Surgical options in vitiligo: skin graft and epidermal suspension diluted in hyaluronic acid gel
Opções cirúrgicas no vitiligo: enxerto de raspado cutâneo e suspensão epidérmica diluídos em ácido hialurônico gel

ABSTRACT
Introduction: Vitiligo is an acquired skin dyschromia characterized by the physical and/or functional reduction of melanocytes. We present two surgical proposals for the treatment of vitiligo.

Case reports: 1) Implant of skin graft diluted in hyaluronic acid gel: We obtained the material through curettage, diluted it in hyaluronic acid gel, and applied it to receptor areas. 2) Epidermal suspension obtained through curettage and diluted in hyaluronic acid gel: After the curettage of the donor area, we treated the material with trypsin-EDTA, centrifuged it, and diluted it in hyaluronic acid gel. The receptor area received the graft.

Conclusion: These are safe, easy, and satisfactory surgical procedures for the presented cases.

Keywords: Vitiligo; Keratinocytes; Ambulatory surgical procedures

INTRODUCTION
Vitiligo is an acquired, idiopathic cutaneous dyschromia characterized by physical and functional reduction of melanocytes. The global prevalence is around 0.5% to 1%. Clinically, it presents achromatic macules and patches of different sizes and forms.¹ Stable vitiligo cases resistant to clinical treatment are candidates for surgical treatment, including non-cultured epidermal suspension grafts treated enzymatically with trypsin 0.25%; thin dermo-epidermal skin grafts; suction blister epidermal grafts (SBEG); total punch grafting; epidermal grafting or in vitro isolation; and culture of melanocytes.² ³
Additionally, Machado (2000) demonstrated the feasibility of obtaining material for grafting through simple epidermal curettage of the donor area to be implanted in a recipient area, also curetted. The obtained graft is humidified with physiological saline to obtain a “paste”; it is applied to the recipient area and fixed with an adhesive semi-permeable membrane.

The present article describes two surgical techniques for treating vitiligo, considering variations of the curettage technique used for obtaining the graft from the donor area.

**Implant of epidermal curettage graft diluted in hyaluronic acid gel**

The technique’s characteristic is the dilution of the obtained graft in a gel of hyaluronic acid. This biocompatible and hygroscopic substance provides greater viability and adhesion to the receptor area. The curettage of the donor area to the papillary dermis obtains the material (Figure 1A). It is then diluted in 1-2 ml of hyaluronic acid gel at 0.5-2% (Figure 1B). The recipient area is also curetted reaching the papillary dermis and obtaining the same size as the donor area. Finally, the graft is applied over the recipient area (Figure 1C) and covered with a dressing of porous membrane of cellulose, maintained in site for seven days. Topical medications and phototherapy are reintroduced 14 days after the procedure. Satisfactory results are observed after 90 days (Figure 1D and 1E).

**Non-cultured melanocyte-keratinocyte cells suspension obtained by curettage and diluted in hyaluronic acid gel**

It corresponds to the association of the techniques curettage grafting and uncultured epidermal suspension methods. Mulekar (2003 and 2005) and van Geel (2001) initially described the use of hyaluronic acid in epidermal suspensions. After curettage of the donor area until the onset of the papillary

**Figure 1:** Implant of curetted skin graft diluted in hyaluronic acid gel. A - Curettage of the donor area. B - Graft diluted in hyaluronic acid gel. C - Post-grafting recipient area. D - Preoperative. E) Postoperative (90 days)
dermis’s punctate bleeding, the collected graft is exposed to a proteolytic solution (Trypsin EDTA 0.025% – LGC Biotechnology™ – Brazil) and incubated for 20 minutes at 98.6° Fahrenheit. After incubation, a pipette aspirates the trypsin. The sample is then washed with 0.9% saline solution and transferred to a centrifuge tube containing the DMEM culture medium (LGC Biotechnology™ – Brazil). After six minutes of centrifuging at 1500 rpm, the supernatant of epidermal cells is discarded, and the pellet is suspended in 1-2 ml of hyaluronic acid gel 0.5-2% (Paulista Center for Pharmaceutical Development™ – Brazil). The suspension concentration, which can vary according to the clinical case of vitiligo, generates a donor to receptor area ratio ranging between 1:10 to 1:20. The recipient area is curetted or dermabraded to the papillary dermis. After applying the epidermal graft, it is occluded with a porous cellulose membrane, which should remain on the site for seven days. The topical medications and phototherapy should be reintroduced after 14 days. Satisfactory results are observed after 90 days and improve 180 days after the procedure (Figures 2A and 2B).

CONCLUSION
In conclusion, curettting the skin until the papillary dermis is an affordable procedure, easy to perform, which provides a satisfactory sample for grafting. When this technique is associated with hyaluronic acid, it allows greater viability and adherence of the graft to the recipient area. Nowadays, there are not indexed publications of those described techniques, and future studies are necessary for further elucidation and improvement of these treatment modalities.

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Figure 2: Uncultivated suspension obtained from curettage with subsequent dilution in hyaluronic acid gel.
A - Preoperative.
B - Postoperative (180 days)
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