer of AuIB, it is more active than the native isomer. It appears that once bound, the ribbon isomer is able to have a highly complementary fit with the receptor site.

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REFERENCES
