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BOTULINUM NEUROTOXIN TYPE B USES A DISTINCT ENTRY PATHWAY MEDIATED BY CDC42 INTO INTESTINAL CELLS VERSUS NEURONAL CELLS

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ABSTRACT

Botulinum neurotoxins (BoNTs) are responsible for severe flaccid paralysis by inhibiting the release of acetylcholine at the neuromuscular junctions. BoNT/B most often induces mild forms of botulism with predominant dysautonomic symptoms. In food borne botulism and botulism by intestinal colonization such as infant botulism, which are the most frequent naturally acquired forms of botulism, the digestive tract is the main entry route of BoNTs into the organism. We previously showed that BoNT/B translocates through mouse intestinal barrier by an endocytosis-dependent mechanism and subsequently targets neuronal cells, mainly cholinergic neurons, in the intestinal mucosa and musculosa. Here we investigated the entry pathway of BoNT/B using fluorescent C-terminal domain of the heavy chain (HcB) which is involved in the binding to specific receptor(s) and entry process into target cells. While the combination of gangliosides GD1a/GD1b/GT1b and synaptotagmin I and to a greater extent synaptotagmin II constitutes the functional HcB receptor on NG108-15 neuronal cells, HcB only uses the gangliosides GD1a/GD1b/GT1b to efficiently bind to m-ICcl2 intestinal cells. HcB enters both cell types by a dynamin-dependent endocytosis which is efficiently prevented by Dynasore, a dynamin inhibitor, and reaches a common early endosomal compartment labeled by EEA1. In contrast to neuronal cells, HcB uses a Cdc42-dependent pathway to enter intestinal cells. Then, HcB is transported to late endosomes in neuronal cells, whereas it exploits a non-acidified pathway from apical to basal lateral side of m-ICcl2 cells supporting a transcytotic route in epithelial intestinal cells.
INTRODUCTION

*Botulinum* neurotoxins (BoNTs) are potent toxins that inhibit the release of spontaneous and nerve evoked neurotransmitters and more specially the release of acetylcholine (Ach) at the neuro-muscular junctions and other cholinergic endings of the peripheral nervous system. Botulism results in a flaccid paralysis and reduction of secretions which lead to skeletal muscle paralysis, respiratory insufficiency, and death in the most severe forms (Sobel, 2005, Poulain et al., 2008). BoNTs are divided into seven toxinotypes (A to G) according to their immunological properties and in numerous subtypes based on amino acid sequence variations (Hill et al., 2013, Peck et al., 2017). A new BoNT type called H has been reported but was characterized as an A/F hybrid (Barash et al., 2014, Maslanka et al., 2016). In natural conditions, botulism is acquired by ingestion of preformed BoNT in food (foodborne botulism), or after intestinal or more rarely wound colonization and subsequent in situ BoNT production (Sobel, 2005). Indeed, *C. botulinum* can contaminate and synthesize BoNT in foods, and under certain circumstances this bacterium can grow and produce toxin in the intestine. This is the case in babies and young children (infant botulism) or in adults with predisposing factors such as intestinal surgery, antibiotherapy, chronic inflammation, necrotic lesions of the intestinal mucosa, and anatomical or functional abnormalities of the intestine. An immaturity or perturbation of the resident intestinal microbiota seems to be the main factor which facilitates the implantation of *C. botulinum* in the digestive tract (Arnon, 1989, Fox et al., 2005, Brook, 2007, Fenicia et al., 2009, Rosow et al., 2015). Therefore, most often the digestive tract represents the entry pathway for botulism. In food intoxication and during intestinal colonization with *C. botulinum*, the ingested or locally produced BoNT has to cross the digestive tract epithelium prior to have access to the target neurons.

BoNT/B is commonly associated with numerous cases of food borne botulism or infant botulism in humans throughout the world (Sobel, 2005, Popoff et al., 2013). Although all BoNT types induce a descending flaccid paralysis, differences in the clinical picture have been observed according to the BoNT type. Thereby, type B botulism is more often less severe and less lasting than type A botulism (Woodruff et al., 1992, Cherington, 1998). Dysautonomia (diplopia, blurred vision, dry mouth, dysphagia, constipation) is more frequent, and the motor symptoms are less pronounced in type B botulism than in type A (Hughes et al., 1981, Merz et al., 2003, Potulska-Chromik et al., 2013). BoNT/B seems to have a higher affinity to cholinergic autonomic than motor-neuron endings (Brashear et al., 1999). BoNT/B and BoNT/A recognize distinct receptors on neuronal cells, synaptotagmin...
(Syt) and synaptic vesicle (SV2) protein, respectively, which might account for the differential recognition of neurons by the two toxins (review in (Rummel, 2013)). In addition, BoNT/B and BoNT/A might use distinct trafficking pathways or mode of dissemination from the digestive tract to the target neurons.

BoNTs associate with non-toxin proteins (ANTPs) produced by *C. botulinum* to form complexes of variable sizes (300 to 900 kDa). ANTPs encompass a non-toxic and non-hemagglutinin component (NTNH), and either several hemagglutinin components (HAs) or other non-hemagglutinin proteins called OrfX1, OrfX2, OrfX3 or P47 (Oguma *et al.*, 1999, Popoff *et al.*, 1999, Quinn *et al.*, 2001, Sharma *et al.*, 2003). One of the main role of ANTPs is to protect BoNT from degradation by the stomach acidic pH and digestive proteases. Indeed, NTNH retains a similar structure than that of BoNT and assembles with BoNT forming a compact interlocked complex resistant to acidic pH and protease degradation (Gu *et al.*, 2012, Inui *et al.*, 2012, Sagane *et al.*, 2012). The mode of passage of either BoNT alone or the whole BoNT complexes through the intestinal epithelial barrier is still controversial. HA complexes, but not OrfX proteins, have been shown to bind to epithelial cadherin (E-cadherin) and to disrupt the intercellular junctional complexes between epithelial cells thus facilitating the passage of BoNT complexes through epithelial barrier by the paracellular way (Jin *et al.*, 2009, Sugawara *et al.*, 2010, Sugawara *et al.*, 2011, Lee *et al.*, 2014, Lam *et al.*, 2015). However, it is not clear whether whole BoNT complex or BoNT alone is transported through the intestinal barrier. At acidic pH, BoNT complexes are highly stable, but at the neutral or basic pH of the duodenum and jejunum, BoNT complexes dissociate (review in (Poulain *et al.*, 2008). BoNTs alone have been reported to cross the intestinal barrier upon a transcytotic mechanism without the help of HAs or other ANTPs (Fujinaga *et al.*, 1997, Maksymowycz *et al.*, 1998, Park *et al.*, 2003, Maksymowycz *et al.*, 2004, Nishikawa *et al.*, 2004, Ahsan *et al.*, 2005, Couesnon *et al.*, 2008). Notably, we have shown that BoNT/B free of HAs is able to translocate through mouse intestinal barrier via an endocytosis-dependent process and rapidly targets in 10 min the enteric nervous system, mainly the cholinergic intestinal neurons (Connan *et al.*, 2016). These finding led us to investigate and compared the mode of BoNT/B entry into cell models of intestinal cell and neuronal cells.

**RESULTS**

**HcB enters neuronal and intestinal cells**

The C-terminal domain (Hc) of clostridial neurotoxin heavy chains is involved in the specific recognition and interaction with cell surface receptors. Hc has already extensively
been used to investigate internalization and intracellular trafficking of clostridial neurotoxins in cells as well as toxin translocation through the intestinal epithelium and dissemination in the subjacent tissues (Lalli et al., 2003, Maksymowych et al., 2004, Bohnert et al., 2005, Roux et al., 2006, Harper et al., 2011, Restani et al., 2012, Lam et al., 2015). Notably, we used fluorescent Hc domains of BoNT/A and BoNT/B to analyze the neurotoxin passage and dissemination in mouse intestinal mucosa (Couesnon et al., 2008, Connan et al., 2016). Thereby, recombinant Hc from BoNT/B (HcB) labeled with Cy3 was used to monitor BoNT/B entry into cells. First, fluorescent HcB was incubated with several cell types at 37°C for 15 min. HcB nicely decorated the extensions and plasma membrane of primary rat cortical neurons and was visualized in intracellular compartments (Fig. 1). HcB-Cy3 also labeled the membrane of the neuronal cell line NG108-15, but to a lower extent than in primary neuronal cells. Primary neuronal cells likely retain a larger number of HcB receptors or higher affinity receptors compared to cultured neuronal or intestinal cell lines. When incubated at 37°C for 15 min, the largest part of HcB-Cy3 was localized in intracellular compartments indicating that HcB was efficiently internalized into the NG108-15 cell line (Fig. 1). In most cells, HcB-Cy3 was distributed in perinuclear compartments. Since HcB-Cy3 internalization was more easily monitored in NG108-15 cells due to the larger volume of NG108-15 versus that of primary neuronal cell bodies, NG108-15 cells were used in the subsequent experiments.

HcB-Cy3 entry was tested is several intestinal cell lines (CaCo-2, T84, mouse intestinal crypt cells (m-IC_{cl2})) with approximately the same efficiency (data not shown). m-IC_{cl2} were used as they were found to be a relevant model for analyzing the entry of BoNTs into intestinal cells (Couesnon et al., 2008). After 15 min incubation, HcB was localized in intracellular compartments, which were more scattered through the cytoplasm than in neuronal cells (Fig. 1).

HcB-Cy3 entry was investigated in polarized m-IC_{cl2} cells grown on filters. HcB-Cy3 (2.5 μg/ml) was added in the apical chamber of the cell cultures which were incubated at 37°C for 3 min. Then, the cells were washed and further incubated with culture medium at 37°C for the indicated times (Fig. 2A and 2B). The HcB-Cy3 spots were monitored in the apical, center, and basolateral side of the polarized cells over time (Fig. 2B). Thus, a progressive passage of HcB-Cy3 was observed from apical brush border binding to basolateral endocytic compartments of m-IC_{cl2} cells within 10-20 min. HcB-Cy3 was also visualized in cultured NG108-15 over time (Fig. 2C and 2D). It is noteworthy that the
internalization level of HcB-Cy3 was lower in intestinal cells compared to neuronal cells (Fig. 2).

**HcB enters neuronal and intestinal cells in a dynamin-dependent manner**

BoNTs enter target cells via a receptor-mediated endocytosis (Beard, 2014, Rossetto et al., 2014), but the precise mechanism of endocytosis according to the cell types is still elusive. BoNT/A has been found to enter neuronal and intestinal cells by a dynamin-dependent pathway in both cell types (Coesnon et al., 2009). Dynasore, an inhibitor of dynamin-mediated endocytosis (Macia et al., 2006) blocks BoNT/A entry into cultured hippocampal neurons (Harper et al., 2011). Here, we investigated the inhibitory effect of Dynasore in the entry of HcB-Cy3 in neuronal and intestinal cells (Fig. 3). Cells were pretreated with Dynasore (80 µM) for 30 min and were then incubated with HcB-Cy3 for additional 10 min. As shown in Fig. 3A, Dynasore efficiently prevented HcB-Cy3 entry in both neuronal NG108-15 cells and intestinal crypt m-IC_{cl2} cells. However, HcB-Cy3 still decorated the plasma membrane of NG108-15 neuronal cells indicating that HcB binding to surface cell receptor was unaffected by Dynasore treatment. To further support the dynamin-dependent HcB-Cy3 entry into cells, NG108-15 and m-IC_{cl2} cells were transfected with the dominant negative dynamin2 K44A as previously described (Coesnon et al., 2010) and then exposed to HcB-Cy3. Dynamin2 K44A prevented HcB-Cy3 internalization in both neuronal and intestinal cells (Sup. Fig. 1).

The HcB entry pathway in NG108-15 and m-IC_{cl2} was further investigated using potassium depletion followed by hypotonic shock which disrupts the formation of clathrin-coated vesicles and prevents clathrin-mediated endocytosis (Carpentier et al., 1989, Kovbasnjuk et al., 2001). HcB-Cy3 internalization was significantly prevented in NG108-15 pretreated in K+-depleted buffer. The fluorescent HcB molecules mainly decorated the cell periphery. In contrast, K+ depletion did not significantly alter the HcB-Cy3 entry into m-IC_{cl2} indicating that HcB mainly used a clathrin-mediated endocytosis in NG108-15 and an alternative pathway in m-IC_{cl2} cells (Fig. 4).

**HcB uses gangliosides GD_{1a}, GD_{1b}, GT_{1b} but not GMI as receptors on intestinal cells**

The specific receptors of BoNT/B on neuronal cells have been identified as gangliosides GT1a, GD_{1b}, GT_{1b} associated to SytI and SytII (Nishiki et al., 1994, Nishiki et al., 1996a, Jin et al., 2006, Rummel et al., 2007). Does BoNT/B recognize the same receptors
on intestinal cells than on neuronal cells for its entry into epithelial cells? Gangliosides are components of plasma membrane of all cell types, but they are predominantly concentrated in nerve endings. Firstly, we investigated the role of gangliosides in HcB entry into neuronal and intestinal cells. Competition experiments were performed between HcB-Cy3 and gangliosides in m-IC\textsubscript{cl2} cells versus NG108-15 cells. When HcB-Cy3 was preincubated with a preparation (gg) enriched in gangliosides GD\textsubscript{1a}, GD\textsubscript{1b}, and GT\textsubscript{1b} (250 µg/ml), HcB-Cy3 internalization was strongly reduced in m-IC\textsubscript{cl2} cells and to a lower extent in NG108-15 cells (Fig. 5). In contrast, GM1 (50 µg/ml) did not significantly impair the binding and entry of HcB-Cy3 into intestinal and neuronal cells (Fig. 5). GM1 was already reported to not be included in the BoNT/B receptor ganglioside moiety on neuronal cells (Nishiki \textit{et al.}, 1996a, Dong \textit{et al.}, 2003).

To further evidence the role of gangliosides in HcB entry into m-IC\textsubscript{cl2} cells, the cells were treated with DL-threo-1-phenyl-2-palmitoyl-amino-3-morpholino-1-propanol (PPMP), an inhibitor of glucosylceramide synthase. HcB-Cy3 internalization in cells exposed to PPMP was significantly lower compared to untreated control cells, whereas subsequent loading with exogenous gangliosides enriched in GD\textsubscript{1a}, GD\textsubscript{1b}, and GT\textsubscript{1b} restored the entry into cells (Fig. 6) further supporting the BoNT/B receptor role of gangliosides.

\textit{HcB uptake in intestinal cells does not require synaptotagmin}

Syt is a Ca\textsuperscript{++} sensor anchored in synaptic vesicle membrane of neuronal cells, and the isoforms SytI and SytII have been identified as BoNT/B receptors (review in (Rummel, 2013)). SytI and SytII are also expressed in intestinal cells (Sup Fig. 2). Competition experiments were performed by preincubating HcB-Cy3 with a 10 fold excess of SytI or SytII and then incubating the mix with cells for 10 min (Fig. 7). SytII induced a significant but partial decrease in HcB-Cy3 entry into NG108-15 cells. Indeed, preincubation with SytI or SytII decreased HcB-Cy3 entry into neuronal cells by 30 and 50\% (Fig. 7), respectively, in agreement with previous report (Nishiki \textit{et al.}, 1996a). In contrast, no significant inhibitory effect of SytI or SytII was observed in HcB-Cy3 entry into m-IC\textsubscript{cl2} cells indicating that SytI and SytII are not required for HcB uptake in intestinal cells.

Since the functional BoNT/B receptor on neuronal cells consists of an association of gangliosides and Syt (Nishiki \textit{et al.}, 1994, Nishiki \textit{et al.}, 1996a, Jin \textit{et al.}, 2006, Rummel \textit{et al.}, 2007), we performed competition experiments with combinations of gg and SytI or SytII. A stronger decrease (80\%) in HcB-Cy3 internalization into NG108-15 cells was obtained by
preincubation of HcB-Cy3 with gg associated with SytI or SytII (Fig. 8), compared to incubation with SytI or SytII alone, 30 and 50 % inhibition, respectively (Fig. 7). This further confirms that HcB requires both gg and Syt for efficient internalization into neuronal cells. In contrast, preincubation with SytI or SytII did not significantly prevent HcB-Cy3 entry into m-ICcl2 cells, and the combination of gg and Syt did not induce additional inhibitory effect compared to gg alone (Fig. 5 and 8) indicating that gangliosides, mainly GD1a/GD1b/GT1b, were sufficient to mediate HcB entry into m-ICcl2 cells.

**HcB enters neuronal and intestinal cells via a common early endosomal compartment and then follows a distinct non-acidic pathway in intestinal cells**

Entry pathways of HcB into m-ICcl2 and NG108-15 cells were investigated by colocalization of HcB-Cy3 with markers of specific endosomal compartments. After 10 min incubation at 37°C with both cell types, HcB-Cy3 was distributed in vesicles containing the Rab5 effector EEA1 (early endosome antigen 1) (Fig. 9). Then, endocytosed cargos in early endosomes (EE) can be sorted and transported in distinct intracellular pathways. A major pathway includes the late endosomes (LE)/lysosomes. Rab7 is classically considered as associated with LE/lysosomes. However, Rab5 replacement by Rab7 occurs in EE, and Rab7 seems to have an earlier role in the trafficking between EE and LE/lysosomes (review in (Girard et al., 2014)). As shown in Fig. 10, HcB-Cy3 was found in vesicles containing Rab7 in both m-ICcl2 cells and NG108-15 cells after 10 min incubation at 37°C. However, HcB-Cy3 colocalized differently with Lamp1 (lysosomal-associated membrane protein 1) which is a specific marker of LE/lysosomes, in m-ICcl2 cells and NG108-15 cells (Fig. 10). In NG108-15 cells, HcB-Cy3 was progressively associated with Lamp1 compartments over time from 5 to 20 min (Fig. 11A, 11C). At 20 min, most of HcB-Cy3 was localized in NG108-15 LE/lysosomes, whereas in m-ICcl2 cells only a low level of the fluorescent protein was detected in this intracellular pathway (Fig. 11B, 11C). These results support that the cellular trafficking of HcB differs in neuronal and epithelial intestinal cells.

**HcB enters intestinal cells via Cdc42-dependent pathway**

RhoGTPases are key regulators which control various cellular functions including diverse endocytosis pathways (Etienne-Manneville et al., 2002). Notably, Cdc42 was found to control a clathrin-independent endocytic process termed the Clathrin-Independent Carrier (CLIC)/GPI-anchored protein-enriched Early Endocytic Compartment (GEEC) pathway and was associated with the endocytosis of GPI-anchored proteins, fluid markers and toxins such
as *Helicobacter pylori* VacA (Sabharanjak et al., 2002, Gauthier et al., 2005, Chadda et al., 2007). Transfection of a Cdc42 negative mutant (Cdc42T17N) fused to GFP into m-IC<sub>cl2</sub> cells significantly inhibited the entry of HcB-Cy3. None or only a few endocytic vesicles stained with HcB-Cy3 were observed in transfected cells compared to untransfected cells (Fig. 12). In contrast in NG108-15 cells, HcB-Cy3 entry was either unimpaired or only weakly compared to untransfected cells (Fig. 12). Transfection of the dominant negative forms of RhoA (RhoA T17N-GFP) or Rac1 (Rac1 T17N-GFP) in m-IC<sub>cl2</sub> cells and NG108-15 cells did not significantly prevent HcB-Cy3 entry (Sup. Fig. 3).

**DISCUSSION**

As monitored with HcB-Cy3, the fluorescent protein bound to the periphery of neuronal cells as well as of intestinal cells and then entered into endocytic vesicles. This raises the question whether BoNT/B recognizes the same specific receptors on both neuronal and intestinal cells? SytI/SytII in combination with gangliosides, mainly GD<sub>1a</sub>/GD<sub>1b</sub>/GT<sub>1b</sub>, were identified as specific BoNT/B receptors on neuronal cells (Nishiki et al., 1994, Nishiki et al., 1996a, Nishiki et al., 1996b, Chai et al., 2006, Jin et al., 2006, Rummel et al., 2007). In addition, SytI/SytII were shown to mediate BoNT/B entry into PC12 cells (Dong et al., 2003). We also confirm that HcB interacts with NG108-15 through gangliosides GD<sub>1a</sub>/GD<sub>1b</sub>/GT<sub>1b</sub> and SytI or SytII by using binding competition experiments (Fig. 4, 5, 6, 7 and 8). The combinations of gangliosides with the receptor protein part (SytI or SytII) were more effective than the individual preparations of gangliosides or Syt in binding inhibition of HcB to NG108-15 cells. This supports that the high affinity functional HcB receptor on neuronal cells requires both the ganglioside and protein parts. In contrast, HcB binding to m-IC<sub>cl2</sub> cells was fully prevented by gangliosides alone without the help of a protein part such as SytI or SytII. Thereby, SytI or SytII were not involved in HcB receptor on intestinal cells, and gangliosides of the GD<sub>1a</sub>/GD<sub>1b</sub>/GT<sub>1b</sub> series seemed efficient and sufficient receptors for BoNT/B. SytI and SytII are mainly expressed in neuronal cells where they are localized on synaptic vesicle membrane and play a critical role in the evoked release of neurotransmitter (Sudhof, 2013). In addition, SytI has been found to be involved in endocytosis of some compounds such as Na<sup>+</sup>-H<sup>+</sup> exchanger in intestinal epithelial cells (Musch et al., 2010). We also observed that intestinal epithelial cells expressed SytI and SytII using Western blot (Sup. Fig. 2). Moreover, SytI and SytII were visualized in m-IC<sub>cl2</sub> by cell imaging with specific antibodies (Sup. Fig. 2). It is noteworthy that m-IC<sub>cl2</sub> mainly retains a stem cell profile and can express a wide variety of proteins (Bens et al., 1996). However, SytI and SytII were not
detected by cell imaging in mouse intestinal epithelial cells (Connan et al., 2016), indicating that these proteins are probably weakly expressed in mouse intestinal epithelial cells or only in some cell subpopulations. This further excludes that SytI and SytII have a role in BoNT/B endocytosis in epithelial cells.

The role of gangliosides in HcB entry into intestinal cells was further supported by inhibiting the ganglioside synthesis in m-IC_{cl2} cells with PPMP (Fig. 6). However, this does not preclude the possibility that membrane proteins of intestinal cells might be used in combination with gangliosides for higher BoNT/B affinity receptors. Indeed, BoNT/B did not localize in all cell types of the mouse intestinal epithelium but only in specific cells which represent a preferential passage of the toxin through the intestinal barrier (Connan et al., 2016). In contrast, BoNT/A which uses SV2C as protein receptor on neuronal cells (review in (Rummel, 2013)) enters intestinal cells in a SV2C-dependent manner albeit this molecule has not been evidenced as a BoNT/A receptor in epithelial cells (Elias et al., 2011, Couesnon et al., 2012). This suggests that SV2C might facilitate endocytosis into epithelial cells in a non-specific manner. However, competition assays with SV2C L4 luminal domain and HcB-Cy3 in m-IC_{cl2} cells showed that HcB entry into intestinal cells was not mediated by SV2C (Sup. Fig. 4).

HcB-Cy3 internalization into endocytic vesicles was shown in both neuronal and intestinal cells. However, the distribution of endocytic vesicles containing HcB-Cy3 in both cell types was slightly different. In NG108-15 cells, HcB-Cy3 entered endocytic compartments which migrated in the perinuclear area, whereas in m-IC_{cl2} cells the vesicles containing HcB-Cy3 had a more scattered distribution through the cytoplasm (Fig. 1 and 2). This supports a different trafficking of BoNT/B in intestinal cells versus neuronal cells resulting in a delivery of the whole and active neurotoxin to the basolateral side of intestinal cells. In contrast, in target neuronal cells the neurotoxin is transported in acidic compartments which trigger the translocation of the L chain into the cytosol and its subsequent proteolytic activity towards the SNARE (Soluble N-ethylmaleimide-sensitive-factor Attachment Receptor) synaptobrevin ((Simpson et al., 1994, Keller et al., 2004, Poulain et al., 2008).

HcB-Cy3 uptake was efficiently blocked by Dynasore which is a specific inhibitor of dynamin (Macia et al., 2006) in both neuronal NG108-15 and intestinal m-IC_{cl2} cells (Fig. 3). However, Dynasore not only blocks dynamin-dependent endocytosis but also induces dynamin-independent effects such as reduction of membrane cholesterol, disorganization of lipid rafts, and prevention of actin filament formation involved in membrane ruffling (Preta 2015). Thus, Dynasore could also impair dynamin-independent endocytosis such as
macropinocytosis or fluid phase endocytosis (Preta et al., 2015). The dynamin-dependent entry of HcB-Cy3 into both cell types was further supported by using transfection with the dominant negative dynamin2 K44A (Sup. Fig. 1). A dynamin-based endocytosis has also been evidenced in BoNT/A entry into cells (Couesnon et al., 2009, Harper et al., 2011, Meng et al., 2013). Moreover, K+ depletion experiments indicated that HcB preferentially used a clathrin-independent endocytosis into m-IC12 cells, contrarily to NG108-15 in which HcB uptake was clathrin-mediated (Fig. 4). Indeed, BoNT/B, like BoNT/A, enters neuronal cells, notably at the neuromuscular junctions, via a clathrin-mediated endocytosis hijacking, at least in part, the synaptic vesicle recycling (Poulain et al., 2008, Meng et al., 2013, Rossetto et al., 2014). Thereby, HcB likely exploited a clathrin-mediated entry into neuronal cells and mainly a clathrin-independent pathway into intestinal cells, both mechanisms of uptake being regulated by dynamin. Interestingly, toxins which recognize glycolipid receptors, use mainly clathrin-independent endocytosis. Toxin binding to membrane glycolipids such as Shiga toxin to globothriaosylceramide or cholera toxin to GM1 induces lipid reorganization (Watkins et al., 2014, Pezeshkian et al., 2016), leading to membrane curvature without clathrin (Romer et al., 2007, Ewers et al., 2010). Membrane invaginations containing toxins are then detached from the plasma membrane through a dynamin-dependent process. This endocytic process is also shared by endogenous proteins such as galectins (Lakshminarayan et al., 2014). BoNT/B possibly uses a similar mechanism of binding to gangliosides GD1a/GD1b/GT1b to trigger a clathrin-independent endocytosis in intestinal epithelial cells.

Cargo molecules attached to membrane receptor are internalized into EE, and then are sorted into multiple endocytic pathways such as recycling to the plasma membrane, transcytosis to the opposite side, or sorted to degradative compartments (LE/lysosomes). Endocytic vesicles are dynamic structures which mature by different ways including fusion with homotypic vesicles or other compartments, or budding and fission in smaller vesicles. Rab proteins are key players in fusion of homotypic endosomes. EEA1 binds to Rab5-GTP and additional effectors such as phosphatidylinositol-3-phosphate (PI3P) and syntaxin-6 in sorting EE where it has a tethering/docking role before membrane fusion. In polarized epithelial cells, EEA1 is distributed in basolateral sorting endosomes but not in apical endosomal compartment suggesting that EEA1 is involved in specific directional endocytic pathway (Simonsen et al., 1998, Christoforidis et al., 1999, Wilson et al., 2000, Shin et al., 2005, Wandinger-Ness et al., 2014). Molecules endocytosed by the typical clathrin-dependent pathway as well as by clathrin-independent pathways are internalized into EE labeled with Rab5 and EEA1 (Kalia et al., 2006, Mayor et al., 2007, Grant et al., 2009,
Howes et al., 2010, Maldonado-Baez et al., 2013). Subsequent maturation of EE into LE is mediated through replacement of Rab5 by Rab7 (Rink et al., 2005, Poteryaev et al., 2010). Thus, Rab7 is found in EE and its activity seems to be required in the late trafficking step from LE to lysosome (Feng et al., 1995), but also in early endosomal sorting in a cargo-dependent manner (Girard et al., 2014). Here, we showed that HcB-Cy3 enters an early endosomal compartment labeled with EEA1 and Rab7 at 10 min incubation in both neuronal and intestinal cells, although HcB was internalized by a clathrin-dependent pathway in NG108-15 cells, and by a clathrin-independent pathway in m-ICcl2 cells. Then HcB was transported in distinct intracellular pathways according to the cell type. Overtime from 5 to 20 min, HcB-Cy3 migrated from EE to LE as monitored with Lamp1 in NG108-15 cells, whereas HcB used a distinct pathway in m-ICcl2 cells avoiding the passage in LE/lysosomes (Fig. 11 ABC). Therefore, our findings support the hypothesis that the transport in the LE pathway in neuronal cells where the endocytic vesicles are acidified, facilitates the translocation of the L chain into the cytosol and its subsequent inhibitory activity on the neurotransmitter exocytosis machinery. In contrast to neuronal cells, the whole neurotoxin transits through non-acidified vesicles in epithelial cells and is released to the baso-lateral side in an un-dissociated form. Similar findings were observed with BoNT/A (Couesnon et al., 2009).

HcB transport in intestinal cells was further investigated by analyzing the effects of RhoGTPases which are involved in various cellular functions notably in the control of the actin cytoskeleton and also in different endocytic pathways (Hall, 2012). In the epithelial intestinal m-ICcl2 cells, HcB-Cy3 was transported in a Cdc42-dependent and Rho-Rac-independent pathway, whereas RhoGTPases were not involved in HcB trafficking in neuronal NG108-15 cells (Fig. 12 and Sup. Fig. 3). Cdc42 is a GTPase of the Rho family which controls certain cortical actin formations like filopodia, cell polarity in all cell types, as well as several intracellular vesicle trafficking and endocytosis pathways including caveolae endocytosis (Tapon et al., 1997, Etienne-Manneville et al., 2002, Cheng et al., 2010, Chi et al., 2013). In addition, Cdc42 controls transcytosis from apical to basolateral side in polarized epithelial cells (Rojas et al., 2001, Harris et al., 2010) such as transcytosis of Escherichia coli Shiga-like toxin through intestinal epithelial cells (Malyukova et al., 2009). Moreover, Cdc42 mediates clathrin-independent endocytosis of GPI-anchored proteins, fluid phase markers and toxins in non-neuronal cells such as H. pylori VacA, BoNT/A, and cholera toxin subunit B (Sabharanjak et al., 2002, Gauthier et al., 2005, Chadda et al., 2007, Couesnon et al., 2009, Malyukova et al., 2009, Zachos et al., 2014, Chen et al., 2016). The HcB Cdc42-dependent
entry pathway is distinct from that of GEEC (GPI-anchored protein enriched early endosomal compartment) which is associated with GPI-anchored protein uptake and lipid bound ligands, since in contrast to the HcB pathway, GEEC is dynamin-independent and is acidified (Zachos et al., 2014). However, HcB was not transported in the acidified route leading to LE/lysosomes in intestinal cells contrarily to neuronal cells. Cdc42 might regulate several clathrin-independent pathways. Thus in contrast to neuronal cells, HcB similarly to BoNT/A (Couesnon et al., 2010) used a Cdc42-dependent transcytosis in intestinal cells which still remains to be defined.

In conclusion, HcB used distinct entry pathways in neuronal and intestinal cells. HcB recognized gangliosides (GD1a, GD1b, GT1b)/synaptotagmins on neuronal cells, whereas gangliosides alone seemed sufficient to facilitate the toxin entry into intestinal cells. In both cell types, HcB uptake was dynamin-dependent, but clathrin-dependent in neuronal cells and clathrin-independent in intestinal cells, and led to a common early endosomal compartment labeled with EEA1. Then, HcB was driven into an acidic pathway leading to LE/lysosomes, allowing the translocation of the L chain into the cytosol of neuronal cells, whereas the neurotoxin transcytosed in a Cdc42-dependent manner and released in an undissociated form at the basolateral side of epithelial intestinal cells.

EXPERIMENTAL PROCEDURES

Reagents

Endocytosis assay were performed with Dynasore Hydrate (Sigma, 80µM). Competition assay were performed using a mixture of 250 µg ml⁻¹ gangliosides (18% GM1, 55% GD1a, 15% GD1b, 10% GT1b and 2% others, Calbiochem) or 50 mg ml⁻¹ GM1 (Calbiochem). PPMP was from Sigma. Cells were stained with Hoechst 33342 (Invitrogen, 1:1000 dilution) for the nuclei. The primary antibodies used recognized E-cadherin (Invitrogen, rat, 1:250 dilution), EEA1 (BD Pharmingen™, mouse IgG1, 1:400 dilution), Rab7 (Cell Signaling, Rabbit, 1:200 dilution), and and Lamp1 (BD Pharmingen™, rat, 1:500). The secondary antibodies were Alexa fluor 488 goat anti rat, (Invitrogen, 1:500 dilution). Rat synaptotagmin I (amino acids 1-421) and II (amino acids 8-61) were produced as recombinant proteins with N-terminal 6His-tag from pET28 vector. Rabbit anti-SytI (ab68853) and SytII (ab113545) were from Abcam, mouse monoclonal antibodies against SytI (3F10A) and SytII (8G2b) were provided by C. Leveque (Nishiki et al., 1996a, Charvin et al., 1997).
Recombinant HcB

Recombinant His-tag Hc fragment of BoNT/B was produced and purified from pET28a vector containing DNA encoding for HcB cloned into BamHI and SalI sites, as previously described (Tavallaie et al., 2004). HcB His-tag was labeled with Amersham Cy3 Mono-Reactive Dye Pack (GE Healthcare) according to the manufacturer's recommendations. Free dye was removed from labeled protein using the de Zeba Spin Desalting columns according to the manufacturer's recommendations (Thermo Scientific).

Cell cultures

NG108-15 neuronal cells were obtained from the ATCC, and m-ICcl2 cells were kindly provided by A. Vandewalle (URMS 1149, Paris). Cells were cultured at 37°C in a 5% CO₂ atmosphere and medium was changed every second day. NG108-15 cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM, Invitrogen) supplemented with 10% fetal calf serum (FCS, Invitrogen). m-ICcl2 cells were cultured as described previously (Bens et al., 1996).

For m-ICcl2 cells grown on filters, cells were seeded (1 x 10⁵) on the upper chamber of Transwell Inserts (Corning) and allowed to differentiate for one week. Cells were incubated with HcB-Cy3 (2.5 µg/ml) for 3 min at 37°C and were rinsed three times with culture medium containing BSA 0.1% and were further incubated at 37°C for the indicated times. Cells were washed with cold PBS and fixed with 4% paraformaldehyde (PFA) before phalloidin labeling.

Primary cortical neurons were obtained from fetal mice. Cortices from E17 mouse embryos were dissected, cut into small pieces, and treated with 0.3% protease for 10 min before trituration in PBS–0.1% DNAse (Roche). Dissociated cells were centrifuged through 4% bovine serum albumin (BSA) solution for 4 min at 1000 rpm. Cells were resuspended in Neurobasal medium supplemented with 2% fetal calf serum, 2%B27 supplement, 25M -mercaptoethanol, 2 mM glutamine, and 50g/mL gentamicin. Cells were plated at a density of 5x 10⁴cells/cm² on round coverslips or glass-bottomed microwell dishes (MatTek, Ashland, US) previously coated with 50 µg/mL poly-ornithine followed by 5 µg/mL laminine. After 1 day, cells were treated with 25 µg/mL uridine and 10 µg/mL fluorode-oxyuridine to eliminate proliferating glial cells (Chen et al., 2006).

Binding and internalization studies
Cells (5 X 10^4) were plated on coverslips and incubated in HBSS/BSA 0.5% (HBSS/BSA) containing 5 µg/ml of HcB-Cy3. For internalization experiments, cells were incubated at 37°C for 3 min, washed with HBSS, and either fixed with 4% PFA in PBS for 20 min, or further incubated at 37°C in HBSS before fixation. After washing in PBS and quenching with 50mM NH₄Cl in PBS, cells were permeabilized with 0.2% Triton X-100 in PBS and incubated with Alexa647-labeled phalloidin (Interchim) and Dapi.

For competition studies, cells plated on coverslips (5 x 10^4) were incubated with HcB-Cy3 (2.5 µg) for 10 minutes or HcB-Cy3 (2.5 µg) pre-incubated with gangliosides mix (250 µg ml⁻¹), or Syt I (25 µg) or Syt II (25 µg) or a combination of gangliosides mix and Syt I or Syt II for 30 minutes at room temperature.

Ganglioside depletion in cell membrane with PPMP was performed as previously described (Couesnon et al., 2008) by incubating m-ICel2 cells with 7.5 µg/ml PPMP in culture medium for 48 h. Ganglioside complementation was performed by incubating the cells with the ganglioside mixture (250 µg/ml) in culture medium after 24 h of PPMP treatment.

K+ depletion was performed as previously described (Couesnon et al., 2010). Briefly, cells were incubated in K+ depletion buffer (NaCl 140 mM, CaCl₂ 1 mM, MgCl₂ 1 mM, Hepes 20 mM, D-glucose 1 g/l⁻¹, pH 7.4) for 30 min at 37°C followed by hypotonic shock (potassium depletion buffer diluted 1:2 in distilled water) for 5 min at 37°C. Control cells were incubated in HBSS containing 10 mM KCl. Cells were then incubated with HcB-Cy3 for 10 min at 37°C and processed for microscopy.

**Cell transfections**

Eukaryotic plasmid vectors used in this work are as follows: GFP-Cdc42T17 N (C. Lamaze, Institut Curie, Paris, France). Cell transfections were performed using X-tremeGene HP (Roche) according to the manufacturer’s instructions. Briefly, cells were grown over night on coverslip. DNA- lipid complexes were formed for 10 minutes at 37°C and added dropwise to the cells and further incubated for 4 h at 37°C. The cells were then washed in PBS and grown overnight to allow expression of proteins from plasmid-borne gene. Cells were then incubated for 10 minutes with HcB-Cy3 before PFA fixation.

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REFERENCES


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Mouse primary cortical neurons, NG108-15, or m-IC\textsubscript{cl2} cells plated on glass slides were incubated with HcB-Cy3 (2.5 \( \mu \)g/ml) for 10 min at 37\(^\circ\)C. Then, the cells were fixed and mounted for microscopy observation. Cells were counterstained with Dapi to visualize nuclei, and m-IC\textsubscript{cl2} cells were counterstained with phalloidin (green) to visualize the actin cytoskeleton.
White arrows indicate neuronal cell bodies with internalized HcB-Cy3 in intracellular compartments. Internalized HcB-Cy3 appeared as numerous vesicular spots in m-IC\textsubscript{cl2} cells.
Figure 2. HcB-Cy3 passage through polarized intestinal crypt m-IC_{cl2} cells.
(A) HcB-Cy3 (2.5 μg/ml) was added to the apical chamber of polarized m-IC_{cl2} cells grown on filters for 3 min at 37°C. Then, the culture medium was replaced with Dulbecco's medium. The cells were further incubated at 37°C for the indicated times, fixed and processed for confocal microscopy.
Intracellular HcB-Cy3 was quantified and expressed as the number of HcB-Cy3 spots in apical, center, or basolateral side per cell. Values are means from three independent experiments with three replicates each.

(C) HcB-Cy3 (2.5 µg/ml) was added to NG108-15 grown on cover slides for 3 min at 37°C. Then, the culture medium was replaced with Dulbecco's medium. The cells were further incubated at 37°C for the indicated times, fixed and processed for confocal microscopy.
Figure 3. Inhibition of HcB-Cy3 entry into NG108-15 cells and intestinal crypt m-IC_{cl2} cells by Dynasore.

(A) Cells plated on glass slides were pretreated with Dynasore 80 μM for 30 min at 37°C. Then, HcB-Cy3 (2.5 μg/ml) was added and cells were further incubated at 37°C for additional 10 min before being processed for microscopy.
(B) Quantification of HcB-Cy3 entry into control cells and cells pretreated with or without Dynasore. Fluorescence of intracellular HcB-Cy3 spots (in at least 20 cells from each of three independent experiments) was determined and the results were expressed as relative amounts of intracellular HcB-Cy3 in cells pretreated with Dynasore versus control cells. The level of internalized HcB-Cy3 in control NG108-15 or m-IC_{el2} cells without Dynasore was arbitrarily assigned to 1. Data are mean values ± SD. Differences between means were tested using Student’s t-test, and p-values <0.001 were taken to indicate significance (***).
Figure 4. K+ depletion impairs HcB-Cy3 entry into NG108-15 and not in m-IC_{cl2} cells. (A) Cells were incubated in K+ depleted buffer or in HBSS containing 10 mM KCl for 30 min at 37°C followed by a hypotonic shock of 5 min, and then exposed to HcB-Cy3 for additional 10 min. (B) Quantification of HcB-Cy3 entry in control cells and cells incubated in K+ depleted buffer as described in Fig. 3. The level of internalized HcB-Cy3 in control m-IC_{cl2} cells was arbitrarily assigned to 1. Data are mean values ± SD from at least three independent experiments. Differences between means were tested using Student’s t-test, and p-values <0.001 were taken to indicate significance (***).
Figure 5. HcB-Cy3 entry into m-IC_{cl2} cells is dependent of the gangliosides GD_{1a}, GD_{1b}, GT_{1b} but not of GM1.

HcB-Cy3 (2.5 µg/ml) was preincubated with (A) a preparation (gg) enriched in gangliosides (18% GM1, 55% GD_{1a}, 15% GD_{1b}, 10% GT_{1b}, 2% others; 250 µg/ml), or (C) GM1 (50 µg/ml), for 30 min at room temperature. Then, the mix was added to NG108-15 cells or m-IC_{cl2} cells plated on glass cover slides, and the cells were incubated for additional 10 min at 37°C before being processed for microscopy.

Quantification of HcB-Cy3 entry in untreated cells and cells pretreated with gg (B) or GM1 (D) was performed as described in Fig. 3. The level of internalized HcB-Cy3 in control NG108-15 or m-IC_{cl2} cells was arbitrarily assigned to 1. Data are mean values ± SD from at least three independent experiments. Differences between means were tested using Student’s t-test, and p-values <0.001 were taken to indicate significance (***).
Figure 6. Inhibition of ganglioside synthesis impairs HcB-Cy3 entry into m-IC\textsubscript{c2} cells. (A) Untreated cells and cells treated with PPMP (7.5 \( \mu \)g/ml) for 48 h were exposed to HcB-Cy3 (2.5 \( \mu \)g/ml) for 10 min. For ganglioside complementation, the cells treated with PPMP for 24
h were supplemented with the mixture of gangliosides enriched in GD$_{1a}$, GD$_{1b}$, and GT$_{1b}$ (250 µg/ml) (PPMP + gg). (B) Quantification of HcB-Cy3 entry in untreated cells and cells pretreated with PPMP or PPMP + gg was performed as described in Fig. 3. The level of internalized HcB-Cy3 in control m-IC$_{cell}$ cells was arbitrarily assigned to 1. Data are mean values ± SD from at least three independent experiments. Differences between means were tested using Student’s $t$-test, and $p$-values <0.001 were taken to indicate significance (***)

Figure 7. HcB-Cy3 entry into m-IC<sub>cl2</sub> cells is not dependent of synaptotagmin (Syt) I and II.

HcB-Cy3 (2.5 μg/ml) was preincubated with the recombinant domain of (A) SytI or (C) SytII (25 μg/ml) for 30 min at room temperature. Then the mix was added to NG108-15 cells or m-IC<sub>cl2</sub> cells plated on glass cover slides and the cells were incubated for additional 10 min at 37°C before being processed for microscopy.

HcB-Cy3 entry was quantified in control cells and cells pretreated with SytI (B) or SytII (D) as described in Fig. 3. Data are mean values ± SD from at least three independent experiments. Differences between means were tested using Student’s t-test, and p-values <0.001 were taken to indicate significance (***).
Figure 8. Both gangliosides (GD$_{1a}$, GT$_{1b}$) and synaptotagmin are required for HcB entry into neuronal cells.

HcB-Cy3 (2.5 μg/ml) was preincubated with both a preparation (gg) enriched in gangliosides (18% GM1, 55% GD$_{1a}$, 15% GD$_{1b}$, 10% GT$_{1b}$, 2% others; 250 μg/ml) and the recombinant domain of (A) SytI or (C) SytII (25 μg/ml) for 30 min at room temperature. Then the mix was added to NG108-15 cells and m-IC$_{502}$ cells plated on glass cover slides, and the cells were incubated for additional 10 min at 37°C before being processed for microscopy. HcB-Cy3 entry was quantified in control cells and cells pretreated with gg and SytI (B) or gg and SytII (D) as described in Fig. 3. Data are mean values ± SD from at least three independent experiments. Differences between means were tested using Student’s $t$-test, and $p$-values <0.001 were taken to indicate significance (***)
Figure 9. HcB-Cy3 enters NG108-15 and m-IC_{cl2} cells through early endosomes. A) HcB-Cy3 (2.5 µg/ml) was incubated with NG108-15 cells or m-IC_{cl2} cells plated on glass slides for 10 min at 37°C. Then, the cells were fixed, permeabilized, and stained with antibodies against endogenous EEA1.

B) Quantification of colocalization between HcB-Cy3 and anti-EEA1. Colocalization index was determined as Pearson's coefficient. The distribution of cell structures recognized by HcB-Cy3 was expressed as the ratio of the number of cell structures co-stained with HcB-Cy3 and anti-EEA1 to the total of cell structures labeled with anti-EEA1. The mean values ± SD of the Pearson's correlation coefficient of 20-30 cells from at least three independent experiments are shown.
**Figure 10.** HcB-Cy3 enters NG108-15 cells and m-IC₁₂ cells through an early Rab7 compartment. A) HcB-Cy3 (2.5 μg/ml) was incubated with NG108-15 or m-IC₁₂ cells plated on glass slides for 10 min at 37°C. Then, the cells were fixed, permeabilized, and stained with antibodies against Rab7. B) Quantification of colocalization between HcB-Cy3 and anti-Rab7. Colocalization index was determined as Pearson's coefficient. The distribution of cell structures recognized by HcB-Cy3 was expressed as the ratio of the number of cell structures co-stained with HcB-Cy3 and anti-Rab7 to the total of cell structures labeled with anti-Rab7. The mean values ± SD of the Pearson's correlation coefficient of 20-30 cells from at least three independent experiments are shown.
B

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C

Colocalisation index

- NG108-15
- m-ICd2

Time (min)

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Figure 11. Differential HcB-Cy3 trafficking through late endosomes in NG108-15 cells compared to m-IC<sub>cl2</sub> cells. A) NG108-15 cells plated on glass slide were incubated with HcB-Cy3 (2.5 μg/ml) for 3 min at 37°C. Then, the culture medium was replaced with Dulbecco's medium. The cells were further incubated at 37°C for the indicated times, and were fixed, permeabilized, stained with antibodies against Lamp1, and processed for confocal microscopy.

B) HcB-Cy3 (2.5 μg/ml) was incubated in the apical chamber of polarized m-IC<sub>cl2</sub> cells grown on filters for 3 min at 37°C. Then, the culture medium was replaced with Dulbecco's medium. The cells were further incubated at 37°C for the indicated times, and were fixed, permeabilized, stained with antibodies against Lamp1, and processed for confocal microscopy. Representative plans from apical to basolateral side overtime are shown.

C) Quantification of colocalization between HcB-Cy3 and anti-Lamp1. Colocalization index was determined as Pearson's coefficient. The distribution of cell structures recognized by HcB-Cy3 was expressed as the ration of the number of cell structures co-stained with HcB-Cy3 and anti-Lamp1 to the total of cell structures labeled with anti-Lamp1. The mean values ± SD of the Pearson's correlation coefficient of 20-30 cells from at least three independent experiments are shown.
Figure 12. HcB-Cy3 entry is Cdc42 dependent in intestinal m-IC<sub>cl2</sub> cells but not in neuronal NG108-15 cells. A) m-IC<sub>cl2</sub> cells were transfected with a plasmid encoding GFP-Cdc42T17N and plated on glass slide. The cells were incubated with HcB-Cy3 (2.5 μg/ml) for 10 min at 37°C, and then were processed for confocal microscopy. Arrows indicate transfected Cdc42T17N cells.

B) Quantitative analysis of HcB-Cy3 entry into cells overexpressing Cdc42 dominant negative (Cdc42T17N) was performed by counting the total number of vesicles containing HcB-Cy3 in several cells (n > 40) and in different fields obtained from at least three independent experiments. The mean values were determined for both transfected cells and non-transfected (control) cells. Data are expressed as the mean ratio ± SD between non-transfected or transfected cells versus total number of cells. Differences between means were tested using Student’s t-test, and p-values <0.001 were taken to indicate significance (***).