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Novel small molecule inhibitors of botulinum neurotoxin A metalloprotease activity

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Abstract

Botulinum neurotoxins (BoNTs) are among the most lethal biological substances to have been weaponized and are listed as biodefense category A agents. Currently, no small molecule (non-peptidic) therapeutics exist to counter this threat; hence, identifying and developing compounds that inhibit BoNTs is a high priority. In the present study, a high-throughput assay was used to identify small molecules that inhibit the metalloprotease activity of BoNT serotype A light chain (BoNT/A LC). All inhibitors were further verified using a HPLC-based assay. Conformational analyses of these compounds, in conjunction with molecular docking studies, were used to predict structural features that contribute to inhibitor binding and potency. Based on these results, a common pharmacophore for BoNT/A LC inhibitors is proposed. This is the first study to report small molecules (non-peptidics) that inhibit BoNT/A LC metalloprotease activity in the low μ M range.

Keywords: Bioterrorism; Botulinum neurotoxin; Drug discovery; High-throughput screen; Inhibitors; Molecular modeling; Pharmacophore; Three-dimensional database search; Metalloprotease

Botulinum neurotoxins (BoNTs) are produced by spore forming anaerobic bacteria *Clostridium botulinum* and are among the most lethal of biological poisons [1,2]. Seven immunologically distinct BoNT serotypes (designated A-G) have been identified [3]. Accidental exposure to BoNTs, for example through contaminated food, can result in life threatening flaccid paralysis [4]. Furthermore, BoNTs have been weaponized in highly toxic aerosol form, and consequently pose a significant "dual threat"—i.e., both to civilian and military populations [5,6]. As a result, there is an urgent need for therapeutic countermeasures against BoNTs [7].

BoNT is secreted as a holotoxin composed of two peptide chains that are linked by a disulfide bridge [8]. The heavy chain is responsible for: (1) targeting and binding to surface receptors on nerve terminals; (2) translocation into the neuronal cytosol via the formation of a low pH endosome; and (3) protecting the substrate binding cleft of the light chain prior to neuronal internalization [9,10]. The light chain, which dissociates from the heavy chain in the low endosomal pH, is released into the cytosol where it acts as a zinc metalloprotease that cleaves SNARE (soluble NSF-attachment protein receptor) proteins: SNAP-25 (synaptosomal-associated protein of 25 kDa), synaptobrevin, and syntaxin [9,10]. BoNT serotypes A, C, and E cleave SNAP-25 [11]; serotypes B, D, F, and G cleave synaptobrevin [12-15]; and serotype C can also use syntaxin as substrate [16]. Without functional

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SNARE complexes, acetylcholine is not released into neuromuscular junctions, leading to paralysis.

Research to identify peptide and small molecule inhibitors of BoNT serotype A (BoNT/A) has targeted both holotoxin translocation and light chain (BoNT/A LC) metalloprotease activity. Sheridan et al. [17] and Deshpande et al. [18] have shown that a number of antimalarial agents interfere with BoNT/A translocation into nerve cytoplasm. Specifically, it has been shown that several antimalarial compounds act subsequent to toxin binding to cell-surface receptors [17], and it has been hypothesized that these agents inhibit BoNT/A cytosol entry by raising endosomal pH (an endosomal pH of 5.5 or lower is needed for release into the cytoplasm) [17,18]. Hayden et al. [19] have found that BoNT/A LC is inhibited by mM concentrations of known protease inhibitors: captopril, lysinopril, and enalapril. In the same study, it was also reported that a number of short peptides, from specific "hinge" libraries, inhibit BoNT/A LC activity by as much as 51% at concentrations as low as 0.5 µM [19]. Using a chromatographic method, Schmidt et al. [20] identified the peptide motif CRATKML as a potent inhibitor. In a subsequent study, the Cys residue of CRATKML was replaced with thiol containing organic moieties and it was found that a 2-mercapto-3-phenylpropionyl containing derivative was the most effective ($K_i = 0.3 \,\mu\text{M}$) [21].

Presently, no small molecule (non-peptidic) inhibitors of BoNT/A LC metalloprotease activity, which are effective in the low µM range, have been reported. In this report, a high-throughput assay was used to facilitate BoNT/A LC inhibitor identification [22]. Initially, the National Cancer Institute (NCI) Diversity set was screened, and several compounds possessing >50% inhibition (at 20 µM concentration) were identified. Based on molecular modeling studies, common structural features and binding modes in the BoNT/A LC substrate binding cleft were identified for these leads. Adscreening of 7-chloro-4-aminoquinoline derivatives also resulted in the identification of leads that substantially inhibit BoNT/A LC at 20 µM concentrations. Analyses of these compounds indicate that they share common structural/functional group characteristics with inhibitors identified during the screen of the NCI Diversity set, and that they may also bind to the enzyme's substrate binding cleft in a similar manner. Based on our analyses, a common pharmacophore for BoNT/A LC inhibitors is proposed.

Materials and methods

Compounds. The NCI Diversity set was obtained in 96-well plate format from the National Cancer Institute (http://dtp.nci.nih.gov/branches/dscb/diversity_explanation.html). The syntheses of N, N-bis(7-chloroquinolin-4-yl)alkanediamines and N,N-bis(7-chloroquinolin-4-yl)heteroalkanediamines used in these studies have been

described previously [23,24]. Amodiaquine, chloroquine, quinacrine, quinidine, and quinine were obtained from Sigma–Aldrich (USA).

High-throughput assay of BoNT/A LC activity. The high-throughput assay used to screen BoNT/A LC proteolytic activity has been described previously [22]. Briefly, liquid transfers were done with a SciClone automated pipettor (Zymark, Hopkinton, MA). Recombinant BoNT/A LC [25] was diluted to 0.6 µg/ml in 40 mM Hepes, and 0.5 mg/ml BSA, pH 7.4. Test compounds was diluted to 0.2 mM with methyl sulfoxide. Ten microliters of each compound were transferred into the corresponding wells of another 96-well plate containing immobilized fluorescent-labeled peptide substrate, specific for the protease activity of BoNT/A LC [22]. Ninety microliters BoNT/A LC was then added and mixed, for final concentrations of 20 µM compound and 0.5 µg/ml BoNT/A LC. Blanks (buffer only) and controls (buffer and BoNT/A LC), eight wells each, were included on each assay plate. After incubation at 30 °C for 60 min, 70 µl was transferred from each well to an opaque-wall plate and fluorescence was measured in a Wallac 1420 multi-label counter (Perkin-Elmer, Gaithersburg, MD).

HPLC-based assay of BoNT/A LC protease activity. Compounds exhibiting ≥ 40% inhibition in high-throughput assays were retested in a HPLC-based assay as described previously [1,26].

BoNT/A LC refinement and inhibitor docking. The structure of BoNT/A LC was obtained from the deposited crystallographic coordinates of the holotoxin (PDB refcode = 3BTA) [27]. The 3.2 Å holotoxin includes the heavy chain, light chain, and a protective residue belt that covers the substrate binding site of the light chain. The heavy chain and the protective belt were removed from the holotoxin, and the light chain was energy refined using the Discover (Accelrys, San Diego, CA) program's cff91 force field. During the refinement, the zinc ion and residues coordinating the zinc ion were fixed in their original coordinates. Our strategy entailed applying 2000 kcal/mol per Å² of force that was stepped off the structure in 100-kcal/mol decrements by minimizing with conjugate gradients, until the norm of the gradient was 0.01 kcal/Å [28]. This process was repeated until all applied external force was removed. The resulting coordinates of the final model were within the experimentally determined X-ray crystallographic resolution. The optimized structure had an rms deviation of 2.6 Å across all backbone atoms from the reported crystallographic coordinates of the holotoxin light chain.

Using InsightII molecular modeling software (Accelrys, San Diego, CA), identified inhibitors were manually docked into the BoNT/A LC substrate binding cleft, and van der Waals violations of 0.25 Å were removed by small adjustments to side-chain torsion angles and inhibitor positioning. The inhibitor-enzyme structure coordinates were subsequently tether minimized in the same manner as described above and the final structure was subjected to hydropathic analysis using the program HINT (eduSoft, Richmond, VA).

Conformer generation. Conformational models of inhibitors were generated using Catalyst 4.7 software (Accelrys, San Diego, CA). Inhibitors were imported into Catalyst and energy minimized to the closest local minimum using the generalized CHARMM-like force field, as implemented in the program. Following, the "best quality" conformational search option was employed to generate conformers within 20 kcal/mol from the global energy minimum. In this study, it was found that 100 conformers per compound ensured adequate coverage of conformational space.

Results and discussion

Screening of the NCI diversity set

A high-throughput fluorescence-based assay was initially used to screen the NCI diversity set: a collection of 1990 molecules that were selected to cover a wide range

of conformational space, and at the same time provide pharmacophore diversity and structural rigidity. Following the initial high-throughput screen, a HPLC-based assay [1,26] was used to eliminate false positives resulting from fluorescence quenching by some of the compounds. The final set of inhibitors, which were tested at $20\,\mu\text{M}$ concentration in the presence 0.1 mM substrate, consisted of 21 compounds with potencies ranging from 14% to 100% BoNT/A LC inhibition. Two-dimensional structures for compounds possessing >40% inhibition are shown in Table 1.

Additional testing of NSC 625324 (silver sulfadiazine), initially the most potent of the identified compounds, indicated that this molecule's efficacy was entirely mediated by the silver ion: no inhibition was observed when the unionized, organic form of sulfadiazine was tested in the HPLC-based assay. Furthermore, BoNT/A LC activity was examined in the presence of silver acetate, and 100% inhibition of protease activity was found at concentrations $\geqslant 5\,\mu\text{M}$ silver ion (the IC50 was $1.5{-}2.0\,\mu\text{M}$). Cesium and rubidium salts did not inhibit BoNT/A LC protease activity at concentrations

Table 1 NCI diversity set inhibitors of BoNT/A LC

Compound	NSC No.	% Inhibition ^a	
H_2N \longrightarrow $\stackrel{\circ}{\longrightarrow}$ \longrightarrow	NSC 625324 (silver sulfadiazine)	100	
HO OH HO OH	NSC 661755 (michellamine B)	62	
N H NH ₂	NSC 357756	57	
Соон	NSC 119889	56	
N OH OH	NSC 86372	51	
	NSC 130796	48	
ĊH ĊH	NSC 402959	40	

^a Compounds tested in the HPLC-based assay at 20 μM concentration in the presence of 0.1 mM substrate.

up to $20\,\mu M$. The presence of zinc (5–50 μM) in assays had no effect on inhibition by $3\,\mu M$ silver ion, but addition of 1 mM DTT immediately reversed inhibition. These observations suggest that silver neither displaces zinc from the active site, nor causes irreversible denaturation by reacting with a residue important for conformational stability. It is possible that this ion might bind to polar residues in or near the substrate binding cleft, blocking access to the catalytic site.

Inhibitors NSC 86372 and 130796 both contain an 8hydroxyquinoline moiety (Table 1), which is known to chelate metal ions [29]. Based on our analyses, both of these molecules do chelate the BoNT/A LC catalytic zinc. Support for a chelation mechanism is based on: (1) assay data showing that several congeners of NSC 86372 possess equivalent inhibitory potency to that of the original lead molecule; (2) addition of 20 µM zinc immediately and quantitatively reversed inhibition caused by up to 10 µM NSC 86372; and (3) NSC 86372, its congeners, and NSC 130796 inhibit other zinc metalloproteases (including anthrax lethal factor and BoNT serotype B light chain (BoNT/B LC)). Consequently, the 8-hydroxyquinoline motif is not suitable for further development as a component of BoNT/A LC inhibitors; however, as will be shown, structural similarities between NSC 86372 and other identified compounds do appear to fit a general model for BoNT/A LC inhibition.

Screening of 7-chloro-4-aminoquinoline derivatives

Congeneric series of *N*,*N*-bis(7-chloroquinolin-4-yl)alkanediamines [23] and *N*,*N*-bis(7-chloroquinolin-4-yl)heteroalkanediamines [24] (collectively referred to as BQs for the remainder of the text) were examined for BoNT/A LC inhibition. The compounds were tested in the HPLC-based assay at 20 µM concentration in the presence of 0.1 mM substrate [1,26]. Inhibition of BoNT/A LC by these compounds was not affected by added zinc. Furthermore, when tested against BoNT/B LC protease activity, no significant inhibition was found. Taken together, these observations suggest that these compounds exhibit specificity for binding to BoNT/A LC and are not zinc chelators. Table 2 shows the two-dimensional structures of these compounds, along with percent inhibition of BoNT/A LC.

Results from our studies on BQs led us to hypothesize that other quinoline-based compounds might also inhibit BoNT/A LC enzymatic activity. One method for fast tracking the evolution of therapeutics against BoNT/A LC would be to identify inhibitors of this enzyme that are already approved drugs (and subsequently already possess pharmacological and toxicological data from clinical trials). Subsequently, we tested five readily available antimalarial drugs—amodiaquine, chloroquine, quinacrine, quinidine, and quinine—that share similar structural characteristics with the BQs. These

compounds were obtained and examined in our HPLC-based assay; structures of these compounds and percent inhibition of BoNT/A LC protease activity are shown in Table 2.

Identification of structural similarities and potential binding sites for inhibitors from the NCI diversity set

Superimpositions of several of the most potent inhibitors identified during the NCI Diversity set screen are shown in Fig. 1. An evident feature to emerge from superimposing these compounds is that they all possess biaryl and triaryl heterocyclic scaffolds (for example, quinoline, phenazinium, or xanthene). The exception to this observation is michellamine B—the central naphthalenes of this molecule do not possess heteroatoms. However, like aromatic heterocyclic nitrogens or oxygens, hydroxy and methoxy substituents on the naphthalene rings may participate in electron donating interactions. The superimposed structures in Fig. 1 also indicate that possessing hydrophobic moieties at either end of the aromatic components may be important for potency.

To gain a better understanding of how these inhibitors interact with BoNT/A LC, molecular docking studies were performed. Michellamine B (Table 1) was the first inhibitor to be examined. This molecule was the best lead for initially probing the steric constraints of the substrate binding cleft, as it is an atropisomer that results from hindered rotation of about single bonds [30–33]; it possesses restricted conformations, which reduces the number of potential binding modes that it may assume in the enzyme's substrate binding cleft. Furthermore, unlike NSC 86372, inhibition of BoNT/A LC by michellamine B is not affected by added zinc, suggesting that inhibition is not simply the result of michellamine B acting as a chelator.

Fig. 2 shows michellamine B docked in the BoNT/A LC substrate binding cleft. There is a good complement between the steric space of the binding cleft and the twist planar conformation of michellamine B. Hydropathic analyses of favorable and unfavorable contacts in several docked models indicated that the best binding mode results when the naphthalenes (labeled A and B for descriptive purposes, Fig. 2) extend down the length of the catalytic cleft. In this model, the two hydroxyl moieties of the naphthalenes exist in close proximity to the catalytic zinc and form an intramolecular hydrogen bond. Either of the hydroxyl moieties may displace the water molecule that is used by the enzyme's catalytic engine during peptide lysis. As seen in Fig. 2, the methoxy and methyl groups of naphthalene A point toward a hydrophobic pocket (referred to as binding subsite 1 for the remainder of the text) that is formed by the aromatic side-chains of residues Phe 162, Phe 177, and Phe 193, as well as the side-chain methyl of Thr 219. The methoxy

Table 2 Quinoline-based inhibitors of BoNT/A LC

Bisquinolines (BQs)

	Compound	% Inhibition ^a		Compound	% Inhibition ^a
RHN	Q1-21	24	RHN	Q3-81	32
RHN	Q3-87	7	•		
H RHN N NHR	Q2-59	26	NHR	Q3-79	27
RHN	Q1-19	5	v		
RHN N NHR	Q2-61	50		Q2-97	15
RHN N NHR	Q2-15	60	RHN NHR		
RHN N NHR CH ₃	Q2-43	52	NHR NHR	Q2-11	8
RHN O NHR	Q1-3	12	MIK		
Antimalarial drugs ^b					
HN	N	20	HN Am	odiaquine	30
CI N (r:	acemic)		CIN		
HN	N		HO.*		
CI N	Quinacrine racemic)	30	اد الاحا	ne (% = r) ine (% = s)	9

^a Compounds tested in the HPLC-based assay at 20 μM concentration in the presence of 0.1 mM substrate.

moiety of naphthalene B packs into a deep pocket (referred to as binding subsite 2 for the remainder of the text) that is located behind His 226 of the catalytic engine, and surrounded by residues Met 164, Thr 175, Arg 230, and Pro 238. The methyl moiety of naphthalene B packs into space behind the side-chain pyrrolidine of Pro 238.

The tetrahydroisoquinoline attached to naphthalene A binds in such a way that the 6,8-diol moieties point toward the solvent, while the 1,3-dimethyl moieties point toward hydrophobic residues of the substrate binding cleft. The ionizable amine of the tetrahydroisoquinoline attached to naphthalene B points towards a cluster of polar residues, including Glu 55, Gln 161, Glu

^b Antimalarial drugs tested in the HPLC-based assay to 50 μM concentration in the presence of 0.2 mM substrate.

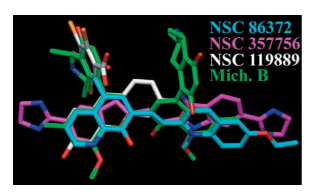


Fig. 1. Overlay of BoNT/A LC inhibitors from the NCI Diversity set. For all structures, nitrogen atoms are blue, oxygen atoms are red, and iodine atoms are orange. Carbon atoms for michellamine B are green, for NSC 86372 are light blue, for NSC 357756 are magenta, and for NSC 119889 are white. There are structural trends in: (1) the superimposed aromatic scaffolds and (2) the positioning of hydrophobic moieties.

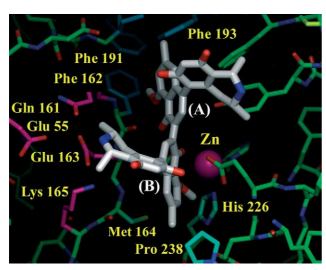


Fig. 2. Michellamine B docked in the BoNT/A LC substrate binding cleft. Nitrogen atoms are blue and oxygen atoms are red. Carbon atoms of michellamine B are white, and the naphthalene scaffolds of the inhibitor are labeled A and B (as referenced in the text). Carbon atoms of residues in predicted binding subsites are light blue (binding subsite 1 in the text), light green (binding subsite 2 in text), and magenta (referred to as the polar region in the text). All other enzyme carbons are green. For michellamine B, the methoxy and methyl substituents of naphthalene A pack into binding subsite 1, while the methoxy of naphthalene B packs into binding subsite 2. The ionizable nitrogen of the tetrahydroisoquinoline attached to naphthalene B points toward a cluster of polar residues: Glu 55, Gln 161, Glu 163, and Lys 165.

163, Lys 165, and Arg 176, and may either engage directly in an ionic bond with one of the Glu residues in this area, or participate in a water mediated hydrogen bond with any of the indicated polar residues (Fig. 2). Along the same lines, the ionizable amine may also serve to solubilize the tetrahydroisoquinoline ring. The 6,8-diol substituents of this tetrahydroisoquinoline are oriented toward the solvent interface.

Subsequent docking studies with other BoNT/A LC inhibitors from the NCI Diversity set indicated that these compounds may use binding modes similar to that of michellamine B. Fig. 3 shows a two-dimensional schematic that matches inhibitor substituents with proposed binding subsites for three inhibitors from the NCI Diversity set. Our analyses indicate that the most potent inhibitors engage in contacts with residues in both binding subsites 1 and 2, and in each case, either an oxygen or a nitrogen (in an aromatic heterocyclic ring) is positioned in close proximity to the catalytic zinc. For example, the 8-hydroxyquinoline of NSC 86372 (Fig. 3), when optimized in the substrate binding cleft, is positioned in such a way that zinc chelation is possible. Additionally, the methyl substituent of the compound's quinolinium substituent inserts into binding subsite 2 (similar to the methoxy moiety of michellamine B naphthalene B), while the ethoxy moiety of the quinolinium fills space located behind Pro 238 (similar to the methyl moiety of michellamine B naphthalene B).

In another example, the most favorable binding mode of NSC 357756 indicates that the dihydro-imidazolyl (attached to the benzofuran) inserts into binding subsite 1 (Fig. 3) (the same pocket that is occupied by the methoxy and methyl moieties of michellamine B naphthalene A (Fig. 2)). The oxygen of the benzofuran moiety is positioned in close proximity to the zinc ion, and the 3-amino substituent on the indole ring engages in a hydrogen bond with Glu 163. In binding subsite 2, a nitrogen in the dihydro-imidazolyl (which is attached to the 6 position of the indole) engages in a hydrogen bond

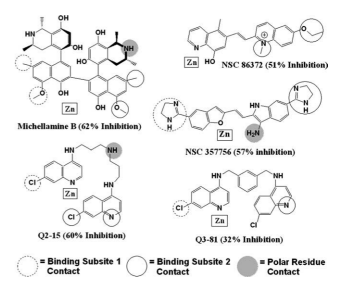


Fig. 3. A two-dimensional schematic relating compound substituents to predicted binding subsites in the BoNT/A LC substrate binding cleft. Dashed circles indicate substituents that engage in favorable contacts in binding subsite 1, solid circles indicate substituents that engage in favorable contacts in binding subsite 2, and gray spheres indicate substituents that engage in favorable interactions with polar residues.

with the side-chain guanidinium of Arg 230, while the methylenes of this ring pack behind the pyrrolidine ring of Pro 238.

Structural evaluation and molecular docking of BQs and antimalarial drugs

BQs possessing a flexible linker between the 7-chloro-4-aminoquinoline moieties are more potent inhibitors than conformationally constrained derivatives (Table 2). In addition, it is also evident that an ionizable amine in the flexible linker increases potency (for example, Q2-61 versus Q1-19). Compound Q2-15 shows that a seven atom linker, with a central ionizable amine, is favored over shorter linkers that also contain an ionizable amine.

Conformational analyses of the most potent BQs show that these compounds favor a folded conformation, with the 7-chloro-4-aminoquinoline moieties positioned front to back and slightly offset relative to one another (Fig. 4A). Based on these observations, it appears that these congeners may all bind in the BoNT/A substrate binding cleft in a similar manner.

To gain a better understanding of how the folded conformations of these congeners might bind in the BoNT/A LC substrate cleft, Q2-15 was fit onto michellamine B (in its docked conformation). The superimposition shown in Fig. 4B indicates a good overlap between: (1) the 7-chloro-4-aminoquinoline components of Q2-15 and the naphthalene moieties of michellamine B and (2) the ionizable nitrogen of the Q2-15 flexible linker and the ionizable nitrogen of one of the michellamine B isoquinolines.

These observations led to docking studies to determine if the BQs might also bind to predicted subsites 1 and 2 in the BoNT/A LC substrate binding cleft. For the most potent compounds, the ring nitrogen of one of the 7-chloro-4-aminoquinoline moieties sits in close proximity to the catalytic zinc, while the 7-chloro substituent packs into binding subsite 1. The ionizable amines in compounds with flexible linkers engage in potential ionic bonds with the polar side-chains of either Gln 161 or Glu 163. Another explanation for the importance of the ionizable amine may be that it is necessary for solubilizing the otherwise hydrophobic linkers found in this series of compounds.

Docking studies also indicate that the second 7-chloro-4-aminoquinoline moiety (of the most potent derivatives) binds in subsite 2 such that the nitrogen of the heterocycle points toward the solvent and engages in a hydrogen bond with the side-chain guanidinium of Arg 230 (as was also observed for a nitrogen in one of the dihydro-imidazolyl rings of NSC 357756), and the 7-chloro substituent points into the hydrophobic space behind His 226. For comparison, molecular docking of compounds with rigid linkers (for example, Q2-11 and Q2-97) indicates that steric constraints prevent these

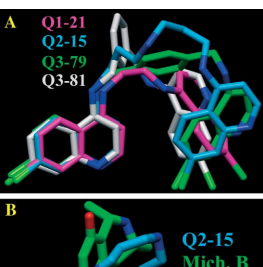
compounds from adopting binding modes in subsites 1 and 2 that are as favorable as those observed for congeners with flexible linkers. In general, these studies provide evidence that the BQs interact with the BoNT/A LC substrate binding cleft in a manner similar to inhibitors from the NCI Diversity set. Fig. 3 shows two-dimensional schematics matching substituents from two of the most potent BQs (Q2-15 and Q3-81) with their corresponding binding subsites in the BoNT/A LC substrate binding cleft.

Of the five examined antimalarial drugs (Table 2), amodiaquine and quinacrine were the most potent (30% inhibition of protease activity). However, the fact that these two drugs inhibit BoNT/A LC with approximately half the potency of Q2-15 provides further evidence that a second aromatic component, such as found in the BQs or michellamine B, is necessary for enhanced potency, and supports our hypothesis that binding subsite 2 (in the enzyme's substrate binding cleft) is an important contact region for inhibitors. Furthermore, as quinacrine contains an acridine scaffold, these studies have revealed a new aromatic component that may be used to identify and/or synthesize new BoNT/A LC inhibitors.

The antimalarial agents that we found to inhibit BoNT/A LC protease activity have previously been shown to increase the time to BoNT/A holotoxin induced muscle paralysis [17,18]. In the earlier work, it was hypothesized that these compounds delayed paralysis by interfering with toxin translocation into the nerve cytoplasm and therefore they were not tested for specific inhibition of BoNT/A LC protease activity. The percent inhibition of BoNT/A LC that we observed for these drugs was not equivalent to the level of protection that they afforded against muscle paralysis in the previous studies [17,18]. For example, quinacrine, which we report to inhibit BoNT/A LC protease activity by 30% at 50 µM (0.2 mM substrate), was found, in an earlier study [17], to delay the time to 50% muscle paralysis by at least 30% (over control) at 3.3 µM [17]. Thus, the results from the earlier studies, combined with the findings from our investigation, suggest that these compounds possess a dual mechanism of action—interfering with both BoNT/A entry into the cytoplasm and with the protease activity of the light chain. This dual mechanism of action is appealing from a drug design standpoint, as one molecule can be used to inhibit two separate functions of the same target. Indeed, if the BQs are found to also inhibit BoNT/A cytoplasm entry, then these studies will be the basis for the development of a new family of bifunctional inhibitors.

A common pharmacophore for BoNT/A LC inhibitors

Based on conformational sampling and molecular docking studies, a common pharmacophore for BoNT/A LC inhibitors is proposed (Fig. 5A). An example of



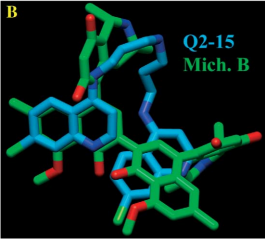


Fig. 4. (A) Overlay of folded conformers of BQs: Q1-21, Q2-15, Q3-79, and Q3-81. For all compounds, nitrogen atoms are dark blue and chlorine atoms are light green. The carbon atoms of Q1-21 are magenta, of Q2-15 are light blue, of Q3-79 are green, and of Q3-81 are white. A striking feature of the superimposed compounds is the close correspondence of their chloro substituents, which provides evidence that they may bind within the enzyme's substrate binding cleft in a similar manner. (B) Overlay of michellamine B and Q2-15. Carbons of michellamine B are green and carbon atoms of Q2-15 are light blue. The superimposed conformers show a good correspondence between: (1) the ionizable amine in the flexible linker of Q2-15 and the ionizable amine in one of the tetrahydroisoquinolines of michellamine B and (2) the chloro moieties of Q2-15 and the hydrophobic methoxy/methyl moieties of michellamine B.

how one of the inhibitors (Q2-15) fits the pharmacophore, using Catalyst 4.7 (Accelrys, San Diego, CA) as the query engine, is shown in Fig. 5B. Specifically, our studies show that the BoNT/A LC binding cleft can accommodate up to two biaryl/triaryl aromatic components, which are defined by planes A and B in the pharmacophore (Fig. 5A). These planes serve as scaffolds for positioning other components of the pharmacophore, provided that the distance between centroids of the two planes only varies from approximately 6.5–9.5 Å (Fig. 5A). An additional requirement of plane A is the presence of a heteroatom (Fig. 5A) that may serve to either directly engage in an interaction with the catalytic zinc, or potentially replace the water molecule used by zinc during substrate lysis. Examples of structural

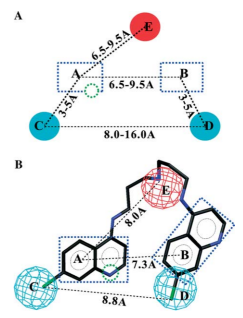


Fig. 5. (A) Common pharmacophore for BoNT/A LC inhibition. Planar components A and B are blue dashed rectangles. The dashed green circle in plane A represents a heteroatom. Hydrophobic components of the pharmacophore (C and D) are shown with light blue spheres. The positive ionizable component of the pharmacophore (E) is shown with a red sphere. (B) Q2-15 mapped onto the common pharmacophore (shown in (A)) in Catalyst 4.7 (Accelrys, San Diego, CA). Carbon atoms are black, nitrogen atoms are blue, and chlorine atoms are green. Planar components A and B are outlined with blue dashed rectangles. The heteroatom in plane A is indicated with a dashed green circle. Mapping of the hydrophobic components (C and D) is indicated with light blue mesh spheres; mapping of the positive ionizable component (E) is indicated with a red mesh sphere.

components of inhibitors that occupy planes A and B include: the two quinoline rings of BQs (a quinoline nitrogen would be the electron donor) (Table 2, Figs. 3, 4, and 5B); the two naphthalene rings of michellamine B (a hydroxyl oxygen would be the electron donor) (Table 1, Figs. 2 and 3); and the indole and benzofuran rings of NSC 357756 (the benzofuran oxygen would be the electron donor) (Table 1 and Fig. 3).

Two hydrophobic moieties (C and D) are key components of the common pharmacophore (Fig. 5A). In relation to the BoNT/A LC substrate binding cleft, C and D correspond to binding subsites 1 and 2, and facilitate hydrophobic collapse between inhibitors and the enzyme. The $C \rightarrow D$ distance range of the common pharmacophore shown in Fig. 5A reflects the extent of the potential binding space that inhibitors may occupy in subsite 2 of the BoNT/A LC substrate binding cleft. Examples of C and D in inhibitors, as they map to the common pharmacophore, include: the two methoxy substituents of michellamine B (separated by a distance of 8.6 Å) (Table 1 and Figs. 2 and 3); the two chlorines of the BQs (separated by an average distance of 9.8 Å for all examined BQs) (Table 2 and Figs. 3, 4, and 5B);

and the two dihydro imidazolyl moieties of NSC 357756 (separated by a distance of 15.8 Å) (Table 1 and Fig. 3).

The final component (E) of the common pharmacophore is a positive ionizable substituent (Fig. 5A). While inhibitor docking studies have shown that an ionizable moiety may participate in electrostatic interactions with enzyme residues, the positive charge may also aid in solubilizing inhibitors bound within the BoNT/A LC substrate binding cleft; unlike a traditional binding pocket, the BoNT/A LC substrate binding cleft is large and almost completely solvent exposed. Hence, it is plausible that a hydrogen bonding network involving the positively charged moiety(ies) of the inhibitors and surrounding water molecules also serves to stabilize compound binding. Examples of inhibitor components that map to E include: the ionizable secondary nitrogen in one of the michellamine B tetrahydroisoquinolines (Table 1 and Figs. 2 and 3); the secondary nitrogens in the flexible linkers of BQs (Table 2 and Figs. 3, 4, and 5B); and the tertiary nitrogens in amodiaguine, chloroquine, and quinacrine (Table 2). Interestingly, the determined BoNT/A LC peptide cleavage site of the natural substrate, SNAP-25, is between a glutamine and an arginine [34]. Furthermore, the distance between the side-chain guanidinum of arginine (which is ionized at physiological pH) and the backbone Cα of this residue corresponds to the distance between A and E in the common pharmacophore. Hence, it is possible that the ionizable substituent of the inhibitors may be mimicking contacts made by the side-chain guanidinium of the indicated Arg residue of SNAP-25.

Summary

These experiments have, for the first time, identified small organic (non-peptidic) lead therapeutics that inhibit BoNT/A LC protease activity in the low μ M range. Based on the structures of these lead therapeutics, molecular modeling studies have been used to propose a common pharmacophore. This pharmacophore will serve as the basis for directing future database mining studies and synthetic organic chemistry projects to identify and develop BoNT/A LC inhibitors with enhanced potency.

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