Treatment of facial lipoatrophy, morphological asymmetry, or debilitating scars with the hyaluronic acid dermal filler Princess® FILLER

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Purpose: The aim of this study was to evaluate the effectiveness and safety of hyaluronic acid (HA) dermal filler when used in the face for medical reconstructive purposes.

Patients and methods: Adult patients with moderately severe facial lipoatrophy (FLA), morphological asymmetry (MA) of the face, or debilitating scars (DS) on the face were included in a prospective, noncomparative, multicenter, postmarket clinical follow-up study. All patients were treated with an HA filler (Princess® FILLER), which was injected intradermally on study Day 1, with optional touch-up 2 weeks later. The effectiveness of the treatment was evaluated at Weeks 4 and 24, using a six-grade scale ranging from “excellent” to “worsening”. The assessments were conducted by both the investigator and the patient and, at Week 4, by the independent photography reviewer as well. Adverse events were collected at each visit.

Results: Fifty-three patients were included in the study (FLA 23, MA 17, and DS 13), and 46 patients completed a 6-month follow-up (FLA 20, MA 15, and DS 11). At Week 4 (primary endpoint), the overall treatment success rate was 100% (FLA), 100% (MA), and 94% (DS), based on assessments made by the investigator, patients, and the independent reviewer, respectively. In most patients (~95%), the effect was sustained over 6 months. Treatment-related adverse events were reported in five patients (9%) and included injection site hematoma, injection site pain, and headache.

Conclusion: Dermal filling with HA gel is a viable treatment option for the correction of various deformaties of the face resulting from FLA, MA, or DS.

Keywords: clinical study, hyaluronic acid, dermal filler, Princess® FILLER, effectiveness, safety

Introduction

Soft tissue fillers are nowadays widely used in cosmetic dermatology and esthetic surgery for the facial rejuvenation or redefinition of facial contours. Although various fillers are available on the market, hyaluronic acid (HA) gels are most popular. In USA only, they were used in almost 2.5 million esthetic procedures in 2016, representing over 90% of procedures involving any soft tissue filler.1 Such wide-spread use of HA fillers is driven by their versatility, ease of administration, predictable effectiveness, safety track record established during decades of use, and nearly no downtime after the treatment.2,3

Esthetic procedures are typically undertaken to remove or reduce the signs of aging or to increase the attractiveness of an otherwise healthy face. There is, however, a range of pathological conditions, which may result in an “abnormal”, asymmetrical appearance of the face with consequent emotional suffering of the patient, psychosocial...
dysfunction, and reduced quality of life.2–6 Some of these, such as facial lipoatrophy (FLA) and atrophic skin scars, are characterized by reduced tissue volume, which can be restored by injecting a soft tissue filler.7,8 Similarly, in case of morphological asymmetry (MA) of the face, increasing the volume of visually hypoplastic areas with a filler is a simple and minimally invasive solution for harmonizing the appearance. With the exception of HA use for the correction of FLA in patients with HIV infection, which was studied in several clinical trials,9–13 literature data on the effectiveness and safety of HA fillers in the treatment of other facial pathologies are fairly limited. Furthermore, available HA fillers differ in physical properties and tissue residence time,14 parameters that may influence the overall performance.

The present study was undertaken to evaluate the effectiveness and safety of Princess® FILLER, a marketed HA dermal filler, when used in routine practice for medical reconstructive purposes in patients with FLA, MA, or DS. This filler is a low viscosity gel of cross-linked HA, designed for injection into the mid-to-deep dermis. The expected lifetime upon implantation is 6–9 months.

Patients and methods

This was a prospective, noncomparative, multicenter, postmarket clinical follow-up study of a European conformity-marked dermal filler Princess® FILLER (ClinicalTrials.gov, NCT03050723). The study consisted of four visits, performed at baseline and 2, 4, and 24 weeks thereafter. Up to 60 patients with FLA, MA, or DS were to be included to obtain 24-week follow-up data from 10 to 20 patients per indication.

The trial was conducted at three dermatology clinics in Austria, from December 2016 to July 2017, in line with the ISO 14155 standard and ethical principles of the Declaration of Helsinki. The protocol and relevant study documents were reviewed and approved by the ethics committee of the Medical University of Graz and the ethics committee of the city of Vienna. Written informed consent was obtained from each patient prior to inclusion in the study. Individual participant data that underline results reported in this article, after deidentification, are contained within this article.

Study population

Adult patients suffering from FLA of moderate severity (grades 2–3 according to the Ascher’s scale15 and unrelated to aging), MA of the face, or at least one DS on the face were included. The defect had to be correctable by a dermal filler in the investigator’s opinion, the skin on the face healthy (with the exception of visible atrophic acne scars), and the patient willing to take part in the study and abstain from any other esthetic or surgical procedures in the treatment area over the following 6 months.

Exclusion criteria comprised pregnancy, breastfeeding, or unwillingness to use contraception throughout the clinical investigation (for women of child-bearing potential only); previous allergic reaction or hypersensitivity to HA; a history or presence of hypertrophic scars, keloids, or pigment disorders; any autoimmune or connective tissue disease, or current treatment with immune therapy; a silicone implant or nonabsorbable tissue filler in the area targeted for intervention, or treatment with any HA filler in the previous 6 months; presence of infectious, inflammatory, or proliferative lesions in the area targeted for intervention, including a history of herpes zoster or recurrent herpes simplex; current treatment with anticoagulant or antiplatelet drugs, unless the risk of bleeding has been discharged; current participation in another clinical trial, or treatment with any investigational drug or device within 30 days prior to enrolment; and any other disease or condition, which in the investigator’s opinion represented a safety risk for participation in the study. Institutionalized persons with legally limited civil rights were excluded as well.

Intervention

All patients were treated with an HA dermal filler, containing 23 mg/mL of HA produced by Streptococcus equi bacteria and cross-linked using 1,4-butanediol diglycidyl ether (Princess® FILLER, Croma-Pharma GmbH, Leobendorf, Austria). Pretreatment with a topical anesthetic was optional.

At the baseline visit, the filler was injected into the mid to deep dermis, in amount needed to achieve a desired level of correction. A touch-up was allowed after 2 weeks and was performed at the investigator’s discretion. The total volume of the filler was limited to 10 mL per patient.

Outcome assessments

At Weeks 4 and 24, the effectiveness of the treatment was evaluated by the investigator and separately by the patient using a six-grade scale presented in Table 1. At Week 4, the effectiveness was also evaluated by an independent reviewer using patient photographs and the same scale. The primary effectiveness endpoint was the proportion of patients with successful treatment outcome at Week 4 based on the investigator’s assessment, where success was defined as an excellent, good, or moderate correction of the defect. Treatment success rates derived from other assessments described earlier, overall...
Table 1  Treatment outcome grading

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Description</th>
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<tbody>
<tr>
<td>Excellent</td>
<td>Significant improvement in patient’s appearance vs baseline, with complete or almost complete correction of the treated defect</td>
</tr>
<tr>
<td>Good</td>
<td>Significant improvement in patient’s appearance vs baseline, with achieved correction in line with expected result</td>
</tr>
<tr>
<td>Moderate</td>
<td>Obvious improvement in patient’s appearance vs baseline, with achieved correction slightly below the expectation</td>
</tr>
<tr>
<td>Slight</td>
<td>Minimal improvement in patient’s visual appearance vs baseline</td>
</tr>
<tr>
<td>No effect</td>
<td>No change in patient’s visual appearance vs baseline</td>
</tr>
<tr>
<td>Worsening</td>
<td>Deterioration of patient’s visual appearance vs baseline</td>
</tr>
</tbody>
</table>

Note: Treatment success is defined as an excellent, good, or moderate correction of the defect.

and stratified by indication, were secondary endpoints. The tolerability and safety were evaluated based on the occurrence of adverse events, which were collected at each visit.

Statistics

The sample size of 60 patients was arbitrarily selected, based on the anticipated feasibility of recruitment, and was sufficient to provide robust estimation of the primary endpoint. With a sample size of 60, a two-sided 95% CI for a single proportion using the large sample normal approximation extended 0.076 from the observed proportion for an expected proportion of 0.900. The primary population for data analysis was the safety population (SP), defined as all patients who received the investigational device and had at least one posttreatment assessment. The effectiveness was also analyzed in the per-protocol population (PP), defined as all patients who received the investigational device and have completed the study without a major protocol deviation. All data were analyzed using descriptive statistics. For the primary endpoint and other treatment success rates, percentages were calculated based on the number of patients treated (intent-to-treat approach with missing data accounted as nonsuccess) and 95% CIs were derived using the Pearson–Clopper interval. In addition, for individual grades of treatment outcome, percentages were calculated based on the number of observations.

Results

Patient baseline characteristics and disposition

A total of fifty-three patients were included in the study: 23 patients with FLA, 17 patients with MA, and 13 patients with DS. All patients were white (non-Hispanic or Latino), and all but one with DS were women. The mean age of the patients was 48.1 years (range 23–76 years) and was slightly higher in those with FLA (mean 48.2 [range 26–75] years) or asymmetry (51.7 [34–76] years) than those with scars (43.0 [23–75] years). Among patients with FLA, 11/23 (48%) patients suffered from grade 2 and 12/23 (52%) patients suffered from grade 3 according to Ascher’s scale (grades 1–5, with 5 being the most severe). In patients with MA, the severity of volume deficit on the left side, compared to the respective area on contralateral side, was mild in 4/17 (23%) cases, moderate in 11/17 (65%) cases, and severe in 2/17 (12%) cases; on the right side, it was mild in 8/17 (47%) cases and moderate in 9/17 (53%) cases. The severity of scars was not graded. In 11/13 (85%) patients, the scars were hypotrophic, and in 2/13 (15%) patients, the scars were hypertrophic; none was calcified.

In a total of nine (17%) patients, including 3/23 (13%) patients with FLA, 5/17 (29%) patients with asymmetry, and 1/13 (8%) patients with scars, the defect had previously been treated with an HA filler. In two of these patients, it could not be ruled out that the treatment had occurred within 6 months prior to screening due to an incomplete treatment date (month and year only).

Of the 53 patients treated, all attended Week 2 visit, 51 (96%) patients returned for Week 4 visit, and 46 (87%) patients completed a 24-week follow-up. One patient was withdrawn from the study because of pregnancy, two patients withdrew consent, and four patients were lost to follow-up after Week 4 visit. All patients who took part in the study were included in the SP, and 44 (83%) patients met all criteria for the PP. Major protocol deviations were related to missed visits due to premature termination, missed visit windows, and potential violation of the wash-out period for previous HA treatment as described earlier.

Treatment details

In most patients, the filler was applied bilaterally and into more than one area of the face (Table 2). At the initial application, 4.3 mL was injected on average, most commonly by serial puncture (in 30 patients; 57%), followed by linear threading (14; 26%), cross-hatching (5; 9%), and serial threading (4; 8%). Serial puncture was a leading injection technique across the indications, used in 11/23 (48%), 9/17
(53%), and 10/13 (77%) patients with FLA, asymmetry, and scars, respectively. A touch-up was performed in 37 (67%) patients, and a total of 6 mL of the filler was administered on average (Table 3). The treatment was generally performed without local anesthesia; anesthetic cream was applied only in one patient who suffered from FLA.

### Treatment effectiveness

In the SP, the treatment success rate at Week 4 was 96.2% (51/53; two-sided 95% CI 87.0–99.5); exactly the same result was obtained with both the investigator and the patient assessments. In the independent reviewer’s opinion, the treatment was successful in 91% of patients (48/53; 95% CI: 79.3–96.9). At Week 24, success rates were lower (83% by the investigator [44/53; 95% CI: 70.2–91.9] and 81% by patients' self-assessment [43/53; 95% CI: 68–90.6]), primarily because drop-outs were accounted as treatment failures. When success rates were calculated based on the number of actual observations, these reached 94%–100% at Week 4, depending on the evaluator, and 94%–96% at Week 24 (Figure 1). Treatment results are also illustrated in Figures 2–4, depending on the evaluator, and 94%–96% at Week 24 (Table 2–3). The treatment was generally performed 2–3 days after initial or touch-up treatment, respectively, and resolving within 3 weeks. Two additional patients had moderate injection site pain, one of them also reported a headache, and further one experienced headache only. These events occurred after touch-up and resolved within 2–4 days.

### Discussion

This study has shown that various soft tissue defects of the face can be corrected successfully with the Princess® FILLER, an HA dermal filler. Princess® FILLER was approved in Europe in 2008 for lip augmentation and the correction of moderate-to-severe facial wrinkles and folds. In 2015, the approval was extended to medical indications discussed here.

The three indications studied are diverse in terms of pathogenesis and clinical presentation but share a common denominator—a deficit in tissue volume. In FLA, the volume loss is due to the atrophy of facial adipose tissue and the disease is primarily manifested on cheeks, temples, and in the preauricular, orbital, and perioral regions. The underlying cause is known in some patients (eg, antiretroviral therapy in HIV-positive individuals, autoimmune connective tissue diseases, various lipodystrophy syndromes, and trauma).7,15 but in many, it remains unclear why slow physiological involution of facial fat pads takes a more aggressive course, resulting in pathological appearance. In contrast to FLA, volume deficit in depressed scars results from either inflammatory destruction of collagen fibers and subcutaneous fat, which typically occurs in atrophic acne scars,17 or physical loss of skin supporting structures during the insult, which preceded the scar. In MA of the face, volume deficit may be actual due to unilateral hypoplasia of various skin supporting structures,
or virtual due to hyperplasia on contralateral side. While the cause of asymmetry may be known (eg, congenital anomaly, genetic disorder, trauma, infection, neoplasm, and other diseases of the facial structures), in many cases, it remains obscure. It is speculated that factors such as habitual mastication on one side and constant facial pressure exclusively on one side during sleep, deleterious oral habits, or unilateral crossbite may trigger the asymmetric development of facial structures, resulting in disharmony.18,19

Obtained results indicate that pathological deficits in facial or dermal volume can be successfully restored by the tested dermal filler, regardless of the cause or location. In investigator’s opinion, the treatment was successful in all patients who attended Week 4 visit. Importantly, a complete or almost complete correction of the defect was achieved in ~60% of the patients, with no major difference between the indications. In a further 35% of patients, a moderate response was observed in line with the expected outcome. Similar results were obtained when evaluation was based on the patient’s self-assessment, corroborating the investigator’s ratings. High treatment success rate was also confirmed by the independent review of patient photographs. In general, the assessments made by the independent reviewer were slightly more conservative than those made by both the investigator and the patients. This may be partly attributed to genuine limitation of photo review compared to live assessments, since it provides a restricted insight from few preselected perspectives only. With respect to the durability of therapeutic effect, in most patients (~95%), the effect was maintained for at least 6 months. Notably, at 6 months after the intervention, the effect was still graded as excellent or good in around 75% of the patients.
The effectiveness observed in patients with FLA was comparable to the effectiveness of other HA fillers, tested in patients with HIV-related lipoatrophy. The scales used across the studies were slightly different, but in general, the esthetic improvement rates at 6 months ranged from 90% to 100%. The frequency of side effects in the current study was lower than stated in the literature. In the present study, treatment-related adverse events were reported in four patients with FLA (18%), compared to 29%–70% in other trials. Although none of our patients had a history of HIV, a higher incidence of side effects was more likely related to subcutaneous administration of the filler, which was driven by characteristics of the filler used. The average amount of the filler was similar across the studies, around 6 mL per patient, which is why this factor could be excluded as an explanation.

Regarding MA, most faces are asymmetric to a certain extent and the point at which “normal” asymmetry becomes “abnormal” cannot be easily defined. It often depends on the clinician’s sense of balance and, more importantly, the patient’s sense of imbalance. No details were collected on asymmetric anatomic landmarks, but in most patients, the filler was administered bilaterally, into the midface and one or both of remaining areas. Literature data on HA filler used for the correction of facial asymmetry are limited to several case reports and two series of seven to eight patients with upper eyelid asymmetry who were successfully treated with...
Hyaluronic acid filler for facial lipoatrophy, asymmetry, or scars

HA filler was also successfully used in combination with botulinum toxin to correct asymmetric peri orbital features (nine patients), or functional asymmetry caused by hemifacial spasm (18 patients).

The effectiveness and safety observed in patients with DS were consistent with data published by Hasson and Romero on a similar HA filler in 12 patients with atrophic scars of various origin. The effectiveness of HA in the treatment of scars was also conveyed in three small studies (5–12 patients each) on the HA treatment of atrophic acne scars by vertical filler technique, microinjection, or pneumatic injection, in a study in 15 patients with cicatricial ectropion, where HA was injected into the lower lid to stretch the tethered skin and correct the lid and several case reports.

Conclusion

Dermal filling with cross-linked HA is a viable treatment option for the correction of various soft tissue defects of the face resulting from FLA, MA, or depressed scars. The filler used in the present study, Princess® FILLER, was well-tolerated by patients, safe, and highly effective, with an overall treatment success rate of 94%–100%, depending on the evaluator. In most patients, the response was sustained over 6 months after the treatment.

Disclosure

Zrinka Ivezic-Schoenfeld, Ira G Federspiel, and Martin Prinz are employees of Croma-Pharma GmbH, which sponsored the present study. Daisy Kophera, Doris Grablowitz, and Benjamin Gehl served as investigators for Croma-Pharma GmbH. The authors report no other conflicts of interest in this work.

References