EVALUATION OF THE EFFICACY OF LEUKOCYTE-PLATELET RICH FIBRIN VERSUS PLATELET RICH FIBRIN MATRIX ON BONE HEALING

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ABSTRACT

Objective: The aim of this study was to compare the efficacy of leukocyte-platelet-rich fibrin (L-PRF) and platelet-rich fibrin matrix (PRFM) on bone healing.

Methods: Eight rabbits were included in this controlled, prospective, blinded study. Defects with a diameter of 3 mm and 2 mm depth were created in the bone of both tibias, filled with L-PRF (group II), PRFM (group III) or left empty (group I). The bone mineral density (BMD) was measured by dual energy absorptiometry (DXA) and histological examination was performed on the 10th and 21st days.

Results: According to Emery’s histopathological healing criteria, control group (group I) revealed score 0 and score 3 at 10 and 21 days respectively. Group II showed score 2 and score 5 at 10 and 21 days respectively. On the other side group III treated by PRFM showed score 3 and score 7 at 10 and 21 days respectively.

Conclusion: Both L-PRF and PRFM showed increased bone formation in surgically created bone defects in rabbits, however, the effect of PRFM was more pronounced. The use of PRFM and L-PRF may contribute to the success of bone regeneration.

KEYWORDS: Bone regeneration, L-platelet-rich fibrin, platelet-rich fibrin matrix.

INTRODUCTION

Bone tissue regeneration is a complex process involving a number of cellular functions and mineralization of the defect followed by an eventual remodeling of the defect site to reconstruct the original structure (1). Several studies have shown that bone regenerative procedures may be enhanced by the addition of specific growth factors (2,3) and the use of biomaterials for guided tissue/bone regeneration (4).
Platelet-rich fibrin (PRF) was introduced by Dohan et al. as a second-generation platelet concentrate (5). In a simple preparation technique, blood is collected without anticoagulant or thrombin and immediately centrifuged only once. The growth factors present in PRF are the same as those in platelet rich plasma (PRP). Kawase et al. (6) suggested that a fibrous network of insoluble fibrin provides a scaffold for cells and serves as a substrate for the sustained release of growth factors. A previous study demonstrated that fibrin provides a matrix for cell growth and differentiation and provide a preferable environment for osteoblastic differentiation, during which cells contact fibrin molecules and exhibit three-dimensional cell–cell interactions (7). Moreover, PRF has played a role in space making for bone regeneration. PRF in a compressed membrane-like form has also been used as a substitute for commercially available barrier membranes in guided-tissue regeneration (GTR) treatment. However, the PRF membrane is resorbed within 2 weeks or less at implantation sites; therefore, it can barely maintain sufficient space for bone regeneration (8).

Lucarelli et al. (9) showed that the mechanical properties of PRF can be improved by inducing extensive fibrin network formation through increased gravitational force through high-speed centrifugation of intact platelets and fibrin in the absence of exogenous thrombin. With this procedure, platelet rich plasma (PRP) can be produced in the form of platelet-rich fibrin matrix (PRFM). The PRFM preparation process creates an ideal scaffold that contains high concentrations of nonactivated, functional, intact platelets, contained within a dense cross-linked fibrin matrix. Carroll et al. (10) have demonstrated, in vitro, that the viable platelets in PRFM released 6 growth factors in about the same concentration for the 7-day duration of their study. The PRFM membrane could be of great potential benefit in both soft tissue and bone repair.

Therefore, the aim of this experimental study was to evaluate and compare the osteogenic effects of L-PRF and PRFM in surgically created bone defects of rabbit tibia.

METHODS

The study was conducted between January 2017 and March 2017 in accordance with the Declaration of Helsinki of 1964, revised in Tokyo 2004 and was approved by the institutional ethics committee. The study included eight rabbits (weight, 2.5-3 kg). Two preoperative intravenous samples of 4.5 cm³ of blood were taken from the veins of the ear of each rabbit and the blood samples were centrifuged to produce autologous L-PRF and PRFM.

L-PRF Preparation

PRF was prepared according to the technique described by Dohan et al. (5). Immediately after 4.5 cc of blood had been drawn from the veins of the ear of each rabbit, the blood sample is centrifuged in a 5-mL sterile tube (without anticoagulant) at 400 g for 10 min. L-PRF layer is located between the red corpuscles at the bottom of the tube and acellular plasma at the top of the tube [Fig. 1 (a)]. The L-PRF was thus removed from the tube using sterile tweezers and pressed between two pieces of surgical gauze to obtain the membrane form.

PRFM Preparation

PRFM was prepared according to the manufacturer’s instructions (FIBRINET®, Cascade Medical Enterprises, and Wayne, NJ, USA). Briefly, 4.5 cm³ of blood was collected into sterile supplied citrate containing tubes. After gentle mixing, the tubes were centrifuged at 1100 g for 6 minutes to obtain PRP. Under sterile conditions, PRP was transferred into a calcium chloride-containing tube (0.25mL CaCl2 1M). The tube was gently swirled and then immediately centrifuged at 4500g for 25 minutes at room temperature. A translucent, yellow-white PRFM was recovered at the bottom of the tube (9) [Fig. 1 (b)].
Experimental groups

Eight rabbits were included in this blinded prospective study. The animals were divided into two experimental groups (Group A and Group B) \( n=4 \) animals/group. Group A animals was sacrificed after 10 days and Group B animals was sacrificed 21 days after surgery. The animals were operated under intravenous general anaesthesia. Both tibias were shaved and cleansed with 10% povidone iodine before manipulation. After a skin incision in the tibial area, the subcutaneous tissue was dissected down to the periosteum and a periosteal incision was made. In both groups, two defects with a diameter of 3 mm and 2 mm depth were osteotomized in the right tibial bone, and one defect with the same dimensions was osteotomized in the left tibial bone in the same rabbit. Sterile saline at 1500 rpm was irrigated over the drilling site during the procedure. The upper defect of the right tibia received L-PRF. The other defect on the same tibia left empty (control group) [Fig. 2 (a)]. The defect of the left tibia received PRFM [Fig. 2 (b)]. After obtaining adequate hemostasis, the periosteum and the muscle were sutured using 4/0 polyglycolic acid sutures, and the skin with 3/0 silk sutures. Antibiotics (Gentamycin 5 mg/kg, Sigma Chemicals, USA) were injected intramuscularly to all rabbits to prevent infection. All efforts were made to minimize pain or discomfort of the animals.

Bone Mineral Density Analysis

In order to measure bone mineral density (BMD), we used specimens harvested at 10 and 21 days (4 animals at each time interval). A total body dual-energy X-ray (DXA) scanner (DPX, Lunar, Madison, WI, USA) and software which was tailored for small animals (Norland, Small Animal Software, 1.0c, Lunar) were used. The operated tibia were examined by the DXA scanner. Regional BMD measurements were obtained by placing rectangular
(18 mm × 10 mm) regions of interest (ROI) on the scan images. Two ROI were positioned in the left tibia and one ROI was positioned in the right tibia. Results of BMD were expressed as grams per square centimeter.

**Histopathological examination:**

The animals were sacrificed at 10 and 21 days (4 animals at each time interval) via pentobarbital overdose (100mg/kg). The tibias were carefully dissected free from soft tissues, and hard-tissue samples were fixed in 10% neutral buffered formalin for 48 h then decalcified in 10% ethylene diamine tetra-acetic acid (EDTA). Tissue processing, including dehydration, clearing, impregnation, and embedding, was done through graded ethanol, xylol, and paraffin. Histologic sections with a thickness of 6 μm were prepared from each defect containing an intact border of the bone, and then the samples were routinely stained with hematoxylin and eosin (H&E) staining. For histological evaluation of healing process, Emery’s histopathological healing criteria were used, which are explained in Table 1. Histological examinations were performed by a single blinded investigator.

**TABLE (1) Emery’s histopathological healing criteria:**

<table>
<thead>
<tr>
<th>Tissue present</th>
<th>Score (point)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Empty cavity</td>
<td>0</td>
</tr>
<tr>
<td>Fibrous tissue only</td>
<td>1</td>
</tr>
<tr>
<td>More fibrous tissue than fibrocartilage</td>
<td>2</td>
</tr>
<tr>
<td>More fibrocartilage than fibrous tissue</td>
<td>3</td>
</tr>
<tr>
<td>Fibrocartilage only</td>
<td>4</td>
</tr>
<tr>
<td>More fibrocartilage than bone</td>
<td>5</td>
</tr>
<tr>
<td>More bone than fibrocartilage</td>
<td>6</td>
</tr>
<tr>
<td>Bone only</td>
<td>7</td>
</tr>
</tbody>
</table>

**RESULTS**

**Clinical evaluation**

All the surgical sites healed well. No signs of inflammation or infection were observed at any time point in any of the samples.

**BMD Analysis**

The PRFM group showed the highest BMD score at 10th and 21st days and the BMD score in the control group was the least (Fig. 3).

![Fig. (3) DXA analysis of tibia bone.](image_url)

**Histopathological findings**

Regarding to bone specimens collected after 10 days of surgery; control group (group I) revealed empty space (score 0) fig. 1-a. Group II treated by PRF showed activation of mesenchymal cells in the soft tissues. Bone surrounding the fracture gap differentiates into chondroblasts that make fibrocartilage and hyaline cartilage. More fibrous tissue than fibrocartilage (score 2) were seen fig. 1-b. On the other side group III treated by PRFM showed more fibrocartilage than fibrous tissue which characterized by well developed cartilaginous tissue with prominent basophilic matrix (score 3) fig. 1-c.

Histopathological findings of bone gap after 21 days of surgery; control group I showed more fibrocartilage than fibrous tissue (score 3) fig.2-a.
In group II treated by PRF revealed small foci of chondrocytes proliferated in the bony tissues. In this group, the gap was filled with trabecular bone characterized by acellular osteocyte lacunae and fibrocartilage tissues (score 5) fig.2-b. Group III treated by PRFM showed that the newly formed cartilaginous tissue in the fracture gap undergoes enchondral ossification (score 7). The healed bone gap inhabited osteocyte lacunae and delimited by osteoblasts that forming a network of bone connected to the reactive trabeculae deposited elsewhere in the medullary cavity and beneath the periosteum fig. 2-c.

**DISCUSSION**

Fibrin-based membranes could be used as scaffolds for proliferation of periosteal and osseous cells. However, their clinical beneficial effects on bone regeneration need to be thoroughly explored. Although fibrin-based membranes have longer resorption time than the other platelet concentrates (13), it has features to similar to those of natural clotting. Therefore, experimental evaluations investigating the early stages of healing would be beneficial.

![Fig. (4) Histopathological section of induced bone gap: control group showing empty gap (a), group treated by L-PRF showing more fibrous tissue than fibrocartilage (b), group treated by PRFM showing more fibrocartilage than fibrous tissue (c) (H&E X200)](image)

![Fig. (5) Histopathological section of induced bone gap: control group showing more fibrocartilage than fibrous tissue arrow (a), group treated by L-PRF showing small foci of chondrocytes proliferated in the bony tissues arrow (b), group treated by PRFM showing enchondral ossification arrow (c) (H&E X200).](image)
Histological samples at early steps in the bone regeneration process are needed to demonstrate the hypothetical beneficial effect of Fibrin-based membranes on bone regeneration. For these reasons, it is relevant to use a standardized animal. In the present study, we evaluated L-PRF in comparison to PRFM in surgically created experimental bone defects. The results showed more new bone formation in defects filled with PRFM than L-PRF and empty defects. Moreover, the BMD analysis showed higher bone density in PRFM and L-PRF groups than in the control group at 10 and 21 days evaluation period. The better results obtained with PRFM may be attributed to structural differences between PRFM and L-PRF which may be partially attributed to their differences in polymerization as a result of inducing extensive fibrin network formation through increased gravitational force during centrifugation.

The present study shows beneficial effects with adding fibrin based –membranes to surgically created defects in animal model. Better results was obtained with PRFM, however, the study had some limitations. The sample size was not large enough (4 animals in each group) to show statistically significant differences between groups and longer evaluation periods is needed to evaluate the effects on all stages of bone regeneration.

REFERENCES