ABSTRACT

Objectives: The present study was designed to evaluate the clinical, radiographic and histochemical significance of using the mandibular tori as autogenous bone graft for treatment of intraosseous defects in patients with chronic periodontitis.

Materials and methods: Twenty-eight sites from 14 patients with chronic periodontitis were included in this study. Each patient was treated with split mouth design; one site received torus mandibularis bone graft and the other site received a full-thickness flap alone. Histopathologic assessment was evaluated on removal of torus mandibularis to evaluate its histologic structure and by the end of the study 9 month later. Clinical and radiographic parameters were re-evaluated at 3 months interval for 1 year.

Results: The results of the present study revealed significant gain in the clinical attachment level (CAL) (88.4%, 4.53 ± 0.06 mm) for torus mandibularis sites compared to (39.7%, 2.01 ± 0.04 mm) for full-thickness flap. Moreover, there was a reduction in the probing pocket depth (PPD) of (75.4%, 5.75 ± 0.12 mm) for torus mandibularis sites and (49.6%, 3.73 ± 0.14 mm) for sites treated with a full-thickness flap only; CAL and PPD differences were significant at p-value ≤0.01. Concomitantly, significant radiographic increase in the bone height and density were recorded in the test group.

Conclusion: The use of mandibular tori as autogenous bone graft could provide benefits as a periodontal therapeutic modality and enhance regenerative potential of periodontal intraosseous defects.

Keywords: Autogenous bone grafts, Chronic periodontitis, Mandibular tori.

INTRODUCTION

The optimal goal of periodontal therapy is to resolve inflammation and control disease progression where successful regeneration of a new attachment apparatus replaces the destroyed one. Unfortunately, this has long been an elusive goal for more than a century. Regeneration is defined as the reproduction or reconstitution of the lost structures of the periodontium restoring its architecture and function. Histological evidence of periodontal regeneration must include new alveolar bone, reestablishment of periodontal ligament and cementum. Bone grafts and synthetic substitutes have been utilized to achieve these therapeutic goals.

Different treatment modalities have been attempted to restore the lost periodontal apparatus with varying outcomes. Extaoral or intraoral autogenous bone grafts showed the best regenerative results histologically. Demineralized freeze-dried bone showed similar results. Consequently, autogenous bone grafts have been widely used for regenerative procedures. A variety of intraoral sites have been utilized for harvesting, including extraction sockets and bony exostoses. Several studies have reported on the clinically successful use of autogenous bone grafts harvested from the intraoral sites in the treatment of intrabony defects. These authors show that regenerative procedures including autogenous bone grafting was superior to surgical debridement alone. A resolution of more than 50% has been shown when using intraorally harvested bone.

Bone exostoses, are localized bony protuberances that arise from the cortical plate and sometimes from the
spongy layer, with no pathological significance. Torus palatinus and torus mandibulae are two of the most common intraoral exotoses. The other types of exotoses as buccal or palatal exostoses are less commonly encountered. Torus mandibulae, usually found bilaterally, are localized on the lingual side of the mandible in the premolar region superior to the mylohyoid ridge, first described in 1908 by Fürst, 1908.23

Mandibular tori may limit tongue space occasionally causing phonetic problems. Therefore, surgical removals of tori are occasionally required for prosthetic reasons. The surface of removed exotoses which are rich with proliferating osteoblasts as well as fibro-cartilaginous capsule that might contain undifferentiated osteoblasts attracted our interest in utilizing it as autogenous bone graft to restore the periodontal bony defects. The present study protocol was designed to evaluate and compare the treatment outcome, clinically, radiographically, and histochemically of the two treatment modalities, full-thickness flap with and without autogenous bone grafting from mandibular tori; in the treatment of intrabony osseous defects.

MATERIALS AND METHODS

Fourteen patients (8 females and 6 males; age range: 37 to 48 years with an average age of 45.3 ± 3.81 years) with chronic periodontitis (localized form, less than 30% involved sites) and having torus mandibularis were selected to participate in this randomized, controlled and parallel clinical design study. Patients were selected from the outpatient clinic, College of Dentistry, University of Dammam, Kingdom of Saudi Arabia. Patients with any systemic illness known to affect the outcomes of periodontal therapy, compromised immune system, pregnancy and/or lactation, smoking or the use of other tobacco products, those taking drugs known to interfere with wound healing, alcohol use habits, allergy or sensitivity to any medication to be used in the study, and those with unacceptable oral hygiene after the reevaluation of phase I therapy were excluded from the study.

The inclusion criteria were the presence of infrabony defect (Vertical bone loss) with loss of CAL of at least 5 mm and PPD of not less than 6 mm. Twenty-eight sites from fourteen patients were selected using a split mouth design for each patient which was randomly determined through a coin toss. One site (group I) received a full-thickness mucoperiosteal flap and the infrabony defect was filled with Torus mandibularis bone graft. The other site (group II) received a full-thickness flap alone. Each patient was prepared for surgery with an initial phase of therapy which included oral hygiene instructions and scaling and root planing. Approximately 4 weeks after the initial therapy, the patients were reevaluated to assess clinical parameters and plaque control. All patients were required to achieve a good oral hygiene (less than 20% O’Leary plaque index) done by scaling and root planing and oral hygiene instructions prior to progressing to the surgical phase of therapy.

Full explanation of the research objective was explained to all patients included in this study, and then they were asked to sign a written consent demonstrating their approval of the procedure and publication of its results. In addition, this research was approved by the ethics committee, College of Dentistry, University of Dammam. The following clinical parameters were assessed at the baseline, 3, 6, 9 and 12 months postoperatively using the same periodontal probe (NUC-15, Hu-Friedy, USA); the gingival index (GI; Löe and Silness, 1963),20 pocket probing depth (PD) and clinical attachment level (CAL; Ramfjord, 1967).22 For 1 year, the clinical and radiographic parameters were re-evaluated at 3 month intervals. The measurements recorded at 6 points (recording the deepest site) around the treated teeth.

Bone Graft Collection

All procedures were performed under sterile conditions. Prior to surgery, all patients rinsed with 0.12% chlorhexidine for 30 seconds. A full-thickness mucoperiosteal flap was elevated for torus mandibularis access and for bone graft collection (Fig. 1A). Vertical releasing incisions (one distal or mesial releasing incision with minimal length was elevated to expose the mandibular tori) were performed if necessary for better access. A surgical bur #701 and 702 (FGOS) in a slow speed handpiece was used under copious irrigation to remove the torus mandibularis. Bone grafts were collected with a bone trap according to the method described by Sivolella et al28 (Omniasurg ASP 100; Omnia srl, Fidenza, PR, Italy). The trap filter was equipped with a removable internal mesh with a pore diameter of 300 µm. Two distinct systems were used for aspiration and bone collection. One system was used for the control of saliva, bleeding, and was kept at a distance of 1.5 cm from the osteotomy site. The other system was sterile and disposable, and comprised a filter for collecting bone chips and a plastic suction tube. The latter was held as close as possible to the osteotomy site in order to collect the bone debris and reduce the aspiration of saliva (Fig. 1B). The collected tissues were placed in a sterile bone well containing sterile saline solution at room temperature and covered with a cap in order to minimize the risk of contamination. After preparing of the receiving site, the excess of saline solution was removed and only the collected materials were used to fill the intraosseous defects.

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in-group I. The flaps were adjusted by gingivectomy and closed with a resorbable type surgical thread (3-0, Manufacture; Johnson and Johnson Intl. XHGHC0) and simple interrupted sutures.

Periodontal Surgical Procedure

Intracrevicular incisions, full-thickness muco-periosteal flaps were raised vestibularly for defect access and granulation tissue removal on both sides of the teeth (Fig. 1A). For a better access or a better closure of the surgical sites, vertical releasing incisions (mesial and distal to the surgical site extended below the mucogingival junction) were performed. All granulation tissues were debrided from the root surfaces and the defects. One site was treated with torus mandibularis as autogenous bone graft and the other site was treated only with a full-thickness mucoperiosteal flap. The graft was delivered to the bony defect with a spatula and added in incremental fashion. Light pressure was used to maintain space between the graft particles to allow revascularization of the site. The defect was filled or slightly overfilled (Fig. 1C) to maximize regeneration without compromising flap closure or vascular supply. The flaps were closed and sutured with interrupted sutures (Fig. 1D). All patients received antibiotics for 1 week (Amoxicillin 500 mg/3 times/day) and rinsed with 0.2% chlorhexidine solution twice daily. The sutures were removed 14 days postoperatively. All patients have been re-called after 3, 6, 9, and 12 months for the clinical and radiographic assessment.

Histochemical Analysis

At the time of torus mandibularis removal, a small sample of the collected materials was fixed in 10% formalin, routinely processed and embedded in paraffin. Serial sections were cut at a thickness of 5 µm. Sections were stained with hematoxylin (HE) and Vankossa histochemical stain as described elsewhere.29 During the 8 month postoperative follow-up period, one case of the tested group involving teeth #2, 3, 4 and 5 demonstrated severe pain in the un-restorable tooth #2 due to deep carious lesion with pulp exposure. After consultation with endodontist it was decided to extract the tooth and approved by the ethics committee, College of Dentistry, University of Dammam. The research team surgically removed tooth # 2 including its proximal alveolar bone (demarcated and removed by Pizotome) and was fixed in 10% formalin, decalcified in 5% formic acid, routinely processed, sectioned at 5 µm thickness and subjected to histochemical staining using HE and

Figs 1A to D: (A) A sulcular incision full-thickness flap reflected, showing intrabony defects at the upper posterior teeth, (B) Torus mandibularis removed with surgical bur, and the bone grafts collected with the bone trap, (C) Torus mandibularis bone graft fills the intraossous defects and (D) Suturing the flap with interrupted sutures
von Kossa that help to demonstrate the histopathologic background of our radiographic findings.

**Standardized Radiographs**

All the standardized radiographs were taken with the Rinn film holder. This device consists of film holder into which the dental radiograph fits, a plastic plate, which the patient bites on, and a rod connected to the film holder that protrudes from the patient’s mouth to allow parallel alignment of the X-ray tube.\(^{30}\) To ensure that the patient’s bite was reproducible in the standardized group of radiographs, an impression material (Henry Schein Blu-Bite Vinyl Polysiloxane Derivative Registration Material: Henry Schein Inc, Melville, NY, USA) was placed on the plastic bite block. The patient was instructed to bite on the bite block while holding his or her teeth in their posterior position (centric relation). Once the impression material set, the film holder was removed from the mouth. The film was then added; the patient set his or her teeth in the registered bite, and the X-ray was taken. Identical exposure parameters were used at all examinations, and the examination film was automatically processed. All periapical radiographs were digitized and saved in a tagged image file format (TIFF). Then, the bone density and marginal bone levels were measured by using Image J software.\(^ {31}\)

**Data Analysis**

Clinical data were expressed as means ± and standard deviation (SD) for each variable and examined intervals. The change in clinical parameters from baseline to 3, 6, 9, and 12 months after surgery within each group were compared using the paired t-test. One-way analysis of variance was used to test the difference between groups. The level of significance for analysis was set at \(p \leq 0.05\).

Change in radiographic findings was done to evaluate increase in alveolar bone level (measured at significant level of \(p < 0.001\)) and bone density (measured at significant level of \(p < 0.05\)), parallel to the clinical evaluation, to estimate the progress in bone gain for each group. For clarification and optimization purposes, baseline radiographs for each case was digitally overlapped to 12-month postoperative one, using adobe Photoshop CS4 software, and the difference was measured as for bone density and height by image J software and the average was calculated and compared.

**RESULTS**

All patients completed the study without complications and recalled for evaluation at 3, 6, 9 and 12 months. The results of the present study showed no statistically significant differences in the clinical parameters measured at the baseline between the treated sites. With regard to the gingival index, statistical significant changes were observed after treatment with torus mandibularis (group I) comparing to the pretreatment period (baseline 2.1 ± 0.21, after 12 months 0.02 ± 0.03). In addition, there was a statistically significant change following treatment with torus mandibularis (group I) compared to sites treated with a full-thickness flap alone (group II) (0.02 ± 0.03, 0.12 ± 0.03 respectively) (Table 1).

The present work illustrated a comparison between both groups with regard to probing depth and clinical attachment level. There was a gain in CAL of 88.4% (mean value at baseline of 5.12 ± 0.52, at 12 months 0.59 ± 0.04 mm) for torus mandibularis sites, and 39.7 % (mean value at baseline of 5.07 ± 0.41, at 12 months 3.06 ± 0.06 mm) for full-thickness flap alone (\(p \leq 0.01\)). With respect to probing depth, there was a reduction in probing depth of 75.4% (mean value at baseline of 7.62 ± 0.42, at 12 months 1.87 ± 0.16 mm) for torus mandibularis sites, and 49.6% (mean value at baseline of 7.51 ± 0.31, at 12 months 3.06 ± 0.31 mm) for sites treated with a full-thickness flap alone (\(p \leq 0.01\)) (Table 1 and Fig. 2).

A radiographic comparison of baseline and 12 months scores (Figs 3D and E) of the experimental group was

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<th>Table 1: Changes of the clinical parameters measurement in the test group (GI) and control group (GII)</th>
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<td><strong>Test group</strong></td>
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<td>GL (scores)</td>
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<td>NS: Nonsignificant differences; *: Significant differences</td>
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done by overlapping the two images (Fig. 3F). Similar procedure was done for the control group (Figs 3A to C). Our results show increase in alveolar bone level and density in experimental group (Fig. 3F) compared to the control group (Fig. 3C). Differences between and the test and control groups for bone density was (17%) as expressed in percentage of difference in pixel/mm² (Fig. 4) that was significant at p < 0.05.

Histopathologically, HE staining supports our hypothesis that the torus mandibularis contains active osteoblasts that were mainly located at the surface of preoperative sample as shown in (Fig. 5A). Higher magnification (Figs 5B and C) demonstrates the active osteoblasts with active production of bone matrix that stain deep violet in HE stain when compared to the main eosinophilic structure of lamellar bone of torus.

The proximal alveolar bone of the extracted tooth #2 at 9th months postoperatively demonstrate clumped osteoblasts surrounding the eosinophilic zone on HE staining (Fig. 6A) interrupted by violet materials presenting active osteogenesis that focially stain black with van Kossa shown in (Fig. 6B) representing mineralization process of the newly formed bone.

DISCUSSION

The ambitious goal of periodontal therapy is to restore the periodontal attachment apparatus to its prediseased state. The proper placement of graft materials for the infrabony defects are one mode of therapy that attempts to restore the lost periodontal attachment apparatus. The bone graft material provides regeneration through inductive or conductive processes. The conductive graft act as scaffold to support the new tissue growth and replaced by the host tissue. The inductive graft stimulates the host tissues to regenerate the lost structures.

All medical devices that come into a direct contact with living tissues must be composed of biologically compatible materials. They must not cause either systemic or local toxicity and should not be carcinogenic or genotoxic. In addition, they must not affect the blood clotting mechanisms and should not induce immune responses.

The present study was designed to evaluate the clinical significance of using mandibular tori as a graft material in the treatment of intraosseous defects in patients with chronic periodontitis.

The present study focused on deep intrabony defects characterized by clinical attachment level and probing pocket depth not less than 6 mm. This is in agreement with the other investigations, which demonstrated a positive interrelationship between defect depth, clinical attachment level gain and osseous fill after the indicated periodontal therapy. In addition, each patient included in the present study served as their own control by the use of split mouth techniques to avoid the individual variations as a vulnerable point in such studies. Moreover, this method is limited to patients only having torus mandibularis.

It is important to note that, several factors might dictate important effects on the outcome of the regenerative therapy. These factors are: age, patient’s gender, oral hygiene, compliance to postoperative instructions and care, surgical manipulation and perfect, defect characteristics and severity. However, in a split-mouth design these factors have a similar influence on both therapeutic modalities to be compared.

A sterile and disposable aspiration set with a plastic suction tube and filter equipped with an internal mesh with a pore diameter of 300 µm was used to collect bone fragments in this study. The results of the present study recorded that there were no problems with filter blockage during collection. This is in agreement with a study
performed by Sivolella et al.,\textsuperscript{28} who demonstrated that 300 µm meshes are capable of catching most of the suctioned material, without the problems of fragment blocking of the aspiration system from obstruction of the filter. In addition, by the use of a two distinct aspiration tubes for bone chip collection and control of salivation and bleeding, the bacterial contamination can be reduced. A study by Etchenson et al\textsuperscript{35} demonstrated that osseous coagulum collected in bone traps had potential for bacterial contamination. They also suggested methods for decontamination. Their results indicate significant bacterial contamination of osseous coagulum. They have shown that bone traps used with stringent surgical protocols that include segregated suction tips and pre-procedural chlorhexidine rinsing, with tetracycline will have a significant, however, incomplete influence on the reduction of bacterial contamination of osseous coagulum. The same protocol as the one used by Etchenson et al\textsuperscript{35} was used in this study. The initial therapy period was 4 weeks. In the Etchenson study, the initial treatment phase was not utilized for procuring graft material. Moreover, Tezulas et al\textsuperscript{36} demonstrated the decontamination of autogenous bone grafts collected during dental implant site preparation by using clindamycin or
Fig. 4: Diagram show, the bone density (in pixel/mm²) following treatment with mandibular tori (group I) comparing to a full-thickness flap alone (group II).

chlorhexidine solutions. Their results showed that both agents effectively decontaminated the collected bone particles. In this study, the protocol was utilized two surgical suction tips: one to aspirate saliva and blood and another to collect osseous coagulum from the osteotomy site.

The gingival index remained reduced when compared to the baseline, throughout the study for both groups, with a statistically significant difference between the groups.

It was suggested that the clinical objectives of periodontal regeneration include: increased bone height, gain in clinical attachment levels, probing depth reduction, improved esthetics and long-term maintenance of teeth in health and comfort.37,38 In the present study, a significant reduction in pocket probing depth and clinical attachment gain was obtained when using of mandibular tori as autogenous bone graft. These results are in accordance with the study performed by Scott and Ganz39 which claimed that mandibular tori could be used as the donor sites of autogenous grafting materials to fill the remaining defects around implants installed in extraction wounds. In addition, the results of the present study showed a significant reduction in pocket depth and attachment level gain when using mandibular tori compared to a full-thickness flap alone.

Contrary to the accepted concept, our results show that mandibular tori can be used successfully in the treatment of advanced periodontal diseases. Histochemical analysis of torus bone indicates the presence of active osteoblasts mainly on the bone surface. A study by Scott...
Mandibular Tori as Bone Grafts: An Alternative Treatment for Periodontal Osseous Defects

In general, successful periodontal therapy means regeneration and/or maintenance of periodontal support. Complete pocket elimination by regeneration or pocket resolution is impossible. The results of the present study showed that it was possible to reduce the probing pocket depth in sites treated with mandibular tori, thus allowing more effective maintenance therapy. The present study suggested that mandibular tori autogenous bone graft could provide benefits as a periodontal therapeutic modality. Furthermore, the radiographic bone gains recorded after treatment with mandibular tori confirmed the clinical observations. The present study show that the use of mandibular tori as bone grafting material may provide additional benefits in the treatment of advanced periodontal disease; and enhance the regenerative potential of periodontal intraosseous defects.

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