Botulinum neurotoxin type-A (BoNT/A) formulations are widely used in clinical practice. Although they share a common mechanism of action resulting in presynaptic block in acetylcholine release, their structure and pharmacological properties demonstrate some similarities and many differences. Bioequivalence has been discussed since the onset of the clinical use of BoNT/A. In this review, we provide an update on the studies and compare the molecular structure, mechanisms of action, diffusion and spread, as well as immunogenicity and dose equivalence of onabotulinumtoxinA, abobotulinumtoxinA and incobotulinumtoxinA.

Key words: botulinum toxin A formulation, pharmacological similarities and differences, abobotulinumtoxinA, onabotulinumtoxinA, incobotulinumtoxinA

Introduction

Botulinum toxins are ‘natural products’ of living bacteria of the genus Clostridium. Particular therapeutics of botulinum toxin, although based on the same serotype A formulations (BoNT/A), have distinct properties. The main three BoNT/A products commercially available worldwide today are derived from Clostridium botulinum Hall strain: onabotulinumtoxinA (ONA-BoNT/A) marketed as Botox/Vistabel by Allergan Inc. (Irvine, CA, USA); abobotulinumtoxinA (ABO-BoNT/A) marketed as Dysport/Azzalure by Ipsen/Galderma (Paris, France); and incobotulinumtoxinA (INCO-BoNT/A) marketed as Xeomin/Bocouture by MerzPharmaceuticalGmbh (Frankfurt, Germany).

New BoNT/A formulations have recently been introduced to the market: prabotulinumtoxinA-xfvś (PRA-BoNT/A) marketed as Jeuveau/Nabota/Nuceiva by Evolus/Daewoong and daxibotulinum toxin A (DAXI-BoNT/A) (formerly RT002) by Revance. Additionally, letibotulinum toxinA (Croma/Hugel with Botulax) is in Phase III trials although results have not been published yet.

Botulinum neurotoxin type-A preparations use in clinical practice is based on presynaptic chemical denervation of cholinergic synapses due to the cleavage of specific synaptic

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proteins. This results in a decrease of acetylcholine release. Despite the common mechanism, these preparations are distinct medications, with many differences in terms of their structure, potency and immunogenicity. These differences may result in differing clinical efficacy and safety as well as pharmacoeconomic profiles, and have been discussed in medical literature over many years.

The aim of this paper was to show pharmacological similarities among, and differences between, the three most widely used BoNT/A preparations: ONA-, ABO-, and INCO-BoNT/A.

**Structure**

ONA-BoNT/A and ABO-BoNT/A are purified neurotoxin complexes including the BoNT/A1 and BoNT/A2 toxin molecules, respectively, and neurotoxin accessory proteins: NAPs - three haemagglutinin (HA) proteins and one non-toxic non-HA protein. It has been suggested that non-toxic HA not only stabilises the biological activity of the product in vivo, but also enables HA-botulinum toxin complex to adhere to muscle tissue [1]. INCO-BoNT/A contains only purified BoNT/A1 [2–4]. Results from a few studies have shown that ~150 kDa BoNT/A protein is mostly linked with NAPs [5, 6] at physiological pH levels, but other studies have suggested that prior to or shortly after injection the NAPs dissociate from botulinum toxin [7, 8]. Another study concluded that ONA-BoNT/A and ABO-BoNT do not contain neurotoxins in complexed form [8]. All commercial products contain an excipient, known as human serum albumin (HSA), which improves toxin stability and diminishes toxin loss during lyophilisation, prevents protein aggregation and surface adsorption, as well as extends shelf life [4].

ONA-BoNT/A is vacuum dried, while ABO-BoNT/A and INCO-BoNT/A are freeze dried. All formulations before clinical use are reconstituted with sterile normal saline buffer, yielding a solution that is slightly acidic [9, 10]. The reconstitution processes of ONA-BoNT/A results in a complete dissociation of 900 kDa complexes and the release of more than 85% of neurotoxins in free form [8].

A comparison of selected characteristics of BoNT/A preparations is set out in Table 1.

**Mechanism of action**

The activity of the botulinum toxin known as 'chemical denervation' refers to the decrease of the pre-synaptic release of acetylcholine, and temporary muscle paresis or inhibition
of glandular secretion. Additionally, BoNT/A inhibits the release of other neurotransmitters and influences inflammatory cells. This is probably the basis of its antinociceptive activity [11].

The mechanism of action of botulinum toxin A includes:
1. binding to nerve terminals; 2. internalisation within an endocytic compartment; 3. translocation into the cytosol; 4. the cleavage of SNARE complex by L chain; and 5. reduction of acetylcholine release from the pre-synaptic terminal.

1. In detail, according to 1., the C-terminal heavy chain (H) contains a translocation domain (HN) and a receptor binding domain (HC). The HC includes an N-terminal subdomain (HCN) of unspecified function and a C-terminal subdomain (HCC) that selectively bonds to dual neuron-specific receptors - ganglioside GT1b, and the protein receptor SV2C on the presynaptic plasma membrane in particular neurons [12, 13]. BoNT serotype A1 and A2 binds to the glycosylated SV2C receptor synaptic vesicle glycoprotein 2C (SV2C) [13], which allows for rapid penetration of toxins, at similar rates, via the same synaptic vesicles. HCA2 has higher affinity for receptor and neurons than HCA1 [14]. Glycosylation Asn559 in SV2C is critical for binding of BoNT/A to presynaptic plasma membrane. Glycosylation patterns in this site vary among adult individuals [15]. Pirazzini et al. have suggested that this feature may be responsible for a different onset and duration of induced neuromyopathy in humans following administration of the same dose of BoNT/A1; probably, different amounts of bound toxin are likely to match different numbers of L chains entering the cytosol in nerve terminal [11].

2. The toxin enters the synaptic vesicles of motor axon terminals by endocytosis. Internalisation of BoNT/A is mediated by receptor of a polysialoganglioside (PSG), the glycosylated luminal domain of a synaptic vesicle protein and unique N-glycans attached to synaptic vesicle (SV) glycoproteins [16], as well as to E-cadherin [17–19], fibroblast growth factor and vanilloid receptors [20, 21]. The increased endocytosis rate of BoNT/A and a frequent exposure of the SV lumen was observed during stimulation of nerves. It has been demonstrated [18] that nontoxic HA protein (present in ONA- and ABO-BoNT/A) sequesters E-cadherin in the monomeric state, disrupts the intercellular epithelial barrier, and facilitates paracellular absorption of BoNT/A [20, 22].

3. The L chain (L) of toxin is translocated across the vesicle membrane into the cytosol. Acidification of the synaptic vesicle lumen triggers HN to form a channel to L translocation. Next, the L chain is released from H chain by reduction of the interchain disulfide bond.

4. Next, L chain cleaving the soluble N-ethylmaleimide-sensitive factor attachment protein receptors (SNAREs), particularly synaptosomal associated protein of 25 kDa, SNAP-25.

5. Finally, SNAP-25 prevents the docking and exocytosis of acetylcholine from pre-synaptic vesicles at neuromuscular junctions. Detailed mechanism of nerve paralysis by BoNT/A is described in a review by Pirazzini et al. [11].

The study by Grando and Zachary [23] presented the non-neuronal and non-muscular mechanisms and effects of BoNT/A in many normal and cancer cell lines. Differentially altered genes expression by BoNT/A involved in signal transduction, immunity and defence, protein metabolism and modification, neuronal activities, intracellular protein trafficking and muscle contraction [24] show the huge range of mechanisms and possible effects of botulinum toxin. For example, ONA-BoNT/A injection markedly reduced, by 53%, urothelial ATP release in patients with spinal cord injury [25], increased nitric oxide (NO) release from the urothelium in the bladder detrusor [26], decreased expression of purinergic receptors (P2X3) in the bladder mucosa [27], inhibited the evoked release of CGRP from afferent nerve terminals in the bladder reducing pain [28], reduced bladder inflammation by decreasing urothelial apoptosis and the expression of vascular endothelial growth factor in the bladders of patients with interstitial cystitis (IC)/bladder pain syndrome (BPS) [29]. Recent studies have revealed that BoNT/A has antinociceptive peripheral effects by blocking synaptic transmission of glutamate, dopamine, ATP and gamma-aminobutyric acid regulation and serotonin [30].

The results of analysis of over 40,000 BoNT/A treatment reports indicate that patients who received BoNT/A in a broad range of injection sites had a significantly lower number of depression reports compared to patients undergoing different treatments for the same conditions [31]. Such results have allowed for the introduction of ONA-BoNT/A in Phase III studies designed so as to obtain FDA indication for major depression [32, 33].

The main mechanism of the ability of BoNT/A to weaken hyperactive secretory cells and relax tense muscles is the same for currently registered preparations, while the detailed additional effects mentioned above are described for ONA-BoNT/A only.

**Diffusion and spread out**

Distant effects of BoNT/A formulations may be the result of haematogenous spread defined as migration in local and regional muscles [34, 35], or as distant migration in areas non-contiguous with the injection [36–39]. Diffusion is characterised as microscopic movement of a soluble molecule's dispersion by passive transport to local and distant tissues [40] away from the intended area to nearby anatomical structures. The local spread of all BoNT/A formulations after injection depends on dilution, needle size, dose and volume, as well as injection technique [41]. Kutschenko et al. [42] suggested that the volume of injection is one of the major factors influencing the degree of muscle paralysis. The volume-dependent
reduction of paresis in a wheel-running test was observed in mice injected with INCO-BoNT/A. Kutschenko et al. suggested that larger volumes induce more intense paresis [42]. Pirazzini et al. suggested that the amount of toxin needed for a certain application should be diluted according to the size of the muscle/area [11]. Based on this data, it may be considered that the diffusion of BoNT/A from the injection site is increased by its gradual dilution in increasing volumes of extracellular fluids thus diminished binding to the presynaptic membrane. Additionally, different degrees of paresis after ONA-BoNT/A, ABO-BoNT/A and INCO-BoNT/A were presented after injecting identical volumes (10 µL) containing the same number of mouse units of BoNT/A into both hind leg muscles. Based on this experiment, the conversion ratio of INCO-BoNT/A and ONA-BoNT/A was estimated as being between 1:0.75 and 1:0.5. ONA-BoNT/A has shown a two-fold greater potency than ABO-BoNT/A [42].

Aoki et al. [36] proposed that protein complex size and pharmacological properties influence the diffusion of BoNT/A. That study showed that high-molecular-weight toxin complex of ONA-BoNT/A limits tissue distribution compared to ABO-BoNT/A [36]. More recent studies in which the size of anhidrotic halos was measured have shown different results. A comparison of ONA-BoNT/A and ABO-BoNT/A (using dose ratios of 1:2.5, 1:3, and 1:4, and identical injection volumes) presented a larger area of anhidrosis after ABO-BoNT/A [43]. Kerscher et al. obtained different mean maximal areas of the forehead anhidrosis of patients at 6 weeks after injection of BoNT/A formulations: comparable spread to ONA-BoNT/A and INCO-BoNT/A, and significantly greater to ABO-BoNT/A [44]. In another study, no significant differences between the mean size of halos produced by ONA-BoNT/A and ABO-BoNT/A were observed [45]. Similarly, no differences in diffusion of ONA-BoNT/A and INCO-BoNT/A injected to forehead at the same dose and using the same technique were demonstrated after 6 weeks and 6 months [46]. In other study, similar, limited to a distance of 30–45 mm [41], diffusion from the site of injection has been well documented by N-CAM staining and characterised ONA-BoNT/A, INCO-BoNT/A and ABO-BoNT/A when they were used in a ratio of 1: 1: 4 and in the same toxin injection volume (25 µL) [47].

Results from the study by Brodsky et al. showed that the presence of complexing proteins in ONA-BoNT/A and ABO-BoNT/A does not reduce migration of the neurotoxin [48]. The diffusion for all formulations of BoNT/A is similar in the majority of studies, but the dose and volume of injection may be the most important factors in differentiating diffusion efficiency.

The retrograde axonal transport of BoNT/A to spinal motor neurons, followed by anterograde transport to the other motor units, has also been suggested [49]. Caleo et al. [50] showed that BoNT/A physically leaves the motoneurons to enter second-order neurons. After injection of ONA-BoNT/A into the nasolabial musculature of rats and mice, catalytically active ONA-BoNT/A was transported to the facial nucleus. The authors suggested that these findings highlight cell-specific, direct central actions of BoNT/A, which are important to fully understand its mechanisms of action and therapeutic effectiveness in movement disorders and pain treatment.

A few studies have shown that BoNT/A, injected intramuscularly, is transported both anterogradely along sensory axons and retrogradely by central neurons and motoneurons axons to the motoneuron soma in the spinal cord [51–54]. Autophagosomes undergo dynein-dependent retrograde axonal transport to the neuronal soma [55]. Moreover, Antonucci et al. observed SNAP-25 cleavage in the contralateral hemisphere after unilateral BoNT/A delivery to the hippocampus [51]. Harper et al. and Restani et al. showed that BoNT/A-HC is internalised in synaptic vesicles and undergoes retrograde trafficking [56, 57]. The retrograde axonal transport and transcytosis to second-order nociceptive neurons explains mechanisms of action of ONA-BoNT/A in migraine [58]. ONA-BoNT/A is the only one approved for the treatment of chronic migraine. Selected papers have presented retrograde transport for ONA-BoNT/A only.

According to the Simpson et al. study, botulinum toxin accesses the perineuronal fluid compartment and does not cross the blood–brain barrier [59]. These authors suggested that BoNT/A is a large molecule and it is not able to cross the blood–brain barrier.

**Immunogenicity**

Antibody formation against the accessory proteins was observed in patients after injection of BoNT/A formulation with associated proteins, but they did not interfere with the biological activity of the toxin (‘non-neutralising’) [60]. Results from a preclinical study suggest that the NAPs may physically secure neurotoxin against the immune system and finally against the formation of toxin-neutralising antibodies interfering with clinical response [61]. However, antibodies formed against the heavy chain may or may not prevent its biological activity. The immunological response of humans to BoNT/A is very low, ranging from 0% to 3%: 0% was reported for ONA-BoNT/A [62, 63] and for ABO-BoNT/A used in glabellar lines [63], 1.2% for ONA-BoNT/A [62, 63], less than 3% for ABO-BoNT/A in cervical dystonia [63], and 1.1% for INCO-BoNT/A in upper limb spasticity. Each patient injected with INCO-BoNT/A was previously treated with a botulinum toxin A product which contained complexing proteins [7, 64]. A comprehensive meta-analysis of 61 studies by Fabbri et al. [65] analysed the frequency of antibodies among 8,525 patients receiving all registered types of BoNT/A across several clinical indications. Generally, the prevalence of antibodies among clinically responding patients was lower (3.5%) than in secondary nonresponse patients (53.5%). The frequencies of antibody formation independent of clinical responsiveness
to BoNT/A formulations across all analysed clinical indications were 1.5% for ONA-BoNT/A, 1.7% for ABO-BoNT/A, and 0.5% for INCO-BoNT/A. The results of this analysis indicate the lowest frequency of antibody formation after INCO-BoNT/A. The prospective, single-arm, dose-titration TOWER study showed that no patient with spasticity with a cerebral cause developed secondary nonresponse due to neutralising antibodies after administration of INCO-BoNT/A in a range of doses between 400 and 800 [66].

Based on the results of the aforementioned studies, the presence of complexing proteins in BoNT/A formulations may increase the risk of the formation of neutralising antibodies.

The immunogenicity of the BoNT/A formulations depends on some factors that differ in the manufacturing process, mainly the source of toxin and the antigenic protein load and the presence of inactive or denatured toxin acting as a toxoid. Treatment-related factors such as the toxin dose, frequency of injections, as well as prior exposure via other routes (intradermal or distant to the target muscle), different formulations (e.g. first application of ONA-BoNT/A or ABO-BoNT/A and second of INCO-BoNT/A) and site of anatomical region (especially near lymph nodes) seem to play a role in the immunogenic response. Based on this knowledge, clinical practice suggests the use of the lowest effective doses and to maintain 12 weeks of minimal interval treatment [67]. On the other hand, shorter, less than half as long, intervals of injection of INCO-BoNT/A have been described as well tolerated and free of antibodies [68].

Doses

A dose equivalence of BoNT/A formulations is still being discussed. The potency of BoNT/A preparations is expressed as Units (U) and 1U corresponds to one LD50 in mouse bioassay [69, 70]. Different diluents for LD50 testing have used by manufacturers: Allergan uses saline [71]; Ipsen uses gelatin phosphate buffer [72]; human serum albumin as a stabiliser was added by Merz to undisclosed diluent [73]. However, it has been suggested that stabilisers can enhance the activity of BoNT/A products at low concentrations in preclinical tests [74]. It is suggested that the diluent buffer significantly influences biological activity of BoNT/A products. Nonparallel dose-response curves of ONA-BoNT/A and ABO-BoNT/A with different relative potencies can explain a dose conversion ratio between Botox and Dysport [74, 76] or 1:2 [43].

There are no internationally accepted standardised tests for BoNT/A product comparisons. For this reason, different assay methods with different proprietary product-specific reference standards for testing potency units are used. The clinical effect of one unit is not interchangeable between formulations due to differences in the bioassay methodologies used by producers [77]. The clinical literature has reported an equivalent potency between ONA-BoNT/A and INCO-BoNT/A [3], but this was not the case in an animal (mouse) study [42]. The potency of INCO-BoNT/A and ONA-BoNT/A in inducing hind limb paresis in the wheel-running performance test in mice showed a conversion rate of between 1: 0.75 and 1: 0.5 [42]. The Allergan LD50 assay used ONA-BoNT/A and INCO-BoNT/A diluted in normal saline [72] to compare their activity. The obtained results showed that one INCO-BoNT/A vial contained less than 100 Allergan units (i.e. 69–78 units for three different lots) and clearly suggested the non-interchangeability of units in the studied products. Additionally, these results were confirmed in an enzymatic cleavage assay, the Digit Abduction Score assay, as well as replication of the LD50 results [78, 79]. Dressler et al. [73] indicated that assay conditions markedly influence potency measurements. Moreover, dose-response data of BoNT/A formulations is used to determine the therapeutic dose range as the ‘benefit–risk’ rate from acceptable efficacy and safety profiles. Significantly different muscle weakening efficacies identified as 50% maximal (median effective dose -ED50) have been reported for the three main BoNT/A products, and furthermore not equipotent units of the botulinum toxin formulations that are under experimental conditions were presented [36, 79].

Additionally, different quantities of 150 kDa (ng protein/100 U) of BoNT/A in formulation (the lowest in INCO-BoNT/A and the highest in ONA-BoNT/A) were shown by Ferrari et al. [80] and Field et al. [81] (see also Table 1). Calculated analysis shows differences between BoNT/A formulations. The highest amount of neurotoxin per product unit (in pg) and the total amount of active BoNT-A (in ng) injected at the recommended dose for an adult lower limb and an adult upper limb were obtained for ABO-BoNT/A. However, the relative quantity of rBoNTA assessed as a ratio quantity obtained by the EndoPep method to protein quantity tested by ELISA method demonstrated not significant differences in LC activity-tested BoNT/A formulations. This indicates that the 150 kDa neurotoxin molecules in each product are equally active [81].

Some studies have shown the non-interchangeability of units of ONA-BoNT/A and INCO-BoNT/A. The highest total amount of active BoNT/A being found after injection of ABO-BoNT/A to lower and upper limbs may suggest a focus on the conversion rate between ONA-BoNT/A and ABO-BoNT/A or INCO-BoNT/A and ABO-BoNT/A.

Summary

The major difference in structure between BoNT/A formulations concerns the presence or absence of complexing proteins. The effectiveness of BoNT/A preparations is not dependent on complexing proteins, but they may increase the risk of the formation of neutralising antibodies. The mechanism of action of all BoNT/A is similar, but the central effects of ONA-BoNT/A may expand indications for its use in major depression. Diffusion and spread out for all formulations of BoNT/A is similar in most studies. The retrograde axonal transport and transcytosis to second-order nociceptive neurons described for
ONa-BoNT/A only justifies its usefulness in the treatment of chronic migraine. The non-interchangeability of units of ONa-BoNT/A and INCO-BoNT/A was shown in an animal study, and the highest total amount of active BoNT/A after ABO-BoNT/A injection to lower and upper limbs may suggest the correct conversion rate doses between BoNT/A preparations.

Based on the differences in biological assays and the variations of biological activity [82, 83], regulatory agencies in most countries worldwide require a statement of unit non-interchangeability among BoNT/A products.

References


