Skin Mast Cell Promotion in Random Skin Flaps in Rats using Bone Marrow Mesenchymal Stem Cells and Amniotic Membrane

Farzaneh Chehelcheraghi¹*, Abolfazl Abbaszadeh² and Magid Tavafi²

¹Department of Anatomical Sciences, School of Medicine, Lorestan University of Medical Sciences, Khorramabad, Iran; ²Department of Surgery, Lorestan University of Medical Sciences, Khorramabad, Iran

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ABSTRACT

Background: Skin flap procedures are employed in plastic surgery, but failure can lead to necrosis of the flap. Studies have used bone marrow mesenchymal stem cells (BM-MSCs) to improve flap viability. BM-MSCs and acellular amniotic membrane (AAM) have been introduced as alternatives. The objective of this study was to evaluate the effect of BM-MSCs and AAM on mast cells of random skin flaps (RSF) in rats. Methods: RSFs (80 × 30 mm) were created on 40 rats that were randomly assigned to one of four groups, including (I) AAM, (II) BM-MSCs, (III) BM-MSCs/AAM, and (IV) saline (control). Transplantation was carried out during the procedure (zero day). Flap necrosis was observed on day 7, and skin samples were collected from the transition line of the flap to evaluate the total number and types of mast cells. The development and the total number of mast cells were related to the development of capillaries. Results: The results of one-way ANOVA indicated that there was no statistically significant difference between the mean numbers of mast cell types for different study groups. However, the difference between the total number of mast cells in the study groups was statistically significant (p = 0.001). Conclusion: The present study suggests that the use of AAM/BM-MSCs can improve the total number of mast cells and accelerate the growth of capillaries at the transient site in RSFs in rats. DOI: 10.29252/ibj.22.5.322

Keywords: Angiogenesis, Necrosis, Mast cells, Reconstructive surgical procedures, Surgical flaps

INTRODUCTION

Surgical skin flaps are frequently used to heal wounds caused by trauma, congenital defects, tumor removal and other issues. Partial skin flap necrosis is a common difficulty in the clinic, particularly on the distal part of the flap. Flap necrosis is caused mainly by inadequate blood perfusion or ischemia-reperfusion. It promotes several damaging alterations in the tissue and vasculature, such as reactive oxygen species and superoxide dismutase activity[11]. Research has been focused on improving blood flow in the flaps, decreasing ischemia, and minimizing necrosis[12,13]. These studies have used bio-scaffolds and stem cell therapy and produced increased blood flow and neoangiogenesis, decreasing the necrotic areas in the skin flaps[4,5]. Stem cells have a high ability for differentiation, and this ability increases during tissue engineering (TE)[6]. Studies have demonstrated that the transplantation of bone marrow mesenchymal stem cells (BM-MSCs) can improve ischemia and flap survival by promoting neovascularization[7], BM-MSCs promote the secretion of angiogenesis growth factors, such as vascular endothelial growth factor (VEGF) and basic fibroblast growth factor (bFGF), which are critical for neovascularization[7]. However, the squat amount of BM-MSCs in bone marrow has limited its clinical use. The amniotic membrane (AM) is a form of bio-scaffold derived from the human placenta. AM has the
capacity to reduce scarring and inflammation, has anti-
microbial properties and improves wound healing. It
also assists, as a scaffold, in cell proliferation and
differentiation. An AM scaffold is a template of the
extracellular matrix (ECM). The goal of AM is the
application of biological scaffolds in TE. In addition,
AM is a biomaterial that can be easily obtained,
produced and transported. AM can be used either
with the amniotic epithelium (complete AM) or
without it (acellular AM [AAM]). Complete removal
of the cellular constituents from AM is essential for
the denudation protocol to preserve the mechanical
components of the remaining scaffold. A scaffold with
the chosen ECM components and low immunogenicity
is a preferred matrix for TE.

In wounds, mast cells serve as the producer of pro-
inflammatory mediators (cytokines). Cytokines can
preserve inflammation, vascular modifications and
leukocyte penetration. They can generate NGF
(nerve growth factor), PDGF (platelet-derived growth
factor), VEGF, FGF2, histamine, and tryptase. These
growth factors are involved in the proliferation of
epithelial cells and fibroblasts. Cytokines can be
detected in the early inflammation phase of wound
healing. The response of the cytokines to inflammation is stimulation of re-epithelialization, re-
angiogenesis and, eventually, collagen and matrix
remodeling. After severe inflammation in the first
period of wound healing, necrotic tissue is replaced by
granulation tissue, which has numerous capillaries.

Mast cells are key effector cells in inflammation and
wound healing. The aim of the present study was to
evaluate morphological changes such as the total
number and degranulation of mast cells and the number
of type 1, 2, and 3 mast cells in the transitional line
using stereological techniques. This experimental study
was designed to explore the effects of BM-MSCs and
AAM on mast cells in transitional lines in a random rat
skin flap model. Wound healing was assessed by
evaluating angiogenesis and capillary density.

MATERIALS AND METHODS

Cell isolation and labeling
This work was authorized by the Ethics Committee
of Lorestan University of Medical Sciences
of Wistar rats were excised and the bone marrow was
removed by flushing the hollow bone marrow. This
was done using a syringe with a 20-gauge needle filled
with DMEM. The collected BM-MSCs were gently
pipetted to open cell clusterings and to attain a
homogenous cell suspension. The cells were
centrifuged at 1200 ×g for 7 min, and the cell pellet
was then re-suspended in 3 ml of culture medium. The
cell suspension was seeded in 25-cm polypropylene
tissue culture flasks with a 5-ml culture medium and
preserved in a humid atmosphere with 5% CO2 at
37 °C for 15 min. Cultures of the BM-MSCs were
examined every three days, and the results were
recorded. The cultures were passage when the BM-
MSCs had attained about 80% confluency.

The mesenchymal population was isolated on the
basis of its ability to attach to the culture plate. The
cell markers on the BM-MSC culture surface to
differentiate between media were analyzed using flow
cytometry. The cells were marked using field markers
for BM-MSC, including CD90, CD105, CD45, and
CD34. To place and orient the BM-MSCs at the
wound healing flap, the cells were tagged with
fluorescent cell tracker 1,1-diocetyl-3,3',3'-
tetramethylindocarbocyanine perchlorate (DiI) dye
according to manufacturer instructions. Rat BM-MSCs
were fully grown in 75 cm² culture flasks containing
fluorescent Dil dye. On the day of labeling, the media
was exchanged with 50% fresh BM-MSC media and
50% serum-free media. The cells were collected and
re-suspended in serum-free media at a density of 1 ×
10⁶/ml. The cells were then labeled with Dil
fluorescent cell tracker (cell tracker CM-Dil (C-7000,
Molecular Probes Inc., USA) by immersion in 1 ml of
a serum-free medium containing 10 µl of Dil at 37 °C
for 20 min. The cells were cleaned twice with
phosphate buffered saline (PBS) before injection.

Acellular amniotic membrane preparation
Human AMs were acquired from aseptic sites from 10
women who received elective cesarean sections at
Shafa Hospital in Lorestan, Iran. They showed no signs
of immunological disorders, hepatitis B, hepatitis C,
cytomegalovirus, unsophisticated pregnancies, white-
out preterm rupture of membranes, infections, and
streptococcus B on vaginal smears or other issues. The
AM was eroded three times with antiseptic PBS that
included 50 µg/ml of penicillin and 50 µg/ml of
streptomycin. The spongy layer was excised and cut
into 2.5 × 2.5 cm pieces. The epithelial cells were
carefully detached in 0.05% ethylenediaminetetraacetic
acid (EDTA; Invitrogen, Germany) at 37 °C for 2 h
and were lightly debrided with a cell hand tool under a
microscope. The evacuation of epithelial cells was
established by exhausting hematoxylin and eosin dye
(Sigma-Aldrich, Germany) ¹₄,₁₅.

Experimental design
Forty male Wistar rats, weighing 250 to 350 g, were
used in this study. Animal care was provided according
All the rats were placed in a prone position and anesthetized with ketamine (50 mg/kg) and xylazine (5 mg/kg). The animals were randomly assigned to one of the four groups as follows: Group 1, dorsal skin flap without MSC injection and AAM transplant receiving 0.5 ml saline injection from the distal to proximal flap in bed; Group 2, dorsal skin flap with an injection of 1 million MSCs in the distal area of the flap; Group 3, dorsal skin flap in which 1 million cells with AAM transplanted flap in bed; Group 4, dorsal skin flap without MSC injection with AAM transplanted flap in bed (Fig. 1).

Random dorsal skin flap model
After the skin of the rats had been shaved, random skin flaps (RSFs) were prepared, and the full thickness of the skin and skin muscle (panniculus carnosus) were measured. The RSFs were placed at the distal end of the animal on a horizontal line between the iliac crests. The size of the flaps was 30 × 80 mm. After placement, the flaps were immediately renewed using stitched 4.0 nylon thread. The day of surgery was day zero. Directly after surgery, the surface area of the flap was assessed\[^{16}\].

Clinical assessments
Assessment was carried out on day 7. Flap viability was determined by clinical (color and capillaries) and histological examinations. The flaps were sliced at the transitional lines and removed and fixed in 10% formalin (Sigma-Aldrich) in PBS at a pH of 7.6\[^{17}\].

Histological examinations
The mast cells in the slices were stained with 10% toluidine blue. There are three types of mast cells\[^{18}\]. In this work, the formation of the cells (connective tissue) in rats were identified and rested. Considering the grade of formation in each cell, mast cells from the transitional line of the flap were used to epitomize three stages of mast cell suppression. Cells that were entirely stained dark blue were called type 1. Cells in which some granules had been produced (cell outline preserved) were identified as type 2. Cells showing a large and generalized degranulation (total or incomplete impairment of the cell) were identified as type 3 (Fig. 2).

To accurately assess the number of each type of mast cell in the transitional line of the flap area, a linear zoom (1033.3 mm\(^2\)) was used on the area between necrosis and the healthy zone, with a 100× target. Each group (n = 10) was photographed, and stereological techniques were used to obtain information for 3-dimensional organization based on measures from the
two-dimensional sections\textsuperscript{19-21}. In this work, 15 mast cell investigation was performed using a specialized cell investigation grid (Fig. 3).

**Immunohistochemistry**

Six sample sections of the transitional line area in individual groups were deparaffinized in xylene and rehydrated, then transferred to graded ethanol baths. After coating, the sections were sealed with 3\% (v/v) H\textsubscript{2}O\textsubscript{2} and preserved with 10.2 mM sodium citrate buffer (antigen retrieval) at 95 °C for 20 min. They were then blocked with 5\% (w/v) bovine serum albumin and 1\% (v/v) Tween-20 in PBS for 10 min. The sections were immersed in VEGF (1:500; PeproTech, Rocky Hill, USA) at 4 °C overnight. Then the sections were immersed in a suitable horseradish peroxidase-conjugated secondary antibody (Santa Cruz Biotechnology, USA) and counterstained with hematoxylin. The number of VEGF-positive blood vessels for each measurement area (2 mm) were assessed. Six random fields of three random sections from each tissue section were used to compute the positive cells\textsuperscript{22}.  

**Statistical analysis**

The data was evaluated with SPSS (ver. 16), and all information was analyzed using one-way ANOVA and Tukey\textquoteright s post hoc comparisons. The results were considered to be significant at \( p < 0.05 \).
Mean ± SD for the numbers of type 1, 2, and 3 mast cells, and the total number of all types of mast cells in each group in 1033.3 mm² area of full thickness skin of transitional line on day 7, estimated by stereological methods (magnification ×100). Difference between the mean number of mast cells in type 1 (A), type 2 (B) and type 3 (C) in different groups was not statistically significant (p = 0.307, p = 0.536, and p = 0.587, respectively). (D) The mean total number of mast cells in BM-MSCs was higher than AAM group (p = 0.002), AAM/BM-MSCs group (p = 0.001), and the control group (p = 0.001), as well as in AAM, it was higher than AAM/BM-MSCs group (p = 0.044) and the control group (p = 0.003).

**DISCUSSION**

Studies have revealed that BM-MSCs can affect the speed and the quality of wound healing. A variety of stem cells have been used to treat ischemia, among which BM-MSCs, adipose tissue derived-MSCs and human umbilical cord MSCs are the most studied. The current study seeded BM-MSCs on AAM because of the positive effect of stem cells, especially BM-MSCs, on healing in ischemia and the positive effect of mast cells on angiogenesis in wound healing [13]. The combination of these two features can hypothetically improve the effectiveness of either treatment separately on ischemia of skin flaps and can provide a better prognosis after flap transfer. In the current study, the role of AAM and BM-MSCs on mast cells during ischemia in a dorsal RSF was investigated. Angiogenesis of the skin flap is a crucial and a complex process; it involves the proliferation of endothelial cells and the cooperation of various growth factors [23]. Some studies have found that the expression...
of CD31 and the presence of VEGF leads to angiogenesis and vascular progression. Although pure BM-MSCs and AAM do not have the ability to secret VEGF, the effect of this new management was investigated by measuring the types of mast cells and the vascular density in skin flaps. The gradual changes in appearance of mast cell have been the subject of study aimed at finding the key function of these cells in the dynamic physiology of skin healing. Consideration of the number of mast cells and statistical analysis indicates that the use of BM-MSCs notably expanded the number of mast cells in all experimental categories on day 7. It appears that in the AAM/BM-MSCs group, the number of type 2 mast
cells increased compared with control, but the increase was not significant. The number of type 3 mast cells decreased in all categories, but the decrease was not statistically significant. The number of type 3 mast cells decreased on day 7, indicating that AAM/BM-MSCs delayed the conversion of type 2 into type 3 mast cells. In the AAM group, the type 3 mast cells decreased significantly on day 7 compared to the BM-MSCs and AAM/BM-MSCs groups. This event indicates that AAM has a notable effect on decreasing the number of type 3 mast cells, in other words, in converting type 2 mast cells to type 3. It is clear that AAM modulates the production of type 3 mast cells.

Studies have suggested that decreasing the components of inflammation, such as mast cell degranulation, as well as increasing the total number of mast cells can positively affect wound healing[25,26]. This finding indicates that mast cells have similar potential for regenerating flap tissue. Mast cells are sensitive to biochemical signals at sites of injury and inflammation and can be polarized against or in favor of inflammatory phenotypes that can have numerous downstream effects[3,22]. The results of our study showed an increase in type 1 mast cells as well as an increase in the area of necrosis in the control group. The process of converting of type 2 to type 3 mast cells in all treated categories indicates that stages in the control category will change. The process of change was as for the experimental category. It appears that the change type 2 into type 3 mast cells continues in the control and experimental categories.

It was found that although BM-MSCs increased the total number of mast cells in the BM-MSC and AAM/BM-MSC groups compared with the control, this increase is related to type 1 and possibly type 2 mast cells. A previous study has revealed that mast cells contribute to scar tissue during wound repair[17].

When AAM decreases the presence of type 3 mast cells during wound repair, it may prohibit scar formation by its direct and strong influence on fibroblast proliferation[25]. It has been suggested that mast cells contribute to tissue repair through the secretion of soluble factors rather than trans-differentiation[27]. Even though mast cells act as a homeostatic ensemble of skin healing under gradual degranulation, it becomes harmful when it exceeds the threshold of destruction. As a result, the repair stages can be affected by chronic inflammation and changes in the dynamics of proliferation[28].

In the current study, when AAM/BM-MSCs reduced the type 3 mast cells throughout wound healing, it reduced scar formation and influenced fibroblast proliferation. Biological scaffolds such as AM

![Fig. 8. Dil-fluorescence in skin flap. Arrows represent the BMMSCs identified by Dil-fluorescence in skin flap.](image-url)
microvascular builder scaffolds are widely used in wound healing [29]. AM acts instead of a scaffold for TE. The ECM elements of the base membrane on the AM contains collagen, fibronectin, laminin, and other proteoglycans, which is remarkable for cell growth. These elements are integrin ligands and are effective in cell adhesion [8].

The study by Youn et al. [30] revealed that normal wound healing requires mast cells. The results of another study showed that microdistortion of wounds is a healthy stimulant for the onset of cellular mast-cell activity in a complete wound healing model [39]. Research has demonstrated a connection between the surge in mast cell degranulation and micro-distortion wound therapy and other wound-healing factors [30]. It can be concluded that AAM/BM-MSCs increase the number of type 2 mast cells and cause the development of microvessels. However, further research is needed to determine other mechanisms in which AAM/BM-MSCs suppress the presence of mast cells in inflammation.

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CONFLICT OF INTEREST. None declared.

REFERENCES


