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Published paper

Abstract
Soluble N-ethylmaleimide sensitive factor attachment protein receptors (SNAREs) are crucial for exocytosis, trafficking, and neurite outgrowth, where vesicular SNAREs are directed toward their partner target SNAREs: synaptosomal-associated protein of 25 kDa and syntaxin. SNARE proteins are normally membrane bound, but can be cleaved and released by botulinum neurotoxins. We found that botulinum proteases types C and D can easily be transduced into endocrine cells using DNA-transfection reagents. Following administration of the C and D proteases into normally refractory Neuro2A neuroblastoma cells, the SNARE proteins were cleaved with high efficiency within hours. Remarkably, botulinum protease exposures led to cytotoxicity evidenced by spectrophotometric assays and propidium iodide penetration into the nuclei. Direct delivery of SNARE fragments into the neuroblastoma cells reduced viability similar to botulinum proteases’ application. We observed synergistic cytotoxic effects of the botulinum proteases, which may be explained by the release and interaction of soluble SNARE fragments. We show for the first time that previously observed cytotoxicity of botulinum neurotoxins/C in neurons could be achieved in cells of neuroendocrine origin with implications for medical uses of botulinum preparations.

Keywords: botulinum, cytotoxicity, neuro2A, SNARE, syntaxin, transfection reagents.


Original Article

Botulinum protease-cleaved SNARE fragments induce cytotoxicity in neuroblastoma cells

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Vesicular transport, protein trafficking, and exocytosis necessitate membrane fusion of vesicles, which is catalyzed by soluble N-ethylmaleimide sensitive factor attachment protein receptors (SNAREs) (Bock et al. 2001; Jahn et al. 2003; Meunier et al. 2003; Bajohrs et al. 2005; Sudhof and Rothman 2009; Mohrmann et al. 2010; Gao et al. 2012; Risselada and Grubmuller 2012). SNAREs are divided into two categories depending on their cellular locations: v-SNAREs on vesicles (also known as synaptobrevins; or vesicle-associated membrane proteins) and the t-SNAREs located on the target membrane, the synaptosome-associated protein of 25 kDa (SNAP25), and syntaxin (Bajohrs et al. 2005). SNAREs form a highly stable, tetrahedral coil–coil bundle that fuses the vesicle to the target membrane (Fig. 1a). Syntaxin and synaptobrevin each contribute one α-helix, whereas the SNAP25 protein contributes two (Wendler and Tooze 2001; Sudhof and Rothman 2009). There have been over 35 different SNARE isoforms identified, each associated with distinct organelles that participate in the precise and scrupulous vesicle fusion essential for proper cellular function (Bock et al. 2001; Wendler and Tooze 2001; Koticha et al. 2002; Xu and Xu 2008).
Botulinum neurotoxins (BoNT) have been invaluable in the studies of SNARE proteins and exocytosis (Erbguth and Naumann 1999; Sorensen et al. 2002). BoNTs are natively produced by anaerobic bacteria of the genus Clostridium and are responsible for the deadly disease called botulism manifested by neuromuscular paralysis (Erbguth and Naumann 1999; Montecucco and Molgo 2005). The BoNTs carry selective SNARE proteases, which have highly precise substrate recognition sites (Montecucco and Schiavo 1994; Montal 2010). BoNTs have been utilized in many medical and biotechnological applications (Davletov et al. 2005; Chaddock and Marks 2006; Foster et al. 2006; Ferrari et al. 2011; Arsenault et al. 2013). A typical BoNT is expressed as a single-chain precursor protein that is processed into two polypeptide chains. The SNARE protease (50 kDa light chain) located at the N terminus is linked, via a disulfide bond, to the C terminal part (100 kDa heavy chain) which is composed of the translocation domain and the receptor-binding domain (Chaddock and Marks 2006; Binz and Rummel 2009). Among seven commonly known BoNT serotypes (A–G) BoNT/A, C and E cleave SNAP25, BoNT/B, D, F, and G cleave synaptobrevins, and BoNT/C also cleaves syntaxin (Montecucco and Schiavo 1994; Rummel et al. 2004; Dong et al. 2006; Antonucci et al. 2008; Binz et al. 2010). To reach their intraneuronal substrates, BoNTs first bind neuronal surface gangliosides and then a synaptic vesicle protein on the pre-synaptic membrane for subsequent internalization (Montecucco et al. 1994; Rummel et al. 2004, 2009; Dong et al. 2006; Mahrohd et al. 2006; Binz and Rummel 2009). Once the internalized vesicle acidifies, the translocation domain forms a putative protein transduction channel that enables translocation of the protease into the cytosol following reduction in its disulfide bond (Koriazova and Montal 2003; Puhar et al. 2004; Fischer 2013; Pirazzini et al. 2013). BoNTs are potentially lethal as they paralyze muscles, such as the diaphragm, but are thought to be non-cytotoxic to neurons with the clear exception of BoNT/C and possibly BoNT/E (Williamson and
Neale 1998; Berliocchi et al. 2005; Zhao et al. 2010; Peng et al. 2013). The underlying causes of BoNT/C-mediated neurotoxicity are still under investigation. BoNT/C-released syntaxin fragment (aa 1-253) may cause cytotoxicity, but was thought to be rapidly degraded as observed by the loss of syntaxin immunoreactivity (Foran et al. 2003). However, immunoreactivity is not necessarily a measure of protein presence and other studies have been able to identify syntaxin fragments in cell extracts (Tsukamoto et al. 2012). If these fragments are free to diffuse and to engage SNAP25 and synaptobrevin or other non-neuronal SNAREs (Fasshauer et al. 1999), then they may potentially interfere with the scrupulous fusion events. Others have reported that BoNT/C-affected neurons first undergo axonal degeneration, possibly because of trafficking problems, followed by apoptosis (Berliocchi et al. 2005; Zhao et al. 2010; Peng et al. 2013).

BoNT protease-based medicines have recently been proposed for the treatment of endocrine disorders and therefore we investigated the possible cytotoxic effects of SNARE cleavage in endocrine cells. We have utilized model neuroblastoma cells (N2A) that abundantly express all three neuronal SNARE proteins: syntaxin 1, SNAP25, and synaptobrevin 2 (Syb2). N2A cells have been shown to be resistant to native BoNTs because of lack of neuronal gangliosides (Yowler et al. 2002). We thus used transfection reagents to internalize the proteases into the cell interior as it was shown before that polycationic lipid and polymers can enable the endocytosis and translocation of the BoNT type A protease as well as other peptides and proteins into cells (Kuo et al. 2010; Oba and Tanaka 2012). We now show that a selection of transfection reagents can mediate the intracellular entry of the botulinum protease serotypes C and D into neuroblastoma cells allowing investigation of botulinum action. Our results show that the intracellular cleavage of syntaxin and SNAP25 by the type C protease and synaptobrevin by the type D protease can cause severe cytotoxic effects. The soluble SNARE fragments were able to form ternary complexes and when transduced, triggered loss of cell viability mimicking the type C and D protease effects.

Materials and methods

Cell culture

Mouse Neuro2A cells (ATCC: CCL-131; LGC Standards; Teddington, UK) were grown in a 37°C incubator at 5% CO2 in low-glucose Dulbecco’s modified eagle medium (Gibco; Paisley, UK) supplemented with 10% fetal calf serum (HyClone; Fisher Scientific; Loughborough, UK) and 1% penicillin/streptomycin (P/S) (Invitrogen; Paisley, UK). Every 3–4 days, cells were washed with phosphate-buffered saline (PBS) and resuspended in culture medium using flow pressure, then counted by hemacytometer. Min6 and SH-SY5Y cells were cultured as previously described (Arsenault et al. 2013). Rat cortical neuronal cells were isolated as previously described (Arsenault et al. 2013). Sprague–Dawley rats of both sexes were obtained from the Medical Research Council Laboratory of Molecular Biology’s Biological Services Group. Experiments were approved by the Medical Research Council Laboratory of Molecular Biology, Cambridge. Hela cells were cultured in high-glucose Dulbecco’s modified eagle medium with 10% fetal clone 1 calf serum and 1% P/S. Cells were plated at 1 × 106 cells per 9 cm culture dish (BD Biosciences; San Jose, CA, USA), at 8 × 105 cells per well in uncoated 24-well plates (BD Biosciences) with or without cover slips, or at 8 × 105 cells per well in uncoated 96-well plates (BD Bioscience).

Protein and peptide synthesis

BoNT/A1 (1-872), rat SNAP25B (22-206) (all four Cys mutated to Ala), and rat synaptobrevin 2 (25-84) were prepared as Glutathione S-transferase (GST)-tagged proteins cleavable by thrombin and purified as previously described (Darios et al. 2010; Ferrari et al. 2012). Syntaxin 1 (1-226) was obtained from ATgen (Bio Trend, Köln, Germany). The production of serotype C and D proteases was described elsewhere (Vaidyanathan et al. 1999; Sikorra et al. 2006). Rat synaptobrevin 2 (25-52) Ac-RLQQTQAQVEDEVDMRNVN DKVLERD-NH2 and syntaxin 1A (201-245) Ac-EIIELENS IRELHDMFMDMAKLVESQGMIDRIYNEVHAVDYYE-NH2 peptides were prepared as previously described (Darios et al. 2010; Ferrari et al. 2012). Rat syntaxin 1A conjugated to Fluorescein isothiocyanate (FITC) FITC-Ahx-EIIELENSIRELHDMFMDMAKLVESQGMIDRIYNEVHAVDYYE-NH2, rat SNAP25B (22-206) Ac-RLQQTQAQVEDEVDMRNVN DKVLERD-NH2, and syntaxin 1A conjugated to Fluorescein isothiocyanate (FITC) FITC-Ahx-EIIELENSIRELHDMFMDMAKLVESQGMIDRIYNEVHAVDYYE-NH2, rat complexin (31-59) Ac-GGGERKAKYAKRAMEERVMRQGIRDKYGIKKG-NH2, rat synaptobrevin 2 conjugated to FITC (31-55) sequence FITC-Ahx-RQQTQAQVEDVDMRNVN DKVLERD-NH2 and penetratin Ac-RQIKIWFQNRRMKWKK-NH2 were synthesized by Peptide Synthetics (Southampton, UK). Protein and peptide concentration was determined by Pierce bichinonic acid (BCA) protein assay kit (Thermo-scientific; Loughborough, UK) according to manufacturer protocol.

Protein transduction

Transduction was performed 24 h after plating with proteins and peptides at desired concentrations that were pre-incubated for 30 min in 100 μL of Opti-MEM (GIBCO) with 2.5 μL of transfection reagents (per 500 μL of culture medium in a 24-well plate). Lipofectamine (Invitrogen), Lipofectamine LTX (Invitrogen), Transpass P (New England BioLabs; Hitchin, UK), and Fugene HD (Promega; Southampton, UK) were used as received from manufacturer. Pro-Ject (Thermo Scientific) was prepared according to manufacturer protocol. Cells were incubated with proteins and transfection reagents at 37°C in cell culture incubator for 42 h or as otherwise indicated.

Confocal microscopy

Cells were grown on cover slips in 24-well plates and treated with compounds 16 h before fixation. Wells were washed once with PBS, and incubated in 4% paraformaldehyde (Alfa Aesar; Hayesham, UK) in PBS for 20 min at 22°C. Wells were washed three times with PBS for 5 min and incubated 10 min in 10 mM NH4Cl (Sigma-Aldrich; Dorset, UK). The cells were then washed with PBS, and incubated in 4% paraformaldehyde (Alfa Aesar; Hayesham, UK) in PBS for 20 min at 22°C. Wells were washed three times with PBS for 5 min and incubated 10 min in 10 mM NH4Cl (Sigma-Aldrich; Dorset, UK). The cells were then
were transferred on Immobilon-P polyvinylidene fluoride membranes (EMD Millipore), and then incubated for 30 min with Alexa Fluor® 488 goat anti-rabbit IgG (H+L) and/or Alexa Fluor® 594 goat anti-mouse IgG (H+L) (Invitrogen) diluted 1 : 800 in 5% BSA in PBS at 22°C. Wells were washed three times in PBS. Cover slips were over Turned onto Vectashield (Vectorlabs; Oroton Southgate, UK) mounting medium, sealed with nail varnish, and visualized on Zeiss 710 (Cambridge, UK) on 10 or 63X. The fluorescent gains intensities and pinhole size (1 AU) were identical between experimental samples.

### Western immunoblotting

The medium was removed from the wells and cells were incubated for 5 min in 100 µL loading buffer (56 mM sodium dodecyl sulfate (Sigma-Aldrich), 0.05 M Tris-Cl (Bio-Rad; Hemel Hempstead, UK) pH 6.8, 1.6 mM UltraPure EDTA (Gibco), 6.25% glycerol (Fisher Scientific), and 0.00001% bromophenol blue (Fisher Scientific)]. One unit of benzoinase (Novagen; EMD Millipore) supplemented with 1 µL of 1 M MgCl₂, was added to each well and plates were shaken at 1500 rpm for an additional 10 min. Samples were boiled for 1 min at 95°C then run on 12% Bis-Tris sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS–PAGE) gels (Invitrogen). Migrated samples were transferred on Immobilon-P polyvinylidene fluoride membranes (EMD Millipore), and then incubated for 30 min in blotting solution (5% milk, 0.1% TWEEN 20 (Thermo Scientific) in PBS). All primary antibodies, including rabbit polyclonal anti-synaptobrevin 2 (clone 69.1; Synaptic Systems), mouse monoclonal anti-syntaxin 1 (clone HPC-1; Sigma-Aldrich), were incubated for 30 min in a permeabilization solution composed of 1% Triton X-100 (Sigma-Aldrich) in PBS containing appropriate primary antibodies for 90 min at 22°C. Mouse monoclonal anti-SNAP25 (SM181; Novagen; EMD Millipore; Feltham, UK), rabbit polyclonal anti-synaptobrevin 1/2/3 antibody (Synaptic Systems; Goettingen, Germany), and mouse monoclonal anti-syntaxin 1 (clone HPC-1; Sigma-Aldrich), were diluted at 1 : 500. The wells were washed three times in PBS, then incubated for 30 min with Alexa Fluor® 594 goat anti-mouse IgG (H+L) and/or Alexa Fluor® 488 goat anti-rabbit IgG (H+L) (Invitrogen) diluted 1 : 800 in 5% BSA in PBS at 22°C. Cells treated with compounds and transfection reagents were washed twice with PBS and resuspended in 10 mM HEPES (Fisher Scientific), 140 mM NaCl, 2.5 mM CaCl₂, pH 7.4 with 2 µg/mL propidium iodide (PI; Sigma-Aldrich) into 12 × 75 mm round bottom test tube (Scientific Laboratory Supplies; Wilford, UK). After 10 min of incubation with PI, fluorescent intensities of the cell populations were measured using FACScalibur 2 (BD Bioscience). Proper gating was determined by untreated control, and laser intensity was determined by PI- and PI+ controls during each experiment. Data were interpreted using FlowJo version 9.4.4 (Tree Star Inc.; Ashland, USA).

### Flow cytometry

SNARE complex reactions

GST-synaptobrevin 2 (25-84) was incubated with or without the type D protease for 6 h at 22°C. Glutathione Sepharose 4B beads (GE Healthcare; Little Chalfont, UK) were added and incubated overnight at 4°C under constant rotation. Samples were washed four times with buffer A (0.8% w/v n-octyl-β-D-glucopyranoside (Sigma-Aldrich), 100 mM NaCl, 20 mM HEPES, pH 7.4) and divided into two tubes. A 1 : 1 ratio of syntaxin 1A (201-245) and SNAP25B (22-206) was added to one sample while syntaxin and buffer A were added to the other and incubated for 3 h at 22°C. Samples were washed four times with buffer A then heated at 95°C for 1 min and loaded on 10% Bis-Tris SDS–PAGE gel (Invitrogen). For the second assay GST-SNAP25 (22-206) was added to Glutathione Sepharose 4B beads (GE Healthcare) and incubated for 90 min at 4°C under constant rotation. Samples were washed four times with buffer A (0.8% w/v n-octyl-β-D-glucopyranoside (Sigma-Aldrich), 100 mM NaCl, 20 mM HEPES, pH 7.4) and divided into four tubes. Buffer A was added to the first sample, FITC-synaptobrevin (31-55) was added to the second sample. A 1 : 1 ratio of FITC-syntaxin 1A (201-245) and FITC-synaptobrevin (31-55) was added to the third sample and a 1 : 1 ratio of syntaxin 1A (1-226) and FITC-synaptobrevin (31-55) was added to the fourth sample. Samples were incubated for 30 min at 22°C then washed three times with buffer A, heated at 95°C for 1 min, and loaded on 10% Bis-Tris SDS–PAGE gel (Invitrogen). ChemiDoc XRS (Bio-Rad) was used to capture fluorescent and Coomassie images of the gels.

### Statistical analysis

All experiments were performed in at least three independent experiments. Results are presented as mean ± standard deviation (SD). Data analysis was performed using Graphpad Prism 5.0 (La Jolla, CA, USA). The unpaired two-tailed Student’s t-test was used for comparison. A p < 0.05 was considered statistically significant.
Results

Direct delivery of botulinum proteases into neuroblastoma cells

First, we ascertained the presence of the SNARE proteins syntaxin, SNAP25, and synaptobrevin in mouse neuroblastoma N2A cells by confocal microscopy. Fig. 1b shows that the t-SNAREs (red) are located on the plasma membrane, whereas the v-SNARE (green) is located in intracellular vesicular pools (Bock et al. 2001; Koticha et al. 2002; Sudhof and Rothman 2009).

Next, we explored a possibility of delivering botulinum proteases, which cleave membrane-embedded SNAREs, into normally resistant N2A cells. As it has been observed that transfection reagents are capable of causing the internalization of the Botulinum A and E proteases targeting SNAP25 (Kuo et al. 2010), we tried an array of transfection reagents to deliver the type C and D proteases, which, respectively, target t-SNAREs and the v-SNARE. Fig. 1c and d (upper panels) show that Lipofectamine LTX and FuGENE HD were able to deliver the C and D proteases into N2A cells very efficiently as evidenced by changes observed in immunoblotting of the SNARE proteins. TransPass, a bona fide protein transduction reagent, was also efficient, but caused cell death as evidenced by a reduction in total SNAP25 protein in the loading controls at the tested dose (Fig. 1c and d upper panels). We chose to use Lipofectamine LTX for subsequent experiments because of its excellent tolerance by cells. Fig. 1c and d (middle panels) show that a dose of 0.2 μg/mL of botulinum proteases was sufficient to achieve near-complete SNARE cleavage upon 20 h incubation. This dose was used in all subsequent experiments. Fig. 1c (bottom panel) also shows a time course of type C protease activity, where a complete cleavage of the syntaxin substrate can be observed even within 4 h, whereas the SNAP25 cleavage slightly lags behind and is completed within 8 h. We also observed a near-complete cleavage of synaptobrevin by the D protease within 4 h (Fig. 1d, bottom panel). Together, extensive cleavage of all three SNAREs in Lipofectamine LTX-treated cells can be observed in the presence of the respective botulinum proteases at 0.2 μg/mL within 8 h.

Cytotoxic effects of the botulinum protease serotypes C and D

It is well-known that BoNT/C has neurotoxic properties, but botulinum effects on survival of neuroendocrine cells have not been specifically addressed. To test cell viability of neuroblastoma cells, we used a cell counting kit (CCK-8). Fig. 2a (left panel) shows the normalized signals exhibited by N2A cells following treatment with botulinum proteases. The type C protease significantly reduced cell viability after a 40 h exposure (*p < 0.01). Curiously, this effect was enhanced with the addition of the type D protease, whereas the type D protease alone was not sufficient to significantly affect cell viability. As dehydrogenase activity might be affected by botulinum-induced changes in membrane trafficking, we corroborated our observations using the BCA protein assay which measures total protein content present in the wells which is a direct measure of adherent cells. The protein content of adherent cells (Fig. 2a, right panel) precisely correlated with the CCK-8 kit results (*p < 0.01; **p < 0.001). Lipofectamine LTX on its own did not affect cell viability (Fig. 2a). In further experiments, we chose to study the combined effect of type C and D proteases as together they mediated the most potent cytotoxicity. Fig. 2b shows a confocal image of botulinum-treated cells versus control cells, with a substantial reduction in the number of cells evident following the protease treatment. Next, we tested other cell lines of neuroendocrine origin and which carry syntaxin 1, SNAP25, and synaptobrevin 2 (Fig. 2c). We observed a significant cytotoxic trend in the human neuroblastoma cell line SH-SY5Y (*p < 0.01) and substantial cytotoxic effect in mouse insulinoma Min6 cells (*p < 0.01) and neurons (*p < 0.01) compared with their respective untreated controls (100%). The non-neuroendocrine human cell line Hela did not show loss of survival which can be explained by the low levels of botulinum proteolytic substrates in these cells.

We next investigated the time course and features of the N2A cell demise. Fig. 3a shows that the loss in cell viability following botulinum treatments (C and D) continues to occur for at least 40 h. The type C protease showed a significant reduction in viability after 36 h (*p < 0.01), whereas the type C and D protease displayed a significant reduction as early as 16 h (*p < 0.01). When observed under higher magnification (Fig. 3b), the botulinum protease C- and D-treated cells often displayed nuclear abnormalities, as would be expected during programmed cell death (Danial and Korsmeyer 2004). For flow cytometry, cells with normal morphology were gated according to the forward and side scattering patterns using untreated cells as a control and then total percentages of gated cells were calculated (Fig. 3c). There was a significant reduction (p < 0.05) in the percentage of morphologically normal cells in the sample cotreated with C and D proteases. Fig. 3d shows the analysis of these gated cells proportional to their PI labeling, a widely used necrosis and late apoptosis marker (Lecoeur 2002). A significant rightward shift in PI-positive cells suggests increased cell death of the C and D proteases-treated cells (p < 0.01). We could not rely on the Annexin V-FITC labeling of apoptotic cells as lipofection by itself alters lipid balance in the plasma membrane and thus gives artificially high Annexin V binding without affecting cell survival.

Botulinum protease-cleaved SNARE fragments mediate cytotoxicity

As botulinum type C protease cleaves both syntaxin and SNAP25, we next aimed to address whether cleavage of
syntaxin or SNAP25 is responsible for the cytotoxic effects observed above. We used the BoNT/A protease, the main ingredient of the BOTOX and Dysport preparations, which exclusively cleaves SNAP25. Fig. 4a shows the comparative proteolytic activity of the type A and C proteases as seen in western immunoblotting. The major difference which can be observed between the A and C proteases is the cleavage of syntaxin 1. Fig. 4b shows a significant reduction in cell viability exclusively when the C protease is used together with the type D protease, whereas the D protease alone or the A and D proteases in combination exhibit no discernible cytotoxicity. This result indicates that it is the cleavage of syntaxin by the type C protease that is the driving force for the observed cytotoxicity.

Finally, we tested whether transduction of type C protease does result in production of syntaxin fragments, as previous results (Fig. 1) indicated loss of immunoreactivity. We used an antibody directed against the head domain of syntaxin 1 (clone HPC-1) and found a clear appearance of the syntaxin proteolytic fragment within several hours of application of the botulinum protease at a dose of 0.002 μg/mL (Fig. 5a).

As botulinum-cleaved syntaxin and synaptobrevin products are cleaved from their transmembrane anchors, they could be soluble and may potentially form aberrant SNARE complexes. We tested whether short SNARE fragments can form such ternary complexes in bead pull-down assays. We first used GST-synaptobrevin 2 attached to beads. As can be seen in Fig. 5b, the D protease cleaved the bead attached GST-synaptobrevin 2 (22-84) very efficiently generating a cleavage product (GST-Syb2D product) still attached to beads through its N-terminal GST fusion. The BoNT/C protease...
produces the syntaxin 1-253 aa fragment which encompasses the short syntaxin peptide 201-245 (Ferrari et al. 2012). Fig. 5b also shows that GST-synaptobrevin, cleaved or not, was able to pull-down both SNAP25 and the short syntaxin fragment. This suggests that botulinum protease C- and D-released products can still interact to form ternary SNARE complexes. Furthermore, we used GST-SNAP25 bound to beads to investigate SNARE interactions with truncated syntaxin 1-226 and a minimal synaptobrevin further reduced to only 25 amino acids. Fig. 5c shows that the short synaptobrevin and syntaxin fragments, with or without the syntaxin regulatory head domain, can still form SNARE complexes evidenced in the pull-down reactions.

To investigate whether shortened SNARE fragments can trigger cytotoxicity, we treated the N2A cells with syntaxin 1 (201-245, 5 μg/mL) and/or synaptobrevin 2 (25-52, 5 μg/mL) peptides, with or without Lipofectamine LTX. Complexin, a SNARE protein that does not directly contribute to the tetrahelical bundle was used as control. Penetratin, a cell-penetrating peptide, was also used as a negative control to eliminate the possibility that non-specific internalization might contribute to cytotoxicity. Fig. 5d shows that N2A cells treated with the shortened SNARE peptides exhibited decreased cell survival compared with control conditions. The syntaxin fragment alone was very efficient even at low concentrations (Fig. 5d and e), thereby masking possible cumulative effects. No decrease in viability was observed with the same concentrations of complexin or penetratin demonstrating the importance of the SNARE ternary interactions involving syntaxin as the major driving factor in triggering cytotoxicity.

Discussion

Botulinum neurotoxins have been invaluable in the studies of the SNARE machinery. Here, we demonstrate a straightforward and easy method to deliver botulinum type D and C proteases into normally refractory cells to cleave membrane-embedded v- and t-SNAREs. Our results expand upon previous observations of the delivery of SNAP25 proteases type A and E (Kuo et al. 2010). A number of studies highlighted BoNT/C cytotoxic effects on central and peripheral neurons, (Foran et al. 2003; Berliocchi et al. 2005; Zhao et al. 2010; Peng et al. 2013), but possible cytotoxic effects in normally resistant neuroendocrine cells have not been addressed. Our newly discovered ability to introduce botulinum proteases in all cultured cells suddenly revealed cytotoxic effects especially for the type C botulinum protease. We observed increase in nuclear PI labeling and nuclear abnormalities following botulinum type C and D protease application which correspond to necrosis and late apoptosis, respectively. This is consistent with the observed caspase 3 activation and apoptosis of cerebellar neurons upon BoNT/C administration (Berliocchi et al. 2005). Our data
show that botulinum protease-induced cell death can occur independent of axonal changes.

The precise mechanisms of botulinum cytotoxicity in neurons are not fully understood. As one possible mechanism which warrants further investigations, we propose that botulinum cleavage products that still contain SNARE interacting domains are able to compete with normal SNARE interplay and disrupt functionally important trafficking and vesicular fusion events. Our results showing synergistic effects of botulinum proteases on cytotoxicity of non-neuronal cells, as well ternary complex formation by shortened SNARE peptides, suggest that cellular well-being generally relies on proper SNARE localization and precisely regulated stoichiometry of SNARE interactions. Although a recent study suggested that the general presence of neuronal SNARE proteins is important for neuronal survival, this can be explained by the importance of intercellular signaling for the maintenance of neuronal circuitry (Peng et al. 2013). Similarly, it could be argued that cell–cell communication is important for cancer cell survival. However, it is well-known that cancer neuroendocrine cells sometimes lose secretory pathway components and yet they are able to grow and proliferate in a normal way (Pance et al. 2006). As botulinum-released SNARE fragments can form ternary complexes they may contribute to deregulation of precise membrane trafficking necessary for correct cell function (D’Alessandro and Meldolesi 2013).

Whereas botulinum type C protease possesses the most potent cytotoxic effects, our data show that it is specifically cleavage of syntaxin rather than SNAP25 which mediates cell toxicity. Indeed, when we tested the type A protease, which only cleaves SNAP25, no cytotoxic effects were observed. A previous study suggested that BoNT/C-cleaved syntaxin degrades too quickly to interfere with intracellular functions, but this inference was based on loss of immunoreactivity (Foran et al. 2003). When we probed neuroblastoma cells using a well-known syntaxin 1 antibody we observed a persistent accumulation of the syntaxin fragment triggered by the type C protease treatment which is in accord with a previous study (Tsukamoto et al. 2012); evidently the fragments are not completely degraded and may take part in aberrant SNARE interactions. We now show that syntaxin fragments, as short as 45 aa, can drive SNARE complex formation as evidenced by pull-downs with SNAP25 and synaptobrevin, even if the latter cleaved by the botulinum type D protease.

This study raises several questions for future investigations. The mechanism of Lipofectamine LTX-induced transduction of botulinum enzymes and small peptides remains unclear because of proprietary reasons and the active ingredients should be understood considering that this reagent was more effective than bona fide protein transduction preparations. It is also not clear, why generation of syntaxin fragments is more cytotoxic compared with synaptobrevin despite that in both cases botulinum proteases sever the SNARE membrane-embedded parts (Peng et al. 2013). It will be revealing to detect even smaller SNARE degradation products following botulinum treatment as small syntaxin and synaptobrevin fragments can still form irreversible SNARE complexes (Ferrari et al. 2012). This is also important in view of SNARE degradation and deregulation which apparently takes place in neurodegenerative disorders (Garcia-Reitbock et al. 2010). The importance of normal SNARE function was previously suggested by a genetic study where deletion of synaptic vesicle protein cysteine string protein α, a SNARE chaperon, led to massive neurodegeneration in vivo (Sharma et al. 2011).

Our results have implications for the use of BoNT/C in biomedical applications and further validate the safer cytological characteristics of BoNT/A-based preparations such as BOTOX and Dysport. BoNT proteases are currently of high pharmaceutical interest (Chaddock and Marks 2006; Chen and Barbieri 2009; Foster and Chaddock 2010; Davletov et al. 2012; Ferrari et al. 2013; Naumann et al. 2013) and the newly highlighted delivery mechanisms could greatly advance the ongoing botulinum research. Considering the observed cytotoxicity, the botulinum proteases could be utilized as molecular surgery tools for neuroendocrine cancer.

therapies. Targeted delivery of botulinum proteases could even be combined with short syntaxin peptides for cumulative cytotoxic effects. Many neuroendocrine tumors give rise to complicated endocrinopathies that require constant monitoring and problematic surgical interventions (Bangaru et al. 2010; Batcher et al. 2011). The possibility of using certain botulinum proteases/SNARE peptides to destroy cancerous neuroendocrine cells paves the way for devising strategies to both block the pathological release of hormones and concomitantly halt tumor proliferation.

Acknowledgments and conflict of interest disclosure

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All experiments were conducted in compliance with the ARRIVE guidelines. The authors have no conflict of interest to declare.

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