INTRODUCTION

Periodontitis is a chronic inflammatory disorder that degrades the integrity of dental support tissues, which include alveolar bones, periodontal ligament, cementum, and gingiva [1]. Periodontal regeneration therapy is an alternative therapy for periodontal disease that restores tissue to its previous form and function. This technique aims to eliminate granulation and necrotic tissue at the bone defect area and thus induce healing. Bone grafts and guided tissue regeneration using membranes are common regenerative materials used in periodontal therapy [2,3].

Regenerative therapy using common regenerative materials is limited to specific types and small- to moderate-sized bone defects. A bone defect with crater defects or a vertical defect with a three-wall bony defect is suitable for the bone grafting procedure [2]. A one-wall bone defect or horizontal bone defect has less healthy cells and a large defect, causing difficulties in the ability of grafts to produce regenerative cells that are similar to or that resemble the lost tissue [4,5]. To overcome this limitation, several combinations of regenerative therapies and tissue engineering applications have been studied.

Tissue engineering techniques stimulate the regeneration of damaged but alive tissues using cells, scaffolds, and triggers (signaling factor) [6]. Isolated cells were grown in vitro to obtain a sufficient amount, and then, they were transplanted using a scaffold material to the damaged area. Cell sheet technology (CST) is one of the approaches used in tissue engineering techniques [7]. CST uses special surfaces, known as peptide structures of, that can promote cell attachment; this approach affords advantages such as simple synthesis, minimal cost, low immunogenic activity, relative stability, and tight control of conformation. The peptide structure of RGD enhances the attachment of the chitosan scaffold to the defect area or the root surface of tooth in tissue engineering regenerative therapy [10]. No previous studies have compared the application of a chitosan cell sheet and RGD-modified chitosan cell sheet for treating a large defect such as a one-wall bone defect or horizontal bone defect.

This study evaluates the regeneration outcome of CST application with and without RGD modification through a protein that could define the regeneration process. Cementum protein-1 (CEMP-1) is a specific biomarker for cementum regeneration and the main regulator in cementogenesis [11].

EVALUATION OF REGENERATIVE THERAPY USING CELL SHEET THROUGH CEMENTUM PROTEIN-1 EXPRESSION ON MACACA NEMESTRINA

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ABSTRACT

Objective: The main objective of periodontal therapy is tissue regeneration. Previous studies have identified the potential of mesenchymal stem cells to improve major periodontal defect reconstruction in bone tissue engineering. Cell sheet technology (CST), in which a cell culture is obtained from a material coated with a temperature-sensitive substrate, has been developed for the reconstruction of various tissues, including periodontal tissue. Cementum protein-1 (CEMP-1) is a 50-kDa protein that plays a crucial role in cementogenesis by enhancing the combining of cells formed by cell cementoblast.

Methods: The CEMP-1 was analyzed expression in a sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) gel electrophoresis assay.

Results: CEMP-1 expression in gingival crevicular fluid was observed using the SDS-PAGE method every week for 3 weeks. Protein band expressions on SDS-PAGE gel were identified at around 50 kDa with different thicknesses between groups. The chitosan, chitosan cell sheet, and RGD-modified chitosan cell sheet groups showed protein bands of CEMP-1 between 50 and 70 kDa at weeks 1, 2, and 3; weeks 2 and 3; and weeks 1 and 2, respectively.

Conclusion: Our results demonstrated that the application of chitosan and RGD-modified chitosan cell sheets could enhance bone regeneration, as evidenced by CEMP-1 protein expression levels.

Keywords: Periodontitis, Bone defect, Cell sheet, Cementum protein-1.

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It stimulates the attachment of differentiation cells as well as the formation of hydroxyapatite crystals by cementoblast, which is the cell that forms cementum tissue. CEMP-1 expression is limited to cementoblast and progenitor cells, that is, subpopulation cells in the periodontal ligament [11,12].

METHODS

The study design is a descriptive laboratory experiment on M. nemestrina. This study was approved by Animal Care and Use Committee (Pusat Studi Satwa Primata-IPB) Bogor. A split-mouth design was performed on four regions of the lateral incisor of one M. nemestrina. The alveolar bone was intentionally destroyed in each region to imitate a one-wall defect of chronic periodontitis with an open flap procedure. In each region, a different treatment was applied to the destruction site: Chitosan only as a control group, chitosan cell sheet, and RGD-modified chitosan cell sheet.

The cell sheet was prepared before the flap procedure using ligament periodontal cells extracted from the central incisor. A cell sheet filled with mesenchymal stroma cells was adhered to the chitosan and applied to the defect area. The chitosan group was used as a control in this study. After the treatment, gingival crevicular fluid (GCF) was collected from the gingival sulcus of the lateral incisor using some paper points every week. Paper points filled with GCF were placed in 100 µl of PBS in an Eppendorf tube and stored at −80°C in a refrigerator.

A Bradford assay was conducted to determine the protein concentration of each sample. A sodium dodecyl sulfate-polyacrylamide gel electrophoresis assay was performed to detect proteins in the sample based on their molecular weight.

RESULTS

Fig. 1 shows the results obtained using the electrophoresis gel. The chitosan, chitosan cell sheet, and RGD-modified chitosan cell sheet groups showed a protein band between 50 and 70 kDa in weeks 1, 2, and 3; weeks 2 and 3; and weeks 1 and 2 after application, respectively.

In week 1, a CEMP-1 protein band was seen only in the chitosan group and RGD-modified chitosan cell sheet group, and not in the chitosan cell sheet group.

In week 2, the protein band was seen in the chitosan group and RGD-modified chitosan cell sheet group, and it started to be seen as a thin layer in the chitosan cell sheet group. In week 3, the protein band was still seen in the chitosan group, and it was seen to become thicker in the chitosan cell sheet group; however, it was no longer seen in the RGD-modified chitosan cell sheet group.

DISCUSSION

Periodontal regenerative therapy aims to restore oral soft and hard tissues through cells, scaffolds, and/or signaling approaches to functional and esthetic oral tissues. Common periodontal regenerative materials find limited applications when dealing with large bone defects such as a one-wall bone defect. The prognosis for this condition is quite poor owing to the difficulties faced in stabilizing the graft material until regeneration starts [5,13]. An earlier study on M. nemestrina and a human mandible showed the first evidence of improved bone regeneration when using chitosan as a scaffold [4,9]. Furthermore, the use of ligament periodontal cell sheets was found to promote regeneration in animal study [14,15]. The difficulties faced in cell sheet application owing to its fragile structure could be overcome using a scaffold such as chitosan. To increase its adherence to defect areas or to a root surface, another peptide structure containing RGD was applied to chitosan [10,16]. The results obtained in this study show that the application of chitosan, chitosan cell sheet, and RGD-modified chitosan cell sheet can promote regeneration based on the expression of CEMP-1. Cementum-forming cells (cementoblasts) have the primary function of making and secreting the extracellular matrix proteins required for cementum mineralization. The finding that CEMP-1 is synthesized by cementoblast cells and by restricted periodontal ligament cell populations indicated that CEMP-1 may act as a local regulator of cell differentiation and extracellular matrix mineralization. CEMP-1 is a cementum protein that indicates cementoblast action. Its absence in the chitosan cell sheet group was probably caused by a lack of cementoblast cells in the cell sheet taken from the periodontal ligament.

![Fig. 1: Differences in cementum protein-1 band from weeks 1 to 3 in (a) chitosan, (b) chitosan cell sheet, and (c) arginineglycineaspartic acid-modified chitosan cell sheet groups](image1)

![Fig. 2: Differences in cementum protein-1 band between chitosan, chitosan cell sheet, and arginineglycineaspartic acid-modified chitosan cell sheet groups in weeks (a) 1, (b) 2, and (c) 3](image2)
cell. Its appearance in early weeks after application indicated early bone regeneration [12].

**CONCLUSION**

A CEMP-1 band could be detected in the chitosan and RGD-modified chitosan cell sheet groups at different appearance times. The application of a cell sheet may enhance healing and bone regeneration.

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