Filling with hyaluronic acid and transcutaneous blepharoplasty in stable systemic sclerosis

Preenchimento com ácido hialurônico e blefaroplastia transcutânea em esclerodermia sistêmica estável

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ABSTRACT

Systemic sclerosis is a disease of unknown cause, is a rare pathology that, when not treated in time, produces severe damage to the facial aesthetic anatomy causing a significant impact on the quality of life. The case of a 54-year-old woman, with a history of systemic sclerosis of 25 years of evolution, and who was stable from her disease over four years ago. The patient consults for resolution of her facial deformity, which is why she proceeds to perform minimally invasive low-risk procedures such as transcutaneous blepharoplasty and fillings with hyaluronic acid obtaining aesthetically adequate results.

Keywords: Blepharoplasty; Hyaluronic acid; Scleroderma, diffuse

INTRODUCTION

Systemic sclerosis (SS) is a disease of the connective tissue that is characterized by an excessive deposit of collagen and other substances in the extracellular matrix, producing cutaneous sclerosis and damage to the microvasculature of the skin and internal organs. It predominantly affects women, with a women:men ratio of 4:1. The most common symptoms include Raynaud’s phenomenon, polyarthritis, dysphagia, heartburn, edema, skin thickening, and contractures of the fingers. There are three phases of dermal involvement: initially, there is an edematous phase, which often presents stiff and swollen hands and fingers; the second phase, called indurative,
is characterized by thickening and hardening of the skin (sclerodactyly and the classic expressionless face); and finally, there is an atrophic phase.5,6

SS is classified as limited (LSS), diffuse (DSS), and sine scleroderma (without scleroderma - ssSS). In cases of LSS (CREST syndrome: calcinosis, Raynaud’s phenomenon, esophageal dysmotility, sclerodactyly, and telangiectasia), patients develop skin tension on the face and in the distal portion of the elbows and knees. They may also have gastroesophageal reflux disease.6,7 In DSS, there is great diffuse skin involvement; patients present Raynaud’s phenomenon and gastrointestinal complications. This type of SS tends to evolve rapidly. Interstitial lung disease and scleroderma renal crisis are the main complications.8-10 In ssSS, patients have antibodies related to SS and visceral manifestations of the disease, but no skin involvement.9

Its incidence is <10 per 1 million per year, and its prevalence is <150 per 1 million in northern Europe and Japan. In the United States, Canada, southern Europe, and Australia it has an incidence of >10 per 1 million per year and prevalence estimates of >150 per 1 million.11

The term scleroderma was introduced by Gintrac in 1847 and arose from the importance of the skin’s participation in vascular disease and fibrotic changes.12 Currently, the term scleroderma refers to the skin involvement of patients. Every patient with morphea has scleroderma (cutaneous fibrosis), but not every patient with SS has scleroderma, which is why the term scleroderma should not be used as a synonym for systemic sclerosis.5

The exact pathophysiology of scleroderma is unknown. It is considered to be secondary to an autoimmune reaction that causes localized collagen overproduction. In some cases, it has been linked to exposure to chemicals. Genetic and infectious factors were involved as possible causal agents.5

Facial impairment of systemic sclerosis and morphea is associated with oral complications, and aesthetic changes strongly affect the patient’s self-image and quality of life.13-15

CASE REPORT

We present the case of a 54-year-old woman with a history of hypothyroidism, controlled with levothyroxine 50mg, severe depression, treated with escitalopram 10mg, and a diagnosis of SS with 25 years of evolution. She was being treated with tacrolimus 1.5mg every 12 hours, sirolimus 1.5mg per day, and methylprednisolone 4mg per day.

On physical examination, she presented atrophic skin, not very hard, without dyschromia, and also severe atrophy of the malar and zygomatic fats and the cheek area. We also observed, secondary to this loss of facial volume, a great herniation of the lower eyelid fat compartments, and microstomy, resulting in the classic aspect of mask-like appearance in this type of patients (Figure 1).

Procedure: Local anesthesia without vasoconstrictor was performed, followed by a transcutaneous incision 0.5cm in the area of greater fat extrusion in the lower eyelid. We removed part of the exuberant fat bags one centimeter below the eyelid, controlling the almost nonexistent bleeding and suturing the lesion with a single stitch using 6-0 monocryl, which was removed after seven days. We decided on this approach through the skin due to the exuberance of the bags, as it was the easiest and least complicated technique, without risk of ectropion (Figure 2).16,17

One week later, filling with high G prime or high-density hyaluronic acid (HA) (Voluma, Allergan, Guarulhos, Brazil) was applied with a cannula 21g in a linear retrograde fan pattern in the supraperiosteal plane in the malar, zygomatic, and in Bichat’s fat, totaling 4ml of HA per cheek (Figure 3).19-23

Then we apply 1ml of low-density HA (Volbella, Allergan, Guarulhos, Brazil), using a with needle 30G, to perform linear retroinjection in the perioral wrinkles and lip contour. Finally, we applied medium-density HA 1ml (Voliftt, Allergan, Guarulhos, Brazil) in the infraorbital fat compartment, followed by suturing of the skin incision (Figure 2).

**Figure 1:** Physical examination of the patient: (A) At rest. (B) In dynamics

**Figure 2:** Pre and immediate postoperative, skin incision and visualization of infraorbital fat

Guarulhos, Brazil), with a needle in small points in the lip vermilion to improve the turgor. 24,25

A very suitable aesthetic result was obtained, significantly improving the quality of life of this patient (Figures 4 and 5).

**DISCUSSION**

Recent studies cite that injections of HA, a non-sulfated anionic glycosaminoglycan widely distributed in all connective tissues, epithelial and neural, would be a possible treatment for cutaneous fibrosis.18,19 It is believed that hyaluronic acid would be a valid therapy in patients with scleroderma due to its filler properties, in addition to its ability to retain water, smoothing, and moisturizing the skin.20,21 Also, HA has been shown to induce the production of type I collagen in the dermis, which could explain its long-lasting effect.22

Recently, the group led by Guggino included ten women between 18 and 70 years old, with systemic sclerosis. Each patient was treated with three injections of HA and platelet-rich plasma every 15 to 20 days. All patients were assessed monthly, at three and 24 months after the end of the treatment, regarding mouth’s opening, freedom of movement of the lips, and skin elasticity. The group observed that eight patients (80%) showed higher mouth’s opening and increased thickness of the upper lip since the first month of follow-up, maintaining these results after two years of initial control.

Another potential filler could be the autologous fat. Autologous fat lipotransfer for soft tissue filling has been described as a well-established aesthetic, and plastic surgical technique.14 Patients with stable scleroderma who have been injected with autologous fat have been reported.13 It is believed that the fat grafting may have tissue regenerative properties, not only serving as a filler, thus postulating that in the fat there could be stem cells, fibroblasts, and endothelial cells that would decrease fibrosis.13,14

There is no consensus in the literature on the application of fillers and minimally invasive surgery in patients with systemic sclerosis. Thus, we suggest and consider that the stability of the systemic condition for more than two years (clinical and laboratory stability) should be sufficient to allow the use of HA or autologous fat in these patients. Note that, before using any filler, you must confirm the stability of the disease, verify and all medications in use by the patient (immunosuppressants, anticoagulants, etc.), and update the laboratory tests (complete blood count, platelet count, hepatitis C and B, HIV, and quantiferon-TB).16

**CONCLUSIONS**

Minimally invasive procedures, such as transcutaneous blepharoplasty and facial filling with HA, can successfully improve the cutaneous cosmetic complications of SS. The appropriate aesthetic result will depend on the experience, technique, and skill of dermatologists and/or plastic surgeons.●
REFERENCES


