EVALUATION OF DENTAL IMPLANTS IN MANDIBULAR NARROW RIDGES USING BONE EXPANSION AND BENDING TECHNIQUE

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ABSTRACT

INTRODUCTION: Narrow dentoalveolar ridges remain a challenge for the successful placement of implants; success depends largely on the quality and quantity of the available bone in the recipient site. This however may be compromised or unavailable which necessitates the need to manipulate the residual bone by contouring or making dimensional changes to create an intra-bony cavity to form a receptor site for an implant while preserving bone integrity and viability.

OBJECTIVES: Evaluation of the success of dental implants in mandibular narrow ridges (≤ 5mm) after using bone expansion and cortical bone bending technique with a new design of bone expander.

MATERIALS AND METHODS: This clinical study was performed on a total of eight patients with mandibular narrow ridges (≤ 5mm). All implants were followed for 6 months. Clinically, each patient was evaluated for pain, tenderness or discomfort, presence of swelling or infection and mobility of the implant. Also, radiographic investigations were performed for the assessment of marginal bone level, bone width and the bone density around the implant. An implant stability assessment of the osseointegration progress was conducted using the resonance frequency analysis technique (Ostell) immediately after implant placement and after three months and after six months.

RESULTS: There was a statistically significant increase in the bone density around the implants throughout the evaluation period and there was increase in implant stability quotient after six months was statistically significant (p=0.008). There were significant decreases in the mean of marginal bone level changes by time in all cases, the increase in bone width quotient after six months was statistically significant p<0.001.

CONCLUSIONS: using the newly designed star shape bone expanders as a bone expansion and bending technique, showed an adequate clinical and radiographic performance in the mandibular narrow ridges.

KEYWORDS: narrow ridges, bone expansion, implant stability.

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INTRODUCTION

Nowadays, dental rehabilitation of partially or totally edentulous regions through dental implants has become common practice with reliable long-term results (1).

However, due to bone atrophy, periodontal disease and trauma, alveolar ridges are more often compromised with horizontal deficiency. Compromised bone quality and quantity after tooth extraction always occurs. These deformities affect implant fixture placement, stability, and long-term success, preventing a proper implant treatment. Implants placed in poor quality bone are often related to compromised primary stability and poor osseointegration (2).

The foundation of implant success has been attributed to their firm bone anchorage called osseointegration or functional ankylosis. It is a predictable tissue response to the placement of tooth root analogues by direct contact between living bone and implants at the light microscopical level. In order to achieve osseointegration the preparation of the bone must be done so that minimal tissue injuries produced (3).

Various surgical widening techniques have been described (4), including lateral augmentation with or without guided bone regeneration (GBR), ridge expansion osteotomy (5), ridge splitting technique with or without interpositional grafting and horizontal distraction osteogenesis (6,7).

These procedures always give chances of surgical risk, postoperative morbidity and multiple long operating surgeries, exposure of the membrane or using bone grafts that could lead to infection. Thus, surgical alveolar ridge expansion was proposed as an alternative technique, where the two cortical plates are expanded to increase the width and thereby to introduce the implant with an appropriate diameter without the removal of any bone from the patient (8, 9).

Bone expansion can be performed by means of osteotomes and manual expanders; these are special set of instruments that were developed to form or shape bone in preparation for the placement of dental implants. They increase the width of bone for implant placement and allow immediate placement of implants in narrow ridges at the time of expansion (10).

This noninvasive way of widening narrow ridges does not require harvesting of bone, reduces operating time and postoperative morbidity, shortens the rehabilitation time, and eliminates the risk of exposure of the membrane or bone graft that could lead to infection (11).

A New design of bone expanders was introduced recently, these expanders are star shaped and they are designed to allow bone expansion with bending of the cortical bone. They provide more surface area and prevent the fracture of the alveolar bone by reducing condensation forces and load (12).
Inclusion criteria
Patients selection criteria of Dentistry, Alexandria University. Patients had an of the Oral and Maxillofacial Surgery Department, Faculty and past dental history was fulfilled. Inspection and number, past medical history, family history, drug history including name, age, gender, occupation, address, telephone evaluated regarding both dental and medical status, The prior to any treatment approach, every patient was the achievement of primary stability in any type of bone simultaneously condensing the bone, in order to optimize cuts through the bone with no bone chipping while designed to evaluate their efficacy in mandibular narrow ridges. This study proposes that this newly designed bone expanders may have similar results to conventional bone expanders.

MATERIALS AND METHODS

1) Study design
Thirteen patients were selected from the Outpatient Clinic of the Oral and Maxillofacial Surgery Department, Faculty of Dentistry, Alexandria University. Patients had an atrophic mandibular edentulous ridge.

Patients selection criteria
Inclusion criteria
- Patients of both sexes, with age of 30-50 years.
- Patients with narrow mandibular ridges (≤5mm).
- Good oral hygiene.
- Adequate inter-occlusal space at the implant site.
- Sufficient available bone to fully accommodate the implant.

Exclusion criteria
- Presence of any bony or soft tissue pathological condition at the surgical site.
- Presence of any uncontrolled systemic disease.
- Para functional habits.
- Heavy smokers.

Informed consent: All patients received explanations about the planned treatment and its potential risks and complications, and signed a written informed consent form prior to being enrolled in the study. It was also mentioned that the patient had the right of withdrawal from the study anytime without any consequences. Ethical approval for this study was obtained from the research ethics committee, Faculty of Dentistry, Alexandria University before beginning the study.

2) Materials

Magic Implant
All implants IBS (InnoBioSurg Co., Ltd, Korea ) were made of titanium alloy with a resorbable blast media surface.

Implants were available in different sizes, lengths (7 - 9 - 11 - 13mm) and diameters (4 - 4.5 - 5mm).

These implants had the following features; a switched platform, a tapered shape, a conical connection with hex base, single threads with new design "fin threads". They were specially designed for self-tapping which cuts through the bone with no bone chipping while simultaneously condensing the bone, in order to optimize the achievement of primary stability in any type of bone density.

The magic kit
Contains guide drill, magic split and a star shaped bone expander.

A) The pre surgical phase
Prior to any treatment approach, every patient was evaluated regarding both dental and medical status. The preoperative data were collected and recorded in full details including name, age, gender, occupation, address, telephone number, past medical history, family history, drug history and past dental history was fulfilled. Inspection and palpation of the site of implant placement was performed, as well as adjacent and opposing teeth, adjacent structure and occlusion; a study model was casted for pre-operative assessment. A CBCT was performed to all patients to measure the width of the alveolar ridge, Planification of the implant placement and the Approximation to mandibular or mental nerve.

B) Surgical phase (Fig 1)
- All patients were operated under local anesthesia, using infiltration technique (Articaine HCl and Epinephrine 1:20,000).
- All patients were instructed to rinse with 0.12% chlorhexiding for 2 minute.
- Full-thickness flap was performed, and extended to one tooth mesial and one tooth distal to the implant site, and slightly lingual to the crest of the ridge.
- A sharp end periosteal elevator was used to elevate the mucoperiosteal flap.
- 1.6 guide drill was used to guide the initial osteotomy for the Magic Split, Drilling was done with low speed (1000 rpm) high torque with internal irrigation with normal saline.
- The Magic split with angled hand lever was then inserted in the osteotomy site, gentle tapping was performed using the mallet until the planned depth was reached, and then the magic split was used for mesiodistal cortical bone incision.
- The 1.6 guide drill was used again to remove the mesiodistal cortical bone and form an oval shape hole.
- The star-shaped magic expander with angled hand lever was then inserted to the length of the implant, by gentle tapping using mallet until the planned depth was reached, the magic expander was used for the bone expansion with ideal condensation.
- Tapered implants were inserted using a ratchet and its adapter. The implant stability was measured by using Osstell. The SmartPeg was screwed into the internal thread of the implant, using a torque of approximately 4-6 Ncm. Then, the Oststell probe was placed in close proximity to the Smart Peg without touching it, to measure the primary stability of the implant (13). The mucoperiosteal flap was sutured, using 3-0 black silk suture (14).

Figure (1): Case 1: (A) Photograph showing missing mandibular first molar. (B) Photograph showing reflected full mucoperiosteal flap. (C) Magic split. (D) Magic expander. (E) Osteotomy. (F) Implant with cover screw. (G) Photograph showing the mucoperiosteal flap sutured, using 3-0 black silk suture. (H) Photograph showing the final prosthesis cemented after 3 months.
C) Post-operative phase
All patients were instructed to apply cold packs extra-orally intermittently every 10 minute for 2 hours on the first day. Chlorhexidine 0.12% mouth wash started after the day of surgery 3 times daily for 7 days, an Antibiotic every 12 hours for 5 days and Non-steroidal anti-inflammatory drugs. Every 8 hours daily for 3 days were instructed as a post-operative medication the sutures were removed after one week post-surgically.

D) Follow up phase
Clinical evaluations were performed for each patient, immediately after implant placement and at a set of intervals, patients were evaluated for pain daily for the first week, then weekly for the first month through Visual Analogue Scale (15). Patients were evaluated for post-operative complications daily for the first week, and then weekly for the first month. A visual descriptor scale (16) was used to indicate presence/absence of edema and inflammation. any post-operative complications such as peri-implant mucositis, peri-implantitis, infection, numbness, wound dehiscence occurring at the implant site during the entire follow-up time was recorded. The depth of the peri-implant sulcus was performed by using calibrated periodontal probe with light force to avoid undue tissue damage and over extension into the healthy tissue. Measurements were taken after implant placement, at three months and at six months post-operatively (17).

The Osstell was used to measure implant stability after implant placement, at three months and at six months post-operatively (18, 19).

Cone beam computerized tomography (20) was performed immediately and six months post-operatively (Fig 2) to assess: Bone width, marginal bone level, Bone density.

![Figure (2): Radiographic evaluation of case 1: (A) immediate postoperative cone beam CT showing the implant in place. (B) Photograph showing A 6month postoperative CBCT showing the implant.](image)

Buccolingual Bone width was measured at certain points on the CBCT preoperatively and measured immediately post operatively and after 6 months.

For Assessment of the bone density around the implant Exposure was performed using Scanora (Scanora 3Dx-Soredex-Finland) at 10 MA, 90 KV and at a proper field of view.

- Image reconstruction was performed using a special software called “Ondemand 3D” (Ondemand3D: Cybermed, Korea) version 1.0.7. Measurements were taken as follows: The bone density apical, buccal and lingual to the implant was used as a known measurement in Hounsfield Unit (To convert from the normal units found in CT data (a typical data set ranges from 0 to 4000 pixel) you have to apply a linear transformation of the data. The equation is: \( \text{hu} = \text{pixel\_value} \times \text{slope} + \text{intercept} \).

E) Prosthetic phase
After 3 months post operatively, the cover screw was removed and the healing abutment placed from 1-2 weeks.

Then, the healing abutment was removed and the abutment was tightened, a condensation silicone impression material was used to make the impression, and definitive porcelain fused to metal restorations were delivered to all patients (Fig 1).

Statistical analysis
- Statistical analysis were performed using a statistical software package SPSS.
- Data were represented as mean ± standard deviation. Repeated measures analysis of variance (ANOVA) test was used to compare numeric variables within the studied group of patients.
- For categorical data, non-parametric ANOVA (Friedman’s test) was performed to detect significant changes within the studied group of patients.
- Post Hoc test was done if ANOVA or Friedman tests were positive. Pearson’s correlation coefficient was used to identify relations between the studied variables.
- In all tests, result was considered statistically significant if the p-value was less than 0.05.

RESULTS
A total of thirteen implants were placed in eight patients (6 females and 2 males), of mean age 38 years. They were selected from the Outpatient Clinic of the Oral and Maxillofacial Surgery Department, Faculty of Dentistry, Alexandria University. Nine implants with 4 mm diameter ×11 mm length were placed in six patients and four implants with 4 mm diameter ×9 mm length were placed in two patients, and. All patients were followed up both clinically and radiographically for 6 months.

I- Clinical evaluation
Pain was evaluated postoperatively daily for the first week then weekly for the first month using the visual analogue scale. On the first postoperative days, all patients experienced mild to moderate pain at the surgical site scoring between two, and three on visual analogue scale. Postoperative tenderness and discomfort were minimal in all cases. During the follow up period, all patients felt no pain, tenderness, or discomfort after implant placement except one case that felt mild postoperative pain scoring two on visual analogue scale and moderate discomfort in the third week after implant placement. In two cases, peri-implant infection with suppuration and swelling in the operated area was found; the first case was failed after three weeks of implant placement and the other was failed after 2 months. In the other cases, patients continued the follow up period without any complications, clinical signs of inflammation, swelling, numbness or peri-implant infections after implant placement or during the evaluation period.

Implant stability quotient was measured in all patients using the resonance frequency analysis technique by the Osstell TM device immediately after implant placement at three months and at six months. The data was collected and tabulated and statistical analysis was done for all patients, the change in implant stability from one interval to the subsequent visit was statistically insignificant after implant placement to the third month (p1=0.688).
In addition, the increase in implant stability from implant placement to the six month was statistically significant (p=0.015) and finally the increase in implant stability from 3 months to 6 months was statistically significant (p < 0.001). The statistical analysis of implant stability scores was done for all patients in Table (1).

Table (1): Comparison between the three periods according to Ostell (n=11).

<table>
<thead>
<tr>
<th>Periods</th>
<th>Bone density (HU)</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Preoperatively</td>
<td>746.97 ± 160.36</td>
<td>9.861*</td>
<td>0.008*</td>
</tr>
<tr>
<td>6th month</td>
<td>717.40 ± 78.0</td>
<td>6.63</td>
<td>0.02</td>
</tr>
<tr>
<td>12th month</td>
<td>689.35 ± 129.01</td>
<td>7.807*</td>
<td>&lt;0.001*</td>
</tr>
</tbody>
</table>

F: F test (ANOVA) with repeated measures. Pairwise comparison between each 2 groups was done using Post Hoc Test (Bonferroni)

p: p values for comparing between three periods

2- Radiographic evaluation

1-Assessment of bone density of the newly formed bone:

Data were collected regarding mean peri-implant bone density values. The mean was calculated immediately postoperatively as the baseline and after 6 months.

In the immediate post-operative phase, the mean peri-implant bone density was 689.35 ± 129.01 HU with a minimum recorded value of 487.14 HU and a maximum recorded value of 864.30 HU.

In the sixth month, the mean peri-implant bone density was 974.53 ± 160.36 HU with a minimum recorded value of 746.97 and a maximum recorded value of 1295.70 HU. These differences were statistically significant (p < 0.001). The statistical analysis of bone density scores was done for all patients in Table (2).

Table (2): Comparison between the two periods according to bone density (n=11).

<table>
<thead>
<tr>
<th>Bone density</th>
<th>Immediate</th>
<th>6th month</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Preoperatively</td>
<td>846.30 ± 6.15</td>
<td>5.0 – 6.63</td>
<td>4.80 – 7.13</td>
<td>0.02</td>
</tr>
<tr>
<td>6th month</td>
<td>746.97 ± 7.13</td>
<td>5.94 ± 0.59</td>
<td>6.15 ± 0.70</td>
<td>0.02</td>
</tr>
</tbody>
</table>

Paired t-test

p: p values for comparing between Immediate and 6th month

*: Statistically significant at p ≤ 0.05

2-Assessment of the marginal bone level:

The mean value of the marginal bone level was calculated and recorded immediately post-operative and after 6 months:

- Immediate post-operative, mean marginal bone level value was 9.07 mm, with SD ± 0.78 mm.
- After 6 months, mean marginal bone level value was 8.05 mm, with SD ± 1.27 mm.
- The Decrease in marginal bone level was calculated, the minimum decrease was 0.13 mm with mean value 1.09 mm and the maximum marginal bone decrease was 2.50 mm, show that there were significant decreases in the mean of marginal bone level changes by time in all cases.

3-Assessment of bone width (bone expansion):

Buccolingual Bone width was measured from the CBCT; in certain point preoperatively, immediately after implant placement and after 6 months.

The mean bone width quotient pre-surgically was 4.82 ± 0.27 mm. It was an increase immediately after implant placement 5.97 ± 0.59 mm.

The change in bone width from one interval to the subsequent visit was statistically significant between the preoperative period to the immediate period (p1=0.001).

However, between the periods of immediate placement to the 6 months showed statistically insignificant in bone width (p3 < 1.000). The statistical analysis of implant stability scores was done for all patients in Table (3).

Table (3): Comparison between the three periods according to bone width (n=11).

<table>
<thead>
<tr>
<th>Bone width</th>
<th>Preoperative</th>
<th>Immediate</th>
<th>6th month</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Preoperatively</td>
<td>4.25 – 5.32</td>
<td>5.0 – 6.63</td>
<td>4.80 – 7.13</td>
<td>20.035</td>
<td>&lt;0.001*</td>
</tr>
<tr>
<td>6th month</td>
<td>4.82 ± 0.27</td>
<td>5.94 ± 0.59</td>
<td>6.15 ± 0.70</td>
<td>0.02</td>
<td></td>
</tr>
</tbody>
</table>

F: F test (ANOVA) with repeated measures. Pairwise comparison between each 2 groups was done using Post Hoc Test (Bonferroni)

p: p values for comparing between three periods

*: Statistically significant at p ≤ 0.05

DISCUSSION

This study was conducted to evaluate the dental implants in narrow mandibular ridge using bone expansion and bending technique.

In the present study 8 patients (6 females and 2 males) from 30-50 years. Patients were selected free from any systemic diseases such as osteoporosis diabetes mellitus, heart diseases, as such conditions may complicate the surgical procedure or the healing process. Accordingly, in the year 2009, Michaeli et al (21) mentioned that the diabetes remains a relative contraindication for implant therapy.

In addition, in the year 2000, Becker et al (22) documented that osteoporosis has been considered as a risk factor for implant failures due to low bone mass and a micro-architectural deterioration of bone leading to fragility. Patients were selected with no Para functional habits as bruxism or clenching which is the most common cause of implant bone loss, Glauser et al in 2001 (23) identified a higher failure rate in the group with bruxism than among non-bruxers (41% compared to 12%).

In the year 1988, Koth et al (24) and Dhanrajani and Al-Rafee in 2005 (25) proposed the considerations and factors affected osteointegration, the quantity and quality of the residual ridges and the bone width is an important criteria affect the osteointegration, the dimension of residual ridge, surgical access and proximity of vital anatomical structure all affect the osteointegration process.
Therefore, placement of implants surrounded by good quality bone is a prerequisite for osseointegration. As a result, a healing period of at least six months has been recommended between extraction of a tooth and subsequent implant placement (26).

In this study CBCT was performed for all patients preoperatively, corresponding to the studies conducted by Cassetta et al in 2013 (27) and Bornstein et al in 2014 (28), they reported that the use of CBCT in implant dentistry vary from preoperative analysis regarding specific anatomic considerations.

In this study surgical alveolar ridge expansion was proposed in mandibular narrow ridges by using special set of expander instruments that were developed to form and shape the bone in the implant placement site, Chiapasco M, Ferrini F et al (29) documented that Because bone is visco-elastic, the width of the alveolar ridge can be increased by using expanders to create pressure between buccal and lingual cortical plates, in a slow organized manner Without removing of any bone from the patients, That noninvasive way of widening narrow ridges does not require harvesting of bone, reduces operating time and postoperative morbidity, shortens the rehabilitation time, and eliminates the risk of exposure of the membrane or bone graft that could lead to infection.

That agrees with Demetriades et al (30) that reported that Implant placement in atrophic ridge is a challenging procedure and often involves prior bone graft augmentation, resulting in a long time treatment plan and more surgical trauma.

In addition, this technique does not require a waiting period of 4-6 months for bone consolidation prior to implant placement and decreases the morbidity since it avoids a second surgical. However, in the year 2004, Misch (31) reported that this technique presents some limitations. It is not applicable if there is insufficient bone height for implant placement and implementing the technique on atrophic ridges < 3.0 mm wide may result in unfavorable bone fractures that lead to bone resorption, in cases of significant bony ridge defects and unfavorable ridge relationships.

A new design of bone expanders was introduced recently, these expanders are star shaped and they are designed to allow bone expansion with bending of the cortical bone they provide more surface area and prevent the fracture of the alveolar bone by reducing condensation forces and load.

The system is designed to push the alveolar bone to form a placement hole, providing a perfect protection to anatomical structures and reducing the need for bone grafting (12).

A single pilot drill was used, followed by atraumatic expansion of the osteotomy site using manual bone expanders to decrease surgical trauma that is mainly attributed to thermal injury of the bone. Otherwise, thermal injury may lead to osteonecrosis and possible fibrous and granulation tissue encapsulation around the implant (32).

In the year 1984, Eriksson et al (33) stated that overheating the tissues at the bone implant interface could cause bone necrosis and compromise the bone’s ability to survive as well-differentiated tissue. Bone tissues are sensitive to heat at 47°C. Rabbit tibiae heated to 50°C for 1 minute and 47°C for 5 minutes have shown a 30% to 40% bone resorption after 40 days, with bone tissue replaced by fat cells. When bone was heated to 47°C for 1 minute, fat cell injury and inconsistent bone injury were observed. Higher injury was reported after tissue was heated to 53°C for 1 minute, resulting in permanent vascular stasis and irreparable bone tissue necrosis.

Also, this is in accordance with Brisman (34) who stated that overheating of bone during preparation of the implant site can lead to bone necrosis, interfacial formation of connective tissue between the bone and the implant, loss of osseointegration and consequently loosening of the implant.

In this study maximum ridge width was 5.0 mm, postoperative cone beam CT revealed up to 2 mm increase in the ridge width. This accordance with Scipioni et al (35) who stated that wherever dental implants are placed, a minimal thickness of 1 to 1.5 mm of bone should remain on both the buccal and the lingual/palatal aspects of the implants to ensure a successful outcome.

This study showed that the marginal bone loss occurring in the both implants right and left side did not exceed 3 mm during the observation period. The marginal bone loss decreased due to the preparation of the implant site and increased again due to bone remodeling during osseointegration.

Regarding to the bone density around the implant, the diagnosis of bone density around the implant is a key of success of the clinical study, the strength of the bone is directly related to bone density. Factors as the amount of bone contact, the modulus of elasticity, and axial stress contours around an implant are affected by bone density, As a consequence the treatment plan which includes implant number and size, should be modified as stress factors and bone density increases (36).

This study showed a significant change in the measurement of bone density around the implant throughout the period of evaluation which indicating successful integration that agreement with Bergkvist et al (36), Twenty-one patients received 137 implants (87 in maxillae and 50 in mandibles) Bone mineral density (BMD) was significantly correlated with bone quality classification in both arches (P < .001) Mean BMD was also significantly correlated with stability values (P < .001).

Different methods for objectively assessing primary stability have been proposed. Several studies have reported that resonance frequency analysis is a useful tool to analyze primary stability after implantation, as well as the degree of stability after osseointegration, in this study using resonance frequency analysis ostell ISQ presents “almost perfect” reproducibility and repeatability after statistical analysis by means of the intra class correlation coefficient. Therefore It can be concluded that Oststell system measurements are highly reliable regarding repeatability.

These results are in accordance with the studies performed by Le Resche et al in 2000 (38) stated that the pain following the implant placement ranged from mild to moderate scoring between 0 and 3 on visual analogue scale. The peak of pain perception occurred on day one following surgery.

In two cases, peri-implant infection with suppuration and swelling in the operated area was found, the first case was after three weeks of implant placement and the other...
was after 2 months. In the year 2012, Sakka et al (39) stated that peri implant infection seem to be one of the most important causes of early implant failure. Early signs of infection may be an indication of a much more critical result than if the same complications occur later, because of disturbance of the primary bone healing process, early infection around the implant and failure of this case could be attributed to lack of oral hygiene maintenance by the patient in spite of the instructions given to her. Early peri implant infection may have a role in preventing osseointegration.

In the other cases, patients continued the follow up period without any complications, clinical signs of inflammation, swelling or peri implant infections after implant placement or during the evaluation period.

CONCLUSION
From the results of this study we can conclude that using the newly designed star shape bone expanders as a bone expansion and bending technique, showed an adequate clinical and radiographic performance in the mandibular narrow ridges.

CONFLICT OF INTEREST
The authors declare that they have no conflicts of interest.

REFERENCES
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