Botulinum toxin type A for the management of glabellar rhytids

Anne Marie Tremaine
Jerry L McCullough

Department of Dermatology, University of California, Irvine, CA, USA

Abstract: There is an increasing demand for minimally-invasive cosmetic procedures to arrest the aging process. Botulinum toxin type A injections are the most commonly used nonsurgical cosmetic procedures in the United States. There has been research spanning over two decades dedicated to safety, efficacy, dosing, and complications of botulinum toxin type A. There are now two Food and Drug Administration (FDA) approved botulinum toxin type A options in the United States: Botox® and Dysport™, with new advances being made in the field.

Keywords: botulinum toxin type A, glabella, Botox, Dysport, complications, dosing

Introduction

There is an increasing demand for a reversal of the aging process and recently more people are turning to minimally-invasive methods to meet this goal, in lieu of surgery. Botulinum toxin type A injections are the most commonly used nonsurgical cosmetic procedures in the United States, with 4.6 million procedures in 2007.1 Glabellar lines occur naturally with facial animation, a continuous practice that facilitates communication. Contraction of the procerus and corrugator supercilii muscles produces creasing of the glabellar skin and ultimately permanent rhytids develop. Administration of low doses of botulinum toxin type A into bilateral corrugator supercilii and procerus muscles paralyze muscular activity, thus diminishing the appearance of dynamic rhytids of the glabella. Botulinum toxin injection is a minimally invasive procedure with relatively quick onset of action seen within three days to two weeks of administration. The effects of botulinum toxin type A commonly last for three to six months, although there is a report of duration as long as twelve months.2 Botulinum toxin type A is a successful treatment clinically, but more importantly patient satisfaction is consistently high with its use.3 The purpose of this review is to discuss the important topics surrounding and pertaining to the role of botulinum toxin type A in the management of glabellar rhytids, as well as, to compare and contrast the FDA approved commercial botulinum toxin type A products.

Botulinum toxin is a neurotoxic protein produced by Clostridium botulinum, a Gram-negative anaerobic bacterium. In the 1980’s, Dr Alan Scott was the first to utilize botulinum toxin clinically with his research on strabismus and blepharospasm, after using it successfully in experiments using monkeys in the 1970’s. Now botulinum toxin is used to treat many medical conditions including cervical dystonia, hyperhidrosis, strabismus, and blepharospasm. The first study4 indicating the utility of botulinum toxin type A for the treatment of hyperfunctional facial lines occurred in the early 1990’s.
Normal facial muscle cell contraction occurs when acetylcholine released from the nerve terminal, diffuses across the synaptic cleft, and attaches to its receptor on the muscle cell (Figure 1a). After injection of botulinum toxin type A into the muscle, the neurotoxin is taken up by the adjacent nerve terminal. Within the terminal it prevents proper binding of the synaptic vesicle containing acetylcholine. The neurotoxin accomplishes this by cleaving SNAP-25, a protein that is crucial for docking of the vesicle to the nerve ending. Thus, neurotransmitter release into the synaptic cleft is inhibited and muscle contraction cannot occur (Figure 1b).

There are several botulinum toxin type A products available worldwide. In North America, onabotulinumtoxinA, known as Botox® (Allergan, Inc., Irvine, CA, USA) and abobotulinumtoxinA, known as Dysport™ (Ipsen Ltd, Wrexham, UK and Medicis Aesthetics Inc., Scottsdale, AZ) are FDA approved products for aesthetic treatments. Botox® was approved in 1989 for certain medical conditions, but it was not until 2002 that this product received FDA approval for temporary improvement in the appearance of moderate to severe glabellar lines in adults aged 65 or younger. It was just several months ago that Dysport™ gained FDA approval for cosmetic indications although it has been used widely in Europe for several years. The list of ingredients for each of these products is listed in Table 1. Two other botulinum toxin type A products are available outside the United States for treatment of blepharospasm and torticollis: Xeomin® (Mertz, Frankfurt, Germany) and Neuronox® (Medy-Tox Inc., South Korea).

Botox®, Dysport™, and any other botulinum toxin type A products are distinctive and not interchangeable. Thus, the generic term, botulinum toxin type A, will be used when a distinction does not need to be made, but otherwise the specific product will be named throughout the review.

Safety and efficacy of botulinum toxin type A

The safety and efficacy of botulinum toxin type A has been investigated in both small, single-center trials,4-12
as well as, large, multicenter, randomized, double-blind studies.\textsuperscript{13–17} Recently, several trials have addressed the concern of long-term safety and efficacy of botulinum toxin type A and these found sustained safety and efficacy after repeated administration of the neurotoxin.\textsuperscript{15,18–23} There is unequivocal evidence from two decades of research that botulinum toxin type A is safe and effective for the treatment of glabellar lines.

Two interesting clinical pearls have been discovered during the course of the research trials. First, while the data is limited and not statistically significant, it appears that there is a significant decrease in efficacy seen in subjects older than 65 years of age.\textsuperscript{16,24,25} Second, Carruthers et al. found that the effects on patient appearance at rest appear to be sustained longer than the effect at maximal frown, thus suggesting that botulinum toxin type A may have a persistent benefit even once the paralysis has reversed. The authors hypothesize that this may occur for several reasons including dermal remodeling, slight muscle atrophy, and behavior modification.\textsuperscript{13–14}

**Dosing**

Many different doses have been evaluated for the treatment of glabellar rhytids, although the current suggested dose is 20 Units (U) for Botox\textsuperscript{®} and 50 U for Dysport\textsuperscript{™}.\textsuperscript{24,25} It is recommended that there should be at least a three month period between botulinum toxin injections. Carruthers et al. compared 10, 20, 30, and 40 U doses of Botox\textsuperscript{®} for the improvement in glabellar rhytids in females and found that dosing with 20–40 U of botulinum toxin type A was significantly more effective at reducing glabellar lines than 10 U alone.\textsuperscript{2} A study evaluating 20 U compared to 30 U in African American females found that either dose was safe and effective.\textsuperscript{27} Larger doses of botulinum toxin type A are needed to improve the glabellar rhytids in men. In one dose-comparing study, it was shown that 20 U of Botox\textsuperscript{®} was
ineffective and greater treatment outcomes were seen with the use of 40, 60, and 80 U doses. 28

In June of 2009, Carruthers et al published a dose-comparing study with Botox® for the improvement in upper facial rhytids (crow’s feet, forehead, and glabella) and found similar efficacy and safety with 32, 64, and 96 U. Due to the dose-dependent response observed and a higher incidence of brow ptosis with 96 U, the authors concluded it is best to use a more moderate 64 U. 29

There is less published literature evaluating dosing for Dysport™. After a Phase II dose-ranging trial, 26 most subsequent trials evaluated a 50 U dose compared to placebo and found it to be safe and effective.15,17 Another dose-ranging trial comparing 20, 50, 75 U, or placebo found 50 U of Dysport™ to be the optimal dose. 30 Comparatively, another trial found that both 30 U and 50 U were safe and effective, although there was a slight improvement in treatment outcomes with the 50 U dose. 31 Kane et al propose that a standardized dose is not optimal since gender and muscle mass affect the dose required for efficacy. 16 This group carried out a randomized, double-blind, placebo-controlled trial that evaluated variable doses of Dysport™ based on gender and muscle mass. After grading the mass of the procerus and corrugator muscles (small, medium, large) the investigators established the dosing for each subject with options including: 50, 60, or 70 U in women and 60, 70, or 80 U in men. The investigators found the variable doses to be efficacious compared to placebo and did not report an increase in adverse events in the higher treatment doses.

As mentioned, Botox®, Dysport™, and any other botulinum toxin type A product have different potencies and the doses are not interchangeable. The biologic activity of the toxin is measured in mouse units (LD$_{50}$), the median intraperitoneal lethal dose in mice. Despite the fact that both neurotoxins are supplied in units of biologic activity (LD$_{50}$), the units are not transposable due to differences in assays used to determine the units. There have been a few studies attempting to create dose ratios between Botox® and Dysport™ to compare efficacy. Lowe et al found 2.5:1 dose ratio (Dysport™: Botox®) to be comparable in terms of tolerability. The authors reported prolonged efficacy and higher patient satisfaction in the Botox® treatment arm at the week 12 timepoint. 32 Others suggest that at 4:1 dose ratio (Dysport™: Botox®) provides similar clinical efficacy. 33,34 Wohlfarth et al recently carried out a systematic review of preclinical and clinical dose ratio studies of botulinum toxin type A used for various therapeutic indications. The authors found a range of Dysport™: Botox® ratios from 2:1 to 11:1 described in the literature. Their literature review established that randomized, controlled clinical trials indicate that a 3:1 dose ratio is more appropriate than 4:1, and there is no evidence for a ratio greater than 4:1. However, the studies do not prove that a 3:1 dose ratio is equivalent clinically. 35 A previous study by Wohlfarth et al used statistical modeling with compound muscle action potential amplitude of the extensor digitorum brevis to confirm the use of a 3:1 dose ratio. 36

### Dilution

There is disagreement over the effect of dilution volume on treatment efficacy and there is limited clinical trial data to support either argument. Dilutions range between 100 U/cc to 10 U/cc, with most choosing to dilute 100 U of botulinum toxin with 1–3 cc of saline. Using a more concentrated solution, such as 1 U/0.1 cc, may allow for more accurate placement, limit pain, restrict diffusion, and thus decrease risk of side effects. Administering less concentrated doses, such as 4 U/1 cc, may be easier to work with. The increased potential for diffusion with lower concentration, higher volume doses can be advantageous since it allows fewer units to cover a greater area. Dilution may just be a matter of preference, as one comparative study for the treatment of blepharospasm showed no difference in efficacy or incidence of patient-reported complications (eg, bruising, redness, complications of injection) with two different dilutions (10 U/cc vs. 100 U/cc) injected into either ocular area. 37 Similarly, other studies found that concentrations varying from 10 to 100 U/cc had no difference in efficacy and the adverse events experienced were similar across all dilution arms. The more dilute treatment arms did have more subjects with swelling and ptosis.

### Table I List of ingredients for Botox® and Dysport™

<table>
<thead>
<tr>
<th>Product</th>
<th>Active ingredients</th>
<th>Inactive ingredients</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>BOTOX®</strong> (onabotulinumtoxinA)</td>
<td>50 Units of botulinum toxin type A</td>
<td>0.25 mg of human albumin, 0.45 mg of sodium chloride</td>
</tr>
<tr>
<td></td>
<td>100 Units of botulinum toxin type A</td>
<td>0.5 mg of human albumin, 0.9 mg of sodium chloride</td>
</tr>
<tr>
<td><strong>DYSPORT™</strong> (abobotulinumtoxinA)</td>
<td>300 Units of botulinum toxin type A</td>
<td>125 mcg of human albumin, 2.5 mg of lactose</td>
</tr>
<tr>
<td></td>
<td>500 Units of botulinum toxin type A</td>
<td>125 mcg of human albumin, 2.5 mg of lactose</td>
</tr>
</tbody>
</table>

**Note:** Each product is supplied in multiple unit vials.

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[Reference numbers cited in the text.]
although the data was not statistically significant.\textsuperscript{38,39} Hankins et al found no difference in efficacy or adverse events in concentrations ranging from 50 to 200 U/cc.\textsuperscript{7} However, a study conducted by Hsu et al determined that dilution did impact treatment efficacy, as the larger volume injections led to greater diffusion and increased the area affected.\textsuperscript{40}

With the more concentrated dilutions, it is efficient to inject with a 30-gauge needle and a 0.3 cc insulin syringe. The insulin syringe does not have a dead space within the needle hub and this serves to decrease the waste of botulinum toxin within the needle. With more dilute concentrations, it is best to use a traditional 30–32 gauge needle.

**Diffusion**

With injections into the small targets of the glabella and knowledge of the delicate surrounding areas, there is concern of product diffusion leading to muscle paralysis outside of the target site. There is debate about the mechanism of diffusion, as well as, how and if diffusion differs among botulinum toxin type A products.

Some believe the complex size of the botulinum toxin affects the diffusion potential and that larger proteins have less diffusion potential. If this is the case, Botox\textsuperscript{®} (uniform 900 kDa complexes) would be less likely to diffuse outside the target tissue compared to Dysport\textsuperscript{™} (heterogeneous mixture of 500–900 kDa complexes). The molecular weight of each product is reported and refuted in the literature and hence the validity of this data is debatable.\textsuperscript{41}

There are several studies that have found no significant difference in diffusion between Botox\textsuperscript{®} and Dysport\textsuperscript{™}.\textsuperscript{36,42,43} Furthermore, investigators suggest the complex protein size is irrelevant since dissociation of the complex occurs immediately after injection, releasing uniform-sized botulinum toxin. These researchers believe diffusion simply depends on the concentration and volume of product used. In other words, the higher the concentration and the greater the volume, the greater the diffusion potential.\textsuperscript{36,41}

**Reconstitution and storage**

Previously, it was thought that botulinum toxin should be used within 4 hours of reconstitution.\textsuperscript{44} The Botox\textsuperscript{®} package insert instructs to use the product within 24 hours of reconstitution,\textsuperscript{24} while the Dysport\textsuperscript{™} package insert instructs use within 4 hours.\textsuperscript{25} The concern is decreased efficacy and increased bacterial growth the longer the reconstituted neurotoxin remains unused. Many different studies have evaluated longer reconstitution periods including 15 days, 42 days, and 49 days without lessening efficacy or producing evidence of bacterial contamination.\textsuperscript{45–47} Both preservative-free and preserved saline were used in these studies and both were shown to be safe after prolonged reconstitution. Once reconstituted, botulinum toxin must be kept at a refrigerated temperature between 2–8°C.

**Complications**

The most common side effects reported with botulinum toxin type A injections include pain, swelling, erythema, ecchymosis, respiratory infection, headache, nasopharyngitis, sinusitis, flu-like symptoms, nausea, and limited hypesthesia. In clinical trials with Botox\textsuperscript{®}, the incidence of headache, nausea, and flu-like symptoms in the treatment arm was the same as seen in the placebo arm.\textsuperscript{13,14,24}

Upper eyelid ptosis is also a complication in treatment of the glabellar region. Understanding anatomic landmarks and proper technique will decrease the incidence of ptosis. This complication arises when the neurotoxin diffuses through the orbital septum and affects the levator palpebrae superioris muscle. The rate of ptosis appears to be determined by the skill and experience of the injector. In studies with repeated administration of botulinum toxin type A, the rate of ptosis decreased over successive cycles indicating improved technique with experience.\textsuperscript{18} Reported rates of ptosis from large clinical trials are similar between Botox\textsuperscript{®} and Dysport\textsuperscript{™} and range from 0.8% to 5.4%.\textsuperscript{13–15,17–19,26} Ptosis can be treated with \(\alpha\)-adrenergic agonist (apraclonidine 0.5% or phenylephrine hydrochloride 2.5%) ophthalmic drops twice a day to the affected side.\textsuperscript{48}

The complications associated with botulinum toxin injection are most often mild and self-limited. Most complications are related to technique, hence complications decrease with proper training of facial anatomy, dosing, and injection technique.

**Administration technique**

Proper technique for injection of the glabellas is debatable, but variations of procedure will still lead to successful treatment outcomes. The patient should be seated in the upright position and all injections should be aimed away from the eye. The neurotoxin should be injected into the muscle belly of the procerus and corrugator supercilli muscles. Administration of the neurotoxin into the forehead should be superficial, aiming for the subcutis. The neurotoxin will spread down to the frontalis muscle. The toxin may spread up to 3 cm from the injection site and this should be considered during administration. There is increased likelihood of eyelid ptosis if the toxin is injected too inferior on the forehead. Medial corrugator injections should be placed 1 cm above the bony supraorbital ridge to avoid ptosis.
Massage after injection is another debated practice among physicians. Massaging horizontally may facilitate smoothing of the lump after injection, aid with diffusion, and prevent inferior spread of the botulinum toxin.\(^4\) Massage becomes problematic, however, if the botulinum toxin diffuses to other muscles. There are currently no studies to support either argument. Digital pressure after the injection may also decrease the diffusion of the toxin.\(^4\) After the treatment, patients may be instructed to stay upright for at least 4 hours to prevent the diffusion of the product in the wrong direction, although there is no data to confirm the necessity of this action. Some physicians also instruct patients to contract the treated muscles for the first couple hours to distribute the toxin to the entire muscle, but, again, this may be unnecessary.

Several of the complications discussed above can potentially be prevented. For instance, to prevent inducing ecchymosis with superficial injection the patient should be advised to avoid aspirin, nonsteroidal anti-inflammatory drugs (NSAIDS), and high-dose vitamin E\(^49\) for at least a week prior to injection. Likewise, removing makeup prior to injection will allow for better visualization of the treatment area and avoidance of small, superficial vessels.\(^50\) It has been deemed helpful to hold direct pressure and application of ice to the injection site.\(^48,50\) To decrease the amount of discomfort for the patient, use of a topical lidocaine prior to the procedure, and slow injection with small amounts of concentrated botulinum toxin type A using a 30–34 gauge needle is recommended.\(^48,51,52\)

Furthermore, it has been found that patients complain of less pain when the botulinum toxin is reconstituted with preserved saline, rather than sterile, non-preserved saline as directed on the package insert since the preservative in the saline, benzyl alcohol, acts as an anesthetic. This is supported by a finding in a randomized clinical trial where not only did the subjects report less pain with botulinum toxin type A reconstituted with preserved saline, but it was found to be as safe and effective as that reconstituted with non-preserved saline.\(^52\) With preserved saline, there does not appear to be an increased risk of bacterial contamination after prolonged reconstitution and repeated extractions from the bottle.\(^53\) Preparing patients for the possibility of headache after injection and instruction to use over the counter (OTC) analgesics can prevent unnecessary patient panic and worry.

### Contraindications to use

There are several contraindications for the use of botulinum toxin type A, including active infection at the injection site or history of a hypersensitivity reaction to any of the ingre-
dients (ie, human albumin, lactose, saline, botulinum toxin type A). Dysport™ may contain trace amounts of cow’s milk protein and patients known to be allergic to this should not be treated with Dysport™. However, it is safe for patients with lactose intolerance to receive Dysport™. Low doses of botulinum toxin may induce a neuromuscular crisis. Patients with a history of neuromuscular disorders including myasthenia gravis, Eaton–Lambert syndrome, and amyotrophic lateral sclerosis should not receive botulinum toxin injections. Patients taking aminoglycoside antibiotics including, but not limited to, amikacin, neomycin, streptomycin, tobramycin, or gentamicin should avoid botulinum toxin because this combination may potentiate the effect of the neurotoxin. Patients should also avoid botulinum toxin injections if they are taking other drugs that interfere with neuromuscular transmission such as magnesium sulfate, succinylcholine, penicillamine, tetracyclines, calcium channel blockers, lincosamides, polymyxins, or anticholinesterases. Despite the lack of evidence of teratogenicity with Botox® or Dysport™, both are category C drugs and should not be administered during pregnancy. It is not known whether the neurotoxin is excreted in human milk and therefore should not be used in nursing patients. Botulinum toxin type A should not be used in subjects with a bleeding disorder and used with care in patients taking medication that affects clotting. Relative contraindications to use include patients with unrealistic goals and patients with psychiatric disease.\(^24,25\)

### Botulinum toxin and resistance

Botulinum toxin is a potentially immunogenic protein that can cause neutralizing antibody formation with repeated injections. Antibody formation may lead to decreased effectiveness of the neurotoxin. There appears to be a heightened risk of antibody formation with increased doses and frequency of administration. With the first generation of Botox®, it was recommended that injections take place at greater than 1 month intervals and that no more than 100 Units be used in a patient at one time. The protein load of the newer generation Botox® has decreased 5-fold consequently reducing the antigenic potential.\(^48\) Several researchers in the fields of neurology and dermatology have looked into the issue of resistance and established a variety of findings. These findings must be evaluated while keeping in mind that observed incidence of antibody positivity in an assay may be affected by methodology and several other factors. In a four year trial for cervical dystonia, 4 of 326 (1.2%) subjects tested positive for antibodies. Three of these subjects stopped responding clinically to botulinum toxin.\(^54\) Lange et al\(^55\) report that while they had a large number of subjects...
with neutralizing antibodies, many of them still responded clinically to treatment. Secondary non-responders with neutralizing antibodies were seen in higher dose indications (eg, focal spasticity and spasmodic torticollis) and with shorter injection intervals. Also, neutralizing antibody development was independent of the commercial preparation used. Other trials within the scope of cosmetic treatments found no antibody formation with repeated botulinum toxin type A injections. It is possible that the dosage of botulinum toxin type A for cosmetic indications is not significant enough to induce antibody formation.

Evaluating treatment outcomes

In clinical trials, treatment outcomes are measured with investigator global assessments and patient satisfaction rating scales. These traditional global assessments do not evaluate the specific outcomes that are significant to patients. Outside of clinical trials to determine efficacy, patient reported outcomes are truly the most important measure of treatment success. Carruthers et al have proposed two new patient-reported outcome measures: the facial line outcome questionnaire (FLO) and the self-perception of age (SPA). The FLO uses scales to allow patients to rate the extent to which their facial lines impact their self-perceptions. The goal of the SPA is to assess the patient’s current perception of his or her age of appearance. The FLO and SPA assessments provide evidence that botulinum toxin type A injections are improving patients’ perception of themselves, which is the real determinant of treatment success with cosmetic procedures.

Improving treatment outcomes

As discussed above, patient satisfaction is crucial and there are several ways to improve treatment outcomes in this area. The initial visit should consist of a detailed consultation discussing the goals and expectations of each patient. It is advisable to use a mirror to allow the patient to point out what they consider to be the problem area while gently pointing out facial asymmetry, as well. If the patient has grand visions, attempts should be made to reestablish realistic expectations prior to any cosmetic procedures. Providing sufficient pain management should not be overlooked as patients will reflect upon the entire process when determining treatment success. At follow-up it is recommended to demonstrate muscle immobility with a mirror and to utilize before and after photos.

In addition to determining patient-perceived efficacy, the FLO assessment can be used to ascertain treatment goals for a patient. The FLO assessment may help physicians gain a better understanding of a patient’s objectives for treatment success compared to general global assessments. The SPA can be used to show patients the value of treatment. Patient satisfaction appears to improve when multiple facial areas are treated in conjunction since this may lead to a more natural look. Treatments with botulinum toxin type A need to be customized to each individual patient depending on physician assessment of the patient’s needs and the patient’s own treatment expectations.

Emerging science: Topical botulinum toxin and new injectables

There has been increasing discussion about topical preparations of botulinum toxin type A, but there is concern over adequate percutaneous penetration and drug delivery to the muscle since neurotoxin is a large molecule. Consequently, the appropriate vehicle will be crucial to the success of a topical version. A topical product would be a useful, painless option, especially for treating needle-phobic patients. It seems unlikely that there will be as great an effect with the topical compared to injection treatment though.

Azzalure® (Galderma, USA) is a topical product adapted from Dysport® that has been approved in several European countries. Azzalure® is indicated for the temporary improvement in the appearance of moderate to severe glabellar lines in adult patients under 65 years, when the severity of these lines has an important psychological impact on the patient. Revance Therapeutics (Newark, CA, USA) is currently conducting clinical trials with RT001, a botulinum toxin type A topical gel for the reduction of crow’s feet wrinkles. Phase I studies found RT001 to be safe and tolerable when applied to the forearms of 41 healthy subjects. Early Phase II studies found it was safe and effective for improvement in the appearance of lateral canthal lines. Adverse events experienced included mild skin erythema, ocular erythema, and ocular burning/stinging sensation. Currently, RT001 is in a US Phase II B clinical study. Chajchir et al conducted a small, single-center study comparing a topical botulinum toxin cream (CosmeTox) versus placebo cream on upper face wrinkles and found the topical neurotoxin to be effective in terms of subject-perception. In this study, the botulinum toxin type A was homogenized with a novel vehicle (InParT) which is thought to aid in appropriate penetration and delivery of the toxin to the muscle.

Also in pre-clinical trials, is the next generation botulinum toxin type A injection, RT002 (Revance Therapeutics). The manufacturer hypothesizes that the botulinum toxin type A molecule plus the patented TransMTS™ peptide technology...
could improve onset of action, increase duration of effect, and limit diffusion away from treatment site. Large, double-blind, randomized clinical trials need to be carried out to confirm safety and efficacy in each of these new products.

**Conclusion**

When properly used, botulinum toxin type A is effective for the improvement of glabellar rhytids and with little incidence of complications. There are multiple botulinum toxin type A products that are commercially available, although only two are FDA approved. Each product is distinct and, as mentioned above, they are not interchangeable. Hence, clinical research must be carried out for each specific product and conclusions from clinical trials cannot necessarily be applied to any other botulinum toxin type A product. There is still room for growth and research to be done within the discipline of aesthetic use of botulinum toxin type A.

**Disclosures**

The authors report no conflicts of interest in this work.

**References**