A utologous fat transplantation is a common and ideal technique for soft-tissue augmentation and for filling soft-tissue defects due to its minimal tissue reaction and easy availability.

**Background:** The increased resorption and the difficulty of the fat graft take following autologous fat transplantation procedure are associated with reduced fat tissue revascularization and increased apoptosis of adipose cells. We suppose that the lipofilling procedure induces an inflammatory environment within the fat graft mass, whose evolution influences the efficacy of autologous fat graft survival. Erythropoietin (EPO) is a glycoprotein hormone known to exert angiogenetic and anti-inflammatory effects; therefore, our purpose was to investigate its reaction with adipose tissue used in lipofilling.

**Methods:** Fat masses were harvested using manual suction lipectomy and then seeded on dishes in appropriate culture and treated for 3 weeks with 3 doses of EPO. CD31 and CD68 immunohistochemistry was used to identify microvessels and several infiltrating leukocyte cells.

**Results:** Following EPO administration, we have detected an increase in the number of CD31-positive microvessel endothelium cells and CD31-positive small leukocytes and a reduction of CD68-positive cells. These effects were more conspicuous following higher EPO dose.

**Conclusions:** Our findings evidence EPO treatment as a useful strategy to sustain the revascularization of grafted tissue and to reduce its inflammatory state. (Plast Reconstr Surg Glob Open 2015;3:e338; doi:10.1097/GOX.0000000000000305; Published online 30 March 2015.)

However, although this technique follows standardized protocols, there is a relatively high resorption rate of the fat graft, the long-term graft survival is unpredictable, thus the desired effect is not always achieved. It has been observed in long-term follow-up studies that 20–80% of the graft volume is lost.

The fat graft survival depends on microenvironment, and a dynamic remodeling after fat graft was well received by the recipient site, but the mechanisms of tissue revascularization/regeneration and subsequent absorption/fibrosis of the graft are not yet completely understood.

Quality and amount of revascularization are the key points for graft survival. After autologous fat transplantation, the increased resorption and in-
creased apoptosis of adipose cells are direct consequences of poor revascularization. Besides, some clinical and experimental studies have reported continuing volume loss of the grafted adipose cells even after the grafts appeared revascularized.

Several studies have documented the production of inflammatory cytokines by adipose tissue. Their occurrence depends not only on macrophage cells normally present in the adipose tissue mass, but a large amount of this production may directly depend on the preadipocytes, which have been observed to be able also to produce monocyte chemotactic protein-1 and macrophage migration inhibitory factors.

Based on these findings, we suppose that lipofilling procedure induces an inflammatory environment within the fat graft mass, whose evolution influences the autologous fat graft take.

Erythropoietin (EPO) is a glycoprotein hormone that was first characterized by its ability to stimulate erythropoiesis. In following researches, it has been discovered that EPO is also involved in angiogenesis induction: instigating the secretion of angiogenic factors, enhancing the immune response, and exerting anti-inflammatory effects.

Our purpose was to investigate the effect of EPO on adipose mass used in lipofilling, focusing mainly on the effects on tissue revascularization and the influence on immune cell behavior.

**MATERIALS AND METHODS**

Fat graft was harvested using manual suction lipectomy under general anesthesia from 10 women, 41 ± 4 years old, during procedures of reconstructive surgery. The participants gave written informed consent. The fat was harvested using a 14-gauge blunt cannula and centrifuged at 1500 rpm for 2 minutes.

The fat mass was seeded on dishes in appropriate culture on average for 24 hours and then treated for 3 weeks with 3 different doses of EPO (EPREX, Janssen-Cilag, Milan, Italy): 0.15, 0.30, and 0.60 µg/ml. A positive control was performed administering a cocktail of trophic drugs (biotin, T3, pantothenate, hydrocortisone, and insulin; from Sigma, St. Louis, Miss.). The negative control was left without any treatment.

At the end of this phase, the fat mass was fixed in formalin and processed for paraffin inclusion and cutting; the obtained 5-µm-thick slides were stained with CD31 (to identify active vessels and immune system cells) and CD68 (to identify macrophages).

Quantification of cell infiltration in the fat grafts was estimated by counting the number of positive vessels and cells. To avoid counting bias, due to the change of adipocytes diameter, the counting data were referred to tissue fat mass of 80 adipocytes. Parametric statistical analysis was obtained with Prism5.0 (GraphPad Software, San Diego, Calif.).

**RESULTS**

The study of the fat mass cultured under microscope evidenced 3 different zones in respect of adipocytes dimension, extracellular matrix deposition, and vessels density. These zones are identified as follow:

- **Zone 1**: wide presence of extracellular matrix, small adipocytes (686.3 ± 33.2 µm²), presence of large microvessels, and few capillary vessels.
- **Zone 2**: appreciable presence of extracellular matrix, large adipocytes (4573.4 ± 162.3 µm²), and presence of capillary vessels.
- **Zone 3**: poor occurrence of extracellular matrix, very large adipocytes (12359.6 ± 347.6 µm²), and presence of capillary vessels.

Our data were referred specifically to the identified zones.

Vessels count has evidenced no changes in the number of CD31-positive and CD31-negative vessels in zone 1 and zone 3 between the different experimental groups analyzed. Differently, zone 2 that has been treated with a higher dose of EPO showed a higher number of CD31-positive vessels in comparison to other experimental groups (Fig. 1).

The CD31-positive cells count showed the occurrence of 2 cell populations named small cells (22.3 ± 1.1 µm²; morphologically compatible with helper lymphocytes) and intermediate cells (68.5 ± 6.1 µm²; morphologically compatible with cytotoxic and natural killer lymphocytes). The positive control group showed a reduction of the number of CD31-positive cells compared with the negative control group. Differently, EPO treatment showed an increase in the number of small cells and a light decrease of intermediate cell number in all the 3 zones (Fig. 2).

The CD68-positive cells count showed the occurrence of 2 cell populations named very large cells (247.1 ± 15.2 µm²; morphologically compatible with macrophages) and large cells (129.2 ± 9.3 µm²; morphologically compatible with B-lymphocytes). In comparison to the negative control, the positive control group exhibits a reduction of CD68-positive cell number in zone 2 and zone 3, whereas all EPO treatments showed reduction of very large cell number in all zones examined and a reduction of large cells in zone 3. Following higher EPO dose treatment, the
very large cells reduction was proportionally more evident (Figs. 3 and 4).

**DISCUSSION**

There is a wide presence of inflammatory cells in adipose mass extracted for refilling, suggesting the potential of the fat mass to produce an inflammatory response.

EPO is a well-known factor able to induce tissue revascularization. EPO receptors have been found in several human leukocytes; therefore, researchers supposed that the EPO receptors have a role in modulating some signaling pathways.

Our findings clearly remark the provascular properties of EPO. Specifically, we observed EPO to induce an increase in the number of CD31-positive microvessels. CD31 reactivity is a well-known marker, indicating the functional activation of microvessels in the tissue; therefore, several CD31-negative microvessels observed in culture should be interpreted as
Fig. 3. Bar graph showing results of the morphometrical analysis performed on CD68-positive cells of zone 3, detected in the fat mass.

Fig. 4. Microphotograph panels showing the immunohistochemical positive signals occurring in the fat mass. CD68 immunohistochemistry.
Effects of Erythropoietin on Adipose Tissue

quiescent microvessels. In addition, EPO presents a
direct effect on tissue infiltrating leukocytes. The re-
duction of macrophage number is an important find-
ing that may evidence a reduced inflammatory state,
with macrophages being the principal cells involved
in the tissue inflammatory pathways. Other significant
evidence is the increase of CD31-positive small lym-
phocytes. In the absence of a more detailed analysis,
the increase of the number of lymphocytes can be
imputed only to a shift of cellular physiology from a
nonactive state to an active state.

Small lymphocytes widely participate in the in-
flammatory phenomena; on the other hand, they are
also involved in the release of tissue trophic factors.7
These findings may indicate lymphocytes as crucial
cells in supporting the tissue survival and they may
also be at the origin of the observed reparative ef-
fects detected on skin lesions and bone fractures.8,9

CONCLUSION
In conclusion, EPO treatment may be a useful
strategy to reduce the inflammatory state of adipose
grafted tissue and to sustain its revascularization.

Further studies are required to analyze the ex-
tent of graft tissue damage related to inflammatory
response and which pattern of cytokines is involved
in modulating the inflammatory response. We think
that a controlled inflammatory response may be use-
ful to obtain a higher and more predictable survival
rate in fat transplantation.

Optimizing the fat tissue graft take may lead to a
reduction in the number of procedures required to
obtain the desired outcome and thus improving the
cost/benefit rate.