Advances in Facial Rejuvenation: Botulinum Toxin Type A, Hyaluronic Acid Dermal Fillers, and Combination Therapies—Consensus Recommendations

ARTICLE in PLASTIC AND RECONSTRUCTIVE SURGERY · JUNE 2008
Impact Factor: 2.99 · DOI: 10.1097/PRS.0b013e31816de8d0 · Source: PubMed
Advances in Facial Rejuvenation: Botulinum Toxin Type A, Hyaluronic Acid Dermal Fillers, and Combination Therapies—Consensus Recommendations

Jean D. A. Carruthers, M.D.
Richard G. Glogau, M.D.
Andrew Blitzer, M.D., D.D.S.,
and the Facial Aesthetics Consensus Group Faculty

Vancouver, Canada; San Francisco, Calif.; and New York, N.Y.

Background: Facial aesthetics and rejuvenation are evolving rapidly due to changes in products, procedures, and patient demographics. Clinicians can benefit from ongoing guidance on products, tailoring treatments to individual patients, treating multiple facial areas, and using combinations of products and ways to optimize outcomes.

Methods: A multidisciplinary group of aesthetic treatment experts convened to review the properties and uses of botulinum toxin type A (BoNTA) and hyaluronic acid fillers and to update consensus recommendations for facial rejuvenation using these two types of products. The group considered paradigm shifts in facial aesthetics; optimal techniques for using BoNTA and hyaluronic acid fillers alone and in combination; the influence of patient sex, ethnicity, cultural ideals, and skin color on treatment; general techniques; patient education and counseling; and emerging trends and needs in facial rejuvenation.

Results: The group provided specific recommendations by facial area, focusing on relaxing musculature, restoring volume, and contouring using BoNTA and hyaluronic acid fillers alone and in combination. For the upper face, BoNTA remains the cornerstone of treatment, with hyaluronic acid fillers used to augment results. These fillers are central to the midface because of the need to restore volume. BoNTA and hyaluronic acid in combination can improve outcomes in the lower face.

Conclusions: Optimal outcomes in facial aesthetics require in-depth knowledge of facial aging and anatomy, an appreciation that rejuvenation is a three-dimensional process involving muscle control, volume restoration, and contouring, and thorough knowledge of properties and techniques specific to each product in the armamentarium. (Plast. Reconstr. Surg. 121 (Suppl.): 5S, 2008.)

Substantial and rapid change typifies the field of facial aesthetics and rejuvenation as physicians and patients continue their quest for minimally invasive yet highly effective and safe approaches to mitigate the signs of facial aging.1,2 In 2004, the publication Consensus Recommendations on the Use of Botulinum Toxin Type A in Facial Aesthetics provided guidance on the use of the Botox formulation (Botox Cosmetic; Allergan, Inc., Irvine, Calif.) of botulinum toxin type A (BoNTA).3 Statistics for 2006, the latest available, show that BoNTA injection continues to be the single most common aesthetic procedure performed in men and women in the United States, but that its use is accompanied by growth in the use of other modalities, including dermal fillers.4

Disclosures: The faculty’s financial relationships or relationships to products or devices they or their spouse/life partner have with commercial interests related to the content of this continuing medical education activity are listed in an Appendix at the end of this article.
Nearly 3.2 million procedures were performed with BoNTA, and another 1.6 million procedures were performed with hyaluronic acid fillers, constituting about 81 percent of all procedures involving soft-tissue augmentation. Together, BoNTA and soft-tissue fillers comprise approximately 54 percent of all nonsurgical aesthetic procedures. With the fillers now available, such as the newer hyaluronic acid products, it can be anticipated that both demand and use will increase because the variety of options affords clinicians and patients greater versatility in tailoring therapy to individual needs and goals.

Changes have occurred not only in products but also in patient demographics, treatment goals, and the ways in which products are used. For example, in the year 2000, ethnic minorities accounted for approximately 15 percent of all minimally invasive procedures in the United States. In 2006, this had increased to approximately 22 percent. Men had 8 percent of all cosmetic procedures in 2006. Growth rates for BoNTA injection and use of hyaluronic acid fillers are higher in men than in women. Cultural and ethnic differences in aesthetic ideals and treatment goals are likely to have an increasing effect on how aesthetic medicine is practiced.

AN EVOLVING PARADIGM

One of the most important changes in facial rejuvenation has been a shift from a two-dimensional focus on hyperdynamic facial lines and immobilization of corresponding muscles to an increased comprehension and appreciation of the three-dimensional aspects of facial aging, particularly the loss of volume and its effect on treatment approaches. This change manifested itself initially in the idea of using BoNTA for facial shaping, but it has expanded to meld concepts of movement control, recontouring, and volume restoration using multiple modalities.

With changing aesthetics have come modifications in the way BoNTA is used in clinical practice. Although most clinical trials report results of BoNTA treatment on single areas of the face, clinicians tend to treat multiple areas to provide a more natural, relaxed look. Increasingly, too, BoNTA is used in combination with other modalities, including hyaluronic acid dermal fillers. By using fillers in combination with BoNTA, it becomes possible to address facial rejuvenation from a three-dimensional rather than a two-dimensional approach, thereby providing more pleasing, longer-lasting aesthetic outcomes.

CLINICAL IMPLICATIONS

The rapid changes occurring in aesthetic medicine necessitated updated clinical recommendations to guide the practice of more advanced techniques. Specifically, areas to be addressed included the following:

- Understanding the importance of how differences among products of the same class influence clinical use and outcomes
- Treating multiple areas in a single session
- Tailoring treatments to individual needs and circumstances, including ethnic and cultural considerations, such as skin type, facial shape, and aesthetic ideals
- Using combinations of products
- Ensuring patient satisfaction

On April 13 to 15, 2007, a multidisciplinary group of experts in aesthetic treatments convened to review the properties and clinical implications of various BoNTA and hyaluronic acid filler formulations and to develop updated consensus recommendations for facial rejuvenation with more advanced techniques using these two types of products. Specifically, these experts addressed the following topics:

- Optimal techniques for using BoNTA and hyaluronic acid fillers alone and in combination to treat multiple upper, middle, and lower facial rhytides, volume depletion, and folds
- Effect of sex, ethnicity, cultural ideals, and skin color on treatment
- General techniques and patient education/counseling
- Emerging trends and needs in facial rejuvenation

For purposes of discussion, the face was divided into thirds (upper, middle, and lower). The faculty stressed, however, the importance of evaluating the overall patient presentation and assessing the effect of change in one area on others. They concurred that the goals of aesthetic treatments are to help patients feel more attractive and as satisfied with their appearance as possible, by creating and maintaining a harmonious, balanced, and refreshed look. This continuing medical education supplement represents a compilation of their recommendations, continuing the process begun in 2004. The specific recommendations made in this document reflect the consensus of this group based on their clinical experience as well as published data regarding the use of BoNTA and hyaluronic acid fillers. It should be noted that other treatments, including non-
hyaluronic acid fillers, may be suitable for facial rejuvenation in individual patients. Recommendations for optimizing their use may become available in the future.

**BOTULINUM NEUROTOXINS AND HYALURONIC ACID FILLERS IN FACIAL AESTHETICS**

**Botulinum Neurotoxin Formulations**

Botulinum neurotoxins are produced by various strains of *Clostridium botulinum*, resulting in seven known serotypes, of which A and B have been developed for routine clinical use. Although basic mechanisms of action are the same for the various types of toxins, they differ both between and within types in a number of properties that have important implications for clinical use. Of the major toxins available for clinical use, only two formulations of BoNTA have been studied and used extensively for both cosmetic and therapeutic indications (Table 1). The type B formulation, commercially available as Myobloc (Solstice Neurosciences, Inc, Malvern, Pa.), is indicated in the United States for the treatment of cervical dystonia. It was deemed less suitable and practical for cosmetic use by the consensus faculty members because of discomfort on injection due to its acidity (pH approximately 5.6) and its shorter duration of effect, which has also been reported in the literature. Therefore, the remainder of this discussion focuses on the two major formulations of BoNTA. Only one (Botox) is approved for a cosmetic use (to treat glabellar lines) by the U.S. Food and Drug Administration; the other (Dysport, Reloxin; Ipsen Limited, Berkshire, United Kingdom) is undergoing Food and Drug Administration review and is indicated for aesthetic use in other countries.

According to clinical studies, product labeling, and clinical experience, these two formulations are not interchangeable and cannot be substituted one for the other by any simple or fixed-dose conversion ratio. Perhaps one of the most important clinical considerations is the probability of diffusion or migration outside the targeted muscle. From both preclinical and clinical studies, it appears that the Dysport/Reloxin formulation is more likely to diffuse farther from the injection site. In addition, the duration of effect may differ, with the Botox formulation providing a longer response at a dose ratio of 2.5:1 (Dysport:Botox).

The consensus faculty reported on their experience with the two formulations of BoNTA. All have used the Botox formulation, and 87 percent reported experience with Dysport/Reloxin in clinical studies. Approximately 50 percent also have used botulinum toxin type B (Myobloc). On the basis of their experience, most faculty members who have used both Botox and Dysport/Reloxin reported that injection volumes and doses differ, that injection patterns may need to be altered due to differences in migration, and that inexperienced injectors will likely have a steep learning curve to adjust injection techniques, because outcomes can differ substantially as a result of relatively minor differences in injection sites. They recommended that Dysport/Reloxin be used with caution around the eyes, upper lip, and depressor anguli oris and in the mentalis.

**Table 1. Comparisons between Formulations of Botulinum Toxin Type A**

<table>
<thead>
<tr>
<th></th>
<th>Botox (Vistabel), Allergan, Inc.</th>
<th>Reloxin (Dysport), Ipsen Limited</th>
</tr>
</thead>
<tbody>
<tr>
<td>FDA approval</td>
<td>Yes, cosmetic/therapeutic</td>
<td>Pending, cosmetic/therapeutic</td>
</tr>
<tr>
<td>Serotype</td>
<td>A</td>
<td>A</td>
</tr>
<tr>
<td>Complex molecular weight</td>
<td>900 kDa</td>
<td>500–900 kDa</td>
</tr>
<tr>
<td>Units per vial</td>
<td>100</td>
<td>500</td>
</tr>
<tr>
<td>Protein</td>
<td>~5 ng/vial</td>
<td>~5 ng/vial</td>
</tr>
<tr>
<td>Form</td>
<td>Vacuum-dried</td>
<td>Lyophilized</td>
</tr>
<tr>
<td>pH</td>
<td>~7.0</td>
<td>~7.0</td>
</tr>
<tr>
<td>Diffusion</td>
<td>Lower</td>
<td>Higher</td>
</tr>
<tr>
<td>Diluent (saline)</td>
<td>2.5 ml</td>
<td>5.0 to 7.5 ml</td>
</tr>
</tbody>
</table>

FDA. U.S. Food and Drug Administration.

**Hyaluronic Acids**

Naturally occurring hyaluronic acids are present universally in living organisms as part of the extracellular matrix. For many years, forms of hyaluronic acids have been used for intra-articular joint injections and in ophthalmologic procedures such as cataract surgery. More recently, they have become important as dermal fillers and contouring agents. Native hyaluronic acid has a short half-life of only about 1 to 2 days in tissue. The process of cross-linking hyaluronic acid results in larger, more stable molecules that have biocompatibility and viscoelastic properties similar to those of the naturally occurring substance. Cross-linking causes the ordinarily hygroscopic gel to become less water-soluble, thereby increasing product stability in tissue. Properties of hyaluronic acids that are important in determining their clinical per-
formance include the concentration of hyaluronic acid and the degree of cross-linking, which affect longevity and stability; gel hardness, which helps determine flow properties, the extrusion force required, and the structure and stiffness of the finished product; and the degree of gel swelling, or ability to resist dilution, which also influences longevity. 23,26,27

Commercially available hyaluronic acid fillers differ in their specific properties, thus contributing to differences in their clinical applications and performance (Table 2). 26,28–33 These differences allow clinicians to select the most appropriate agent for a given treatment need. For example, one important consideration is longevity of effect, and available dermal fillers offer a wide range of expected durations. 42 In addition, within the category of hyaluronic acid fillers, products differ in expected longevity, and recently, product labeling for the Juvederm fillers (Allergan, Inc., Irvine, Calif.) was revised to note that the treatment effects last up to 12 months. 33 In the clinical experience of the faculty, various fillers differ from each other in their duration of effect.

Some investigators have expressed the opinion that permanent fillers may lead to longer-lasting complications that are difficult to treat. 35 They have noted that such agents may also lead to an aesthetically unsatisfactory outcome that cannot easily be corrected. 34–36 Moreover, areas treated with such substances will not age in the same way as the rest of the face, with potentially unsatisfactory longer-term outcomes. 34–36 Clinical studies on polymethylmethacrylate and bovine collagen have, however, indicated a relatively low rate of late-occurring adverse events. 37 In the current trials reported, late adverse reactions occurred in 2.2 percent of the wrinkles treated, of which two reactions were considered severe in one patient.

On balance, the participants in this conference considered that, at this time, temporary fillers such as hyaluronic acids afford clinicians and patients a long-lasting outcome (Table 2) without the risks associated with permanent fillers and their use can be repeated as needed to maintain a satisfactory outcome and address evolving treatment needs. Hyaluronic acid fillers have an excellent safety profile. Many complications can be avoided by careful injection technique or reversed by injection of hyaluronidase. 38 For patients desiring a more permanent effect, hyaluronic acid fillers can be used initially to provide the patient with an opportunity to evaluate the effects. 35

In summary, the use of minimally invasive products such as BoNTA and hyaluronic acid dermal fillers has expanded significantly from their original applications to encompass mobility restriction, volumizing, and contouring alone and in combination to manage the multidimensional aspects of facial aging (Table 3). Meaningful differences exist among botulinum toxin formulations and hyaluronic acid filler preparations, and practitioners should be familiar with these differences to achieve optimal outcomes.

### Table 2. Comparisons among Selected Commercially Available Hyaluronic Acid Fillers

<table>
<thead>
<tr>
<th>Source</th>
<th>Juvéderm Ultra (24 HV)</th>
<th>Juvéderm Ultra Plus (30 HV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Source</td>
<td>Bacterial fermentation</td>
<td>Bacterial fermentation</td>
</tr>
<tr>
<td>Gelsizing process; product consistency</td>
<td>Sieving; granular</td>
<td>Homogenization; smooth</td>
</tr>
<tr>
<td>Cross-linking</td>
<td>BDDE</td>
<td>BDDE</td>
</tr>
<tr>
<td>HA concentration</td>
<td>20 mg/ml</td>
<td>24 mg/ml</td>
</tr>
<tr>
<td>Duration of effect (FDA label)</td>
<td>~6 months</td>
<td>~12 months</td>
</tr>
</tbody>
</table>

BDDE, 1,4-butanediol diglycidyl ether; HA, hyaluronic acid; FDA, U.S. Food and Drug Administration.

### Table 3. Evolving Aesthetic Uses of Botulinum Toxin Type A and Hyaluronic Acid Fillers

<table>
<thead>
<tr>
<th>Older treatment approaches</th>
</tr>
</thead>
<tbody>
<tr>
<td>Restricting movement to smooth lines (BoNTA)</td>
</tr>
<tr>
<td>● Glabellar lines*</td>
</tr>
<tr>
<td>● Horizontal forehead lines</td>
</tr>
<tr>
<td>● Crow’s feet</td>
</tr>
<tr>
<td>● Bunny lines</td>
</tr>
<tr>
<td>● “Chemical” brow lift</td>
</tr>
<tr>
<td>Filling lines and folds (HA)</td>
</tr>
<tr>
<td>● Nasolabial folds</td>
</tr>
<tr>
<td>● Lips (border and volume)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Newer treatment approaches</th>
</tr>
</thead>
<tbody>
<tr>
<td>Restricting movement, volumizing, contouring</td>
</tr>
<tr>
<td>● Treating multiple areas of the face</td>
</tr>
<tr>
<td>● Using combinations of BoNTA and HA fillers (e.g., brow lifting and shaping)</td>
</tr>
<tr>
<td>● Expanding treatment of the lower face with BoNTA, especially in combination with HA filler</td>
</tr>
<tr>
<td>● Shaping the jaw line with HA filler</td>
</tr>
<tr>
<td>● Using BoNTA for masseter reduction</td>
</tr>
<tr>
<td>● Periorbital volumizing with HA filler</td>
</tr>
<tr>
<td>● Restoring malar projection with HA filler</td>
</tr>
<tr>
<td>● Preventing lines with BoNTA</td>
</tr>
</tbody>
</table>

BoNTA, botulinum toxin type A; HA, hyaluronic acid.

*Botox (Allergan, Inc., Irvine, Calif.) is indicated for temporary improvement in the appearance of moderate to severe glabellar lines associated with corrugator and/or procerus muscle activity in adult patients 65 years of age and older (from Allergan, Inc. Botox Cosmetic [package insert]. Irvine, Calif.: Allergan, Inc., 2005).
FACIAL AGING AND REJUVENATION

General Principles

Facial aging progresses as the result of both intrinsic and extrinsic contributing factors. Over time, loss of subcutaneous fat, gravitational changes due to loss of elasticity, and remodeling of bony and cartilaginous structures lead to visible signs of aging. All portions of the face do not age simultaneously. For example, it has been observed, and confirmed in studies, that the subcutaneous fat of the face exists as separate and distinct anatomical compartments that may age differently with respect to each other. Similarly, the bony changes that occur may affect different parts of the facial skeleton differently. A recent study using computed tomography demonstrated that the skeleton of the midface below the inferior orbital rim undergoes significant changes while the anterior projection of the inferior orbital rim remains relatively fixed at its midpoint.

With age, repetitive muscle activity leads to fine lines that eventually become hyperdynamic wrinkles, which may deepen and ultimately remain apparent even when the muscles are at rest. Photodamage and smoking both accelerate these changes, with photoaging being the single most important contributor to the appearance of aging skin, including its texture and pigmentation. Due to the numerous factors contributing to the visible signs of aging, facial rejuvenation is likely to be best served by a multipronged approach.

Effects of Sex and Skin Color

In addition to environmental influences on aging, sex and skin color contribute to the manifestation of facial aging. Interestingly, although similar environmental factors affect perceived age among men and women, the strength of the association varies between the sexes. In a study of aging twins, for example, a history of smoking, sun exposure, and low body mass index were associated with higher perceived age in males. For females, significant determinants of high perceived age were low body mass index, sun exposure, smoking, and low social class.

In general, muscles of males tend to have a greater mass, which can affect the typical doses of BoNTA used in treatment. Other aspects of muscular anatomy, such as location and insertion point, may vary between individuals without being sex-specific, as indicated by a study of the anatomic position of the corrugator supercili in cadavers. When white males and females were compared within age groups, males tended to have consistently larger dimensions than females. The specifics of growth and aging of their soft tissue also differed, however, based on computerized three-dimensional mesh diagram analyses. It has also been pointed out that the skin of males and females differs substantially, particularly with regard to thickness. The effects of photoaging tend to be less pronounced in males than in females. Female skin is also generally more sensitive to irritants and allergens as well as to procedures and products used in their skin care. Particularly important to treatment planning are sex differences in idealized concepts of facial attractiveness (Table 4), but many of these descriptions are traditionally based on Caucasian ideals, which do not universally apply.

Ethnic groups differ in their facial characteristics with respect to both color and underlying structural or architectural differences. Some of the important and most obvious differences between skin of color and white skin are in melanosome type and size, distribution, and degree of melanization, which determines responses to ultraviolet radiation and thus, to some extent, the effect of photoaging. The darker the skin, the less vulnerable it is to ultraviolet damage. Skin of color also tends to be thicker and more sebaceous and to have more abundant collagen. Consequently, the signs of aging differ among persons of different skin colors and ethnic backgrounds. For

Table 4. Sex Differences in Facial Aesthetic Ideals

<table>
<thead>
<tr>
<th>The Idealized Female Face</th>
<th>The Idealized Male Face</th>
</tr>
</thead>
<tbody>
<tr>
<td>Large, smooth forehead</td>
<td>Overhanging, horizontal brow</td>
</tr>
<tr>
<td>Smaller nose</td>
<td>Larger nose</td>
</tr>
<tr>
<td>Arched or gull wing–shaped eyebrow</td>
<td>Minimal eyebrow arch</td>
</tr>
<tr>
<td>Wide-set eyes for a larger look</td>
<td>Deeper-set eyes, appearing closer together</td>
</tr>
<tr>
<td>Prominent cheekbones</td>
<td>Wider mouth</td>
</tr>
<tr>
<td>Heart-shaped taper in lower face</td>
<td>Squared lower face</td>
</tr>
<tr>
<td>Smaller lower-to-upper face ratio than in males</td>
<td>More equal ratio of lower-to-upper face proportions</td>
</tr>
<tr>
<td>Full, vermillion lips</td>
<td>Beard or coarser texture to lower facial skin</td>
</tr>
</tbody>
</table>

example, darker and thicker skin is not prone to developing the fine lines associated with aging lighter skin types.\textsuperscript{50–52} Both Asian skin and darker skin types have a higher probability of pigmentedary changes with aging as well as dyschromias in response to treatments.\textsuperscript{49,53}

Anthropometric studies conducted in various populations provide quantification of some of the differences among and within ethnic groups and demonstrate how far various populations diverge from Caucasian aesthetic ideals, which have traditionally driven much of facial plastic surgery.\textsuperscript{48} Faculty members discussed the substantial differences in profiles of Asians, African Americans, and Caucasians and their effect on treatment planning.

In Asians, for example, the lips project more than the nose and chin, whereas in Caucasians, the nose and chin project more than the lips. With aging, the lips of Asian patients tend to deflate and become flatter. The treatment goal with filler is to restore volume but not to cause excess protrusion, and faculty members recommended that practitioners separate these two effects conceptually. Asian patients tend to object to a bulging in the lower periorbital area and benefit from small amounts (2 U) of BoNTA just below the inferior ciliary line (approximately 1 to 2 mm). This provides a rounder, more youthful look to the eyes. Note that BoNTA should not be injected into the medial lower lid, because excess weakness in the medial orbicularis oculi may produce epiphora from decreased muscular squeeze toward the canaliculus. These injections may also result in morning edema of the lower eyelid due to decreased squeeze of fluid into the lymphatics.

Adding treatment of the crow’s feet and lateral brow can also help widen the appearance of the eye. In general, faculty members who treat Asian patients cautioned that BoNTA treatment should be conservative. They also noted that combination treatments with BoNTA and hyaluronic acid fillers may provide better results than monotherapy.

Facial aging in African Americans differs somewhat from that in Caucasians. For example, aging in the midface is prominent.\textsuperscript{34} Faculty members who treat African Americans noted that these patients exhibit a significant loss of volume in the midface with age and have heavier cheeks, a prominent tear trough, and fewer fine lines and wrinkles than Caucasians. Treating only the nasolabial folds in patients with this type of presentation can result in a very unnatural look; in contrast, malar augmentation provides more satisfactory outcomes.

Clinical studies have demonstrated that both BoNTA and hyaluronic acid fillers are effective and well tolerated in persons of color.\textsuperscript{55,56} For example, in women of skin types V or VI, BoNTA (20 U, $n = 15$; or 30 U, $n = 16$) safely reduced the severity of glabellar lines, with a high level of subject satisfaction (100 percent at day 30) that lasted for the 4-month duration of the study in 60 percent of subjects.\textsuperscript{55} In another study, the Juvederm formulation of hyaluronic acid provided longer-lasting improvement in the appearance of nasolabial folds in both Caucasians and persons of color compared with bovine collagen.\textsuperscript{56} No differences in efficacy based on skin type were observed throughout the 24-week duration of this study. No increased incidence of hyperpigmentation or hypertrophic scarring occurred in persons of color as compared with Caucasians, and there were no serious treatment-related adverse events. The incidence of adverse events did not differ by treatment or skin type. In clinical practice, faculty members recommend reducing the number of injections in persons of color to reduce the risk of hyperpigmentation.

As faculty members pointed out, treatment planning and goals must take each individual’s presentation into account to provide a harmonious, balanced outcome. Ethnic background and skin type add another dimension to the evaluation: sensitivity and consideration of cultural and ethnic aesthetic ideals and goals, such as maintaining ethnic identification while improving appearance. This is, however, in keeping with an essential guiding principle of aesthetic treatment: tailoring treatment realistically to specific individual goals and needs, which plays an important role in determining timing and use of specific products and ensuring high patient satisfaction.

**General Product Usage Considerations**

Virtually all faculty members stated that they use a variety of products depending on needs, area of the face, patient preferences or request, and product characteristics. With hyaluronic acid fillers, for example, the choice is very dependent on patient presentation. Faculty members generally prefer softer gels for more superficial use and more robust or stiffer products for deeper use, but said that the type of skin may play a role in dictates the choice of product. Faculty members noted that they view all fillers as falling along a continuum from softer to stiffer products. They also stressed the concepts of layering with different products and using enough product to provide
optimal outcomes. In their opinion, complications were in many instances a function of technique rather than solely a result of product characteristics. One important general way to reduce complications with filler injections is to slow the injection down and use finer-gauge needles (e.g., 30 g) that are longer than 1 inch in areas such as the infraorbital hollows. Faculty also recommended avoiding superficial use of stiffer products in areas such as the glabella, where they have the potential to compromise the capillary bed and lead to necrosis. In the experience of the faculty, these adjustments may help minimize swelling. Novice injectors need to understand that each product and syringe set has a different feel to it, that injection technique has to be adjusted for each product and treatment area, and that less may often be more.

**TREATING THE UPPER FACE WITH BoNTA AND HYALURONIC ACID FILLERS**

**Aging Considerations in the Upper Face and Areas of Interest**

Aging changes in the upper face reflect the effects of ultraviolet damage, gravitational changes, and muscles of facial expression on overlying skin. Although volume changes occur in parts of the upper face with age, they are not the most overtly apparent visible signs of aging in this region.

The muscles involved in upper facial expression and the central role of brow position have previously been reviewed. The hyperdynamic lines that form as a result of muscle activity are primarily perpendicular to the contraction direction of the muscles, but with aging, sleep creases also form on damaged skin. Loss of brow elevation is noticeable when the eyebrows descend below the level of the supraorbital ridge. Lateral brow ptosis also becomes apparent as temporal hooding. Overall, the appearance is one of fatigue and/or other negative expressions, such as anger, sadness, or disapproval, that are discordant with reality.

BoNTA has become the standard for nonsurgical upper facial rejuvenation. By reducing the motility of the muscles of facial expression, BoNTA significantly reduces the severity of upper facial rhytides and leads to a smoother, more youthful appearance. Another use of BoNTA in the upper face is to widen the eye. Injection of 2 U of BoNTA subdermally into the orbicularis oculi in the midpupillary line 3 mm below the ciliary margin increased palpebral aperture, especially when combined with injections into the lateral canthus. The authors noted that the outcome was most dramatic in Asian subjects. Moreover, by understanding the interplay of the elevator and depressor muscles of the upper face, clinicians can use BoNTA for managing asymmetries and repositioning, such as eyebrow lifting.

**Literature Update on the Upper Face**

Much of the literature on the use of BoNTA to treat the upper face has focused on line effacement in single facial areas, but in clinical practice, it is common to treat more than one area. One recent placebo-controlled, double-blind study examined the efficacy and safety of treating the forehead, glabellar area, and crow’s feet in a single session. In this study, female subjects received 64 U of BoNTA divided equally among 16 sites in the glabella (five sites), crow’s feet (three sites per side), and forehead (five sites). The Facial Line Outcomes questionnaire was the primary outcome measure on which subjects rated items of importance to their appearance based on their previous 7 days’ experience. Subjects also rated their perceived age of appearance using the single-item Self-Perception of Age. Facial Line Outcomes scores improved significantly after BoNTA but not placebo treatment, and were maintained through the duration of the double-blind period of 12 weeks. In addition, the number of subjects who reported looking younger than their current age increased from 35 percent at baseline to 75 percent at week 4; this was also maintained through week 12. Self-Perception of Age scores did not change with placebo treatment. During the study, no ptosis occurred and any adverse events were transient and mild to moderate in severity. This study confirmed widespread clinical experience that BoNTA can be used safely and effectively to treat multiple areas of the upper face in a single session.

Of increasing interest is the addition of hyaluronic acid filler to BoNTA to expand the options when treating the upper face. The results of a prospective, randomized study of BoNTA in combination with hyaluronic acid filler demonstrated that combination treatment of glabellar rhytides improved outcomes in comparison with hyaluronic acid filler alone. In this study, female subjects with deep resting glabellar rhytides received hyaluronic acid filler either alone (n = 19) or in combination with BoNTA (n = 19). BoNTA (30 U) was injected 1 week before hyaluronic acid filler was used in those receiving combination...
Combination treatment nearly doubled the median duration of response compared with treatment with filler alone. In addition to increasing durability of outcome, combination treatment also increased the percentage of subjects with aesthetic improvement at week 16 (95 percent versus 83 percent). At both rest (through week 32) and maximum attempted contraction (through week 16), responder rates were significantly greater with combination treatment. This study did not include a BoNTA-only arm, so the effect of BoNTA alone compared with combination treatment cannot be assessed from this study.

Consensus Recommendations

What Has Changed?

Many of the changes that have taken place in approaches to facial rejuvenation have come about because of increased understanding of facial aging, expansion in the numbers and types of products available, and associated refinements in aesthetic goals—specifically, providing a balanced, harmonious, and natural look.

Consensus recommendations on use of the Botox formulation of BoNTA alone have previously been published (Table 5). Changes from these original recommendations, based on this current update, are also shown in Table 5. Most importantly, 100 percent of the faculty agreed that they have decreased the number of units of BoNTA they used to treat the forehead.

Changes from Previous Recommendations

To treat horizontal forehead lines in female patients, 57 percent use 6 to 10 U and 43 percent use 11 to 15 U, compared with the typical dose of 15 U previously recommended. For male patients, 73 percent use 11 to 15 U, with the remainder of the faculty evenly divided between higher and lower doses. Previously, the typical range for male patients was 20 to 30 U. Interestingly, the faculty now use somewhat higher amounts of BoNTA to treat crow’s feet in men.

Current Practice in the Upper Face: Staging of Treatment

For the glabellar area, BoNTA is used alone about 75 percent of the time; when it is used in combination with fillers, 73 percent of the faculty noted that they stage the treatments for new patients, treating first with BoNTA and then with filler. This allows for the assessment of residual static lines/folds after BoNTA has taken effect. A minority of the faculty sometimes treat with both BoNTA and filler during a single treatment session. Approximately 43 percent of the faculty split the treatment of the glabella and frontalis into two visits when treating a patient for the first time.

Combination versus Single Treatment

Presently, all of the faculty use BoNTA alone to shape the brow about 50 percent of the time, but more than 90 percent of the faculty also use filler in some patients to replace volume in the brow area. Combination treatment of BoNTA and filler is used in the brow area about 20 percent of the time. Approximately 67 percent of the faculty use filler alone to fill the expression folds of the lateral frontalis immediately above the lateral upper brow of some patients. All faculty members continue to treat crow’s feet with BoNTA, but 80 percent now use it in combination with filler for some patients as part of treatments to enhance malar volume (see Treating the Midface with

Table 5. Consensus Recommendations on the Use of Botulinum Toxin Type A* in the Upper Face†

<table>
<thead>
<tr>
<th>Region/Target Muscle(s)</th>
<th>Usual No. of Injection Points (range); Unchanged from Original Recommendations</th>
<th>Total Starting Dose: Usual Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glabellar complex (procerus, depressor supercilii, orbicularis oculi)</td>
<td>5 to 7; men may require more sites</td>
<td>Women: 20 to 30 U; men: 30 to 40 U</td>
</tr>
<tr>
<td>Horizontal forehead lines</td>
<td>4 to 8; more or fewer may be required based on anatomic and aesthetic evaluations</td>
<td>Women: 15 U, 10 to 20 U; men: 20 to 30 U</td>
</tr>
<tr>
<td>Crow’s feet (lateral portions of the lateral orbicularis)</td>
<td>2 to 5 per side (higher in selected patients)</td>
<td>12 to 30 U; men: 20 to 30 U</td>
</tr>
</tbody>
</table>

*BoNTA injection sites and volumes are for the Botox formulation.
BoNTA and Hyaluronic Acid Fillers). In treating the upper face, faculty members agreed that restoring volume in the temporal region is an emerging treatment area. Faculty reported using a variety of different fillers in this area.

**Order of Treatment When Treating Multiple Upper Facial Areas**

When treating multiple areas of the upper face, 47 percent of the faculty treat the glabellar area first, followed by the forehead and then the crow’s feet. Another 47 percent of the faculty treat crow’s feet first, followed by the glabellar lines and then the forehead. All faculty noted that the most important consideration of the order is to track the treatments that have been performed in terms of injection location and units.

**Using Hyaluronic Acid Fillers in the Upper Face**

Most participants in the consensus meeting reported using more than one product, depending on the specific need. Use of hyaluronic acid fillers was divided evenly among Restylane, Juvederm Ultra, and Juvederm Ultra Plus. The usual amount of hyaluronic acid filler varied by site and ranged from 0.1 ml to 1.0 ml. About 50 percent of the faculty use 0.5 ml or less per site. Specific recommendations for hyaluronic acid filler use in areas of the upper face are listed in Table 6. The faculty noted that loss of volume in the temple area lateral to the tail of the eyebrow is associated with a drop in the tail of the brow. By using filler in the temple area and under the lateral brow and a small amount of BoNTA in the tail of the brow, a very attractive effect can be obtained. Patients may be unaware of the potential for treating this area with combination treatment and may need to be educated about it.

**Avoiding and Managing Complications in the Upper Face**

The most common complications with BoNTA are an overtreated frontalis, dropped brow, asymmetry, and bruising, particularly in the lateral canthus. Understanding the anatomy and evaluating each patient under dynamic and resting conditions can help avoid these complications. Patients with a low brow or evidence of pre-existing ptosis should be evaluated carefully before treatment. Eyebrow ptosis can generally be avoided by injecting no closer than 1 cm above the bony orbital rim in the midpupillary line and using lower doses in the frontalis. Faculty members recommended decreasing the amount of BoNTA used in treating older patients to avoid brow ptosis when trying to elevate the forehead. Many patients may require education about the need for striking a balance between reducing the appearance of forehead lines and the risk of brow ptosis. All patients should be counseled not to manipulate the treated area for several hours after treatment. In injecting the medial aspect of frontalis fibers with BoNTA, without treating the lateral frontalis fibers, can result in a cocked eyebrow in some patients. This can be corrected by injecting 1 to 3 U of BoNTA into untreated fibers that are causing the upward pull. Note also that these recommendations are only for the Botox formulation of BoNTA. Various lines of research, as well as the clinical experience of the faculty, indicate that another formulation of BoNTA (Dysport) exhibits a greater tendency to diffuse from the site of injection, which would be of clinical relevance in the small muscles of the face.

Conference participants agreed that using hyaluronic acid or any other dermal filler in the upper face, particularly the glabella, is an advanced technique that should be undertaken only by very experienced injectors. Although rare, the most serious potential complication is necrosis, which can occur most commonly when any filler is injected intravascularly (Table 7). Injectors should have thorough knowledge of the anatomy and blood supply of the area and must be alert to blanching or sud-

<table>
<thead>
<tr>
<th>Treated Area</th>
<th>Filler Volume</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glabella</td>
<td>&lt;0.25 ml in most patients; up to 0.5 ml</td>
<td>Stay superficial to mid-dermis, and aspirate if necessary; avoid compressing vessels</td>
</tr>
<tr>
<td>Forehead</td>
<td>Depends on length of rhytide and degree of compensation; dictated by product to be used</td>
<td>Use softer products and avoid hard products that bead</td>
</tr>
<tr>
<td>Crow’s feet</td>
<td>0.25 ml</td>
<td>Avoid stiff fillers, and proceed gradually and with caution; place hyaluronic acid fillers deeper than soft collagen</td>
</tr>
<tr>
<td>Bunny lines</td>
<td>~0.10 ml</td>
<td>Use soft, malleable product</td>
</tr>
</tbody>
</table>
den pain, which can indicate blood vessel occlusion or compression.

Summary Recommendations and Key Insights for Treating the Upper Face

BoNTA remains the main treatment option for the upper face, but dermal fillers such as hyaluronic acid are being added to the armamentarium to augment the results achieved by BoNTA alone. Also, doses of BoNTA, particularly in the forehead, are definitely lower than previously recommended to maintain some muscle movement, to minimize adverse outcomes such as brow ptosis, and to maintain a more harmonious aesthetic appearance. Doses of BoNTA greater than 20 U in the frontalis are more likely to lead to suboptimal outcomes. All faculty agreed that a frozen look is not a desirable goal, even though some patients expect to have a high level of immobility. In those cases, patient education is very important, to establish realistic and aesthetically pleasing expectations.

Combination treatment may increase the longevity of the outcomes.65 The consensus of the faculty was that BoNTA treatment should generally be performed first to assess the need for treatment of residual issues, such as static lines and deep folds that can be treated with hyaluronic acid fillers. It is now recognized that shaping of the brow region is a three-dimensional challenge that is well suited to the use of both BoNTA and hyaluronic acid fillers used in the superior orbital rim, especially in the lateral brow. An emerging use for fillers is to address the aging-related loss of volume in the temples. In all cases, it is important to consider patients’ sex and ethnic differences and to keep balance in mind when treating the upper face (e.g., relaxing the frontalis versus the degree of brow ptosis or change in the shape or arc of the brow). Some of these techniques, particularly the use of dermal fillers, should be reserved for use by experienced injectors because of the risk of blood vessel occlusion or compression.

TREATING THE MIDFACE WITH BoNTA AND HYALURONIC ACID FILLERS

Aging Considerations in the Midface and Areas of Interest

An important contributor to aging changes in the midface is the remodeling of underlying cartilaginous and bony structures that occurs, in addition to photodamage and loss of subcutaneous tissue and cutaneous elasticity.40,41 Changes in the structure of and relationships between the inferior orbital rim and the anterior maxilla markedly influence facial appearance, including the appearance of the eyes.40 The key to rejuvenating the midfacial area is volume restoration through re-inflation and recontouring. The specific areas of interest are the malar smile lines, malar projection, infraorbital hollow (arcus marginalis), nasojugal fold (tear trough, which is the medial arcus marginalis), nasolabial folds, and nasal dorsum and tip (Fig. 1). Each of these areas is considered below.

Literature Update on the Midface

Clinical experience using BoNTA and/or dermal fillers to treat various midfacial areas has been reported and reviewed.1,8,71 Several clinical studies and case reports document techniques and outcomes for treating midfacial sites with hyaluronic acid filler. For example, a technique of nonsurgical lower eyelid lift performed by filling the hollows with hyaluronic acid has been described, with the total filler volume reported as ranging from 0.35 ml to 1.40 ml per side, injected in small amounts.72 These investigators cautioned injectors to avoid the infraorbital nerve and periorbital arteries.

In another study, 24 female patients received hyaluronic acid filler to treat tear troughs.73 The filler volume ranged from 0.1 to 0.45 ml per eyelid,
with most patients requiring 0.2 to 0.3 ml. The deepest portion of the medial tear trough was treated first, and multiple passes with small amounts of hyaluronic acid filler were used. At the 10-day follow-up, 22 of the patients reported satisfaction with their appearance.

A larger, retrospective analysis reported on 121 patients who had received hyaluronic acid treatments for periorbital hollows that included the orbital rim hollow, zygomatic hollow, septal confluence hollow, and cheek pad (as well as the eyebrow area). The mean filler volume used to treat the orbital complex was 0.9 ml, and reinjection for maintenance was typically at about 6.5 months. Patient satisfaction was generally high: 86 percent were satisfied after the first injection, 91 percent after the second, and 100 percent after the third. Side effects were mild and transient and included color change, fluid build-up, lumpy or irregular contours, and bruising.

The basis for approving hyaluronic acid fillers for dermatologic use was their utility in treating nasolabial folds. In a multicenter, randomized, comparative study of 137 subjects, the Restylane formulation of hyaluronic acid provided a more durable response at 6 months than did bovine collagen, based on both physician and patient ratings. Similarly, a multicenter, randomized study of Juvederm fillers compared with bovine collagen in 439 subjects demonstrated that Juvederm hyaluronic acid fillers provided a significantly longer duration of effect than did collagen. Subjects also significantly preferred the Juvederm treatments to the collagen treatments. Recently, approval was granted by the U.S. Food and Drug Administration to amend the labeling for Juvederm Ultra and Juvederm Ultra Plus to include the information that the results of treatment with these products have been shown to last to up to 1 year.

Consensus Recommendations

General Principles

All faculty members agreed that one of the most important treatment considerations for the midface is the degree of volume depletion and flattening that occurs in the malar area. Accordingly, 60 percent of the faculty use fillers first followed by BoNTA treatment when a combination approach is indicated, to assess residual volume deficits, lines, and folds. They noted that not all areas of the midface are amenable to combination treatment.

Faculty members agreed that the malar contour should be restored first, followed by the orbital-malar groove and the nasojugal fold, which they noted to be the most complicated. When malar contour is restored, it results in a lift of the face and diminution of the nasolabial folds, affecting how the folds are to be treated. This approach, moreover, sometimes negates the need to treat the nasojugal fold/tear trough, which is a more complicated procedure. Thus, this conservative approach is a very useful way to begin treatment of the midfacial area.

Regarding injection technique, 56 percent reported using an anterograde technique. It is common to use the anterograde technique along the vermilion border, where the material will run along the potential tissue space with little or no additional movement of the needle tip. This can help verify the proper depth of placement for the needle tip. Other advantages of an anterograde approach is that it may yield a softer forward move-
ment through tissues, blunting the impact of the sharp needle tip, pushing vessels out of the way of the advancing needle, and reducing the probability of bruising. It is also possible to inject a small amount of hyaluronic acid filler moving forward to blunt the dissection and to inject additional filler as the needle is withdrawn if a greater volume of correction is needed. Those who prefer the retrograde injection technique believe that injecting slowly as the needle is withdrawn helps avoid intravascular injection of filler. This technique is often chosen for very soft, thin, or vascular areas, such as below the eyes or the malar region. In these areas, it is especially important to avoid tissue trauma; with a fine needle, it is thought that when the material flows as the needle is withdrawn, additional tracks or dissection planes are not created. This may be preferred when injecting areas with named vessels. Regardless of the technique used, it is essential to inject slowly (<0.3 ml per minute) and gently to avoid tissue tears.

Specific Recommendations by Treatment Area

Although it is a key principle to tailor treatment to each patient’s specific presentation, recommendations were undertaken for a typical or average patient, a Caucasian woman approximately 45 years of age (Fig. 2). Filler volumes and treatment recommendations are listed in Table 8. Note that greater or lesser volumes may be suitable for individual patients, depending on their presentation. The recommendations in Table 8 are generally conservative. For treating small malar smile lines, all of the faculty members use hyaluronic acid filler, but none uses BoNTA alone because of the risk of a stroke-like appearance and/or an inability to smile. Fewer than 20 percent use combination treatment in this area. To restore malar projection, faculty use a number of products: 100 percent use Restylane or Juvederm Ultra Plus in some of their patients; another 25 percent use Juvederm Ultra for some patients, depending on needs and goals. For this use, filler may be injected submuscularly as well as subdermally, particularly when using Perlane or Juvederm Ultra Plus.

In treating the nasojugal groove, 67 percent of the faculty use Juvederm Ultra and the remainder use Restylane. For nasolabial folds, treatment of the infraorbital hollow should be undertaken only by experienced injectors, with precautions to avoid injecting deep to the orbital septum. Although not a major application of fillers, a variety of products are also used to contour the nasal dorsum and tip due to surgical depression or atrophic changes caused by aging: 44 percent of the faculty use Restylane, 25 percent use Juvederm Ultra, 19 percent use Juvederm Ultra Plus, and 50 percent also use fillers other than hyaluronic acid, such as calcium hydroxylapatite. Fewer than 10 percent use collagen in this area. Most of the faculty elevate the nose saddle, recontour the nasal tip, and inject filler at the base of the columella to lift the nasal tip as a whole. Asian patients may

![Fig. 2. Commonly treated areas of the midface. Injection sites and patterning for hyaluronic acid fillers are shown.](image-url)
benefit from small amounts of filler in the bridge of the nose, which may also soften the appearance of the epicanthal folds. Some faculty elevate the nasal tip by injecting 3 U of BoNTA and 5 U each into the lower nasalis.

Avoiding and Managing Complications in the Midface

General precautions are the same as for other areas of the face, such as using slow, gentle injections to prevent tissue damage while maintaining steady pressure on the plunger when advancing.

Precautions for treating the midface include the following:

● Avoid using BoNTA in the malar smile lines, to avoid causing a transient stroke-like, frozen appearance or an inability to smile
● Start conservatively with filler to avoid overcorrecting; touch-ups can be performed at follow-up visits in 2 to 4 weeks
● To achieve an optimal aesthetic and avoid clumping, massage filler to distribute it and to permit evaluation of the effect before treating other areas
● Do not inject deeply into the nasal septum
● Do not overfill the nasal area; treat conservatively and re-evaluate and re-treat if necessary

Summary Recommendations and Key Insights for Treating the Midface

The main focus of treatment of the midface is on volume restoration, which dictates that filler use predominate and that BoNTA, if used, play a more limited role. One of the potential first steps is to restore malar projection or volume, which may obviate or modify treatment needs in other areas, such as the tear trough and nasolabial folds. After this, it is important to re-evaluate the remainder of the face and then treat. In treating the midface, it is important not to stint on filler volume, which may require educating patients to understand that undertreatment will yield suboptimal results.

Faculty members suggested marking each side of the face before injecting, which may help treatment of the second side if and when the first swells. Another option is to have the pretreatment photograph(s) available for comparison once the first side of the face undergoes augmentation. It was also suggested that clinicians use half the syringe, and then walk around to the other side of the patient to inject the other side of the face.

TREATING THE LOWER FACE WITH BoNTA AND HYALURONIC ACID FILLERS

Aging Considerations in the Lower Face and Areas of Interest

Aging changes due to photodamage, loss of subcutaneous fat, gravitational changes, and remodeling of the underlying bony and cartilaginous structures are very visible in the lips and surrounding tissues (Fig. 3). The appearance of the lower face is also modified by aging changes in the midface. In addition, dentition changes, resorption of mandibular and maxillary bone, forward rotation and protrusion of the chin, and other aging changes alter the proportions of the lower third of the face relative to the upper two thirds. Gravitational changes result in sagging skin, jowls, and wattles as well. Repeated muscular

---

### Table 8. Recommendations for Treating the Midface with Botulinum Toxin Type A* and Hyaluronic Acid Fillers

<table>
<thead>
<tr>
<th>Treated Area</th>
<th>Filler Volume (% of faculty)</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Malar smile lines</td>
<td>0.2 to 0.4 ml/side</td>
<td>Use deep dermal or subdermal injections and treat conservatively; may combine with resurfacing approaches or very small amounts of BoNTA; layering with different products can be used to address the etched-in superficial lines</td>
</tr>
<tr>
<td>Infraorbital hollow (arcus marginalis)</td>
<td>0.2 to 0.5 ml/side</td>
<td>Undercorrect in this area; do not inject deep to the orbital septum; this should be attempted only by experienced injectors</td>
</tr>
<tr>
<td>Nasojugal fold (tear trough)</td>
<td>0.1 to 0.2 ml/side (80%), 0.25 to 0.4 ml/side (20%)</td>
<td>Treat conservatively and re-evaluate need for additional treatment in 2 to 4 weeks; this should be attempted only by experienced injectors</td>
</tr>
<tr>
<td>Nasolabial folds</td>
<td>0.5 to 1.0 ml/side (81%), 1.1 to 1.5 ml/side (19%)</td>
<td>Visualize the nasolabial folds as a triangular area to be filled rather than as a linear area; hyaluronic acid fillers can be layered with collagen or with each other; massage to distribute and contour fillers and reduce beading/clumping</td>
</tr>
<tr>
<td>Nasal dorsum and tip</td>
<td>≤0.5 ml</td>
<td>Can be used after rhinoplasty for refinement or to treat drooping that occurs with aging; BoNTA has been used to treat this area</td>
</tr>
</tbody>
</table>

*BoNTA injection sites and volumes are for the Botox formulation.
contractions lead to furrows and lines that are superimposed on volume loss and exacerbated by loss of elasticity. Textural changes are also very apparent in the skin of the lower face, which may be amenable to various resurfacing modalities in combination with other treatments.

The orbicularis oris exerts a constant pull on the upper and lower lips that ultimately contributes to the formation of outwardly radiating perioral lines, sometimes called lipstick lines. Downward radiating lines, called marionette lines, form from the oral commissures due to the actions of the depressor anguli oris. The lips flatten and lose fullness, vertical height may increase, and the vermilion area may be reduced. As a result, the amount of dental display may be diminished and the Cupid’s bow may also flatten. Loss of definition occurs around the mandible, and the mentalis area develops a pebbled appearance called peau d’orange, which results from mentalis activity over time in conjunction with volume loss in the chin. Horizontal (necklace) lines form on the neck, and the platysmal bands become more prominent as elasticity is lost with aging.

In treating the lower face, the primary goals are to restore volume, reduce mobility, and resurface where appropriate. Faculty members consider combination treatment to be the standard for rejuvenating the lower face.

Consensus Recommendations

General Principles

The lips are the aesthetic focal point of the lower face. In treating the lips, 100 percent of the faculty use filler to shape the vermilion border and to volumize the lips. Approximately 50 percent each use collagen or hyaluronic acid filler routinely to enhance the vermilion border, and 100 percent use hyaluronic acid filler for volumizing the lips themselves.

For other areas of the lower face, including the depressor anguli oris, prejowl sulcus, mouth corners, and marionette lines, faculty members use BoNTA alone about 25 percent of the time, combination treatment about 50 percent of the time, and filler alone about 25 percent of the time. Some individual faculty members use combination treatment approximately 75 percent of the time. When combinations of BoNTA and filler are used, 87 percent of the participants treat with filler first. Treatments do not have to take place in a single session. One of the benefits of multiple sessions is the opportunity to reassess needs. All faculty members treat platysmal bands with BoNTA, and about 27 percent treat horizontal neck lines with BoNTA. Some practitioners also use hyaluronic acid fillers to manage the horizontal neck lines. They cautioned about the risk of hyperpigmentation when treating horizontal neck lines.

Fig. 3. Changes in the lower face due to aging.
lines in persons of color. As in other areas of the face, participants use a variety of different hyaluronic acid fillers to treat the lower face.

Specific Recommendations by Treatment Area

The use of BoNTA alone in treating areas of the lower face has previously been detailed. In contrast to previous recommendations, the faculty members have tended to reduce the number of BoNTA units for each area treated (Table 8). An important use of BoNTA in the lower face is to treat asymmetrical smiles. It was emphasized that skin landmarks can be used in conjunction with identifying musculature, but not in place of it. The depressor anguli oris can be palpated, and the medial aspects should be avoided. Injecting too high will affect the orbicularis oris and can worsen asymmetry. One tip is not to go higher than halfway between the lip and the mandible.

Participants reported less variability in their treatments than they did previously. In particular, note the substantially lower number of units used to treat platysmal bands so as not to weaken the muscles of the neck. Combination treatment is emerging as the standard for treating the lower face.

In using fillers to treat the lower face, faculty members tend to view the area as a whole rather than as isolated regions (Fig. 4). When treating the vermilion border with hyaluronic acid filler, approximately 80 percent of the faculty use Juvederm Ultra and the remaining use Restylane. For volumizing the lips, Juvederm Ultra, Juvederm Ultra Plus, and Restylane are preferred by 44 percent, 31 percent, and 25 percent of the faculty, respectively. For volume enhancement of the lips and shaping the vermilion border, 67 percent of the faculty use 1.0 ml of hyaluronic acid filler. The remaining 33 percent use 2.0 ml. The faculty cautioned about overfilling the lips and creating excessive protrusion (“duck lips”). It is important to think of shaping the lips, not simply adding volume. For example, when patients request an increase in the vertical height of the vermilion, it must be realized that filler is also going to increase the lips circumferentially. Rebuilding the philtral columns can be effective but must be done conservatively to avoid asymmetry.

For marionette lines (melomental folds) and the prejowl sulcus, 56 percent of the faculty use one syringe of hyaluronic acid filler and the remaining 44 percent use two syringes. Again it was stressed that using an adequate volume of filler is a key determinant of a successful outcome and patient satisfaction. Patients are apt to require education about the potential benefits and limitations of treatment, including the likelihood of suboptimal outcomes if an insufficient volume of filler is used.

Managing Complications in the Lower Face

The previous consensus conference reviewed potential complications with BoNTA in detail. In
the update meeting, faculty members reiterated the critical importance of understanding the anatomy in the lower face and using that, rather than skin landmarks, to direct injection sites. In treating the perioral area with BoNTA, the main precautions are to avoid overtreatment or injecting too close to the mouth, in order to prevent mouth incompetence, drooling, speech problems, and diminished proprioception (Table 9).³

Clinicians should begin conservatively and re-treat at a follow-up visit if necessary. In the first meeting, consensus panel members recommended that injections always be symmetrical and that the midline be avoided to prevent additional flattening. In the update meeting, it was noted that it is sometimes desirable to treat the lower lip area at the same time as the upper lip area to aid in proprioception. When treating the mentalis with BoNTA, it is important to avoid the depressor labii and consequent risk of lower lip depression. Again, injecting too close to the mouth may result in lower lip incompetence and drooling.³

Faculty members suggested that using higher dilutions of BoNTA coupled with more injection points could help avoid asymmetries of the upper lip, some of which result from unequal innervation by the depressor anguli oris. They also noted that injections of BoNTA into the platysma that are too high up on the neck can result in lip asymmetries. For hyaluronic acid fillers, careful injection planning to deliver equal amounts to each side of the face is important in avoiding asymmetries.

The platysma is a very superficial muscle, and injecting BoNTA too deeply in the platysmal bands may affect the strap muscles, causing dysphagia, or the cricothyroid muscle, leading to voice changes.⁷⁸,⁷⁹ It was noted that injecting the pars facialis just below the mandibular margin is safe, and that a danger zone is just at the cervicomental junction because of subjacent muscles involved in swallowing. Complications can be avoided by carefully marking the areas to be injected, injecting only the bands that can be grasped, and/or by using electromyographic guidance if necessary.⁸⁰ In some instances, the reversible, orally active anticholinesterase agent pyridostigmine may be used (60 mg) to counteract some of the effects of BoNTA (J. C. Carruthers, personal experience). This use of pyridostigmine, however, should be balanced against the risk of potential side effects, such as nausea, vomiting, diarrhea, and increased salivation, and pyridostigmine should not be considered for routine administration.⁸⁰ For hyaluronic acid fillers, careful injection planning to deliver equal amounts to each side of the face is important to avoid treatment-induced asymmetries.

Summary Recommendations and Key Insights for Treating the Lower Face

Full rejuvenation of the lower face involves the control of muscle movement as well as the restoration of volume. In many cases, neither BoNTA nor hyaluronic acid filler alone will provide optimal results. It is critical to know the musculature and its complex interactions in this area. Poor treatment planning and reliance on skin landmarks may lead to asymmetries and other suboptimal outcomes. Faculty members stressed that many patients complain about the appearance of

<table>
<thead>
<tr>
<th>Region/Target Muscle(s)</th>
<th>Usual No. of Injection Points (range)</th>
<th>Total Starting Dose: Usual Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Perioral area, orbicularis oris</td>
<td>2 to 6; to start: 4 sites, 1 site per lip quadrant</td>
<td>Original Recommendations: 4 to 10 U evenly divided among sites; Revised Recommendations: 4 to 5 U (75%); Note: treat lower lip in same session</td>
</tr>
<tr>
<td>Dimpled chin (peau d’orange), mentalis</td>
<td>1 to 2 (start with 1 midline or 2 symmetrical, lateral injections)</td>
<td>Women: 2 to 6 U; men: 2 to 8 U; Revised Recommendations: 4 to 5 U (56%); 6 to 10 U (44%)</td>
</tr>
<tr>
<td>Neck, platysmal bands, platysma</td>
<td>Women: 2 to 12 per band; men: 3 to 12 per band</td>
<td>Women: 10 to 30 U; men: 10 to 40 U; Revised Recommendations: 10 U/band (73%); 11 to 20 U/band (27%); 40 to 60 U total per neck per treatment session</td>
</tr>
</tbody>
</table>

their lips, and a substantial degree of education may be needed to help them understand the value of treating the entire lower facial region. In general, the perioral area should be managed conservatively, with follow-up visits scheduled to assess the need for additional treatment. The faculty cautioned against the use of high doses of BoNTA (>50 to 75 U per treatment session) in the platysmal bands because of experience with dysphagia and the need for treatment with a nasogastric tube.

OTHER ADVANCED TECHNIQUES AND EMERGING TRENDS

New and expanding uses for both BoNTA and hyaluronic acid filler continue to emerge rapidly. The faculty members discussed some of the most important techniques, based on their experience and the published literature. They also discussed the most prevalent trends in facial rejuvenation that they are observing in the literature and in clinical practice.

Acne and Other Scars

The faculty agreed that the use of either BoNTA or hyaluronic acid filler for acne scarring is still undergoing investigation. They pointed out that scars differ and that no one treatment is suitable for all. It was also noted that active inflammation is a contraindication to hyaluronic acid filler use and that BoNTA should not be used in infected sites. None of the participants use BoNTA alone to treat scars, whereas all have used hyaluronic acid filler alone. For long-standing, atrophic scars, small volumes of hyaluronic acid filler can be used to soften the appearance. Also, scar appearance may worsen with age, and hyaluronic acid fillers can be used to augment the volume of the area and thereby improve appearance. Repeated treatments may be necessary to provide optimal correction. All faculty members agreed that fractional resurfacing technology is an excellent approach to acne scarring.

Chin Implants

Hyaluronic acid fillers are useful in smoothing the appearance of chin implants, particularly in the transition area between the implant and soft tissue. In addition, the skin over the chin implant may dimple, which can be addressed by injecting approximately 8 to 10 U of BoNTA into the mentalis.

Horizontal Upper Lip Line

A variety of approaches were discussed, and it was agreed that more research into the underlying anatomical basis for this appearance is needed. Some faculty members recommended building up the philtral columns and resurfacing the skin. Others suggested that a small amount of BoNTA (e.g., 0.5 to 2 U) in the infracolumellar area could be effective, particularly in patients whose nasal tip droops when they smile. Some faculty members reported success with 1 U of BoNTA directly under the nose. The treatment is similar to that for nasal tip elevation but depends on the presentation. Another approach is to inject 0.5 to 1 U into each levator labii just lateral to the nasal ala on each side. Understanding and observing the muscle actions in this area are important in determining the treatment.

Risorius

The risorius muscle moves the lips laterally and is responsible for smiling or grimacing. It sometimes results in an unwanted or unintended negative expression if the muscle is hyperdynamic. One unit of BoNTA may reduce this activity if it is excessive. Electromyography-guided injection may increase the accuracy of delivering toxin to this muscle.

Excessive Gingival Display (Gummy Smile)

The use of BoNTA to treat excessive gingival display is a minimally invasive alternative to surgery. In this approach, small amounts of BoNTA are used to treat the muscles associated with raising the upper lip, thereby reducing their activity and lengthening the lip line. The elevator muscles of the upper lip include the levator labii superioris, levator labii superioris alaque nasi, levator anguli oris, zygomaticus major and minor, and depressor septi nasi. In this area, because of the interplay of muscles, conservative treatment is essential. One pilot study recommends 2.5 U per site at the levator labii superioris, 2.5 U per side at the levator labii superioris/zygomaticus minor, and 1.25 U in the orbicularis oris (2 to 3 mm inferior to the nostrils and 2 to 3 mm from the midline). Other recommendations are up to 5 U per site at the levator labii superioris alaque nasi just above the periorbital muscle, which can be identified by placing a fingertip on the pyriform aperture below the nasomaxillary groove. Overtreatment can result in lip ptosis, lower lip protrusion, the possibility of asymmetries, and excessive lengthening of the upper lip. The faculty members who
treat gummy smile tend to inject more conservatively, using about 1 to 2 U per side or 2 to 3 U in the depressor septae at the base of the columella.

Masseter Reduction

Techniques for using BoNTA to treat hypertrophic masseter muscles are well reported in the literature, are particularly evident in Korean-language publications (J. Wu, personal communication; C. Maas, personal communication, April of 2007), and are also reported in articles on treating temporomandibular dysfunctions. English-language publications document the efficacy of approximately 25 to 30 U of BoNTA per side for masseter reduction. It is important that BoNTA be used to treat facial widening only when the masseter is involved and not when mandibular bony prominence is the underlying cause. Faculty members who treat this area generally start with 10 U of BoNTA per side, but increase the dose to at least 20 U per side when needed. It was noted that it may take at least 6 months for optimal results to appear and up to a year for substantial recontouring to become apparent. Retreatments are at 3- to 4-month intervals. To perform the injection, have the patient bite down so that the anterior and posterior borders of the masseter can be felt. Injections should be low, just above the mandible, with one to two sites per side. By starting with 5 U per site, patients can be checked for side effects such as chewing difficulties and nighttime tooth grinding. Doses can be increased to as much as 25 to 50 U as needed and as tolerated.

Earlobe Volume

With age, the earlobes sag and develop creases. Hyaluronic acid fillers can be used to fill the sagging lobe and rejuvenate appearance. Women who have had repairs due to earrings pulling through and tearing the earlobe can benefit from the tissue-expanding effects of hyaluronic acid filler. In the faculty’s experience, the effects are very long-lasting, perhaps because that region is not very active metabolically and is not subject to movement.

Emerging Trends

The shift from a two-dimensional focus to a three-dimensional approach to minimally invasive facial rejuvenation has reinforced the idea of creating overall facial harmony and balance, within the confines of cultural, ethnic, and gender-related goals and ideals. With this approach comes an appreciation of the need for substantial volumes of hyaluronic acid filler to achieve optimal outcomes. This often entails substantial patient education, to ensure realistic expectations and to foster a commitment to longer-term maintenance with sufficient product to provide a high degree of satisfaction. In tandem, it is now recognized that relaxing the muscles of the lower face can play an important role in combination with fillers. In addition, BoNTA treatments, begun at earlier ages, can aid in line prevention. For example, facial lines were compared in a set of identical female twins. One twin had received BoNTA treatments in the upper face regularly for 13 years. The other twin had received only two treatments in the glabella and forehead. The minimally treated twin had visible lines at rest in the forehead and glabella, whereas the regularly treated twin did not. Differences were also apparent in the crow’s feet during smiling. No adverse events were associated with regular treatment. This comparison suggests that long-term treatment with BoNTA can be used safely to prevent the development of lines that are visible at rest.

Faculty members concurred that multimodal approaches to facial rejuvenation and enhancement help clinicians to provide the most satisfactory outcomes for their patients. Resurfacing with light/laser treatments or chemical peels, along with line management, volumizing, and recontouring, has proven to be both safe and effective. For example, the use of nonablative lasers, intense pulsed light, or radiofrequency when used immediately after BoNTA treatment does not reduce the efficacy of BoNTA treatment. Indeed, BoNTA plus laser treatment increases the extent and longevity of improvement compared with laser treatment alone. In addition, BoNTA plus intense pulsed light improves outcomes over intense pulsed light alone when the crow’s feet are treated. Similarly, radiofrequency, lasers, or intense pulsed light can be used safely and effectively in combination with hyaluronic acid filler to treat nasolabial folds and perioral rhytides.

Given their experience, the faculty cautioned that edema from other treatments, such as fractional laser therapy, could distort the anatomy, which could affect filler placement and potential migration of neurotoxin. Therefore, any multimodal treatment plans should take such effects into consideration.

Not to be neglected are topical treatments, such as cosmeceuticals, to protect against photodamage, aid in retexturing of the skin, and serve as an important adjunct to other aesthetic products and procedures. A variety of products are available, and many patients can benefit from...
their clinician’s advice on these products to minimize the risk of hypersensitivity reactions while maximizing benefit. Nearly 70 percent of the faculty members routinely discuss antioxidant supplements and topical treatments with their patients. An effective sunblock is considered one of the most important topical agents that patients can use. In addition to protecting against photoaging, sunblock may also help prevent hyperpigmentation after other aesthetic treatments. Despite general awareness of the need for sun protection, many patients can benefit from education about the types of ultraviolet radiation and how to protect against them most effectively and regularly.

Summary: Other Advanced Techniques and Emerging Trends

In discussing the use of BoNTA and hyaluronic acid fillers, the faculty members noted that both types of products can be used in a broad spectrum of applications. BoNTA, for example, can be used to advantage whenever restricting muscle movement, position, or size is likely to provide aesthetic benefits. These uses include masseter reduction, reduction of excessive gingival display, and prevention of line or scar formation. Hyaluronic acid fillers are advantageous overall for volume restoration and recontouring, including in such new uses as earlobe volume restoration and scar filling. Both BoNTA and hyaluronic acid fillers are integral to multimodal approaches to facial rejuvenation and enhancement that include light and laser techniques as well as topical treatments such as cosmeceuticals. Together, these approaches can help clinicians and patients achieve highly satisfactory outcomes.

GENERAL TECHNIQUES

Anesthesia

Because facial areas differ in their sensitivity to pain, the use of various forms of anesthesia depends on both the product and the area of injection. In the lower face and lips, all faculty members use intraoral blocks (e.g., lidocaine and articaine, with or without epinephrine). All also use ice and many use a Zimmer cooler. Topical anesthetics with agents such as lidocaine are also a useful adjunct. Massagers for vibration anesthesia are used by approximately 30 percent of the faculty. In the midface, none of the faculty members uses a block to treat the malar areas, whereas approximately 40 percent use blocks for treating the nasolabial folds. In the nasojugal groove, 85 percent use a topical anesthetic, 65 percent use ice, and various members of the faculty occasionally use injectable anesthesia, facial cooling, and massage. Only topical agents are used when the upper face is being treated.

Needles and Syringes

The needles and syringes used by faculty members depend on both individual technique/preference and specific application. Faculty members recommended 31-gauge needles with a short hub (30-U insulin syringe), 29-gauge, half-inch to 1-inch needles (0.30-ml syringe; BD Medical, Franklin Lakes, N.J.), tuberculin syringes with 30- or 32-gauge needles, and the hubless, 30/31-gauge system. All faculty members use half-inch needles in their practices, and approximately 30 percent use 1-inch to 1 ¼-inch needles for certain applications. Some faculty members use 32-gauge needles with a tuberculin syringe because their sharpness and thinness decrease injection pain.

Avoiding Complications and Posttreatment Recommendations

Injection-site reactions, such as erythema, edema, and ecchymosis, are the most common side effects of treatment. Before treatment, patients should be cautioned to avoid any agents that may inhibit clotting for several days before treatment (Table 10).

Table 10. General Recommendations to Avoid and Manage Complications

<table>
<thead>
<tr>
<th>Potential Complications</th>
<th>Management Approach</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hypersensitivity reactions</td>
<td>Rare; avoid in patients sensitive to any ingredient in the formulation</td>
</tr>
<tr>
<td>Injection-site reactions (e.g., erythema, edema, ecchymosis)</td>
<td>Patient should avoid nonsteroidal anti-inflammatory drugs, alcohol, and vitamin E for at least several days before treatment; injector should apply firm pressure after treatment; use ice packs</td>
</tr>
<tr>
<td>History of cold sores (herpes labialis, oral herpes simplex infection)</td>
<td>1 g of valacyclovir (orally) on the day of treatment; prescription for valacyclovir to be taken if needed</td>
</tr>
</tbody>
</table>

Copyright © American Society of Plastic Surgeons. Unauthorized reproduction of this article is prohibited.
members have their patients use ice while still in the office. Although it was previously recommended that patients contract the muscles treated with BoNTA to facilitate neurotoxin uptake, none of the faculty members recommended such an approach after combination treatment. All faculty members massage the treatment area after injections, but fewer than 50 percent have their patients massage the area at home.

None of the faculty members advises patients to avoid sun specifically after BoNTA treatment, although sun protection is always important; 28 percent permit their patients to exercise immediately after treatments; an additional 50 percent allow exercise the same day; and the remaining suggest waiting 1 day. Research is needed on the effects of exercise on both BoNTA and hyaluronic acid filler treatments.

Patient Expectations and Concerns

It is prudent to attempt to uncover the patient’s expectations, anxieties, or distress as well as potential misinformation that may derive from the experience of friends or from other sources, such as magazine articles. These discussions also provide an opportunity for the practitioner to identify patients who may not be suitable candidates for treatment and to foster a long-term plan with those who are.

Some practitioners suggested asking patients to bring photographs of themselves dating back 5 to 10 years to help guide the treatment plan and establish realistic goals. One of the most important recommendations is to photograph all areas of the face and not just those undergoing treatment. Having a number of views available can help in the accurate assessment of the face and can be useful to show to patients when they return with questions or issues with their treatment. Photographs can also be used to document outcomes.

Other Practice Management Considerations

All of the faculty noted that their staff members are an integral part of their practices and play an important role in office management, patient comfort and retention, follow-up, and identification of potentially problematic patients. Approximately 65 percent of the faculty employ a business manager to handle many of the office financial matters, including billing, discounts, and payment arrangements, leaving the physician to focus on the practice of medicine.

Patient retention is important, and more than 60 percent of the faculty schedule the next appointment before the patient leaves the office. The vast majority (87 percent) of the faculty and their staff also correspond with their patients by e-mail. Scheduling follow-up telephone calls is another way to improve retention, but in all of these approaches, it is important to maintain confidentiality. All faculty members take strong steps to ensure patient confidentiality and to prevent staff from divulging any patient information outside the office. Typically, any breach of patient confidentiality is grounds for immediate dismissal.

Patient Satisfaction

Most faculty associate patient retention with patient satisfaction. Only one faculty member surveys patients on a routine basis. If validated instruments such as Facial Line Outcomes and Self-Perception of Age are available for more areas of the face, they are likely to be used. Development of such instruments may help track and enhance patient satisfaction with various aesthetic procedures and with a physician’s practice.

OVERALL SUMMARY AND CONCLUSIONS

The number of minimally invasive aesthetic products and procedures has burgeoned in the last several years. In concert, clinicians have continued to expand and refine their techniques to provide their patients with optimal outcomes and a high level of satisfaction. Together, BoNTA and hyaluronic acid fillers are a mainstay of facial rejuvenation, accounting for approximately 54 percent of all nonsurgical aesthetic procedures performed in the United States. Use of these products is increasing among various demographic populations, including among persons of color and those with diverse ethnic backgrounds.

At the same time, concepts of aging and rejuvenation have changed from a somewhat two-dimensional focus to an appreciation of the three-dimensional aspects of aging, including the contribution of volume loss to appearance. A thorough knowledge of the process of facial aging and the anatomy and physiology of the facial musculature is paramount to treat patients successfully. With this understanding has come a paradigm shift from treating wrinkles and lines to relaxing the musculature, restoring volume, andcontouring to provide a balanced, harmonious aesthetic result. Although the various areas of the face lend themselves to different treatments, the overwhelming trend is toward increased combined use
of BoNTA and hyaluronic acid fillers as well as other modalities, such as resurfacing.

Notably, the combination of BoNTA and hyaluronic acid filler appears to increase the longevity of the outcomes, an issue of importance to patients. More than ever, patient education and counseling are essential when developing a comprehensive treatment plan, so that patients can make informed decisions about their treatment choices. Clinicians must also be knowledgeable about the strengths and limitations of various products and how they may be used optimally to individualize treatment for patient needs and goals.

It has also become apparent that, although it is necessary to discuss regions of the face individually, treating areas of the face in isolation does not yield the best possible outcomes for patients. Key insights and recommendations were provided by the faculty for each major region of the face as well as specific areas within those regions. It was stressed that treatment of any one area may have a considerable effect on other areas that should be evaluated as treatment progresses. In addition, the importance of understanding how to avoid or manage complications was addressed in detail. The faculty members emphasized that certain advanced techniques described in the supplement should not be undertaken by inexperienced injectors.

For the upper face, faculty agreed that BoNTA remains the foundation of treatment but that hyaluronic acid fillers can augment results in several ways, including the management of deep resting folds and lines that remain after BoNTA treatment and for the restoration of volume in the eyebrow area. An emerging use of hyaluronic acid fillers is to address the age-related loss of volume in the temple area. One of the BoNTA-specific changes in recommendations since the previous BoNTA consensus meeting was the agreement that lower doses of BoNTA, particularly in the forehead, may help avert a frozen look.

For the midface, the most apparent effect of aging is the loss of volume. Thus, hyaluronic acid fillers play a central role, with BoNTA treatment serving as an important adjunct depending on the specific treatment plan. A key step in facial rejuvenation of the midface is the restoration of volume in the malar region. By treating this area with hyaluronic acid fillers, clinicians can provide a more youthful, rounded area that will affect how other areas of the face, such as the tear troughs and nasolabial folds, are subsequently treated. Importantly, optimal results depend on using sufficient volume, and patients may need to be educated that undertreatment is likely to lead to an unsatisfactory outcome. In treating the midface, faculty members suggested taking any of a number of steps to avoid asymmetries. For example, mark both sides of the face, use half a syringe on one side and then move to the other, and have the pretreatment photograph(s) available for comparison during the procedure.

In the lower face, both BoNTA and hyaluronic acid fillers are important because rejuvenation involves control of muscle movement as well as restoration of volume. Detailed treatment planning is essential, to avoid asymmetries and poor outcomes. Knowledge of the musculature and its complex interactions is crucial. Treating the perioral area is central to the aesthetic outcome of the lower face, but treatment should be initiated conservatively, with follow-up visits for additional treatments. The differences between shape and volume are important in treatment planning and in the optimal use of products. Faculty members stressed that it is valuable to ensure that patients appreciate that not all treatments must be performed in a single session.

The faculty members also provided information and recommendations on other advanced techniques in the use of BoNTA and hyaluronic acid fillers and emerging trends in facial rejuvenation. These techniques include, but are not limited to, the use of BoNTA to recontour the mandible area and to reduce excessive gingival display; combination treatment with BoNTA and hyaluronic acid filler to refine the appearance of chin implants and improve acne and other scars; the use of hyaluronic acid filler to augment earlobe volume; and the use of both BoNTA and hyaluronic acid filler in combination with other modalities, such as laser and light treatments. Specific information on general techniques and practices, such as anesthesia, needles and syringes, avoiding complications, and posttreatment recommendations, is also detailed. Patient selection, education, and counseling, as well as other practice management issues, are considered.

In conclusion, the information presented in this supplement represents the insights and recommendations of medical experts in facial aesthetics on the use of BoNTA and hyaluronic acid fillers in facial rejuvenation. This supplement recognizes that facial aesthetics is a rapidly evolving field of medicine in which physicians and patients continue to have a major interest in the benefits of safe and effective, minimally invasive products and procedures that afford high levels of satisfaction to both clinicians and patients.
APPENDIX: THE FACIAL AESTHETICS CONSENSUS GROUP

The following faculty members comprise the Facial Aesthetics Consensus Group:

Richard L. Anderson, M.D.
Plastic Surgery
Salt Lake City, Utah

Joel L. Cohen, M.D.
Dermatology
Englewood, Colo.

Sue Ellen Cox, M.D.
Dermatology
Chapel Hill, N.C.

Steven H. Dayan, M.D.
Facial Plastic Surgery
Chicago, Ill.

Zoe Diana Draelos, M.D.
Dermatology
High Point, N.C.

Steven Fagien, M.D.
Ophthalmic Plastic Surgery
Boca Raton, Fla.

Ellen C. Gendler, M.D.
Dermatology
New York, N.Y.

Mitchel P. Goldman, M.D.
Dermatology
La Jolla, Calif.

Mary P. Lupo, M.D.
Dermatology
New Orleans, La.

Corey S. Maas, M.D.
Otolaryngology
San Francisco, Calif.

Susan C. Taylor, M.D.
Dermatology
New York, N.Y.

Jessica Wu, M.D.
Dermatology
Los Angeles, Calif.

Steven G. Yoelin, M.D.
Ophthalmology
Newport Beach, Calif.

APPENDIX: FACULTY DISCLOSURES

The Postgraduate Institute for Medicine assesses conflict of interest with its instructors, planners, managers, and other individuals who are in a position to control the content of continuing medical education activities. All relevant conflicts of interest that are identified are thoroughly vetted by the Institute for fair balance, scientific objectivity of studies utilized in this activity, and patient care recommendations. The Postgraduate Institute for Medicine is committed to providing its learners with high-quality continuing medical education activities and related materials that promote improvements or quality in health care and not a specific proprietary business interest of a commercial interest.

The faculty reported the following financial relationships or relationships to products or devices they or their spouse/life partner have with commercial interests related to the content of this CME activity:

Richard L. Anderson, M.D., states that he received consulting fees from Allergan, Inc., and BioForm Medical, Inc., and has received research study support from Allergan, Inc., Ipsen Limited, and Merz Pharmaceuticals. He is a consultant with Medicis Pharmaceutical Corporation.

Andrew Blitzer, M.D., D.D.S., states that he receives royalties from Medronic Xomed, Inc.; consulting fees from Allergan, Inc., and Solstice Neurosciences, Inc.; contracted research funds from Merz Pharmaceuticals; and an unrestricted research grant from Allergan, Inc.

Jean D. A. Carruthers, M.D., states that she and her spouse receive consulting fees from Allergan, Inc., Artes Medical Inc., BioForm Medical, Inc., Lumenis Ltd., Medicis Pharmaceutical Corporation, Merz Pharmaceuticals, Ortho Neutrogena, Solstice Neurosciences, Inc., and Unilever, Canada; and contracted research funds from Allergan, Inc., BioForm Medical Inc., Medicis Pharmaceutical Corporation, Merz Pharmaceuticals, Ortho Neutrogena, Q-Med, Inc., and Richard James Inc.; and have ownership interest in Artes Medical, Inc., and BioForm Medical, Inc.

Joel L. Cohen, M.D., states that he receives consulting fees from Allergan, Inc., BioForm Medical, Inc., Medicis Pharmaceutical Corporation; fees for non-Continuing Medical Education services received directly from a commercial interest or their agents from Allergan, Inc.; and contracted research funds from Allergan, Inc., BioForm Medical, Inc., and Medicis Pharmaceutical Corporation.

Sue Ellen Cox, M.D., states that she and her spouse receive consulting fees and contracted research funds from Allergan, Inc.

Steven H. Dayan, M.D., states that he receives consulting fees from BioForm Medical, Inc., and Medicis Pharmaceutical Corporation; fees for
non-Continuing Medical Education services received directly from a commercial interest or their agents from Aesthera Corp. and Laserscope; and contracted research funds from Aesthera Corp., Allergan, Inc., BioForm Medical, Inc., Contura International A/S, and Medicis Pharmaceutical Corporation.


**Steven Fagien, M.D.**, states that neither he nor his spouse has any financial relationships or relationships to products or devices with any commercial interests related to the content of this Continuing Medical Education activity.

**Ellen C. Gendler, M.D.**, states that she receives consulting fees from Allergan, Inc., and Medicis Pharmaceutical Corporation.


**Mitchel P. Goldman, M.D.**, states that he receives consulting fees from Allergan, Inc., Lumenis Ltd., Medicis Pharmaceutical Corporation, and Mentor Corporation; fees for non-Continuing Medical Education services received directly from a commercial interest or their agents from Allergan, Inc., Lumenis Ltd., Medicis Pharmaceutical Corporation, and Mentor Corporation; and contracted research funds from Allergan, Inc., Lumenis Ltd., Medicis Pharmaceutical Corporation, and Mentor Corporation; and has ownership interest in Lumenis Ltd. and CoolTouch, Inc.

**Mary P. Lupo, M.D.**, states that she receives consulting fees from Dermik Aesthetics and fees for non-Continuing Medical Education services received directly from a commercial interest or their agents from Allergan, Inc.

**Corey S. Maas, M.D.**, states that he receives research/consulting fees, advisory board honoraria, and/or stock interest from Allergan, Inc., BioForm Medical, Inc., Colbar Life Science Ltd., Elan Pharmaceuticals, Inc., Mentor Corporation; fees for non-Continuing Medical Education services received directly from a commercial interest or their agents from Allergan, Inc., Barier Therapeutics, Inc., Dermik Aesthetics, Doak Dermatologics, Galderma Laboratories, L.P., Johnson & Johnson, Medicis Pharmaceutical Corporation, Procter & Gamble, Stiefel Laboratories, Inc.; fees for non-Continuing Medical Education services received directly from a commercial interest or their agents from Allergan, Inc., Barier Therapeutics, Inc., Dermik Aesthetics, Doak Dermatologics, Galderma Laboratories, L.P., Johnson & Johnson, Medicis Pharmaceutical Corporation, Novartis Pharmaceuticals Corporation; and contracted research funds from Amgen, Inc., Atrix, Atrix/Fujisawa, Barrier Therapeutics, Inc., Beiersdorf AG, Doak Dermatologics, Galderma Laboratories, L.P., Genzyme Corporation, Johnson & Johnson, LEO Pharma A/S, Medicis Pharmaceutical Corporation, Palomar Medical Technologies, Stiefel Laboratories, Inc., and Teva Pharmaceutical Industries Ltd.

**Jessica Wu, M.D.**, states that she receives consulting fees from Allergan, Inc., Dermik Aesthetics, Johnson & Johnson, and Stiefel Laboratories, Inc., and contracted research funds from Allergan, Inc.

**Steven G. Yoelin, M.D.**, states that he receives consulting fees from Allergan, Inc., BioForm Medical, Inc., Sanofi-Aventis, and Valeant, Inc.; fees for non-Continuing Medical Education services from Allergan, Inc. and Medicis Pharmaceutical Corporation; and contracted research funds from Allergan, Inc.

The planners and managers reported the following financial relationships or relationships to products or devices they or their spouse/life partner have with commercial interests related to the content of this CME activity:
The PIM clinical content reviewer, Trace Hutchinson, Pharm.D., hereby states that neither he nor his spouse has any financial relationships or relationships to products or devices with any commercial interests related to the content of this Continuing Medical Education activity.

The HLS reviewers, Paula G. Davis, Ph.D., and Ramana Yalamanchili, Ph.D., state that neither they nor their spouses/life partners have any financial relationships or relationships to products or devices with any commercial interests related to the content of this Continuing Medical Education activity.

Jean D. A. Carruthers, M.D.
University of British Columbia
Suite 740-943 West Broadway
Vancouver, V5Z 4E1 British Columbia, Canada
drjean@carruthers.net

ACKNOWLEDGMENT
This activity is supported by an educational grant provided by Allergan, Inc.

REFERENCES


