Emerging aspects in Botulinum Toxin use

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Abstract
With an established number of licensed medical indications, Botulinum Toxin type A (BoNT-A) is also globally acknowledged to effectively refine many of the senescent changes which occur in the ageing face. BoNT-A is widely considered to be an effective first line therapy to eradicate dynamic lines, as well as an adjunctive treatment for a number of aesthetic rejuvenation strategies in softening more established rhytides. The exceptional popularity of the products for aesthetic use and the new potential applications currently under investigation remain unabated. Such demand reinforces the importance of continuous educational updates for the medical practitioner to provide valuable information on the evolving aesthetic and medical modalities, ensuring an optimal understanding of the uses of the product, and safe treatment outcomes for the patient.

Key words: Botulinum Toxin type A, emerging indications, delivery devices, pharmacology, dermatology, pain

Introduction
In this article, we will examine the currently available literature surrounding the emerging indications for Botulinum Toxin type A (BoNT-A), as well as recent pharmacological developments. BoNT remains one of the most extensively researched drugs to date and a very popular treatment for patients. A recent annual survey by the American Society of Plastic Surgeons indicates that there were over over 6 million aesthetic treatments with BoNT-A alone during 2013, reinforcing to the practitioner the importance in maintaining an awareness of current developments. Clinicians should therefore continue to develop their clinical knowledge and practice to ultimately deliver an enhanced aesthetic result.

The history of BoNT in medicine is well known. Initially isolated from an outbreak of food poisoning, the causative organism Clostridium botulinum soon became of intense interest due to the exceptional potency of the neurotoxin produced (Fig. 1). Following the isolation of BoNT, clinical trials and the path to product registration were followed, both in the United States with the product Oculinum (eventually Botox) and Europe as the product Dysport. Since then, the molecule has been isolated and characterized (Fig. 2), but there are still unknown aspects about the mode of action that are elusive and not yet fully determined. For example, why does the activity of the molecule persist for many months when normally, any foreign proteins in the body are rapidly identified and eliminated? Recent evidence indicates that the common phenomenon of phosphorylation may play a role, although this may also significantly reduce enzymatic activity. Currently there are a number of BoNT-A products available worldwide, with regional products and others in development (Fig 3) – a recent number emerging from South Korea in particular. The pharmacology of BoNT-A, when used in aesthetic treatments and for medical indications, has been the subject of considerable discussion in recent years. Attempts have been made, based on pseudo-scientific arguments, to distinguish certain products from the others. However, the arguments used were identified as incorrect some years ago. Originally, these arguments related to the existence of the so-called BoNT complex, a natural formation of the active neurotoxin, produced by the bacteria, that serves to protect the neurotoxin element in the natural environment. The neurotoxin molecule is naturally protected by a family of related proteins. However, unlike certain claims, these accessory proteins were found to dissociate from the neurotoxin molecule during reconstitution of the products in the vial. Apparent product differences relating to, typically, the size of the BoNT-A complex were therefore inaccurate and misleading. Additional features of BoNT-A pharmacology are defined by the injection itself and what occurs subsequently. Diffusion refers to the BoNT complex moving from an area of high concentration to an area of low concentration during the binding to receptors, whereas Spread refers to the physical movement of the product once injected, which may be caused by factors such as dilution volume, needle gauge and injector. The pharmacodynamics of BoNT-A, in particular its uptake into the neuromuscular junction (NMJ), is of clinical importance. Recent reports indicate a rapid uptake with no residual BoNT-A in the injected muscle within 20 minutes. Thus questioning post injection protocols currently being promoted.

The main areas to consider in the use of BoNT represent:
1. Toxin effects on muscles,
2. Effects on glands,
3. Pain relief,
4. Dermatological uses and
5. Methods of delivery.

1. BoNT effect on muscles
The specific target for BoNT is the NMJ, where the nerve synapse connects to the muscle systems. This is a highly complex, universal connection within the body and much about how the NMJ functions is still a subject of investigation. In particular, the existence and role of the neurotransmitters involved in overall signaling processes continues to be an area of discovery. The specific anatomical distribution of the NMJ within a muscle is of critical importance in understanding the effects of BoNT-A on muscle dynamics. Unlike other muscles in the body, facial muscle fibres have single or multiple NMJs which determine the ability to control very fine and intricate muscle movements. However, apart from limited studies to date, we have little information on these target sites in the majority of other facial muscles, even though knowledge of the gross anatomy of these muscles, with respect to use of BoNT-A, has recently been described. A more detailed understanding of the effects of BoNT-A on facial muscle activity with respect to dose, action and recovery, through the use of electromyography, is also being gained. Detailed understanding is essential in optimization of the dose-effect relationship for BoNT-A, in order to gain the best aesthetic results for the patient, with minimal doses. Facial aging is a universal process that affects many different aspects of size, shape, volume and function. However, in addition to ageing effects on muscle size and strength, there is one additional effect that has been overlooked until now – the ageing of NMJs. With clear evidence of such ageing processes,
2. Glandular effects
The effects of BoNT-A on hyperhidrosis are well documented, with over 160 publications covering nearly 20 years of use. The products have widespread acceptability for treatment of the condition, wherever occurring on the body, in both paediatric and adult applications, despite only being approved for axillary hyperhidrosis. BoNT-A is also used to treat gustatory sweating (Frey's Syndrome). Additional recent applications include the use for treatment of salivary glands for control of drooling by reduction of hypersecretion. This continues to be a well-studied area, although a recent Cochrane Review found insufficient evidence to inform clinicians clearly about clinical interventions using BoNT for drooling in children with cerebral palsy.

3. Pain relief
The use of BoNT for the relief of pain has been studied in many different conditions. To date, only the prophylactic treatment of migraines has been approved by the licensing authorities for one of the products. Nevertheless, use of BoNT for routine treatment of many pain conditions has found a widespread, off-label usage. There is little doubt that further, detailed clinical trials, yet to be performed, will bring established use of BoNT in these types of pain conditions. There are close links between certain aspects of the pain indications under investigation following treatment with BoNT, and a number of painful dermatological conditions, as discussed below.

4. Dermatological uses
The dermatological potential for BoNT-A has recently been investigated with early findings in support of treating symptoms associated with rosacea, certain types of psoriasis, facial inflammatory diseases and acne. OnabotulinumtoxinA has been used experimentally to treat erythema and flushing in rosacea and been shown to demonstrate, through the neurogenic component, an ability to actively address vascular dysfunction and inflammation, with potential to reduce hyper seboration. Dayan and colleagues conducted a two-year survey comprising a small number of subjects (13) presenting with rosacea. The authors utilised an intralesional microdroplet injection technique (0.05ml) of onabotulinumtoxinA, which was 100 units in a dilution of 7ml. Multiple injections were performed intradermally, which gave a dose on average from 8-12 units per affected side. Subjects were reviewed at three weeks and again at three months with a visible reduction in erythema reported, with the authors measuring outcomes on the basis of before and after photography, as well as verbal feedback. Some subjects experienced longevity of results beyond three months. The findings indicate a decrease in persistent as well as intermittent flushing, as well as reduced erythema and inflammation. A range of painful facial conditions, often involving inflammation, have also been studied. These include natalgia paresthetica, refractory erythromelalgia, and hidradenitis suppurativa. Other developments include additional knowledge on the effect of BoNT-A on fibroblasts – showing a positive effect with collagen production.

5. Methods of Delivery
Several areas need to be considered in delivering the treatment of BoNT-A:

I. No-needle techniques
II. Pain on injection
III. Accuracy of dosage
IV. No-needle injections

Studies to inject BoNT-A using several delivery devices have looked at the ability to drive the product through the epidermis and dermis.
This can be readily achieved with the electrical technique of iontophoresis.\(^5\)\(^5\)\(^6\) \(^7\)

However, pressure delivery using the Dermojet has also been evaluated and is a technique to deliver the product into different areas, for example, for treating palmar hyperhidrosis. However there is no control over depth of injection and the toxin may be delivered deeper than required, resulting in adverse consequences. More recent research is looking at techniques to modulate the skin barrier and allow penetration of BoNT through the dermis. Here, formulation of the BoNT using various carriers is a key area. The US-based company Revance has completed limited Phase 3 testing of a topical BoNT-A gel. This would allow treatment of conditions such as hyperhidrosis and wrinkles due to superficial muscles – typically Crow’s feet, without a needle to puncture the skin. The company is one of around four in the world examining the potential for topical application of BoNT, but the doses required are known to be much higher than used for injection, and the effects are limited to the more superficial facial muscles; not the stronger and deeper muscles of the glabella complex.

**II Pain**

The volume of solution injected is directly related to the volume used for reconstitution of the product from the powder form, which is provided for each commercial product. Each manufacturer has specific recommendations for the volume of reconstitution of their product, based specifically upon the format used in the registration clinical trials. These can be different for each product. Many clinicians, however, have preferred volumes of reconstitution based on volumes of injection that they are used to. Therefore there is considerable variability between the injection techniques used – with regard to volume from the official recommendations. Trials have been carried out on different volumes of injection, with the same units being administered in each treatment, but the results have demonstrated either no difference in onset or duration of the effects obtained\(^5\)\(^2\) or the opposite, contrary result.\(^5\)\(^3\) Volume of injection has been reported as unlikely to have an effect over the standard range of product reconstitution volumes recommended.\(^5\)\(^4\)

In one study, the volume of injection was found to be related to pain on injection, with a higher volume giving more pain. Unfortunately, there were insufficient patients in the study to conclude significance, but nevertheless a guide was established.\(^5\)\(^6\)

Since the initial use of BoNT-A for aesthetic corrections, clinicians have often chosen to use a diluent containing a preservative (benzyl alcohol) instead of the diluent recommended by the manufacturers: unpreserved saline. To date, nothing has been published directly from the manufacturers when using preserved saline for their products but clinician-led studies are available,\(^5\)\(^6\)\(^5\)\(^8\)\(^5\)\(^9\) and these indicate a reduction in pain on injection when using preserved saline.

There are two possible reasons for this. Firstly, normal (pharmacopoeial) 0.9% saline is incorrectly considered to be a physiological solution.\(^5\)\(^9\) Therefore any change from this state, especially if the pH is nearer to neutrality, could improve the pain perception upon injection. But secondly, benzyl alcohol is also recognized to have a minor anesthetic effect when injected. The net effect is likely to be a combination of these and other factors.

### MAIN BoNT-A PRODUCTS

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<td>HSA 125 ug, Lactose 2.5mg</td>
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<td>Fermentation Precipitation “Crystallisation”</td>
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### BoNT-A PRODUCTS FROM ASIA

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One aspect of injection pain often overlooked is the needle size being employed. Only limited work has been undertaken, but the latest study, from a group of clinicians in Turkey, indicates a smaller 33 gauge needle gives less pain in four areas of the face studied when compared to a 30 gauge needle.10

To further minimize patient discomfort during the injection procedure, Kim and colleagues16 describe findings from their survey comprising 181 participants on their satisfaction after receiving BoNT-A, reconstituted with 1% lidocaine & epinephrine 1:100,000. The addition of lidocaine achieved an immediate paralyzing effect on the treated muscles, with the epinephrine reducing spread to adjacent muscles. The findings presented no decrease in pharmacological potency or increase in adverse effects, with some subjects reporting longer duration of aesthetic effect. The survey established no hypotheses for the possible causes of such finding. The statistical data reflects positively in favour of utilising the combination of lidocaine and epinephrine, since 85.7% (78/95) of the participants felt that such combination was superior, 35.7% (56/157) reported no difference in the result and 6.4% (10/157) stated no improvement. However, this is a single study with limited data and no recommendations on this technique may be made without more supporting evidence.

### III Accuracy of Dosage

With low volumes of solution being injected, it is important to be able to deliver accurate dosages and minimise loss of BoNT due to dead-space in the equipment being used. New devices are being developed to deliver BoNT-A, such as the 3Dose Injection Syringe, which has space in the equipment being used. New devices are being developed to deliver accurate dosages and minimise loss of BoNT due to dead-space loss by 0.08mls compared to the conventional needle and syringe combination. With a 33 gauge needle this reduces product wastage and causes minimal pain from the needle stick. Such devices can enable the practitioner to accurately deliver the required dose to the target muscle, however to achieve optimal results, an accurate knowledge of anatomy is essential.

**Conclusions:**
The advancing science and applications of BoNT strengthen the importance for practitioners to have knowledge of current developments in the field. Further developments, resulting in numerous "off-label" indications, continue to be identified. Knowledge of BoNT is constantly evolving within aesthetic practice, moving from purely treating dynamic lines and facial contouring (masseter) to treating dermatological conditions with significant clinical benefits to patients. BoNT continues to be developed with adjunctive technologies being added to deliver toxim to specific areas with accurate dosage. Intricate and detailed anatomical knowledge, together with these advances, will allow practitioners to achieve optimal results.

**Disclosures**
Andy Pickett is Director and Founder of Toxin Science Limited, Wrexham UK and Adjunct Professor at the Botulinum Research Center, Institute of Advances Sciences, Dartmouth, USA. He is also a Senior Programme Leader in Galderma Aesthetic and Corrective Business Unit. His views and opinions are his own and those of Toxin Science Limited alone.

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