Neuritogenic Actions of Botulinum Neurotoxin A on Cultured Motor Neurons

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ABSTRACT

Botulinum neurotoxins (BoNTs) are extremely potent neuromuscular poisons that act through soluble N-ethylmaleimide-sensitive factor attachment protein receptor (SNARE) protein cleavage to inhibit neurotransmitter release. The ability of BoNT serotype A (BoNT/A) to eliminate localized transmitter release at extremely low doses is well characterized. In the current study, we investigated the less understood characteristic of BoNT/A to induce nerve outgrowth, sometimes referred to as sprouting. This phenomenon is generally considered a secondary response to the paralytic actions of BoNT/A, and other potential factors that may initiate this sprouting have not been investigated. Alternatively, we hypothesized that BoNT/A induces sprouting through presynaptic receptor activation that is independent of its known intracellular actions on the soluble N-ethylmaleimide-sensitive factor attachment receptor (SNARE) synaptosomal associated protein of 25 kDa (SNAP-25). To test this, the effects of BoNT/A application on neurite outgrowth were examined using primary cultures enriched with motor neurons isolated from embryonic mouse spinal cord. In this system, BoNT/A potently stimulated neuritogenesis at concentrations as low as 0.01 nM. The neuritogenic effects of BoNT/A exposure were concentration dependent and antagonized by *Triticum vulgaris* lectin, a known competitive antagonist of BoNT. Similar results were observed with the isolated BoNT/A binding domain, revealing that neuritogenesis could be initiated solely by the binding actions of BoNT/A. In addition, the presence or absence of SNAP-25 cleavage by BoNT/A was not a determinant factor in BoNT/A-induced neuritogenesis. Collectively, these results suggest that binding of BoNT/A to the motor neuronal membrane activates neuritogenesis through as yet undetermined intracellular pathway(s), independent of its known action on vesicular release.

Botulinum neurotoxin (BoNT), notorious as the agent responsible for the potentially fatal paralytic disease botulism, is an extremely potent neuromuscular poison (Arnon et al., 2001). Ironically, the toxin’s ability to virtually eliminate transmitter release within a relatively confined area at extremely low doses makes it highly desirable as a therapeutic agent (Jankovic, 2004; Erbguth, 2008). The seven BoNT serotypes (A–G) are zinc metalloproteases that, once activated, cleave to inhibit neurotransmitter release. The ability of BoNTs selectively cleave specific nerve terminal proteins known as SNAREs (Schiavo et al., 2000). These SNAREs form the core fusion machinery necessary to initiate vesicular neurotransmitter release. Selective proteolytic inactivation of SNARE proteins at the mammalian neuromuscular junction (NMJ) inhibits acetylcholine release causing flaccid muscle paralysis (Kalandakanond and Coffield, 2001a,b).

Although BoNTs have been studied for well over 50 years, their targeting of SNAREs was revealed less than 20 years ago. SNARE protein inactivation is now widely considered the primary intracellular action of the paralytic BoNTs. Since then, only a limited number of other potential intracellular actions have been investigated (Coffield et al., 1994; Ishida et al., 2004). An intriguing corollary to the paralytic activity of the toxins is the observation that after intoxication, some BoNT-poisoned nerve endings extend or “sprout” neurites. A number of studies have examined the importance of sprouting in toxin duration of action and subsequent NMJ remodeling (Angaut-Petit et al., 1990; Comella et al., 1993; Juzans et al., 1996; de Paiva et al., 1999). In addition, a few studies have reported the importance of molecular factors such as neural cell adhesion molecule, laminin, and insulin-like growth factor on BoNT-induced sprouting (Booth et al., 1990; Caroni et al., 1994; Schäfer and Wernig, 1998; Lee et al., 1999). Furthermore, the effects of BoNT on neurite outgrowth are considered a secondarily expressed action of the paralytic toxin. Nevertheless, the possible involvement of BoNT in neuronal remodeling is largely unexplored.

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**ABBREVIATIONS:** BoNT, botulinum neurotoxin; Hc, heavy chain; SNAP-25, synaptosomal associated protein of 25 kDa; BoNT/A, BoNT serotype A; TVL, *Triticum vulgaris* lectin; SNAP-25, synaptosomal associated protein of 25 kDa.
growth in cell culture have been reported by a few groups, with varying results (Bonner et al., 1994; Igarashi et al., 1996; Osen-Sand et al., 1996). None examined these effects on cultured spinal motor neurons. BoNT-induced sprouting is generally considered an indirect response secondary to denervation, and direct presynaptic events mediating this process have been largely ignored. It is possible, however, that the phenomenon is actually a result of intracellular signaling initiated by the direct interaction of BoNT with the presynaptic membrane. Thus, an examination of the potential actions of BoNT on motor neuron neuritogenesis merits further consideration.

In the current study, we examined the effects of intact BoNT serotype A (also known as BoNT/A holotoxin) and its isolated binding domain H2C on neurite outgrowth using motor neuron-enriched primary cultures isolated from the embryonic mouse spinal cord. Both BoNT/A holotoxin and BoNT/A H2C potently stimulated neuritogenesis. These neuritogenic effects were concentration dependent and abolished by the presence of Triticum vulgaris lectin (TVL), a plant protein that has affinity for sialic acid containing membrane components such as polysialogangliosides. TVL was chosen because it is a ligand for complex sialogangliosides (e.g., GTb1) known to act as BoNT receptors (Kozaki et al., 1998).

In addition, TVL has been shown to competitively antagonize the binding of BoNT/A to central nervous system preparations, to antagonize BoNT-induced paralysis, and to inhibit BoNT-induced cleavage of SNAP-25, the SNARE protein target of BoNT/A (Bakry et al., 1991; Blasi et al., 1993; Schiavo et al., 1993; Kalandakanond and Coffield, 2001b). SNAP-25 may also participate in neurite growth, although its exact role remains to be elucidated (Osen-Sand et al., 1996; Morihara et al., 1999; Kimura et al., 2003). Herein, we report that SNAP-25 was expressed in the cultured motor neurons and subsequently cleaved by BoNT/A. It is interesting that neurite outgrowth occurred regardless of whether SNAP-25 was intact or cleaved, indicating that this known intracellular target was not relevant to BoNT/A-induced neuritogenesis in embryonic motor neurons (eMNs). Collectively, these results suggest that binding of BoNT/A to the motor neuronal membrane activates neuritogenesis through undetermined intracellular pathway(s), independent of its known action on vesicle release.

In summary, the findings presented herein support a neuritogenic action for BoNT/A on cultured motor neurons. Further studies are necessary to: 1) clarify the intracellular pathways mediating this function, 2) determine whether this phenomenon holds true in older neurons, and 3) correlate these in vitro findings with BoNT-induced sprouting in vivo.

Materials and Methods

Cell Culture. All procedures involving animals were approved by the Institutional Animal Care and Use Committee. The protocol used was modified from Anderson et al. (2004). In brief, embryos were collected from timed pregnant NIH Swiss mice (gestation days 13–14). After removal from the dam, the embryonic spinal cords were isolated, cleaned of meninges, and then minced with scissors. The minced tissue was transferred to 0.025% trypsin solution for dissociation for 25 min at 37°C, with frequent agitation. The trypsinization step was terminated with the addition of soybean trypsin inhibitor in DNase-containing isolation buffer. After centrifugation, the dissociated tissue was triturated until evenly suspended. Tissue debris was removed by further centrifugation of the dissociated suspension through a bed of 4% bovine serum albumin at 300g for 10 min. The preparation was enriched with motor neurons by loading the cell suspension gently onto a Histodenz gradient (6.7%; Sigma-Aldrich, St. Louis, MO) that was then spun at 500g for 20 min. The motor neuron-enriched interface was collected by gentle aspiration and cleared of cellular debris by centrifugation through another bed of 4% bovine serum albumin at 300g for 10 min. The neuron containing fraction was resuspended in L-15 medium supplemented with 10% glucose, 2 mM L-glutamine, 5% horse serum, 7.5% sodium bicarbonate, 1% penicillin-streptomycin, 5 μg/ml insulin, 100 μg/ml putrescine, 100 μg/ml conalbumin, 3 μM sodium selenite, and 2 μM progesterone.

Neurite Outgrowth Assay. eMNs were plated in 24-well plates coated with poly-n-lysine plus laminin (BD Biosciences, San Jose, CA) at a density of 1 × 10^5 cells/ml. The cultures were maintained in a humidified incubator (5% CO2) at 37°C for 4 h to ensure cell attachment. After 4 h, the cells were treated with intact BoNT/A (holotoxin) or BoNT/A H2C plus or minus TVL and returned to the 37°C incubator for 24, 48, or 72 h. After treatment, the cultures were washed, and the cells were fixed in 4% paraformaldehyde, pH 7.4, for 30 min. After fixation, the cells were washed with phosphate-buffered saline, blocked with 5% goat serum for 30 min to prevent nonspecific binding, and then incubated overnight with a polyclonal antibody to protein gene product 9.5 (1:400; Serotec, Oxford, UK), a protein marker for neurons. Labeled neurons were detected by fluorescence using an fluorescein isothiocyanate-conjugated secondary IgG.

Neurons were visualized using an Olympus (Tokyo, Japan) inverted microscope, and images were captured with a charge-coupled device camera. IPLab Software was used to measure neurite length and to count primary and secondary neurite numbers for each treatment. Total neurite length was defined as the sum of the lengths of all neurites of a given neuron excluding the soma. Primary neurites were defined as neurites that arose directly from the neuronal soma; secondary neurites were defined as branches arising directly from primary neurites. One-way analysis of variance, followed by Bonferroni or Dunnnet post hoc tests, was used to test statistical significance (GraphPad Prism 5; GraphPad Software Inc., San Diego, CA).

Immunobots. Cells were harvested from culture plates at desired time points using T-PER protein extraction reagent (Pierce Chemical, Rockford, IL) plus protease inhibitor cocktail. Cell extracts were sonicated briefly before centrifugation. Supernatants were collected and processed for SDS-polyacrylamide gel electrophoresis followed by Western blot using standard protocols. Protein samples (10 μg/lane) were resolved on 12.5% Tris-HCl gels. Resolved proteins were transferred to polyvinylidene difluoride membranes and probed for SNAP-25 using a rabbit SNAP-25 primary antibody (1:15,000; Sigma-Aldrich). SNAP-25 immunoreactivity was visualized with chemiluminescence.

Biochemicals. BoNT/A holotoxin and BoNT/A H2 were purchased from Metabiologics, Inc. (Madison, WI). TVL (also known as WGA) was purchased from Sigma-Aldrich and used at 1 μM.

Results

BoNT/A Promotes Neurite Outgrowth of eMNs. Neurite outgrowth in vitro is dependent on several factors, including but not limited to cell origin, age, and culture conditions (e.g., plating density, media, substrate, growth factors, etc.). In our initial experiments, we examined neurite outgrowth from day 13 eMNs under control and BoNT/A-treated conditions at three time points (Fig. 1). Neurons were allowed to attach to laminin-coated plates before treatment. Substantial numbers of cells were attached and initiating primary neurite extension by 4 h after plating (Fig. 1B, inset). Between 24 and 48 h in culture, total neurite length in
untreated control neurons more than doubled, with a mean increase of 124%. An additional 76% increase was measured between 48 and 72 h (Fig. 1B, open circles). Altogether, total neurite length in untreated control eMNs increased by 295% between 24 and 72 h.

Initial assessment of neurite outgrowth in eMNs treated with a single concentration of BoNT/A holotoxin revealed a stimulatory effect on neurite outgrowth. In cultures treated with 0.1 nM BoNT/A, total neurite length was significantly greater (p < 0.001) in the BoNT/A-treated eMNs compared with time-matched, untreated control neurons at each of the three time points (Fig. 1A and B, open squares). Furthermore, the rate of neurite outgrowth was greater in the BoNT/A-treated cultures. Total neurite length increased by 156% between 24 and 48 h of treatment and by 174% between 48 and 72 h of treatment. Total neurite length in BoNT/A-treated eMNs increased by 444% between 24 and 72 h.

Neuritogenic Effects of BoNT/A Are Concentration-Dependent. The responses of neurite outgrowth to different concentrations of BoNT/A holotoxin were examined at 24 and 48 h and compared with time-matched untreated control neurons (Fig. 2). In general, the stimulatory effect on total neurite length increased with increasing concentrations of BoNT/A, with the maximal responses observed at 0.1 nM (Fig. 2, A and B). This was followed by a stabilization of the response, with a slight depression in response at the highest concentration (10 nM) at 24 h; however, this response was still significantly greater than untreated controls. These concentration-dependent increases in neurite length ranged from 68 to 152% at 24 h and from 29 to 82% at 48 h.

To examine whether BoNT treatment selectively stimulated primary or secondary neurite growth, primary and secondary neurite lengths were measured separately for two toxin concentrations (data not illustrated). As might be expected, total primary neurite length was greater than secondary neurite length at 24 h, even in untreated cultures. BoNT/A treatment strongly stimulated both primary and secondary neurite growth; however, there was no evidence of a selective effect of the toxin on the two classes of neurites. After 24-h BoNT/A treatment, primary neurite length increased significantly by 75.3 and 54.6% at 0.1 and 1.0 nM BoNT/A, respectively, whereas secondary neurite length increased by 41.8 and 77.2%, respectively. By 48 h, primary neurite length increased by 10.8 and 75.4% at 0.1 and 1.0 nM BoNT, respectively, whereas secondary neurite length increased by 109.9 and 110%, respectively.

In addition to neurite length, the numbers of primary neurite extensions and secondary branches were counted and compared with untreated controls (Fig. 2, C and D). At 24 h, the number of primary neurites in toxin-treated cultures increased by an average of 29% across all four concentra-
tions. The individual concentration responses were significantly different from untreated eMNs and demonstrated a similar pattern of concentration dependence as total neurite length \((p < 0.001)\). The number of primary neurites was also increased at 48 h, albeit to a lesser extent. As previously noted for total neurite length, the primary neurite response observed at the 10 nM concentration was also depressed at 48 h relative to the lower concentrations but was still significantly increased compared with controls. Lastly, BoNT/A treatment also significantly increased the number of secondary branches \((p < 0.001)\). The pattern of this response was very similar to that of the primary neurite response, although the magnitude of the branching response was substantially greater (note that the x-axis for Fig. 2D is 5 times that of Fig. 2C). Collectively, these data suggest that BoNT/A treatment promotes neuritogenesis in eMNs.

**SNAP-25 Is Present in eMNs and Cleaved by BoNT/A.** SNAP-25, the SNARE protein intracellular target of BoNT/A may participate in neurite growth in developing neurons, although its exact role is unclear. In the current study, Western blot analyses revealed the presence of stable SNAP-25 protein expression in the 13-day eMN cultures as early as 4.5 h postplating (Fig. 3, top). It is interesting that exposure to intact BoNT/A (0.1 nM) resulted in cleavage of SNAP-25 in the eMNs, coincident with promotion of neurite outgrowth at 24 and 48 h (Fig. 3, bottom). In this case, cleavage was evident by the appearance of a second band of slightly lower molecular weight \((-24 kDa)\) detectable just below the parent SNAP-25 band. These data demonstrate that neurite outgrowth in these embryonic neurons was not inhibited by cleavage of SNAP-25 and, further, confirm the functional binding and uptake of BoNT/A by the eMNs in culture.

**Effects of BoNT/A Holotoxin Are Mimicked by the Toxin Binding Domain and Blocked by TVL.** To determine whether the stimulatory effect of BoNT/A on eMN neuritogenesis could be mediated by binding of the toxin alone or required internalization of the toxin enzymatic domain, separate experiments were performed using the toxin HC minus the light chain. BoNT/A HC-induced concentration-dependent increases in total neurite outgrowth that were qualitatively similar to BoNT/A holotoxin in response pattern, although somewhat less in response magnitude and sensitivity (Fig. 4A). In this case, the percentage increases in neurite length ranged from 5 to 61% at 24 h and from 3 to 38% at 48 h, with the maximal responses observed at 1 nM. Likewise, BoNT/A \(H_C\)-induced concentration-dependent increases in both the number of primary neurites and secondary branches at both time points (Fig. 4, B and C). The depressed response induced by 10 nM holotoxin relative to the lower concentrations was also evident with the toxin \(H_C\). As confirmed by Western blot (Fig. 3, bottom), the BoNT/A

![Fig. 3. SNAP-25 protein in eMNs. Top, immunoblot of SNAP-25 protein isolated from eMN cultures at 4.5, 5, and 6 h after plating. Bottom, immunoblots of eMN cultures treated with either 0.1 nM BoNT/A holotoxin (left) or 0.1 nM BoNT/A \(H_C\) (right) at 24 and 48 h. The presence of two bands in blots from holotoxin-treated cultures indicates toxin-induced cleavage, with the lower band representing cleaved SNAP-25. Note the absence of the cleaved SNAP-25 bands in the \(H_C\)-treated cultures. Protein samples (10 \(\mu\)g/ lane) were resolved on 12.5% Tris-HCl gels. SNAP-25 immunoreactivity was detected with rabbit SNAP-25 primary antibody and visualized by chemiluminescence.

![Fig. 4. Concentration-dependent effects of BoNT/A binding domain \((H_C)\) on neurite outgrowth. A, comparisons of total neurite length in micrometers \((\text{mean} \pm \text{S.E.M.}, n = 30)\) from the lowest (0 nM) to the highest (10 nM) concentrations of \(H_C\) at 24 and 48 h. B and C, comparisons of the numbers of primary \((B)\) and secondary \((C)\) neurites from 0.01 to 10 nM \(H_C\) at 24 and 48 h. Data are represented as the mean percentage change from control (0 nM) values \(\pm \text{S.E.M.}, n = 30\). A to C, ***, **, and *, values are significantly different from control (0 nM) at \(p < 0.001, p < 0.01, \text{and} p < 0.05\), respectively.\]
Hc lacked enzymatic activity. Thus, binding of the toxin Hc alone was sufficient to induce neurite outgrowth, and cleavage of SNAP-25 did not play any role.

T. vulgaris lectin was used to investigate the potential role of complex ganglioside binding in the enhancement of neurite outgrowth by BoNT/A. The concentration of TVL chosen for this study was well within the range of concentrations shown by Bakry et al. (1991) to competitively antagonize BoNT binding to central nervous system membranes and to antagonize BoNT paralytic action at the NMJ. As illustrated in Fig. 5A, treatment with TVL (1 μM) completely abolished the stimulatory effects of BoNT/A holotoxin on total neurite length at both 24 and 48 h. TVL alone had no effect on neurite outgrowth. In addition, TVL treatment abolished the stimulatory effects of BoNT/A on primary and secondary neurite numbers (data not shown). Finally, treatment with TVL also abolished the stimulatory effects of BoNT/A Hc at 24 h and significantly reduced the response at 48 h (Fig. 5B).

Discussion

Botulinum neurotoxin serotype A, marketed as Botox in the United States, is used currently for an ever-expanding list of neurological disorders characterized by neuromuscular hyperactivity (Jankovic, 2004). The cellular mechanism of action that underlies BoNT/A’s extraordinary ability to eliminate localized transmitter release at extremely low doses is well characterized and will not be further discussed here. Less well appreciated is the inherent ability of BoNT/A to induce axonal sprouting at poisoned nerve terminals (Holland and Brown, 1981; Hopkins et al., 1981; Diaz et al., 1989; Pamphlett, 1989; Angaut-Petit et al., 1990; Holds et al., 1990; Bonner et al., 1994; de Paiva et al., 1999). Such sprouting has also been reported after poisoning by BoNT serotypes D and F (Comella et al., 1993; Meunier et al., 2003). A number of studies have examined the importance of sprouting in toxin duration of action and subsequent NMJ remodeling (Angaut-Petit et al., 1990; Comella et al., 1993; Juzans et al., 1996; de Paiva et al., 1999). Although a few studies have examined this sprouting phenomenon in cell culture with variable results (Bonner et al., 1994; Igarashi et al., 1996; Osen-Sand et al., 1996). However, there has been little study to examine the cellular basis of this sprouting, which has been largely considered a nonspecific response to chemical denervation. In contrast, we hypothesized that this phenomenon may actually be a direct presynaptic neurotogenic effect of the toxin.

In the current study, BoNT/A stimulated neurotogenesis from eMNs grown in culture by increasing the number of primary neurites extended, the number of secondary branches formed, and, thus, total neurite length. These stimulatory effects of BoNT/A were concentration dependent and, more importantly, initiated solely by BoNT/A binding as evidenced by qualitatively similar responses observed after exposure to the isolated toxin binding fragment. The toxin’s actions did not appear to stimulate selectively primary or secondary neurite growth. Rather, the effects seemed to be more widely distributed because both were strongly stimulated. As might be expected, the effects were biphasic, with the stimulatory effects on primary neurites occurring earlier (24-h time point), followed by stimulatory effects on secondary neurites by 48 h.

The stimulatory actions are in general agreement with the work of Bonner et al. (1994) in which chick ciliary ganglion neurons were cocultured with skeletal muscle. In that study, BoNT/A exposure increased both primary and secondary neurite branching frequency; however, in contrast to our work, they did not observe an increase in the numbers of primary neurites. Conversely, our results are at variance with other studies reporting that BoNT/A actually inhibited neurotogenesis because of its cleavage of SNAP-25 (Osen-Sand et al., 1996; Morihara et al., 1999). In fact, in our study, neurotogenesis was promoted despite SNAP-25 cleavage.

Antagonism of the BoNT neurotogenic actions by TVL is in keeping with the previously recognized role of complex poly- gangliosides as universal membrane receptors for the various BoNT serotypes (Bakry et al., 1991; Kozaki et al., 1998; Schiavo et al., 2000). Early studies revealed the importance of gangliosides in BoNT binding, whereas later studies attempted to clarify the nature of ganglioside-BoNT interactions and their relevance to the more selective binding of BoNT to protein receptors. As a consequence, it has been proposed that gangliosides may act to increase toxin receptor affinity or to promote lateral interactions between BoNT coreceptors acting in a receptor array (Montecucco et al., 2004). Gangliosides themselves are reported to mediate neurite outgrowth, possibly through modulation of calcium channels/calcium influx in growth cones (Ledeen and Wu, 2002). Thus, the precise function of gangliosides in the BoNT-induced outgrowth reported here, beyond their known role as toxin receptors, remains unclear. In any event, the fact that the neurotogenic actions of BoNT/A do not require the toxin.

Fig. 5. Antagonism of BoNT/A holotoxin (A) and Hc (B) effects on total neurite length by ganglioside blockade using TVL. For each group, data are presented as mean ± S.E.M. (n = 30). TVL used at 1 μM. *** and * values are significantly different from controls at p < 0.001, and p < 0.05, respectively.
enzymatic domain but can be mediated solely by its binding domain suggests that the interactions of the toxin with gangliosides and/or other potential membrane receptors initiate intracellular signals that ultimately modulate neurite formation. This is somewhat reminiscent of known neurotrophins and is a novel finding because intracellular signaling by BoNT/A has not been recognized previously.

Although the concentration-dependent responses to the toxin binding fragment were qualitatively similar to those of the holotoxin, the magnitude was somewhat less, suggesting potential differences in potency between the intact toxin and the toxin fragment. The reasons for this difference are unclear. One potential explanation may be that separation of the binding fragment from the enzymatic domain alters somewhat the conformation of its binding pocket(s) such that its interaction with membrane receptors is affected. In any event, the stimulatory actions of the isolated binding domain confirm the significance of a receptor-initiated event in the promotion of neuritogenesis by BoNT.

It is interesting that both the holotoxin and the \( H_c \) binding fragment concentration responses displayed inverse U-shaped curves. This biphasic response is reminiscent of hormesis. Hormesis, a biphasic dose-response model characterized by low-dose stimulation and high-dose inhibition, typically displays a U-shaped curve. The inverse or reciprocal \( U \) shape is an alternative way to illustrate this response. Beginning at the low dose, the magnitude of the stimulatory or positive effect increases with dose, until at some high dose the magnitude diminishes, becoming less positive and displaying an inverse \( U \) shape. In fact, this type of response has been demonstrated recently for a variety of neuroactive substances, including neurotoxins, on both neurite outgrowth and neuronal survival (Ca-

In conclusion, the findings presented herein support a neu-
ritogenic role for BoNT/A. Further studies are necessary to: 1) clarify the intracellular pathways mediating this function, 2) determine whether this phenomenon holds true in older neurons and nerve terminals of the peripheral nervous system, and 3) correlate these in vitro findings with BoNT-induced sprouting in vivo.

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References


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