### Citation

### Published Version
doi:10.1038/ncomms14130

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Identification and characterization of a novel botulinum neurotoxin

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Botulinum neurotoxins are known to have seven serotypes (BoNT/A–G). Here we report a new BoNT serotype, tentatively named BoNT/X, which has the lowest sequence identity with other BoNTs and is not recognized by antisera against known BoNTs. Similar to BoNT/B/D/F/G, BoNT/X cleaves vesicle-associated membrane proteins (VAMP) 1, 2 and 3, but at a novel site (Arg66-Ala67 in VAMP2). Remarkably, BoNT/X is the only toxin that also cleaves non-canonical substrates VAMP4, VAMP5 and Ykt6. To validate its activity, a small amount of full-length BoNT/X was assembled by linking two non-toxic fragments using a transpeptidase (sortase). Assembled BoNT/X cleaves VAMP2 and VAMP4 in cultured neurons and causes flaccid paralysis in mice. Thus, BoNT/X is a novel BoNT with a unique substrate profile. Its discovery posts a challenge to develop effective countermeasures, provides a novel tool for studying intracellular membrane trafficking, and presents a new potential therapeutic toxin for modulating secretions in cells.
Botulinum neurotoxins (BoNTs) are a family of bacterial toxins, classified as one of the six most dangerous potential bioterrorism agents (Category A and Tier 1 select agent in the United States)\(^1\). They are also widely used to treat a growing list of medical conditions\(^2\)–\(^4\), including muscle spasms, chronic pain, overactive bladders, as well as having cosmetic applications. There are seven well-established serotypes of BoNTs (BoNT/A–G), traditionally defined based on a lack of cross-neutralization by different antisera raised against each toxin type. All BoNTs share the same structure and function\(^4\)–\(^6\). They are composed of a light chain (LC, ~50 kDa) and a heavy chain (HC, ~100 kDa) connected by an inter-chain disulfide bond. The HC contains two sub-domains: the C-terminal HC\(_5\) that mediates binding to receptors, and the N-terminal HC\(_3\) that mediates translocation of the LC across the endosomal membranes. The LC acts as a protease in neurons to cleave a set of proteins: BoNT/A, C and E cleave at distinct sites on a peripheral membrane protein known as SNAP-25; BoNT/B, D, F and G cleave at different sites on homologous vesicle proteins VAMP1, 2 and 3 (vesicle-associated membrane proteins); and BoNT/C also cleaves the plasma membrane protein syntaxin 1. These proteins are prototypes of the SNARE (soluble NSF attachment protein receptor) protein family, whose members mediate various membrane fusion events in eukaryotic cells\(^7\)–\(^8\). Cleavage of any one of the three neuronal SNARE proteins blocks fusion of synaptic vesicles to plasma membranes, thus preventing neurotransmitter release from neurons.

Recognizing all distinct serotypes of BoNTs is essential for developing effective countermeasures against this family of toxins. BoNT/A and BoNT/B were first identified in 1919 by Georgina Burke\(^9\). The last of the seven serotypes, BoNT/G, was discovered in 1969 (ref. 10), and no new BoNT serotype has been found for the past four decades. Recent progress in genomic sequencing has revealed multiple subtypes (designated with Arabic numbers, for example, BoNT/A1), which can be recognized by the same antisera but contain substantial sequence variations\(^11\)–\(^13\). Furthermore, there are also multiple mosaic toxins. For instance, a ‘type H’ was reported in 2013 but was later designated as a mosaic toxin, as its LC shares ~80% identity with the LC of a BoNT/F subtype, BoNT/F5, and its HC shares ~84% identity with the HC of BoNT/A1 (ref. 14–17). Consistently, this toxin can be neutralized by antibodies against BoNT/A (ref. 16).

The genes encoding BoNTs can be on a plasmid, a phage, or the chromosome, indicating that these genes are mobile and capable of horizontal gene transfer\(^18\)–\(^23\). Some strains contain two or even three different BoNT genes\(^15\)–\(^24\),\(^25\). These strains are usually designated with a capital letter for the toxin type that is expressed at higher levels than the other one, followed with a lower case letter for the second toxin type (for example, BoNT/Af strain). In addition, it has also been reported that some BoNT/A strains contain a complete BoNT/B gene, but only BoNT/A is expressed\(^26\)–\(^28\). Thus, the BoNT/B gene is considered a silent gene and the strains are known as BoNT/A(B) strains. A recent survey of infant botulism cases reported that ~8% isolates are BoNT/A(B) strains\(^29\).

Here we searched published genomic sequences and identified a novel BoNT gene encoded on the chromosome of *Clostridium botulinum* strain 111. This strain was originally identified from an infant botulism patient in Japan in 1996 (ref. 30). The initial characterizations indicated that the toxicity of this strain is due to BoNT/B\(^30\). Later studies confirmed that this strain expresses a subtype of BoNT/B (BoNT/B2) encoded on a plasmid\(^31\),\(^32\). The sequence of this novel BoNT gene was deposited into the GenBank database in February 2015, as a part of the genomic sequence of *C. botulinum* 111. We characterized the protein encoded by this gene at functional levels and established it as a new BoNT serotype with a unique substrate profile.

### Results

**Searching genomic databases revealed a novel BoNT gene.**

In an attempt to survey the evolutionary landscape of BoNTs, we performed iterative Hidden Markov model searches of the Uniprot sequence database. Our search identified all known BoNT subtypes and mosaic toxins, as well as the related tetanus neurotoxin (Fig. 1a; Supplementary Fig. 1). To our surprise, the search revealed a potentially new BoNT, tentatively designated BoNT/X (Fig. 1a, GenBank no.: BAQ12790.1), from the recently reported genomic sequence of *C. botulinum* strain 111. BoNT/X showed the least protein sequence identity with the other BoNTs in pairwise comparisons (Fig. 1b). Furthermore, the low sequence similarity is evenly distributed along the entire BoNT/X sequence (Fig. 1c), indicating that it is not a mosaic toxin. Despite this low sequence identity, the overall domain arrangement of BoNTs is conserved in BoNT/X (Fig. 1c), including a zinc-dependent protease motif HEXXH (residues 227–231, HELVH) in the LC (ref. 33), and a SXWY motif in the HC (residues 1,274–1,277, SAWY), which recognizes the lipid receptor gangliosides\(^34\).

Similar to the other BoNTs, the BoNT/X gene is located in a gene cluster\(^23\). All seven established BoNTs are co-expressed with another 150 kDa protein known as NTNHA (non-toxic non-hemagglutinin protein), which forms a pH-dependent complex with BoNTs and protects them from proteases in the gastrointestinal tract\(^35\). The BoNT/X gene is also preceded by a potential NTNHA gene (Fig. 1d). Besides BoNT and NTNHA, a typical BoNT gene cluster contains genes encoding one of the two types of accessory proteins: (1) the HA cluster encoding three conserved proteins HA17, HA33 and HA70, which form a complex with BoNT/NTNHA and facilitate absorption of toxins across the intestinal epithelial barrier\(^16\)–\(^18\), or (2) the OrfX cluster encoding conserved OrfX1, OrfX2, OrfX3 and P47 proteins with unknown function\(^23\). The BoNT/X gene is located in an OrfX gene cluster, as are BoNT/E, F and members of BoNT/A. Interestingly, the BoNT/X cluster has two unique features (Fig. 1d): (1) there is an additional OrfX2 gene that does not exist in any other BoNT clusters (we designated it OrfX2b); (2) the reading frame of OrfX genes is usually opposite to BoNT/NTNHA genes, but it has the same direction as the BoNT/X gene in the BoNT/X cluster (Fig. 1d). These findings suggest that BoNT/X is a unique branch of the BoNT family.

The LC of BoNT/X cleaves VAMP2 at a novel site. To characterize BoNT/X, we first focused on its LC (X-LC, residues 1–439) and produced it as a His6-tagged protein in *Escherichia coli*. LCS of BoNT/A (A-LC) and BoNT/B (B-LC) were produced and assayed in parallel as controls. Incubation of X-LC with EDTA prevented cleavage of SNARE proteins by X-, A- and B-LCs (Fig. 2a). LCs of BoNTs are zinc-dependent proteases\(^33\). As expected, EDTA prevented cleavage of SNARE proteins by X-, A- and B-LCs (Fig. 2a). Furthermore, incubation of X-LC with the purified recombinant cytosolic domain of VAMP2 (residues 1–93) converted VAMP2 into two lower-molecular-weight bands (Fig. 2b), confirming that X-LC cleaves VAMP2.

To identify the cleavage site, we analysed the VAMP2 (1–93) protein, with or without pre-incubation with X-LC, by liquid chromatography–tandem mass spectrometry (LC–MS/MS, Fig. 2c–e). A single dominant peptide peak appeared after incubation with X-LC (Fig. 2c,e; Supplementary Fig. 2). Its molecular weight is 3,081.7, which fits only the peptide sequence of A67–L93 of VAMP2 (Fig. 2c,e). Consistently, another fragment...
from the beginning of the His6-tag to residue R66 of VAMP2 was also detected (Fig. 2d). To further confirm this finding, we repeated the assay with a different VAMP2 fragment: glutathione S-transferase (GST) tagged VAMP2 (33–86) (Supplementary Fig. 3). Incubation with X-LC generated a single dominant peptide peak with a molecular weight of 2,063.1, which fits only A67–R86 of VAMP2 (Supplementary Fig. 3). Together, these results demonstrate that X-LC has a single cleavage site on VAMP2 between R66 and A67.

R66–A67 is a novel cleavage site on VAMP2, distinct from all established target sites of BoNTs (Fig. 2f). It is also the only BoNT cleavage site located within a region previously known as the SARE motif (Fig. 2f, shaded regions). The VAMP protein family includes VAMP1, 2, 3, 4, 5, 7 and 8, as well as related Sec22b and Ykt6. R66–A67 is conserved in VAMP1 and 3, which are highly homologous to VAMP2. To validate the specificity of X-LC, we expressed HA-tagged VAMP1, 3, 7, 8 and Myc-tagged Sec22b and Ykt6 in HEK293 cells via transient transfection.

Cell lysates were incubated with X-LC. Both VAMP1 and 3 were cleaved by X-LC, whereas VAMP7, VAMP8 and Sec22b were resistant to X-LC (Fig. 2g).

BoNT/X cleaves VAMP4, VAMP5 and Ykt6. Unexpectedly, Ykt6 was also cleaved by X-LC (Fig. 2h). This finding was confirmed using a purified GST-tagged Ykt6 fragment, which shifted to a lower-molecular-weight band after incubation with X-LC (Fig. 2h). The cleavage site was determined to be K173-S174 by mass spectrometry analysis of the intact Ykt6 versus Ykt6 cleaved by X-LC (Supplementary Fig. 4). This site is homologous to the cleavage site of BoNT/X on VAMP2 (Fig. 2i). Among VAMP family of proteins, VAMP4 contains the same pair of residues (K87–S88) at this site as Ykt6. We found that X-LC cleaved both purified GST-tagged cytoplasmic domain of VAMP4 (Fig. 2i), as well as native VAMP4 in BDE (Fig. 2j). As a control, Sec22b was not cleaved by X-LC in BDE. In addition, the GST-tagged cytoplasmic domain of VAMP5 was also cleaved (Fig. 2i). The cleavage sites were determined by mass spectrometry analysis to be K87–S88 in VAMP4 and R40–S41 in VAMP5 (Supplementary Fig. 5). Both sites are homologous to the cleavage site of BoNT/X on VAMP2 (Fig. 2i), demonstrating that the location of the cleavage site is conserved across different VAMPs. The ability of X-LC to cleave VAMP4, VAMP5 and Ykt6 is highly unusual, as their sequences are substantially different from VAMP1/2/3. BoNT/X is the first and the only BoNT known that can cleave VAMPs beyond the canonical targets VAMP1, 2 and 3 (ref. 40).

Proteolytic activation of BoNT/X. We next examined the linker region between the LC and the HC, which must be cleaved by bacterial or host proteases to convert the toxin to an ‘active’ di-chain form. We produced a recombinant X-LC-H2 fragment (residues 1–891) in E. coli and subjected it to limited proteolysis by endoprotease Lys-C. Samples were analysed using Tandem Mass Tag (TMT) labelling and tandem mass spectrometry. TMT labels free N-termini (and lysines). Limited proteolysis by Lys-C

![Figure 1](image_url) **Identification of BoNT/X.** (a) A phylogenetic split network covering all BoNT serotypes, subtypes, mosaic toxins and related tetanus neurotoxin (TeNT) illustrates their potential evolutionary relationships, as well as conflicts arising from e.g. chimerisms, based on their protein sequences. BoNT/X is highlighted in red. An enlarged version of this panel is shown in Supplementary Fig. 1, with the sequence access number for each toxin gene noted. (b) A phylogenetic tree of the protein sequence alignment for BoNT/A-G, TeNT and BoNT/X, analysed by the ClustalW method. The percentages of sequence identity between each toxin and BoNT/X are noted. (c) Upper panel: a schematic drawing of the three domains of BoNT/X, with conserved protease motif in the LC and the ganglioside binding motif in the HC noted. Lower panel: analysis using a sliding sequence comparison window demonstrated that the low identity between each toxin and BoNT/X are noted. (d) A schematic drawing of the orfX gene cluster that hosts the BoNT/X gene (upper panel), which has two distinct features compared with other known orfX clusters (middle and lower panels): (1) there is an additional orfX2 protein (designated orfX2b) located next to the BoNT/X gene; (2) the reading frame of orfX genes has the same direction as the BoNT/X gene.
Figure 2 | The LC of BoNT/X cleaves VAMPs at a unique site. (a) X-LC was incubated with BDE. Immunoblot analysis was carried out to detect syntaxin 1, SNAP-25 and VAMP2. Synaptophysin (Syp) served as a loading control. A-LC and B-LC were analysed in parallel. Cleavage of VAMP2 by B-LC results in two smaller fragments. (b) VAMP2 (1–93) was incubated with X-LC. Samples were analysed by mass spectrometry (LC–MS/MS) to determine the molecular weight of cleaved fragments. Eluted peptide peaks from the HPLC column are plotted over running time (RT, X axis). The mass spectrometry data for the two cleavage products are colour-coded, with mass-to-charge ratio (m/z) noted. The molecular weight is deduced by multiplying m with z, following by subtracting z. The protein sequences for the two cleavage products are colour-coded and listed in e. (f) Sequence alignment between VAMP family members, with the cleavage sites for BoNT/B, D, F, G and X marked in red, and the two SNARE motifs in blue shade. (g) HA-tagged VAMP1, 3, 7 and 8, and Myc-tagged Sec22b and Ykt6 were expressed in 293T cells via transient transfection. Cell lysates were incubated with X-LC and subjected to immunoblot analysis. Actin is a loading control. (h) GST-tagged Ykt6 was incubated with X-LC (100 nM). Samples were analysed by SDS–PAGE and Coomassie Blue staining. (i) GST-tagged VAMP2 (33–86), VAMP4 (1–115) and VAMP5 (1–70) were incubated with X-LC (100 nM). Samples were analysed by SDS–PAGE and Coomassie Blue staining. X-LC cleaved both VAMP2 and VAMP5. We note that VAMP5 protein contains a contaminant band that runs close to the cleavage product. (j) Experiments were carried out as described in a, except that VAMP4 and Sec22b were detected. Synaptotagmin I (Syt I) is a loading control. X-LC cleaved native VAMP4 in BDE. One of two (b,g,j) or three (a,h,i) independent experiments is shown.

produced one new free N terminus mapped to residue N439 in the linker region (Fig. 3a; Supplementary Data 1), confirming that the linker region is susceptible to proteases.

We then examined whether proteolytic activation enhances the potency of BoNT/X. It has been shown that incubation of high concentrations of LC-H₉ of BoNTs with cultured neurons results in entry of LC-H₉, likely through non-specific uptake into neurons⁴¹. Similarly, X-LC-H₉ entered cultured rat cortical neurons and cleaved VAMP2 in a concentration-dependent manner (Fig. 3b). Activation by Lys-C increased the potency of X-LC-H₉: 10 nM activated X-LC-H₉ cleaved similar levels of VAMP2 as 150 nM intact X-LC-H₉ (Fig. 3b). Activated X-LC-H₉...
These results confirmed that the inter-chain disulfide bond is essential for the activity of X-LC-HN, and this inter-chain disulfide bond can be formed via block disulfide bond shuffling. Samples were then analysed by SDS–PAGE and Coomassie Blue staining. A majority of WT X-LC-HN exists as a single band cleaved their substrates. X-LC-HN activated by Lys-C showed a greater potency than non-activated X-LC-HN. X-LC-HN was more potent than B-LC-HN and A-LC-HN, neither of which or X-LC-HN mutants. C423S mutation abolished the activity of X-LC-HN, whereas mutating C461 or C467 did not affect the activity of X-LC-HN. disulfide bond in WT and three cysteine mutants of BoNT/X. X-LC-HN formed aggregates, marked by an asterisk, which disappeared with DTT. The majority of activated WT X-LC-HN separated into two 100 kDa bands without DTT and separated into two ~50 kDa bands with or without DTT. C423/C461/C467 are the only three cysteine residues mutated in the presence of DTT. C423S mutation abolished the activity of X-LC-HN, whereas mutating C461 or C467 did not affect the activity of X-LC-HN. These results confirmed that the inter-chain disulfide bond is essential for the activity of X-LC-HN, and this inter-chain disulfide bond can be formed via either C423-C461 or C423-C467. One of two (b) or three (b,c,f) independent experiments is shown.

The inter-chain disulfide bond in BoNT/X. Like other BoNTs, the linker region of BoNT/X contains two conserved cysteines, but there is also an additional cysteine (C461) unique to BoNT/X (Fig. 3a). To determine the cysteine residues that form the essential inter-chain disulfide bond, we generated three X-LC-HN mutants, each with one of the three cysteine residues mutated (C423S, C461S and C467S). These mutants, as well as wild-type (WT) X-LC-HN, were subjected to limited proteolysis with Lys-C and then analysed via SDS–PAGE and Coomassie Blue staining, with or without the reducing agent dithiothreitol (DTT) (Fig. 3c). Mutating the only cysteine on the LC (C423S) is expected to abolish the inter-chain disulfide bond. Consistently, C423S mutant separated into two ~50 kDa bands without DTT. In contrast, both C461S and C467S mutants showed as a single band at 100 kDa in the absence of DTT and separated into two ~50 kDa bands in the presence of DTT. These results suggested that C423 on the LC can form the inter-chain disulfide bond with either C461 or C467 on the HC. We also found that Lys-C treatment degraded a significant portion of C423S mutant as compared with C461S or C467S mutants (Fig. 3c, +DTT), suggesting that losing the inter-chain disulfide bond makes the molecule more susceptible to proteases. We noticed that a portion of WT X-LC-HN formed aggregates at the top of the SDS–PAGE gel (Fig. 3c, marked by an asterisk). These aggregates disappeared in the presence of DTT. C423/C461/C467 are the only three cysteine mutants of BoNT/X. (f) Experiments were carried out as described in b, except that neurons were exposed to WT or C423/C461/C467 mutants. C423S mutation abolished the activity of X-LC-HN, whereas mutating C461 or C467 did not affect the activity of X-LC-HN. These results confirmed that the inter-chain disulfide bond is essential for the activity of X-LC-HN, and this inter-chain disulfide bond can be formed via either C423-C461 or C423-C467. One of two (b) or three (b,c,f) independent experiments is shown.

Figure 3 | Proteolytic activation and inter-chain disulfide bond in BoNT/X. (a) Sequence alignment of the linker between the LC and HC of the seven established BoNTs plus BoNT/X. The Lys-C cutting site was identified by mass spectrometry analysis (see Method and Supplementary Data 1). (b) Cultured rat cortical neurons were exposed to X-LC-HN for 12 h. Cell lysates were harvested and immunoblot analysis carried out to examine syntaxin 1, SNAP-25 and VAMP2. Actin is a loading control. Trypsin-activated A-LC-HN and B-LC-HN were analysed in parallel. X-LC-HN entered neurons and cleaved VAMP2. X-LC-HN activated by Lys-C showed a greater potency than non-activated X-LC-HN. X-LC-HN was more potent than B-LC-HN and A-LC-HN, neither of which cleaved their substrates. (c) WT and mutant X-LC-HN were activated by Lys-C and analysed by SDS–PAGE and Coomassie Blue staining, with or without DTT. C461S and C467S mutants showed as a single band at ~100 kDa without DTT, and separated into two ~50 kDa bands with DTT. A portion of WT X-LC-HN formed aggregates, marked by an asterisk, which disappeared with DTT. The majority of activated WT X-LC-HN separated into two ~50 kDa bands without DTT. This is due to disulfide bond shuffling as described in the following panel. (d) Lys-C-activated WT X-LC-HN was incubated with NEM to block disulfide bond shuffling. Samples were then analysed by SDS–PAGE and Coomassie Blue staining. A majority of WT X-LC-HN exists as a single band at ~100 kDa without DTT after NEM treatment, indicating that native WT X-LC-HN contains an inter-chain disulfide bond. (e) Schematic drawings of the disulfide bond in WT and three cysteine mutants of BoNT/X. (f) Cultured rat cortical neurons were exposed to X-LC-HN for 12 h. Cell lysates were harvested and immunoblot analysis carried out to examine syntaxin 1, SNAP-25 and VAMP2. Actin is a loading control. Trypsin-activated A-LC-HN and B-LC-HN were analysed in parallel. X-LC-HN entered neurons and cleaved VAMP2. X-LC-HN activated by Lys-C showed a greater potency than non-activated X-LC-HN. X-LC-HN was more potent than B-LC-HN and A-LC-HN, neither of which cleaved their substrates.
increased degradation by Lys-C compared with C423S mutant (Fig. 3c, + DTT). One possible explanation is that WT X-LC-HN contains an inter-chain disulfide bond under native conditions, but this bond can rearrange into intra-chain C461–C467 pair under denaturing conditions in the SDS buffer. This phenomenon is known as disulfide bond shuffling, which often occurs among adjacent cysteines. To test this hypothesis, we utilized an alkylating reagent, N-Ethylmaleimide (NEM), which permanently blocks free cysteines and prevents disulfide bond shuffling. As shown in Fig. 3d, WT X-LC-HN pretreated with NEM showed as a single band at 100 kDa in the absence of DTT, and separated into two ~50 kDa bands in the presence of DTT. These results confirm that WT X-LC-HN contains mainly an inter-chain disulfide bond, but it is susceptible to disulfide bond shuffling due to an extra cysteine in the linker region (Fig. 3e).

We further examined the activity of the three X-LC-HN cysteine mutants on cultured neurons. As expected, C423S mutant was inactive, whereas C461S and C467S mutants both showed similar levels of activity as WT X-LC-HN (Fig. 3f). These results confirm that the inter-chain disulfide bond is critical for the activity of BoNT/X.

**Generating full-length BoNT/X via sortase-mediated ligation.** We then sought to determine whether full-length BoNT/X is a functional toxin. As no antisera against BoNT/X are available, we decided to avoid generating the full-length active toxin gene. Instead, we developed an approach to generate a limited amount of full-length BoNTs in test tubes by enzymatic ligation of two non-toxic fragments of BoNTs. This method utilizes a transpeptidase known as sortase, which recognizes the peptide motif LPXTG, cleaves between T-G, and concurrently forms a new peptide bond with other proteins/peptides containing N-terminal glycine (Fig. 4a). We produced two non-toxic fragments of BoNT/X: (1) LC-HN with a LPETGGS fusion and a His6-tag fused to the C terminus; and (2) the HC of BoNT/X (X-HC) with a GST tag, thrombin cleavage site, and an additional glycine residue at its N terminus. Cutting by thrombin releases X-HC with a free glycine at its N terminus. Incubation of these two fragments with sortase generated a small amount of ~150 kD full-length BoNT/X (X-FL, Fig. 4a). We note that X-HC showed poor solubility and a strong tendency towards aggregation, which might be the reason for the low ligation efficiency (Fig. 4b). In contrast, ligation of X-LC-HN with the HC of BoNT/A (A-HC) achieved a better efficiency, with the majority of X-LC-HN ligated into a XA chimeric toxin (Supplementary Fig. 6a). To ensure biosafety, the amount of precursor fragments in the reaction is strictly limited to generate the minimum amount of ligated toxin necessary for functional assays.

We first analysed the activity of ligated BoNT/X using cultured rat cortical neurons. Neurons were exposed to the sortase ligation mixture and control mixtures in culture medium. As shown in...
Full-length inactive BoNT/X. Finally, we developed an inactive mutant of BoNT/X as a potential reagent for generating neutralizing antibodies. Mutations at two residues (R362A/Y365F) in BoNT/A inactivate the protease activity of the LC and abolish the toxicity of BoNT/A in vivo.45,46. These two residues are conserved in BoNT/X. We introduced the corresponding mutations (R360A/Y363F) in BoNT/X and generated a full-length inactive form, designated as BoNT/XRY.

**Discussion**

BoNT/X is the first serotype of BoNTs identified by genomic sequencing and bioinformatics approaches. It remains unknown whether BoNT/X is ever produced in C. botulinum strain 111. BoNT/X could be a silent gene, or it may not be expressed at detectable toxicity levels under culture conditions in the lab. Thus, this toxin was revealed only by sequencing C. botulinum 111. This illustrates the importance of genomic sequencing and bioinformatics approaches for understanding microbial virulence factors. Whether BoNT/X could be expressed and exhibit toxicity under certain environmental conditions remains an intriguing question.

A remarkable feature of BoNT/X is its unique ability to cleave VAMP4 and Ykt6. VAMP4 is widely expressed and is known to mediate vesicle fusion between the trans-Golgi network (TGN) and endosomes, as well as homotypic fusion of endosomes.47,48 Ykt6 is an atypical SNARE without a transmembrane domain.49 It is anchored to membranes via lipidation, which allows dynamic regulation of its membrane association. Ykt6 is an essential protein in yeast and implicated in multiple membrane fusion events including ER-Golgi, intra-Golgi, endosome-Golgi-vacuolar, and autophagosome formation. Its function in mammalian cells remains to be established. BoNT/X is the first and only BoNT to cleave these SNAREs that mediate various intracellular membrane-trafficking events.

Interestingly, both VAMP4 and Ykt6 are enriched in neurons. Recent studies suggested that VAMP4 contributes to asynchronous synaptic vesicle exocytosis, enlargeosome exocytosis and activity-dependent bulk endocytosis (ADBE) in neurons.50–52 The role of Ykt6 in neurons remains to be established, but it has been shown to suppress the toxicity of α-synuclein in Parkinson’s disease models.53,54 The other substrate of BoNT/X, VAMP5, is mainly expressed in muscle cells and its function remains to be established.55 BoNT/X will be a useful tool to investigate the function of VAMP4, Ykt6 and VAMP5, as well as related membrane trafficking events. In addition, because VAMP4 has been implicated in granule release in immune cells, BoNT/X may have the potential to modulate inflammatory secretion in immune cells.
The X-LC-H₃ fragment showed a higher level of activity in neurons than either A-LC-H₃ or B-LC-H₃, suggesting that its membrane translocation and/or protease activity might be more efficient than the corresponding fragments in BoNT/A and BoNT/B. X-HC is functional for targeting cells, as its presence enhanced cleavage of VAMP2 in neurons over LC-H₃ alone (Fig. 4c). When present without the translocation and LC domains, X-HC is prone to aggregation. This solubility issue is likely due to separation of X-HC from X-LC-H₃, as full-length BoNT/XKY remains stable at high concentrations. The X-FL generated by sortase-mediated ligation has a rather low toxicity in vivo in mice. It remains unknown whether this low in vivo toxicity was intrinsic to BoNT/X. It is also possible that the sortase linking method resulted in an attenuated toxin, as the H₃ and LC-H₃ folded separately and there is an additional linker between the H₃ and H₃ in ligated toxins. Nevertheless, X-FL produced by sortase-mediated ligation is active on neurons and induced typical flaccid paralysis when injected locally in mice, produced by sortase-mediated ligation is active on neurons and induced typical flaccid paralysis when injected locally in mice, demonstrating that BoNT/X is a functional toxin (Fig. 4f). It will be necessary to introduce native BoNT/X to characterize its in vivo potential and determine its biosafety risk. It will be important to neutralize antisera against BoNT/X before producing any native toxin.

Introducing a full-length active toxin gene into any expression system/organism is always a significant biosafety concern. Sortase-mediated ligation assembles a small quantity of full-length toxin from two complementary and non-fragmented expressed and purified individually. The amount of the precursor toxins in the reaction can be strictly controlled, so the amount of ligated toxin is precisely limited to ensure biosafety. The ligated toxin should possess the same mode of action as native toxins, but may exhibit lower toxicity in vivo, possibly due to the addition of the sortase linker and/or reduced compactness of the molecule. Thus, the ligated toxin could be used for functional studies, but may not be suitable for estimating the lethal dose of native toxin. On the other hand, this reduced toxicity from the sortase linking method could be an advantage to create attenuated toxins for research use. It might be also possible to deliberately attenuate the toxicity of ligated toxins in vivo by including additional linkers, such as peptide sequences that are sensitive to serum proteases, to further mitigate biosafety concerns.

Methods

Materials. Mouse monoclonal antibodies for syntaxin 1 (HPC-1, dilution 1:5,000), SNAP-25 (C1712, 1:5,000), VAMP2 (C169.1, 1:2,000), Syf (C72, 1:2,000) and Syt I (mAb848, 1:1,000) were generously provided by E. Chapman (Madison, WI, USA). Most are available from Synaptic Systems (Goettingen, Germany), with the exception of Syt I (mAb848), which is available from the Developmental Studies Hybridoma Bank. Rabbit polyclonal antibodies against VAMP4 (Cat. no. 136002, 1:1,000) and Sec22b (Cat. no. 186003, 1:1,000) were purchased from Synaptic Systems. The following mouse monoclonal antibodies were purchased from the indicated vendors: actin (Sigma, AC-15, 1:2,000); anti-HA (Covance, 16B12, 1:2,000); anti-Myc (Millipore, 9E10, 1:1,000). Equine polyclonal antisera against BoNT/X (BIPV/NT/DC, BoNT/A-F, and BoNT/G) were generously provided by S. Sharma (FDA). Goat polyclonal antibody against BoNT/D was purchased from Fisher Scientific (NB10062469). BoNTs were produced by Metabiologics (Madison, WI, USA). Antibody validation is available on the manufacturers’ websites. 293T (ORL-3216) cells were originally obtained from ATCC, which were negative for mycoplasma contamination but have not been authenticated.

cDNA and constructs. The cDNAs encoding X-LC (residues 1–439), X-H₃ (residues 803–1306), A-LC-H₃ (residues 1–874, GenBank no. M30196), and B-LC-H₃ (residues 1–860, GenBank no. AB229227) were synthesized by GenScript (New Brunswick, NJ, USA). The cDNA encoding X-H₃ was generated in-house using the Gibson assembly method. X-LC, A-LC (residues 1–425) and B-LC (residues 1–439) were cloned into pET22b vectors with His6-tag on their N termini. These proteins were expressed and purified at 11,000 r.p.m. for 12 h. The pellet was collected and solubilized for 30 min in 15 ml Tris-buffered saline (TBS: 20 mM Tris, 150 mM NaCl) for 20 min at room temperature before adding to BDE. The pellet was collected and solubilized for 30 min in 15 ml Tris-buffered saline (TBS: 20 mM Tris, 150 mM NaCl) plus 2% of Triton X-100 and a cocktail of protease inhibitors (Roche, CA). Samples were subsequently centrifuged at 17,000 r.p.m. for 20 min to remove insoluble materials. The final BDE concentration was ~2 mg ml⁻¹. VAMP2 (1–784) was expressed and purified using nickel-chelation chromatography (见NATURE COMMUNICATIONS) and dialyzed against 50 mM HEPES pH 7.2, 500 mM NaCl, 100 mM imidazole, 5% glycerol, 1 mM TCEP, and lyzed with an EmulsiFlex-C3 (Avestin, Mannheim, Germany) at 20,000 psi. Lysates were ultra-centrifuged at 200,000×g for 45 min. Supernatant was loaded onto a 15 ml Ni-NTA agarose column and washed with wash buffer (50 mM HEPES pH 7.2, 500 mM NaCl, 100 mM imidazole, 5% glycerol, 1 mM TCEP). Proteins were eluted with elution buffer (50 mM HEPES pH 7.2, 500 mM NaCl, 250 mM imidazole, 5% glycerol, 1 mM TCEP), and then dialyzed overnight in 50 mM HEPES, 500 mM NaCl, 5% glycerol, and 0.5 mM TCEP. Dialyzed fractions were further desalted with a PD-10 column (GE, 17-0851-01).

Large-scale production and purification of BoNT/XKY. cDNA encoding BoNT/XKY was assembled into a pET22b vector (K360A/Y363F), X-H₃ and X-H₃. It was cloned into a pET22b vector, with the His6-tag on its C terminus. The corresponding plasmid was transformed into E. coli BL21 (DE3). Cultures for expression were first grown using a LEX Bioreactor (Epphcyte3, Ontario, Canada) at 37°C in 1.5l of medium until OD₆₀₀ reached 0.8. The temperature was then reduced to 18°C for induction of expression with 1 mM IPTG, and grown for 16–17 h. Bacteria were harvested, re-suspended in HEPES buffer (50 mM HEPES pH 7.2, 300 mM NaCl, 25 mM imidazole, 5% glycerol, 250 mM NaCl, 5% glycerol, 1 mM TCEP), and then dialyzed overnight in 50 mM HEPES, 500 mM NaCl, 5% glycerol, and 0.5 mM TCEP. Dialyzed fractions were concentrated before being loaded on a Superdex200-16/60 column equilibrated in the same buffer used for dialysis. Elution peak corresponding to BoNT/X was collected and concentrated to ~10 mg ml⁻¹. Sample was aliquoted and flash-frozen in liquid nitrogen for storage at ~80°C.

Cleavage of SNARE proteins in rat BCE. Rat brain was homogenized in 15 ml 320 mM sucrose buffer, followed by centrifugation at 5,000 r.p.m. for 2 min at 4°C. Supernatants were collected and centrifuged at 11,000 r.p.m. for 12 min. The pellet was collected and solubilized for 30 min in 15 ml Tris-buffered saline (TBS: 20 mM Tris, 150 mM NaCl) plus 2% of Triton X-100 and a cocktail of protease inhibitors (Roche, CA). Samples were subsequently centrifuged at 17,000 r.p.m. for 20 min to remove insoluble materials. The final BDE concentration was ~2 mg ml⁻¹. VAMP2 (1–784) was expressed and purified using nickel-chelation chromatography (见NATURE COMMUNICATIONS) and dialyzed against 50 mM HEPES pH 7.2, 500 mM NaCl, 250 mM imidazole, 5% glycerol, 1 mM TCEP, and then dialyzed overnight in 50 mM HEPES, 500 mM NaCl, 5% glycerol, and 0.5 mM TCEP. Dialyzed fractions were further desalted with a PD-10 column (GE, 17-0851-01).

Cleavage of recombinant VAMPs by XLC. VAMP2 (1–93) was expressed and purified as a His6-tagged protein. VAMP2 (33–86), VAMP4 (1–115), VAMP5 (1–70) and Ykt6 (1–192) were expressed and purified as GST-tagged proteins. These proteins were pre-incubated in TBS buffer at 37°C. Samples were either analysed by SDS–PAGE gels and Coomassie Blue staining, or subjected to mass spectrometry analysis. Cleavage of VAMPs in cell lysates: Full-length HA-tagged VAMP1, 3, 7, 8 and Myc-tagged Sec22b, and Ykt6 were transfected into 293T cells using PolyJet transfection reagents (SignaGen, MD) following the manufacturer’s instructions. Cell lysates were harvested 48 h later in RIPA buffer (50 mM Tris, 1% NP40, 150 mM NaCl, 0.5% sodium deoxycholate, 0.1% SDS, 200 μl per 10-cm dish) plus a

Bioinformatics. The Uniprot database was searched with Jackhammer on the HMMER web server, using a BoNT/A sequence as the seed (Uniprot accession number ASH7Z9) until convergence. Returned sequences were aligned with Clustal Omega and a NeighborNet phylogenetic network estimated with SplitsTree.

Protein purification. E. coli BL21 (DE3) was used for protein expression. In general, induction of expression was carried out with 0.1 mM IPTG at 22°C overnight. Bacterial pellets were disrupted in lysis buffer (50 mM Tris pH 7.5, 130 mM NaCl) by sonication, and supernatants were collected after centrifugation at 20,000 g for 30 min at 4°C. Protein purification was carried out using AKTA Prime FPLC system (GE), and purified proteins were further desalted with a PD-10 column (GE, 17-0851-01).

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protease inhibitor cocktail (Sigma-Aldrich). Cell lysates (250 μl) were incubated with X-LC (0.5 mM) for 1 h at 37 °C. Samples were then analysed by immunoblot.

Identification of cleavage sites in VAMPs by LC–MS/MS. Samples were analysed at Taplin Biological Mass Spectrometry Core Facility at Harvard Medical School. For VAMP2, whole-protein samples were loaded onto a 100 μm internal diameter C18 reverse-phase HPLC column packed with 3 cm of beads off-line using a pressure cell. The column was re-attached to an Accela 600 Pump (Thermo Fisher Scientific). A rapid gradient of increasing acetonitrile was used to elute the peptide/protein from the HPLC column. As peptides eluted, they were subjected to electrospray ionization and then placed into an LTQ Orbitrap Velos Pro ion-trap mass spectrometer to acquire a high-resolution FTMS scan at 60,000 resolution, a second scan at low resolution in the ion trap, and a final scan to perform data-dependent MS/MS. The charge state envelopes were de-convoluted manually to obtain mono-isotopic masses when possible or average masses for the proteins. Peptide and protein identity were determined by matching protein databases with the acquired fragmentation pattern using the software program Sequest (Thermo Fisher Scientific). All databases include a reversed version of all the sequences, and the data were filtered to 1–2% peptide false-discovery rate.

For Yk6, VAMP4 and VAMP5, samples were first separated on SDS–PAGE. Protein bands were excised and cut into ~1 mm² pieces. Gel pieces were incubated with 50 mM ammonium bicarbonate solution containing 12.5 ng μl⁻¹ modified sequencing-grade chymotrypsin (Roche Diagnostics). Samples were digested overnight at 25 °C. Peptides were then extracted and separated with reverse-phase HPLC. As peptides were eluted, they were subjected to electrospray ionization and transferred into an LTQ Orbitrap Velos Pro ion-trap mass spectrometer (Thermo Fisher Scientific). Eluted peptides were detected, isolated and fragmented to produce a tandem mass spectrum of specific fragment ions for each peptide.

Identification of the protease cleavage site between LC and Hn. Histagged recombinant X-LC-Hn fragment (residues 1–891) was purified in E. coli and subjected to limited proteolysis by proteopin (Sigma P22889, 100:1 (toxin:trypsin) molar ratio) at 25 °C in 25 mM NaPO₄, 125 mM NaCl, 1% Triton X100, pH 7.4, for 1 h at 25 °C. The reaction was stopped by adding 50 μl of 2× SDS loading buffer. The samples were resolved on a 12% SDS–PAGE gel, and the proteins were transferred to a nitrocellulose membrane using a semi-dry method. The membrane was incubated with mouse anti-X antibody as primary antibody, and with goat anti-mouse IgG conjugated to horseradish peroxidase as secondary antibody. The membrane was developed under the ECL method. The BoNT/X sample was composed of X-LC-HN and X-HC at 1:1 molar ratio. Identification of cleavage sites in VAMPs by LC–MS/MS. Samples were treated with Lys-C were labelled with the heavy TMT. Both samples were protease inhibitor cocktail (Sigma-Aldrich). Lysates were centrifuged for 10 min at maximum speed using a pressure cell. The column was re-attached to an Accela 600 Pump (Thermo Fisher Scientific). Eluted peptides were detected, isolated and fragmented to produce a tandem mass spectrum of specific fragment ions for each peptide.

Cysteine alkylation by NEM. Lys-C-activated X-LC-Hs fragment was diluted into sodium phosphate buffer (10 mM, pH 6.5) at a final concentration of 0.3 mg ml⁻¹, with or without NEM at indicated concentrations (20, 10 and 5 mM) and incubated for 10 min at room temperature. NEM was freshly prepared in sodium phosphate buffer. Samples were mixed with 3× neutral loading dye (200 mM Tris pH 6.8, 30% glycerol, 6% Lithium Dodecyl sulfate, 10 mM NEM, and 0.06% BPP) at room temperature for 10 min, heated for 10 min at 55 °C, and then analysed by SDS–PAGE and Coomassie Blue staining.

Neuron culture and immunoblot analysis. Primary rat cortical neurons were prepared from E18–19 embryos by a papain dissociation kit (Worthington Biochemical) following the manufacturer’s instructions. Neurons were exposed to either BoNT/X fragments or sortase ligation mixtures in culture medium for 12 h. Muscle paralysis and the spread of hind paw in response to a startle stimulus were observed 12 h after injection as previously described.14

Biosafety and biosecurity. All procedures were approved by the Institute of Biosafety Committee at Boston Children’s Hospital and at Stockholm University. To ensure biosafety and biosecurity, no active full-length toxin was produced in any form. The amount of sortase linking reaction is strictly controlled to ensure that only a minimal amount of ligated toxins was produced, which was immediately utilized for functional studies. The loss of toxicity of BoNT/X₁₅₋₋ and reduced toxicity of sortase-linked X-Fl were confirmed using the mouse lethality assay. Inactive BoNT/X₁₅₋₋ was prepared and is available for developing neutralizing antibodies.

Data availability. The data and materials that support the findings of this study are available from the corresponding authors upon request.

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Acknowledgements

We thank members of the Dong and Stenmark labs for discussion, suggestions and comments. We would like to acknowledge Daniel Nilsson for his technical assistance. We thank Edwin R. Chapman (Univ. of Wisconsin—Madison), Shashi Sharma (FDA), Chuan Hu (University of Louisville), Bradley L. Pentelute (MIT) and Jesse C. Hay (University of Montana) for providing antibodies, cDNAs and other reagents. This study was supported in part by NIH IR1FNS808033 (to M.D.), the Swedish Research Council (2014-5667), the Wenner-Gren Foundation, and the Swedish Cancer Society (to P.S.). M.D. also holds the Investigator in the Pathogenesis of Infectious Disease award from the Burroughs Wellcome Fund.

Author contributions

P.S. identified BoNT/X and analysed its gene cluster. P.S. and D.L. carried out the bioinformatics analysis. P.S., G.M., M.M.-C. and L.H. purified the full-length BoNT/Xavr, P.S., S.Z. and M.D. designed and performed all other experiments and analysed data. J.Z. performed the DAS assay. Y.S. and S.-I.M. synthesized the cDNA for X-LC-HN of BoNT/Xavr and analysed its gene cluster. M.D. also holds the Investigator in the Pathogenesis of Infectious Disease award from the Burroughs Wellcome Fund.

Additional information

Supplementary Information accompanies this paper at http://www.nature.com/.

Competing financial interests: Boston Children’s Hospital has filed a provisional patent application (U.S. 62/380,239) for medical use of BoNT/X, with P.S., S.Z. and M.D. as inventors.

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How to cite this article: Zhang, S. et al. Identification and characterization of a novel botulinum neurotoxin. Nat. Commun. 8, 14130 doi: 10.1038/ncomms14130 (2017).

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