INTRODUCTION

Aesthetic procedures with hyaluronic acid (HA) dermal fillers have been rated the second most popular nonsurgical procedure.1 They are favored for their ease of administration and achievement of the desired aesthetic improvement.2 Despite the renowned safety of the procedure, rare adverse events have been reported in the literature.2–7

The author is an ophthalmologist and a laser eye surgeon with over 20 years of experience in HA use, and with over 5,000 cosmetic injection procedures. Over these years, only transient swelling and bruising during injection have been observed as procedure-related expected side effects. The objective of this article was to discuss these and offer a formal protocol for treatment.

Methods: This article presents 5 clinical cases of late-onset inflammatory response occurring at least 3 months after uneventful injection of HA dermal filler.

Results: Inflammation appeared spontaneously, usually 4–5 months after the last injection, but in 1 patient, almost 14 months later. One patient was injected at the same time with fillers manufactured by 2 different technologies. In this case, all areas treated with the same filler showed diffuse swelling of inflammatory nature, whereas the lips, treated with the second filler brand, remained unaffected. Four patients reported a flu-like illness or gastrointestinal upset a few days before the onset of dermal filler inflammation.

Conclusion: Late-onset inflammatory reactions to HA fillers may be self-limiting but are easily and rapidly treatable with oral steroids, and with hyaluronidase in the case of lumps. It is likely these reactions are due to a Type IV delayed hypersensitivity response. Delayed inflammation associated with HA fillers is nonbrand specific. However, the case where 2 different brands were injected during the same session, but only 1 brand triggered a hypersensitivity reaction, suggests that the technology used in the manufacturing process, and the subsequent differing products of degradation, may have an influence on potential allergic reactions to HA fillers.

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as the injection technique (eg, filler volume, repeated treatments, and intramuscular implantation) and different properties of the filler.

CASE 1

A 44-year-old female Asian patient presented with a diffuse swelling and tenderness without lumps 4 months after receiving 1.6 mL injection of Hydrafill Softline® (Inamed Aesthetics, Wicklow, Ireland) in the labiomial corners and nasolabial folds Table 1. The injection was performed in the author’s clinic with a needle, in a deep dermal/subdermal plane, with a retrograde linear thread technique. As the filler was lidocaine-free, a topical EMLA® cream was applied before the treatment.

The patient’s medical history included 7 injections of products from Juvéderm® (Allergan Inc., Pringy, France) and Hydrafill® ranges administered over the previous 3 years (total volume = 5.6 mL) for treatment of the same areas. No known allergies or history of autoimmune diseases were reported. Approximately 1 week before the reported reaction onset, the patient had suffered a flu-like illness.

The reaction began with redness and firm swelling at the corners of the mouth. The patient massaged the area in an attempt to resolve the symptoms. Within 24 hours, the swelling has spread to the inferior nasolabial folds and was characterized by redness, tenderness, and inflammation. The patient self-administered antihistamines (oral cetirizine hydrochloride, 10 mg) for 2 days without improvement. The reaction resolved completely in 1 week with a 5-day treatment with oral steroids (soluble Prednisolone®) in reducing doses of 60, 40, 20, 10, and 5 mg. Hyaluronidase was not required. At the time of the reaction, the use of hyaluronidase as a reversal agent for HA was not widely established in the United Kingdom.

CASE 2

A 48-year-old female Caucasian patient presented to her treating practitioner with a localized redness and swelling without lumps in the nasolabial folds 5 months after receiving 1 mL injection of Restylane® (Q-Med AB, Uppsala, Sweden) in the same area. The injection was performed in a different clinic with a retrograde linear thread technique Table 1. The filler was lidocaine-free and the use of additional anesthetics is unknown.

A couple of weeks before the reaction onset, the patient reported a cold sore on the lip, which had fully healed before the reaction. No further medical history, including any previous treatments with HA fillers and known allergies, is available.

The patient had been treated with oral antihistamines for 5 days without improvement. The treating practitioner called the author, who advised to prescribe a 5-day course of steroids, commencing at 60 mg, with reducing doses, and to schedule a review appointment. The injecting practitioner did not call the author back after 5 days, but subsequent confirmation of full resolution was obtained.

CASE 3

A 54-year-old female Caucasian patient presented with a diffuse, reddish, painful swelling without lumps 4 months after receiving injections of 2.4 mL of Teosyal® Puresense Ultra Deep (Teoxane S.A., Geneva, Switzerland) in the cheeks, chin, and marionette lines and 1 mL of Belotero® Intense (Anteis S.A, Geneva, Switzerland; a wholly owned subsidiary of Merz Pharmaceuticals GmbH) in the lips on the same day Table 1. The injections were performed in the author’s clinic. Teosyal® Puresense Ultra Deep was injected with the supplied needle and a 27G cannula suprapieriesostically and in the deep dermis. Belotero® Intense was injected with the supplied needle by subdermal and submuscular retrograde linear thread technique. The products contained lidocaine, and no additional anesthetics were applied.

The patient’s medical history included a 1 mL injection of Teosyal® Puresense Deep Lines to the nasolabial folds in the preceding year. No known allergies or history of autoimmune diseases were reported. A few days before the reaction onset, the patient had experienced a gastrointestinal upset.

The patient is a health care professional, so she was able to describe her reactions correctly over a phone consultation with the author. The patient was unaware of the injected filler brands, but unprompted, reported swelling in all treated areas, except for the lips. The photographs of her reactions were unfortunately lost over time. The patient administered oral antihistamines for 7 days and reported a gradual improvement. She refused intake of steroids and reported full resolution of her symptoms over a 2-week period.

Two months later, the patient experienced a second gastrointestinal upset, and subsequently the same facial areas flared up with red, tender swelling, but again the lips remained unaffected. The patient reported that the second episode was not as severe as the first one, and she self-administered antihistamines for 7 days with slow recovery of her symptoms.

CASE 4

A 46-year-old female Caucasian patient presented with a diffuse swelling with hard lumps on the forehead and a diffuse swelling of the labiomial corners 5 months after receiving a 1 mL injections of Teosyal® Deep Lines in the glabellar area and corners of the mouth (Fig. 1) Table 1. The injection was performed in a different clinic, to subdermal/deep dermis with a retrograde linear thread technique. The filler was lidocaine-free.

This was the patient’s first experience with HA injections. No known allergies were reported. During the week before the reaction onset, the patient had been abroad on holiday, where she had been systemically unwell and had suffered a gastrointestinal upset.

Before being referred to the author, the patient had taken oral antihistamines for 2 days without improvement. The author treated the affected areas with an injection of hyaluronidase (Hyalase®, Wockhardt UK Ltd.) 1,500 units in 1 mL, and the patient was given oral steroids (soluble
Fig. 1. Photographs of patient 4 (46 years old) taken at first presentation to the author after unsuccessful treatment with antihistamines (5 months after injection). A–D, Immediate reaction onset; A, B, diffuse swelling with hard lumps on the forehead; C, D, diffuse swelling on the labiomental corners.

Fig. 2. Full resolution at labiomental corners and improvement in the glabellar area following treatment with steroids and the first dose of hyaluronidase. A–D, Photographs of patient 4 taken at full resolution at labiomental corners and improvement in the glabellar area following treatment with steroids and the first dose of hyaluronidase.
Prednisolone®) for 5 days in reducing doses of 60, 40, 20, 10, and 5 mg. The erythema and swelling resolved, but a small palpable lump remained in the glabellar area, despite the overall improvement (Fig. 2). Three weeks following the first hyaluronidase administration, the patient was reviewed, and a further injection of Hyalase 1,500 units dissolved in 4 mL was injected into the residual glabellar lump. The patient subsequently called to report full resolution of her symptoms.

CASE 5

A 46-year-old female Caucasian patient presented with an asymmetrical bilateral swelling on the outer lid-cheek margins 14 months after receiving a 1 mL injection of Teosyal® Puresense Global Action in the outer cheeks and lateral tear troughs, as well as 1 mL of the same filler in the labiomental triangles (Fig. 3) Table 1. The injections were performed in the author’s clinic with subdermal retrograde linear technique with needle and cannula. A supraperiosteal bolus was deposited around the cheeks and the lid-cheek junction with a needle. A deep dermal bolus was deposited in the labiomental triangles with a cannula. The filler contained lidocaine, and no additional anesthetics were applied. The reaction was seen only in the lateral tear trough/lid cheek margin area; the labiomental triangles were unaffected.

The patient’s previous medical history included 6 injections with Juvéderm® 18, Succell® One (Sanofi Aventis, Paris, France) and Teosyal® Puresense Global Action administered over the previous 7 years (total volume = 4.8 mL). The patient reported a long-term history of hay fever and atopy, without mentioning any prior systemic illness.

At the reaction onset, the patient did not relate the symptoms to the filler, and thus first contacted her general practitioner. Upon prescription, the patient administered oral antihistamines for a week without improvement. Then the patient presented to the author and was treated with oral steroids (soluble Prednisolone®) for 5 days in reducing doses of 60, 40, 20, 10, and 5 mg leading to full resolution of the symptoms (Fig. 4).

DISCUSSION

Late-onset inflammatory reactions are rare complications, which may occur following injection of HA dermal fillers. Their cause may be infectious or immune-mediated in origin, and their outbreak can be triggered, for example, by a flu-like illness.2,3,5,6,13,14 Nevertheless, the latter events may be coincidental.

Fig. 3. Photographs of patient 5 (46 years old) taken at first presentation to the author after unsuccessful treatment with antihistamines (14 months after injection). A–C, immediate reaction onset. The larger swelling on the left lid-cheek margin, where the reaction is more pronounced due to a slightly larger injected volume, and the smaller swelling on the right.

Fig. 4. Full resolution following steroid intake. A, B, Photographs of patient 5 at full resolution following steroid intake.
Table 1. Patient Data

| Patient Number | Age at the Time of Injection (Years) | Total Volume Injected (mL) | Injectable HA Product (Manufacturer, Cross-Linking Technology) | Anatomic Area of Injection | Injection Depth for Retrograde Thread Technique | Time to Reaction (Months) | Symptoms | Treatment Performed in the Author’s Clinic (Yes/No) | Onset | Outcome |
|----------------|-------------------------------------|---------------------------|---------------------------------------------------------------|---------------------------|-----------------------------------------------|--------------------------|----------|---------------------------------------------------|-------|---------|
| 1              | 44                                  | 1.6 mL                    | Hydratone Softline® (24 mg/ml)                                | Labiomental corners       | Deep dermal/ subdermal                      | 4                        | Diffuse swelling and tenderness without lumps in the labiomental corners | Yes   | Resolved |
| 2              | 48                                  | 1 mL                      | Restylane®                                                  | Nasolabial folds          | Unknown                                      | 5                        | No       | No                                                |       | Resolved |
| 3              | 54                                  | 2.4 mL                    | Teosyal® Puresense Ultra Deep (25 mg/g)                       | Cheeks, chin, and marionette lines | Supraperiosteal/ deep dermal/ subcutaneous | 4                        | Yes      | Yes                                               |       | Diffuse swelling, localized redness and swelling without lumps in the cheeks and labiomental corners | 4     | Resolved |
| 4              | 46                                  | 1 mL                      | Teosyal® Deep Lines | Glabella-area and corners of the mouth | Subdermal/ subcutaneous/ submuscular | 5                        | No       | No                                               |       | Resolved |
| 5              | 46                                  | 2 mL                      | Teosyal® Puresense Global Action (25 mg/g)                    | Outer cheeks/ lateral tear troughs/ labiomental triangles | Subdermal/ supraperiosteal/ subcutaneous/ submuscular | 14                       | Yes      | Yes                                               |       | Asymmetrical bilateral swelling on the lateral tear troughs/ labiomental triangles margin area | 14    | Resolved |

If an infection is suspected, steroids should not be prescribed. All presented cases are believed to be immunological in nature, specifically as all injected areas were affected simultaneously, except in cases 3 and 5 discussed below. In the event of infection, symptoms would be localized or restricted to a discrete area. Moreover, at presentation, the patients were systemically well and had been asymptomatic in the injection area for months between the last treatment and reaction onset. Even the associated lump reported in patient 4 was hard and nonfluctuant and atypical of infection. Therefore, there was no need for an empirical antibiotic treatment. Microbiological analysis for detection of a quiescent biofilm was not performed.

Delayed type IV hypersensitivity following HA implantation is the most likely explanation of the observed late-onset events. This rare systemic response is initiated by T lymphocytes and mediated by CD4+ cells. The reaction manifests as a persistent facial edema in the treated area, at times accompanied by inflammatory nodules. A foreign body granuloma can be suspected in patient 4 due to the presence of a nonfluctuant lump. Based on data from early 2000s, the rates of delayed hypersensitivity vary between 0.02% and 4%. As the number of fillers, performed procedures, and the associated complications have increased in the last decade, a newer estimate would be of interest.

Late-onset hypersensitivity may manifest from weeks to many months after HA injection. It is impossible to predict, and it may occur in both previously injected and first time patients. Several case reports have been published attempting to understand the etiology in relation to HA dermal fillers. Suggested influencing factors include previous infections and trauma, as well as the injection technique (eg, filler volume, repeated treatments, and intramuscular implantation) and different properties of the filler.

Although severe adverse events may occur with any HA filler, the rates seem to vary among different products. The fillers cited in this report are characterized as non-immunogenic, biocompatible, and nontoxic implants, composed of sodium hyaluronate from nonanimal origin cross-linked with 1,4-butanediol diglycidyl ether. The technology of the cross-linking varies among different manufacturers. Based on preclinical data, it is advisable not to modify the HA molecule to such a large extent that it would no longer be recognized as HA, and thus potentially lead to foreign body reactions. The limit of accepted modifications remains unknown. Among the cited fillers, Restylane® products have the lowest and Teosyal® the highest degree of modification. Additional filler-related factors suggested to influence inflammatory activities include presence of impurities from the cross-linking and biofermentation processes, higher concentration of HA, and characteristics of the filler particles (eg, size, surface, and charge). However, as the manufacturing procedures remain confidential, one can only speculate on the technology-related factors associated with filler complications.
onset. The proposed hypothesis involves macrophages remembering stimulations such as a severe systemic infection, and their activation then triggering giant cell formation and foreign body granuloma. Belzay et al. proposed a different mechanism, by which VYCROSS technology HA may contribute to the inflammatory activities and thus exhibit immunologic properties. The suggested mechanism involves release of proinflammatory low molecular weight HA fragments during an accelerated breakdown of HA gels, triggered by a systemic inflammatory response to an unknown antigen.

In all true type IV granulomatous processes, all sites that were originally injected with filler material would be expected to be adversely affected simultaneously, as observed in this report. Patient 3 is a particularly interesting case because all sites injected with 1 brand were inflamed with the exception of the lips, which were injected with another brand. The fact that 2 different brands of filler were injected simultaneously in the same patient, but only 1 brand appears to have triggered a hypersensitivity response, suggests that the technology used in the manufacturing process of these brands can have an influence on possible side effects in the patient, even months after treatment. Published clinical data show that products from the second range of fillers (Belotero®) preserve the structural integrity of the surrounding tissues and have a favorable safety profile.

In case of patient 5, inflammation was observed approximately 14 months postinjection of the lid-cheek margins, but the labiomental triangles were unaffected. Normally, by this time, it would be expected that the dermal filler would have degraded and been eliminated, but published literature suggests that fillers injected in the tear troughs do not degrade as quickly as in other areas. This evidence may explain why the delayed reactions were observed only in the 2 tear troughs, after the other areas were free of injected HA.

Delayed complications are particularly difficult to diagnose and treat due to the time lapse from the last procedure. As observed in the described cases, type IV hypersensitivity reactions are unresponsive to antihistamines. Steroids are required to alleviate the inflammatory signs. In case of patient 3, if steroids had been used when the first reaction occurred, the second flare-up might have been avoided. Type IV foreign body reactions are however generally self-limiting until the foreign body is destroyed. Therefore, patient 3 achieved slow and full recovery even without steroids and attributed it to the antihistamines. Hyaluronidase may be injected to remove the allergen in case of lumps, as shown in patient 4.

Inflamed areas should not be massaged, as done by patient 1. The patient felt the need to massage, as the reaction appeared as if the filler was freshly injected and needed to be dispersed. However, manipulation aggravated the condition, induced tenderness and increased the edema.

The main limitation of this article is the absence of histological analysis for detection of macrophages and their activation then triggering giant cell formation and foreign body granuloma. Biopsies were not performed due to the patients’ desire to have minimally invasive resolutions to their symptoms as quickly as possible. As this is a retrospective data review, and 2 patients were referred to the clinic by different practitioners, the medical history is at times incomplete and exact doses of prescribed medication are also unknown. Only available nonstandardized patients’ photographs are presented in this article.

**CONCLUSIONS**

Late-onset inflammatory response occurs at least 2 months after HA injection, and presents as diffuse, firm, red, nonfluctuant inflammation of all areas containing the dermal filler. Patients are otherwise systemically well. Very late presentation, over a year after the last injection, can occur in some cases, depending on the location of the injected product and the speed of degradation in that area. Such reactions may occur with any HA dermal filler, but their incidence may vary depending on the manufacturing technology.

Prompt identification and correct treatment allow successful resolution of inflammatory symptoms within a few days. In the absence of lumps, the reactions may settle over time without intervention, but will need oral steroids in most cases for rapid and sustained improvement, and to reduce the risk of recurrence. Treatment of persistent lumps additionally requires injection of hyaluronidase for optimal resolution. Patients should be correctly informed of all possible rare adverse reactions before treatment to avoid fear, disappointment, or litigation and to ensure that they seek prompt and correct medical intervention when necessary.

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