Maxillary Sinus Augmentation Using a Titanium Mesh: A Randomized Clinical Trial

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Abstract

BACKGROUND: Various attempts have been implemented using different materials and techniques to augment the maxillary sinus floor for prospect dental implant positioning.

AIM: This contemplate was conducted to assess the osteogenic capability of the maxillary sinus in a two-step sinus membrane elevation using titanium mesh to keep the formed space to place dental implants in atrophic ridges.

MATERIALS AND METHODS: Titanium micromesh was customized and positioned into the sinus on one side to preserve the elevated membrane in position. On the other side xenograft was applied. Instant and 6-months postoperative cone beam computed tomography (CBCT) was done to assess the gained bone height and density. Bone core biopsies were obtained during implant placement for histological and histomorphometric evaluation.

RESULTS: The average bone height values increased in both groups. Meanwhile the average bone density value was higher at the graft group than the titanium mesh group. Histological and histomorphometric evaluation presented the average bone volume of the newly formed bone in the graft group which is superior to that of the titanium mesh group.

CONCLUSION: The use of the titanium micromesh as a space-maintaining device after Schneiderian membrane elevation is a trustworthy technique to elevate the floor of the sinus without grafting.

Introduction

Dental implants have been used widely in partially or completely restoring missing teeth. Maxillary sinus pneumatization and defective bone quality and quantity are amid the factors that limit maxillary ridge rehabilitation using dental implants [1]. Various adjunctive surgical techniques have been developed to improve the quality of bone and the surrounding soft tissues and conquer the bone volume deficiency for successful placement and support of dental implants [2].

Maxillary sinus augmentation has been first presented by Tatum meanwhile Boyne and James were the first to utilize autologous bone grafts with reported long-term follow-up [3]. Sinus augmentation is performed by grafting the surgically produced partition between the superiorly repositioned sinus membrane and the bony sinus floor with proper materials [4]. Numerous grafting materials that can be implemented are autogenous bone, alloplastic and allogenic materials [5]. Autogenous bone is regarded the most superior material for sinus floor augmentation in terms of histological performance. However, donor area morbidity and graft volume loss are amid the major disadvantages of autogenous bone, which directed the majority of efforts towards using bone substitutes and new grafting techniques [6]. There after evolved the idea of maxillary sinus membrane lifting without the application of any bone grafts; which was first presented by Lundgren and co-authors [7].

During the past decade, several contemplates presented the cautious elevation of the Schneiderian membrane followed by concurrent placement of the root form implants to act as tent poles under the membrane [8-14]. Long-period studies using similar technique showed high implant survival rates with marginal bone resorption around the positioned
implants within the acceptable range [15-17].

Attempts were done at introducing space-making devices under the elevated maxillary sinus membrane when no implants could be concurrently placed. Recently, titanium mesh has received growing attention as many documented predictable and consistent results with the use of this material [18]. Many authors of the use of titanium mesh have been recommended. Titanium mesh provides superior space preservation, a basic prerequisite for any bone regeneration procedure. In addition, the pores within the titanium mesh are thought to play a significant role in maintaining and preserving blood supply to the grafted defect. However, the process of new bone formation using titanium mesh is not fully explained. In addition it has also been suggested that the use of titanium mesh could provoke the flap dehiscence and subsequent graft failure. The lateral window approach has been regarded as the classical technique for augmenting the maxillary sinus floor especially when the initial alveolar bone height does not insure primary stability of the implant. It can be done either in a single step with immediate implant placement or in two stages with delayed implant placement, depending on the available quantity and quality of remaining bone at the atrophic ridge site [19, 20].

Therefore the rationale of this study is to evaluate the osteogenic capability of the maxillary sinus in a two-step sinus membrane elevation using titanium mesh to maintain the created gap after membrane elevation without the use of any graft or space-filling material for future site preparation for the positioning of dental implants in severely atrophied posterior maxillary ridge.

**Materials and Methods**

The study was a double blinded study using a split-mouth design. Four patients (2 females and 2 males), with a total of eight sinuses which were augmented for future implant placement. The age range of the patients was between 33-55 years with a mean of 40.5 ± 6.65 years. All patients were recruited from the outpatient clinic of the Periodontology Department, Faculty of Oral and Dental Medicine, Cairo University and the outpatient dental clinic at the National Research Centre, Giza. The study protocol was approved by the Ethical Committee at the National Research Centre and an informed consent after all the study procedures where explained to each patient was obtained. The selected patients were systemically free with complete absence of any infection or pathosis in the bone or covering mucosa at the edentulous site or major bony sepsa as evident on the panoramic radiograph. The distance between the crest of the ridge and the floor of the sinus in areas planned for future implantation had to be less than 5 mm with proper antro-posterior, transverse and vertical dimensions at centric occlusion. Good oral and periodontal health is also required in the selected patients. Patients who had previous surgery done in the maxillary sinus were excluded. Also pregnant females and smokers were not included. Each patient was interviewed in order to obtain a comprehensive history, including full medical and dental history.

Patients were assigned to one of the two treatment groups using computer generated randomization table (Research Randomizer computer software, version 4.0, Pennsylvania, USA) with a drop out ratio 10-15 %.

Allocation concealment was obtained using sealed coded opaque envelopes containing the treatment to the specific subject. The sealed envelope containing treatment assignment was opened at time of the surgery.

Four patients were selected indicated for bilateral external sinus elevation procedure, so that a total of 8 sinuses were enrolled in the present work. A study of a continuous response variable from matched pairs of study subjects with sample size calculation was planned. Prior data indicate that the difference in the response of matched patients was normally distributed with standard deviation (SD) 2.47. If the true difference in the mean response of matched patients is 0.2, we needed to study 8 sides to be able to reject the null hypothesis that this response difference was zero with probability (power) 0.09. The Type I error probability associated with this test of this null hypothesis was 0.05.

**Preoperative procedures**

A comprehensive intraoral examination of the remaining teeth and soft tissue condition at the edentulous ridge was performed as well as evaluation of the inter-arch space, the width of the alveolar bone and the thickness of the covering tissue. Preoperative digital panoramic radiograph was done for each patient as a primary survey with a magnification 1:1.

For the selected patients, preoperative cone beam computed tomography (CBCT) scans were also performed while the patients were wearing a radiographic/surgical stent to precisely measure the bone height and density at the area considered for surgery and future implantation. Location of the exact mesio-distal dimension of the lateral-window osteotomy during surgery and the bone thickness was done in order to determine the amount of drilling to avoid perforation of the Schneiderian membrane.

An alginate impression was taken for the chosen patients, and then a diagnostic wax up was made. A maxillary partial denture was fabricated prior to the surgery, to point out accurately the proposed area that requires sinus elevation. This denture was
checked properly with the opposing teeth to check on the occlusion.

All patients received a single dose of IV antibiotic (ampicillin/sublactam 1500 mg) in the form of Unasyn 1500 mg vial, Pfizer, USA) and another IV corticosteroids injection (Dexamethasone sodium phosphate 8 mg/2ml, Organium laboratories, UK) half an hour before the operation as prophylactic measures.

**Surgical procedures**

The surgery was done under local infiltration anesthesia consisting of 2% lidocaine hydrochloride and 1: 100000 epinephrine administrated in the buccal vestibule and the palatal mucosa opposite to the site of surgery. A lateral sinus floor elevation was performed according to Boyne and James [5] (Figures 1-3). After sinus elevation, maintenance of the newly created space was accomplished using a xenogenic bone grafting material for the control group or graft group (Group A) as seen in Figure 4 and a titanium (Ti) mesh for the intervention group or the Ti mesh group (Group B) as shown in Figure 5 without adding any bone filling material.

**Figure 1: Mucoperiosteal flap reflection**

The mucoperiosteal flap was performed using a mucoperiosteal elevator in the form of three lines in order to expose the lateral wall of the maxilla. It started from the crestal incision positioned slightly palatal to the crest of the ridge and then extended mesially and distally to facilitate the flap reflection. This was done to delineate the outline of the rectangular osteotomy and was guided by the radiographic surgical stent. Exact measurements were taken from the preoperative CBCT radiograph and then transferred to the bone to determine exactly the site of the surgery.

**DASK (Dentium Advanced Sinus Kit, California, USA) drill #4 or #5 was used to prepare a lateral sinus window using light pressure (800-1200 rpm) and rotating strokes.**

**Figure 2: Drilling into the bone**

The inward reflection of the osteotomy and...
the elevated membrane resulted in an empty space with the wall of the sinus as its medial border and the antral membrane, while the upper side with the elevated lateral wall and antral membrane, and inferiorly bordered by the bony floor of the sinus. Before moving to the next step, the integrity of the sinus membrane was ensured using the Valsalva maneuver, which is performed by pinching the patient's nose, thus forcing a moderate exhalation, where the sinus inflates and deflates thus ensuring no perforations in its walls [21].

For the intervention group or Ti mesh group (Group B), the width of the osteotomy was measured and a foil template was trimmed to fit exactly into the created space in an L-shape figure so that one side was resting on the lateral wall of the bone and the other side pushed inside the cavity supporting the membrane at its new level. A 0.1 mm thickness micro titanium mesh (Leibinger, Stryker Co., Geneva, Switzerland) was then cut and trimmed the same size as the template, and its sharp ends were smoothed to avoid tearing the sinus membrane. The titanium mesh was tailored in an L-shape so that the long arm is inserted inside the sinus cavity to support the Schneiderian membrane at its new level and the short arm is fixed to the lateral sinus wall with two micro screws (1.5 mm) above the site of the osteotomy. The osteotomy was then covered with a collagen membrane and the flap was readapted and sutured in its original position using also the 3-0 silk suture in an interrupted pattern.

**Postoperative instructions**

Patients were instructed to apply ice packs for 10 minutes every 30 minutes for the first 24 hours. Meticulous oral hygiene measures were pertained, including: tooth brushing using soft brush carefully and chlorhexidine gluconate 0.1 % mouthwash (every 8 hours for two weeks). Avoidance of any negative or positive pressure like blowing the nose, drinking with a straw or even spitting hardly, for the first 24 hours were explained to the patients. Medical regimen included: a single dose of long acting corticosteroids: Methylprednisolone acetate 80 mg/ml I.M. immediately postoperative. An anti-inflammatory analgesic: Ketolactromethamine 30 mg/amp/2ml I.M. injection every 12 hours for the first 24 hours. Oral antibiotics: Ampicillin/sulbactam 375 mg tablets (every 8 hours for 10 days) + Clindamycin 150 mg capsules (every 6 hours for 4 days). Nasal decongestants: Oxymetazoline HCl 0.25% nasal drops (every 8 hours for 7 days). Systemic decongestants: Pseudoephedrine HCl 60 mg + Triprolidin HCl 2.5 mg tablets (every 8 hours for 7 days). Chlorohexidine Gluconate 0.1 % mouthwash (every 8 hours for two weeks) was prescribed. Sutures were removed after 7-10 days following the surgery.

**Clinical assessment**

Regular clinical assessment was done for the patients after 48 hours, 1 week, 2 weeks, 1 month and then monthly until 6 months postoperatively evaluating the wound for signs of hematoma, bleeding, infection or even membrane exposure. The patients were examined also for signs and symptoms of sinusitis. Each patient was evaluated immediately after the surgery, 3 and 6 months post-operatively for the following: discomfort, tenderness and pain.
Radiographic assessment

Radiographic follow up was done using CBCT preoperative and immediately postoperative as shown in Figure 6 and 6 months postoperatively as presented in Figure 7 and 8 to evaluate the formation of new bone. The patients submitted to this study were evaluated using Next Generation Scanora 3D CBCT scanner (Soredex, Tuusula, Finland). Images were taken prior to surgery, immediately postoperative and 6 months postoperatively. After acquisition, data were exported and transferred in a special format to be secondarily re-evaluated for height and density measurements, where dental software (version 5.3; Anatomage, San Jose, CA, USA) was used for that purpose. The software was used to determine the pre- and post-operative bone height after bone grafting and titanium mesh application as well. Height was measured in mm.

The density of bone was measured in Hounsfield Units (Hu) at the graft site and the area inferior to the titanium mesh application were evaluated in reference to the native bone density in the alveolar ridge. Series of steps were followed to guarantee the standardization of measurements for both bone graft and titanium mesh position. At the software, section module was chosen; which presented the data in the axial, coronal and sagittal views.

The measurements were oriented in a certain position to ensure standardization as the sagittal line was made parallel to the alveolar ridge; hence, coronal line was automatically obtained in the buccopalatal orientation. At the coronal view, the mesial end of the bone graft was determined, and then the height of graft material and alveolar bone as well, were taken, and then, at each slice till the distal end of the graft material posteriorly same measurements were taken. The slice thickness was set at 0.25 mm in thickness and 1 mm interval between each slice. The mean height of the bone graft was calculated, and the mean height of the alveolar ridge as well. For titanium mesh side, same steps were repeated with additional orientation step, where the axial line was made parallel to the mesh to ensure recording of the true vertical height inferior to the mesh at the coronal view.

Histological Assessment

At the time of implant placement, core biopsies were retrieved, guided by the same radiographic-surgical stent used in the surgical procedures. The specimens obtained were immediately fixed in 10% buffered formalin for 1 week, then decalcified and processed according to a standardized protocol of Ethylene diaminetetraacetic acid (EDTA) formic acid combination. Then, specimens were embedded longitudinally into paraffin blocks and oriented in a standardized way for labeling.
and differentiating the newly formed bone from the native bone end. Blocks were cut into longitudinal 5 mm-thick sections using a manual rotary microtome (RM 2135 microtome, Leica, Heidelberger Straße, Nussloch, Germany) and stained with Mayer's hematoxylin and eosin stain (H&E) and Masson Trichrome stain for histological and histomorphometric analysis.

**Histomorphometric Analysis**

All the stained sections were examined with an Olympus CX20 (Olympus, Shinjuku-ku, Tokyo, Japan) microscope attached to a camera and a computer. For each of the native and newly formed bone specimens, the most representative five fields per specimen were captured using magnification (x100). Images of the slides were taken and saved as figure files; the image analysis was done with an image analyzer (Leica Qwin 500, LEICA Imaging Systems Ltd, Cambridge, England) computer system using the Image J software (v. 1.45e, National Institutes of Health, Bethesda, MD, USA) at the Pathology Department, National Research Centre, Cairo. The bone volume (bone area fraction) was measured for each image. For each sinus, the mean bone volumes of the native and newly formed bone were calculated for data analysis. Area fraction (µm²) of native and newly formed bone specimens were measured in five random fields per group and data obtained as mean area and standard error.

**Statistical Analysis**

The mean ± standard deviation (SD) values were calculated and presented for each group. Data showed parametric (normal) distribution. Paired-wise sample t-test was used to compare between related samples while independent-sample t-test was used to compare between unrelated samples. The significance level was set at $p \leq 0.05$. Statistical analysis was done using IBM® SPSS® Statistics Version 20 for Windows (SPSS Inc., Chicago, IL, USA).

**Results**

The current contemplate was carried out on a total of 4 patients with 8 sides indicated for open sinus elevation surgery for future implant placement using the split mouth study design. Patients were randomly allocated into two groups. Their ages ranged from 35 to 55 years old with mean of (40.5 ± 6.65) years. The study included 4 males and 4 females. (Group A) the control or graft group which consisted of 4 sides indicated for external sinus elevation surgeries with xenogenic bone graft augmentation. (Group B) the intervention or Ti mesh group consisted of the other 4 sides indicated for external sinus elevation surgeries using the titanium mesh for space-maintenance. All the height and density measurements were taken from the CBCT with special software preoperative, immediately postoperative and after 6 months.

**Clinical results**

Wound healing was normal in all patients and the post-operative follow up period went uneventful without any signs of infection, suppuration, mucositis, abnormal bleeding, significant hematoma or flap dehiscence. Pain and minor swelling of gingival mucosa was noted in all patients that were completely resolved by the fifth day post-operatively. The membrane rise and mesh fixation procedures were done with no sinus membrane tears. After the second step surgery, a whole number of 14 implants were positioned, with primary steadiness gained in all the operated sinuses.

**Radiographic results**

The postoperative CBCT showed opacifications below and surrounding the titanium mesh obvious in both the cross-sectional and panoramic views as seen in Figure 7 and axial view in Figure 8, which indicates the formation of a blood clot inside the formed space.

Six months postoperatively the CBCT showed substantial amounts of radio-opacities indicating new bone formation. In various cuts the bone did not plug the whole volume below the mesh, forming voids. The line of separation between the newly formed bone and the native bone could be recognized in nearly all the examined views.

**Bone height results (in mm)**

The difference between pre and postoperative bone height in both control and intervention groups was statistically significant ($p = 0.005$). For group (A) a statistically significant difference ($p = 0.001$) was found between preoperative and postoperative bone height showing a mean bone height of $3.39 \pm 0.89$ preoperatively and a mean bone height of $12.58 \pm 2.01$ postoperatively. In group (B) the mean bone height was $3.53 \pm 1.02$ preoperatively and $11.46 \pm 2.78$ postoperatively. Although the postoperative bone height of the control group was higher than the intervention group, no statistically significant difference was seen between both groups ($p = 0.54$), where results of the mean bone heights were $12.58 \pm 2.01$ and $11.46 \pm 2.78$ in the control and intervention groups respectively as seen in Table 1. Calculating the percentage of change revealed that the amount of bone height gain in the control group (73.05%) was
Bone density results (in Hu)

There was a statistically significant difference between the native and postoperative mean bone densities in both control and intervention groups ($p = 0.03, 0.04$) respectively. The mean bone density of native bone before titanium application was $265.00 \pm 14.14$ and following titanium application the density at the site reached $182.00 \pm 37.31$, while the mean bone density of native bone before bone graft application was $345.25 \pm 51.86$ and reached $246.75 \pm 43.29$ at the site of bone graft insertion. There was no statistically significant difference in bone density between (Post-Titanium) ($182.00 \pm 37.31$) group and (Post-Bone Graft) ($246.75 \pm 43.29$) group where ($p = 0.06$). The amount of bone density gain in relation to the native bone density values in group (A) was 71.5% and in group (B) it was 68.7%.

Histological results

Clinical explanation of the obtained core biopsies showed that the color of the newly formed bone was coral pink, as compared with the white color of the native bone. The length of the cores was nearly of the same length as expected from the 6 months postoperative CBCT.

Osteocytes were unevenly oriented within the bony matrix. Osteoblastic rimming could be seen delineating the narrow spaces. Small to moderate areas of trabecular bone with clearly seen lamellae adjoining narrow marrow spaces were seen, with few inflammatory cells. Immature formed bone was also observed.
In the Ti mesh cases, the native bone showed mature trabecular bone, evidently seen lacunae of osteocytes and newly formed bone.

**Histomorphometric analysis results**

The bone area of the native bone of grafted cases ranged from 26.68% to 41.85% with a mean value of 36.7 ± 5.36%, meanwhile the bone area of the newly formed bone ranged from 8.11% to 26.96% with an average value of 13.59 ± 3.53%

In the Ti mesh cases the bone area of the native bone ranged from 28.38% to 44.2% with a mean value of 34.81 ± 2.51%, meanwhile the bone area of the newly formed bone ranged from 8.11% to 8.31% with a mean value of 5.81 ± 0.73%.

![Figure 12: Photomicrograph 6 months postoperative in the Ti mesh group showing the newly formed bone with lacunae of osteocytes (a) and granulation tissue (Masson Trichrome stain x 100)](http://www.idpress.eu/mjms/)

No significant difference (NS) of native bone results was recorded in Ti mesh cases with p < 0.05 as compared with grafted cases. Significant (S) difference of newly formed bone results was seen in Ti mesh cases with p < 0.05 as compared with the new bone in the grafted cases which was higher in this group as recorded in Table 2.

| Case      | Type of bone | Area    | Area%     |
|-----------|--------------|---------|-----------|
| Grafted   | Native       | 4368.11 | 0.36      | 36.77 ± 5.36 |
|           | Newly Formed | 1299.33 | 0.13      | 13.59 ± 3.53 |
| Ti        | Native       | 4136.16 | 0.34      | 34.81 ± 2.51 (NS) |
|           | Newly Formed | 691.12  | 0.058     | 5.81 ± 0.73 (S) |

NS: Non significant; S: Significant.

**Discussion**

Several studies have concluded that sinus membrane elevation accompanied by simultaneous implant placement resulted in new bone formation without placement of any grafting material. However it is not yet clear how bone is formed in such non-grafted sinuses. Previous research has highlighted the osteogenic potential of cells isolated and cultured from the lining of the maxillary sinus and the ability of these cells to form bone in ectopic conditions [22, 23]. On the other hand it was debated that the walls and septa of the sinus are responsible for new bone formation similar to the mechanism of bone formation in extraction sockets after clot formation [24, 25].

Since the height or quality of the residual alveolar crest may not always be enough to provide the initial stability for simultaneous implant placement, studies have attempted the insertion of a space maintaining device after the process of sinus membrane elevation to evade non-supported sinus membrane crumple to maintain the gained space [16].

Thus the aim of the current contemplate was to assess the capability of a micro titanium mesh, acting as a space maintaining device after Schneiderian membrane elevation, to enhance new bone formation in the sinus without the need to use of any grafting material.

The titanium mesh chosen as a space maintaining device in this study is characterized by its excellent biocompatibility. It is easily shaped and adapted on the bone surface, and rigid enough to maintain the sinus at the new level. The holes of the mesh allow direct contact between the blood clot in the newly formed space and the Schneiderian membrane with its osteogenic property, and its 0.1 mm thickness preserves all of the formed space allowing the placement of the longest possible implant [26].

The xenograft used as a positive control in the present study is an ideal scaffold for osseoinduction and can resist resorption caused by increased pneumatization of the sinus as a result of increased sinus pressure [27, 28].

The shape of the osteotomy in this study was chosen to be rectangular in shape with rounded corners to easily accommodate the titanium mesh during insertion and fixation [29]. A study of the arterial architecture of the maxillary sinus' area explained that the vascularization of the grafting material used in sinus elevation procedures occurs through the endosseous and extraosseous arterial anastomosis. In the present study, the osteotomies were large enough to make sure that the titanium mesh covers all the area planned for the bone formation and future implant placement [30].

The issue of the real need to cover the osteotomy window with a collagen membrane or not is still arguable. Some authors preferred to cover up the osteotomy to rule out non-osteogenic connective tissue infiltration and avoid the escape of the graft particles, while others say there’s no need for the
collagen membrane coverage. In the present study the lateral window was enclosed with a membrane to avoid the infiltration of non osteogenic connective tissue into the cavity [31].

Tawil and Mawla noticed a lack of cortication of the graft surface together with encleavage through the sinus window, which was due to the fibrogenic nature of the periosseum once it has been elevated from the bone surface [32]. In the present study minor encleavations were seen in most of the cases (radiographically and histologically); however it neither affected the bone density nor the width of the future implant placement sites.

The lateral window approach for maxillary sinus elevation used here is a predictable and simple technique and offers good and clear access to the Schneiderian membrane [13]. It was reported that the lateral approach for sinus elevation showed more bone formation than the bone added osteotome sinus floor elevation technique [33].

Results of the present study demonstrated a postoperative increase in bone height and bone density in both graft and titanium mesh groups which was bigger in the graft group but with no statistically significant difference.

Although the xenograft used here acted as a good scaffold for osseointegration it did not resorb completely as evident histologically. The xenograft was shown to undergo slow or no resorption for up to 6 years, hence failing to remodel and adapt to the surrounding bone [34].

Both radiographic and histological results demonstrated the effectiveness of the osteoinductive allograft in forming new bone which is consistent with previous studies using similar graft materials, demonstrating the osteoinductive ability of xenograft when used as the sole grafting material in maxillary sinus augmentation [35, 36]. Histologically xenograft was shown to form new trabecular bone and gain satisfactory results in terms of bone height that prove that it is a suitable augmenting material for atrophic maxillary rehabilitation [37]. Another research demonstrated that long-term outcome after sinus augmentation were in favor of xenograft when compared with autogenous graft where only 4.2 % of 47 cases showed 50 % reduction of augmented height in the xenograft group compared to the 8.7 % cases in the autogenous group [38].

On the other hand, the titanium mesh proved to maintain the space below the Schneiderian membrane as demonstrated by the 6 months postoperative CBCT. Previous studies on localized alveolar reconstruction for guided bone regeneration purposes recommended the use of titanium mesh to maintain the space that will be filled with materials like bone morphogenetic proteins, platelet rich plasma or platelet rich fibrin [39].

The same idea of space maintenance beneath the sinus membrane without the placement of any grafting material was implemented in a former study using a bioresorbable tenting device with simultaneous implant placement. The results were however quite disappointing revealing new bone formation only at the implant surface. The authors attributed these results to several factors which included lack of stabilization of the device inside the sinus cavity, disturbed healing process by the rigid material of the device, making its reshaping difficult, or extensive healing that might have caused small initial perforations to be widened. Moreover, the surface of the used device being smooth and broad in width might have increased the contact area with the Schneiderian membrane, thus hindering its osteogenic properties [16].

Thor et al studied the amount of bone gain by the tenting technique revealing a mean bone height gain of 6.51 mm with an average residual bone height of 5.5 mm [14]. Leblebicigolu reported less gain in bone height than the present study with an average bone gain of 3-4 mm. In the current study the bone height was nearly doubled 6 months after the operation. The tenting technique in Leblebicigolu’s study [40] resulted in bone formation around the apex only while in the current study both the reformatted cross-sections and panoramic views CBCT showed bone formation over the area beneath the titanium mesh.

Results in the present study demonstrated that the bone gain in the titanium mesh group was still not complete as it did not reach the titanium mesh in some areas as shown on the CBCT, implying that more time is needed for the bone to fill all over the created chamber. However the x-rays demonstrated that in the areas for planned future implant placement the bone reached up to 11 or 12 mm which is enough for implant placement with proper primary stability.

The present results are in accordance with previous case series study results where the elevated sinus membrane was maintained by fixing a titanium plate to the lateral wall of the sinus, where only 40.2 % of the space below the plate was maintained after 6 months [2]. In this case series study 6 months postoperative CBCT demonstrated a small gap between the titanium mesh and the newly formed bone. In this contemplate the average residual ridge height reached 9.63 ± 1.47 mm after 6 months which is comparable to the results of the present study demonstrating 11.46 ± 2.78 mm mean bone height 6 months postoperatively.

Bone formation inside the sinus cavity requires the migration and differentiation of osteogenic cells into osteoblasts, where they start to synthesize and deposit collagenous matrix for mineralization. The bone marrow is the main source of such cells and it is thought that the mesenchymal stem cells (MSCs) migrate from the bone marrow into the blood filled sinus cavity using the fibrin network as
a scaffold. Lifting the periosteum initiates bone resorption, exposure of the bone marrow and access of stem cells to the sinus cavity as mentioned in a previous study [9].

Kim et al carried out an in vitro study where it was observed that the source of bone forming cells is the periosteum of the lifted sinus membrane that contributed mainly in the process of bone formation. The presence ofMSCs with osteogenic properties in the maxillary sinus membrane was also confirmed. Another source of the osteogenic cells might be from the periosteum of the osteogenic layer covering the lateral window. The titanium mesh surface texture might have a certain role in bone formation as it stimulates thrombin formation on the surface of titanium and together they both activate stimulation and inhibition of the apoptotic process of osteoblasts [29].

In conclusion, the titanium mesh can be used as a space maintaining device to support the Schneiderian membrane without using any grafting material. Future studies using larger sample size with longer follow up intervals are recommended.

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