SURVIVAL RATE, PERI-IMPLANT SOFT TISSUES AND CRESTAL BONE LOSS OF IMPLANTS INSERTED BY FLAP OR FLAPLESS SURGERY IN TYPE 2 CONTROLLED DIABETIC PATIENTS AND IMMEDIATELY LOADED WITH MANDIBULAR FIXED PROSTHESIS. ONE YEAR RANDOMIZED CLINICAL TRIAL

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ABSTRACT

Objectives: The aim of this short-term randomized trial was to evaluate survival rate, peri-implant soft tissues and crestal bone loss of implants inserted by flap or flapless surgery in type 2 controlled diabetic patients and immediately loaded with mandibular fixed prosthesis.

Materials and methods: Twelve completely edentulous patients with controlled type II diabetes mellitus (HbA1c ranged from 5 to 6%) were randomly assigned into two equal groups: Group I (flap surgery) received 6 implants in the mandible using standardized full-thickness mucosal flap surgery and free hand implant placement; Group II (flapless surgery) received 6 implants in the mandible using flapless surgical approach and computer guided implant placement. Implants were immediately loaded with acrylic fixed restoration. Three months later, fixed ceramo-metal restoration was delivered. Implant survival rate, peri-implant soft tissues (plaque index, gingival index, pocket depth and keratinized mucosal width) and crestal bone loss were evaluated at time of implant loading (baseline), 3 months, 6 months and 12 months after loading.

Results: The implant survival rate was 94.5% and 91.6% in the flap and flapless groups respectively. Without difference between groups (p=0.652). For both groups, plaque scores, pocket depth and crestal bone loss significantly increased with time. Gingival scores significantly increased from baseline to 3 months, then significantly decreased thereafter. Keratinized mucosal width significantly increased from baseline to 3 months for flap group and insignificantly decreased for flapless group. Flap group showed significant higher plaque scores, gingival scores, pocket depth, width of keratinized mucosa and crestal bone loss than flapless groups.

Conclusion: Computer guided flapless surgical approach may be recommended for implants placed in edentulous type 2 controlled diabetic patients and immediately loaded with mandibular fixed prosthesis as it was associated with favorable peri-implant soft tissue response and crestal bone loss compared to implants placed with flap surgery after one year.

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INTRODUCTION

Diabetes Mellitus is a group of metabolic disorders characterized by an increase in plasma glucose levels\(^1\). There are mainly two types of diabetes: 1) type I diabetes (insulin-dependent) which characterized by a lack of insulin production and 2) type II diabetes (noninsulin dependent) which is caused by the body’s ineffective use of insulin\(^2\). Diabetic patients have several complications that may affect the healing potential following endosseous implant treatment such as microvascular disease, susceptibility for infection, and delayed wound healing \(^3\). Moreover the implants in these patients subjected to mechanical overload resulting from diabetes-induced lower percentage of bone-to-implant contact, immature bone, and incorrectly formed bone\(^4\). Therefore, patients with diabetes were not considered suitable for implant treatment when the treatment was introduced. However, over the past 2 decades, diabetes has been regarded as a relative (not absolute) contraindication for implant therapy related to the stability of the diabetic’s blood sugar level\(^5\). Measurements of glycated hemoglobin (Hba1c) levels is considered an accurate method to determine glycemic control in diabetic patients as it reflected the blood glucose level over the last 2-3 months\(^5\). Good glycemic control is essential to maintain the function of osteoblast and the decrease the rate of crestal bone loss\(^6\). Recommendations for strict glycemic control for persons with diabetes have targeted maximal Hba1c levels ranging from 6.5% up to 7.0% \(^5\). Implant success rate in patients with type II controlled diabetic patients was reported to range from 9 to 100%\(^10\)-\(^12\).

The traditional used full-thickness flap surgical approach provides good visualization of anatomical structures, ridge shape and landmarks at implant placement which reduce the risk of bone fenestration and perforation but the technique is usually associated with increased postoperative pain, discomfort, bleeding and edema as well as increased marginal bone resorption\(^13\),\(^14\). In contrast, the flapless surgical approach is minimally invasive, is associated with reduced surgical trauma, short observation period, reduced postoperative pain, and swelling, increased patient satisfaction and rapid healing \(^15\),\(^16\). Moreover, it preserves blood supply to the periosteum of bone and decrease bone loss\(^17\). However, flapless surgical approach is a blind technique which may increase the risk of bone perforation, over or under preparation of implant osteotomy and inability to manipulate peri-implant soft tissue to increase keratinized mucosal width\(^18\),\(^19\).

Immediate loading of the implants by professional restoration achieves higher patient satisfaction as it provides immediate restoration of mastication and aesthetics without need to wear dentures during osteointegration. However, uncontrolled loading may induce micromotions at implant/bone interface and cause marginal bone loss \(^20\). Reviewing the literature, the effect of immediate loading on the success rate of implants inserted in patients with controlled type 2 diabetes was scarce and limited to single implants installed in premolar and molar regions and supported single restorations\(^21\) or implants inserted in maxillary anterior region (esthetic zone)\(^5\). However, evaluation of immediate loading of full arch fixed restoration in edentulous controlled diabetic patients was not a concern. Moreover, the effect of the surgical approach (flap Vs flapless) on clinical and radiographic outcomes of implants inserted in controlled type 2 diabetes was limited to delayed loading protocol only which was used either with full arch\(^10\) or single tooth\(^22\) prosthesis and immediate loading protocol was not evaluated. Furthermore, there is still little information in the literature regarding the implant survival and peri-implant tissue outcomes of implants inserted with flapless surgery in controlled diabetic patients\(^10\).

Accordingly, the purpose of the present study was to evaluate the survival rate, peri-implant
soft tissues and peri-implant crestal bone loss of implants inserted by flap or flapless surgery in type 2 controlled diabetic patients and immediately loaded with mandibular fixed prosthesis. The null hypothesis of the study was that there will be no significant difference in the studied outcomes between the 2 surgical approaches.

**MATERIALS AND METHODS**

**Patient enrollment and study design**

Twelve edentulous patients (6 males and 6 females, mean age of 58±4.6 years) with controlled type II diabetes mellitus were selecting from the patients attending regularly at the outpatient clinic of the oral and maxillofacial department who had a desire to replace their missing teeth with fixed restoration. The inclusion criteria are; 1) Completely edentulous maxillary and mandibular ridges, 2) Patients medically diagnosed with type II-controlled diabetes (noninsulin dependent) with a history of well-controlled plasma glucose levels (fasting plasma glucose ≤140 mg/dL, 2-hour postprandial glucose ≤200 mg/dL and glycosylated hemoglobin (HbA1c) from 5 to 6%) as verified by patients’ physicians and patients underwent oral hypoglycemic drugs, 3) patients had sufficient bone height and width (class III-V according to Cawood and Howell) and density in anterior and posterior regions of the mandible to receive six implants of at least 11 mm in length and 3.7 mm in width. This was checked by preoperative cone beam computerized tomography (CBCT), and 4) Implant sites should had at least 4 months of healing following tooth extraction prior to implant placement. The exclusion criteria include 1) patients with history of aggressive periodontitis, 2) Smokers, 3) patients with malignancy who underwent radiotherapy or chemotherapy, 4) patients with harmful habits such as clenching or bruxism, 5) bleeding disorders or other systemic diseases that may contraindicate implant surgery, 6) persons with a history of microvascular or macrovascular complications, and 7) patients with immunosuppressive drugs. Patients were informed about the steps of the study and the need for frequent recalls, then informed consents were obtained from participants. The study was conducted according to the ethical principles stated in the Helsinki Declaration and the study protocol was approved by the local ethical committee of the faculty of Dentistry. CONSORT guidelines for randomized controlled trials were followed.

Each patient was provided a number and entered into a spread Excel sheet. Using RAND command, the patients were randomly distributed. The patients were then randomly assigned into one of two groups using balanced randomization procedures to ensure equal age, and sex distribution between groups. Comparison of baseline criteria between groups was performed after randomization to ensure that there was no significant difference in age and sex between groups before the start of study. Randomization and allocation of participants was performed by blinded dentist. Group I (flap surgery) included six patients (3 males and 3 females) received 6 implants in the mandible using standardized full-thickness mucosal flap surgery and free hand implant placement, group II (flapless surgery) included six patients (3 males and 3 females) received 6 implants in the mandible using flapless surgical approach and computer guided implant placement. Implants in both groups will immediately loaded with provisional acrylic fixed restoration on the same day of surgery. Three months later, fixed porcelain fused to metal restoration we are delivered to all participants.

**Surgical and prosthetic protocol**

For both groups, maxillary and mandibular conventional dentures were constructed using the conventional steps of denture construction and bilateral balanced occlusal concept was used to enhance the stability of maxillary dentures. For both groups, 6 bone-level platform-switched implants (Tiologic, Dentarum, Germany) were
planned to be inserted in the mandible (4 implants equally distributed in the interforaminal area and two implants posterior to the mental foramina). The implant diameters ranged from 3.7 to 4.8 and lengths ranged from 11 to 15mm were installed. Implants were installed with at least 35 Ncm insertion torque to provide good primary stability of the implants which is prerequisite for immediate loading.²

Surgery was performed under local anesthesia for both groups. For flap group (Group I), gutta percha radiopaque markers were attached to mandibular dentures at proposed implant sites to convert the denture to the radiographic template. Cone beam computerized tomography (CBCT, i-CAT Vision, Hatfield, PA, USA) was performed while patients wearing the dentures to evaluate implant sites and the proximity of implants to vital structures. Moreover, CBCT was used to detect the proper implant length and diameter. The denture was then converted to surgical stent by attaching metal tubes at proposed implant sites. For flapless group (Group II), Radiopaque gutta percha markers are added to the polished surface of the mandibular denture at labial, buccal and lingual flanges. Dual scan protocol was followed using cone beam CT (CBCT, i-CAT, Imaging Sciences International ISI, Pennsylvania, USA).

Firstly, the patients were scanned while wearing their mandibular dentures, then the mandibular dentures were scanned alone. The two data sets of the double scans were overlapped then the acquired images were loaded into 3-D image treatment planning software (OnDemand). According to the CT scan, the implants were virtually planned, then an individualized stereolithographic surgical guide was constructed using prototyping technique (fig1). A mucosal supported stereolithographic surgical template with 6 sleeves positioned over proposed

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Fig. (1). Cone beam computerized tomography to evaluate implant sites, plane implant positions and construct stereolithographic guide.
implant sites was constructed using 3D printing technology (In2Guide).

For flap group, a mid-crestal incision was made from first molar area on one side to first molar area on the other side. Then a full-thickness mucoperiosteal flap was raised. Using the surgical guide, 6 implants were inserted (4 implants equally distributed in the interforaminal area and two implants posterior to the mental foramina) parallel to each other's using successive drilling. Healing abutments were connected to the implants and the flap was closed with interrupted sutures (Vicryl, 0000) (fig 2). For flapless group, implants were inserted according the flapless surgical protocol using mucosal supported stereolithographic surgical template and the universal surgical kit (In2Guide, Universal Kit Cybermed Inc) supplied with the template to be used during osteotomy preparation. This kit includes hand drill sleeves with successive increasing diameters that fit the template sleeves.

The mucosal supported template was stabilized in the patient’s mouth by a rubber base interocclusal record and fixed to the mandibular bone using anchor pins. Implant osteotomies were prepared using successive drills of increased diameters that fit into metal sleeves of the template (fig 3). Postoperative panoramic radiograph was made to verify implant position (fig 4).

For both groups, Open tray impression was made on the implant level. Long transfer copings were threaded to the internal hex of the implants. A stock plastic tray was perforated over implant sites. Rubber dam sheets were adapted over the copings to protect the wound and the sutures during impression making. Orthodontic ligature wire was used to connect the transfer copings, then Duralay resin (Reliance, USA, with no dimensional changes) was adapted over the wire and the copings to splinted the transfer copings and to avoid movement of the impression copings during impression making (Fig 2).
Light body rubber base impression material was injected around the impression posts and the overall impression was completed using putty material (Zhermack®, Badia Polesine, Rovigo, Italy). Implant analogues were attached to the impression posts and the impression was poured using hard stone. Recorded bases were constructed record jaw relations and facial support. Straight titanium abutments were connected to the impression analogues, and fixed provisional screw retained tooth coloured acrylic resin was constructed in the laboratory. The prosthesis was screwed to the implants to allow immediate loading of the implants in the same day of surgery (fig 6).

For both groups necessary occlusal adjustments were performed to ensure homogenous occlusal contact in centric and eccentric relations. Each patient was prescribed Antibiotics (amoxicillin and clavulanic acid (Augmentin 1gm/ twice daily) started before surgery and continued for 10 days postoperatively. Mouth wash (chlorhexidine digluconate 0.2%) started before surgery and continued daily for 7 days postoperatively. Pain was controlled by analgesics (600 mg ibuprofen one every 8 h) as long as needed. Also, Nonsteroidal anti-inflammatory drugs (Alphintern) were prescribed 3 times daily for 7 days. Instructions were given to the patient to use ice bags to reduce post operative edema. The
patients were instructed to eat soft diet and perform adequate oral hygiene. 3-months regular recall visits for data collection were scheduled for all participants over the study period.

Three months later (after osseointegration), the provisional acrylic prosthesis was unscrewed, then open tray impression was performed again to construct screw retained metal ceramic fixed prosthesis for both groups. On the resultant cast, Ti-base abutments (Tiologic) were threaded to the implant analogues and the cast was scanned using CAD/CAM device (Ceramill, Austria), then a fixed screw-retained hybrid prosthesis was designed with 12 teeth (from first molar on one side to first molar on the other side) then printed into castable resin and tried in patient mouth. The resin bridge was invested, cast with Cobalt chromium alloy, tried in for passive fit using single screw test. The porcelain powder (VITA, Germany) was mixed, applied onto the metal over the opaque layer, fired, finished and glazed. Pink porcelain was used to replace lost bone and gingival tissues when needed (fig 7). Fixed prostheses were delivered to all patients and panoramic radiographs were made to ensure passive seating of the prosthesis.

Study outcomes

Implant success rate was calculated using the criteria proposed by Albrektsson et al.\textsuperscript{25} which include; Absence of implant mobility; absence of pain, foreign body sensation and/or dysesthesia), absence of peri-implantitis and suppuration, absence of radiolucency around the implants and vertical bone loss was not exceed 1.5mm in the first year. Implant was considered survived if it fulfills function, and does not require removal. Plaque index was measured according to Mombelli et al.\textsuperscript{26} and gingival index was measured using Loe and Silness\textsuperscript{27} Pocket depth was measured by plastic periodontal probe which inserted in the peri-implant sulcus to measure the distance between gingival...
margin and the most apical probing depth. Plaque index, Gingival Index, pocket depth were measured at mesial, distal, buccal, and lingual surface of each implant. The width of keratinized mucosa around each implant was measured in mm using a graduated periodontal probe as the distance between the gingival border to the muco-gingival junction on the buccal side of each implant\(^2\).

Crestal bone loss was measured at mesial, distal, buccal, and lingual surface of each implant using CBCT (i-CAT Vision, Hatfield, PA, USA). For standardization of exposure conditions, all images we are performed with the same acquisition time (14.7 second), voxel size and slice thickness. In the panoramic window of the CBCT software (OnDemand3DApp), marginal bone height was measured at mesial and distal surface of each implant. In the cross-sectional window of the CBCT software, marginal bone height was measured at buccal and lingual surface of each implant. To estimate marginal bone height at all surfaces, the distance from implant abutment junction (point A) to the bone contact with implant (point B) was measured using the ruler measure tool of the software to give bone level\(^2\). Bone loss was calculated by evaluation of bone height changes from base line to 3 months, 6 months and 12 months. The mean marginal bone loss for all surfaces was used.

All clinical parameters were measured at immediate loading with provisional acrylic dentures (baseline), 3 months, 6 months, and 12 months after loading.

**Statistical analysis**

Data was analyzed with SPSS program version 25. Shapiro wilk test of normality was used to evaluate normal distribution of the data. Kaplan Meier analysis and Log rank test were used to calculate implant survival rates. Friedman test, followed by Wilcoxon signed ranks tests were used to compare plaque and gingival indices between different time intervals and Mann-Whitney test was used to compare plaque and gingival indices between groups. Repeated measures ANOVA followed by Bonferroni test was used to compare pocket depth, implant stability, width of keratinized mucosa, and crestal bone loss between time intervals and groups. P is significant if it was less than 5%.

**RESULTS**

In this study, the patients attended the regular follow-up and there were no dropouts. Two implants failed to integrate in the flap group (in 2 patients) and 3 implant failures (in 2 patients) occurred in the flapless group resulting in 94.5% and 91.6% implant survival rate in the flap and flapless groups respectively. Implant failures in both groups occurred within the first three months after implant loading with provisional acrylic restoration. No implant failures occurred later. The failed implants were occurred as a result of implant overloading and implants were associated with mobility, crestal bone loss, gingival inflammation and suppuration. The failed implants were removed and the remaining implants were used to support the final ceramometal prosthesis. The study was conducted according to intention to treat principal. Kaplan Meyer analysis of implant survival rate in both groups is presented in fig 9. There was no significant difference in implant survival rate between groups (log rank test, \(p = .652\)).
Comparison of plaque and gingival scores between flap and flapless groups and between time intervals is presented in table 1. Plaque scores significantly differ between observation times for both groups (p=.014 and .031 for flap and flapless groups respectively). For both groups plaque scores significantly increased with time. Multiple comparison between each two observation times is presented in the same table. For flap group, there was a significant difference between each two observation times. For flapless group, no significant difference in plaque scores between 6 months and 12 months was observed. At baseline, no significant difference in plaque scores between groups was detected. At all other observation times, flap group showed significant higher plaque scores than flapless groups.

Gingival scores significantly differ between observation times for both groups (p=.032 and .037 for flap and flapless groups respectively). For both groups gingival scores significantly increased from baseline to 3 months, then significantly decreased thereafter. Multiple comparison between each two observation times is presented in the same table.

**TABLE (1)** Comparison of plaque and gingival scores between flap and flapless groups and between time intervals

<table>
<thead>
<tr>
<th></th>
<th>Baseline Median (Minimum-maximum)</th>
<th>3 month Median (Minimum-maximum)</th>
<th>6 month Median (Minimum-maximum)</th>
<th>12 month Median (Minimum-maximum)</th>
<th>Freidman (p value)</th>
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<td><strong>Medians of plaque scores</strong></td>
<td></td>
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<tr>
<td>Flap group</td>
<td>.000a (.00-.00)</td>
<td>2.0b (1.0-3.0)</td>
<td>1.50c (1.5-2.0)</td>
<td>1.400d (0.5-1.5)</td>
<td>.014*</td>
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<tr>
<td>Flapless group</td>
<td>.000a (.00-.00)</td>
<td>1.5b (1.00-2.00)</td>
<td>1.1c (0.6-1.4)</td>
<td>1.200c (1.00-1.5)</td>
<td>.031*</td>
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<tr>
<td>Mann-Whitney</td>
<td>1.00</td>
<td>.018*</td>
<td>.021*</td>
<td>.040*</td>
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<tr>
<td>(p value)</td>
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<tr>
<td><strong>Medians of gingival scores</strong></td>
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<tr>
<td>Flap group</td>
<td>.50a (0.25-1.00)</td>
<td>1.50b (1.00-2.00)</td>
<td>1.00c (.50-1.50)</td>
<td>1.10c (.60-1.5)</td>
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<td>Flapless group</td>
<td>.10a (.00-.40)</td>
<td>1.00b (1.0-1.5)</td>
<td>.60c (.30-.80)</td>
<td>0.50c (0.20-0.75)</td>
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<tr>
<td>Mann-Whitney</td>
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<td>.002*</td>
<td>.007*</td>
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</tbody>
</table>

*p is significant at .05. Different small letters in the same raw show significant difference between time intervals (Wilcon test, p<.05), while similar letters show no difference.
table. For both groups, no significant difference in gingival scores was observed between 6 months and 12 months. For all observation times, flap group recorded significant higher gingival scores than flapless group.

Comparison of probing depth, keratinized mucosal width, and marginal bone loss between flap and flapless groups and between time intervals is presented in table 2. For both groups, there was a significant difference in pocket depth between observation times (p=.013, and .015 for flap and flapless group respectively). Pocket depth increased significantly with advance of time in both groups. Multiple comparisons of pocket depth between each two observation times are presented in the same table. For both groups there was a significant difference in pocket depth between each two observation times. At baseline, there was no significant difference in pocket depth between groups. For all other time intervals, flap group recorded significant higher pocket depth of the flapless group. Mean keratinized mucosal width significantly different between observation times for both groups. For flap group, keratinized mucosal width significantly increased from baseline to 3 months. For flapless group, keratinized mucosal width decreased insignificantly with time. For flap group, no significant difference in width of keratinized mucosa between 3 months, 6 months and 12 months was observed. There was a significant difference in mean marginal bone loss between observation times (p=.001 and .007 for flap and flapless group respectively). For both groups marginal bone loss significantly increased with time. Multiple comparison of marginal bone loss between each two observation times is presented in table 2. For both groups, there was a significant difference in marginal bone loss between each 2 time intervals. For observation times, flap group recorded significant higher marginal bone loss than flapless group.

### TABLE (2) Comparison of probing depth, keratinized mucosal width, and marginal bone loss between flap and flapless groups and between time intervals

<table>
<thead>
<tr>
<th></th>
<th>Baseline mean± standard deviation</th>
<th>3 month mean± standard deviation</th>
<th>6 month mean± standard deviation</th>
<th>12 month mean± standard deviation</th>
<th>Repeated ANOVA (p value)</th>
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<tr>
<td><strong>Flap group</strong></td>
<td>1.12±.35a</td>
<td>1.32±.27b</td>
<td>1.58±.31c</td>
<td>1.82±.40d</td>
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<td><strong>Flapless group</strong></td>
<td>.91±.29a</td>
<td>1.00±.29b</td>
<td>1.16±.37c</td>
<td>1.33±.45d</td>
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<td>t-test (p value)</td>
<td>.324</td>
<td>.024*</td>
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**Mean Keratinized mucosal width**

<table>
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<tr>
<th></th>
<th>1.69±.41a</th>
<th>1.82±.33b</th>
<th>1.81±.38b</th>
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<td><strong>Flap group</strong></td>
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<td><strong>Flapless group</strong></td>
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<td>.005*</td>
<td>.002*</td>
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<tr>
<td>t-test (p value)</td>
<td>.155</td>
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**Mean marginal bone loss**

<table>
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<tr>
<th></th>
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<th>.62±.12a</th>
<th>.84±.32b</th>
<th>1.2±.33c</th>
<th>&lt;.001*</th>
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<td><strong>Flap group</strong></td>
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<td>.40±.13a</td>
<td>.63±.24b</td>
<td>.91±.33c</td>
<td>.007*</td>
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<td>t-test (p value)</td>
<td>.011*</td>
<td>.014*</td>
<td>.005*</td>
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*p is significant at .05. Different small letters in the same raw show significant difference between time intervals (Bonferroni test, p<.05), while similar letters show no difference.
DISCUSSION

Three-Dimensional cone beam computerized tomography was used for evaluation of crestal bone height changes around implants as it provides information about buccal and lingual bone resorption in addition to mesial and distal bone resorption which is not provided by the two-dimensional periapical radiography. Also Cone beam CT is easily used with edentulous patients had elevated floor of the mouth which makes periapical radiographs difficult to perform. The use of cone beam CT in evaluation of marginal bone resorption around implants was previously described in other studies.

The implant survival rate was 94.5% and 91.6% in the flap and flapless groups respectively without difference between groups. Similarly, Agrawal et al. found a similar survival rate for implants inserted with flap (95.7%) and flapless (93.5%) surgical approaches and conventionally loaded in controlled diabetic patients. They also found no significant difference in implant survival rate between flap and flapless groups. The high survival rate in both groups could be attributed to the good bone density in the mandible which provide higher implant stability that can withstand micromotions of immediate loading. Another explanation could be attributed to the good glycemic control in the included participants patients.

In this study, plaque and gingival scores increased significantly with time in both groups. The large median plaque and gingival scores obtained after 3 months in both groups. This could be attributed to the presence of provisional acrylic bridge which have spaces around the implants that may hinder adequate cleaning by the patients. When professional restoration was replaced by ceramo-metal restoration, plaque scores decreased after six months due to the smooth convex surface of prosthesis and the high adaptation of the prosthesis to the abutments. The increased gingival scores after 3 months are attributed to the increased plaque scores which cause gingival inflammation. The increased plaque and gingival scores was in line with Peled et al. who observed plaque accumulation and peri-implant gingivitis in controlled type II diabetic patients with advance of time. Decreased plaque scores after six months were similar to finding of another study in which a significant decrease in plaque around implants supporting fixed immediate prostheses was noted after 6 months. The increased plaque and gingival index with advance of time for flap and flapless groups was in line with finding of another study in which the authors reported a progressive increase of plaque index in sulcular bleeding index after one year. The flap group recorded significant higher plaque and gingival index than flapless group. In agreement with this finding, Agrawal et al. reported that flap group had significant increase of plaque index and the mean sulcular bleeding index compared to the flapless group when implants are conventionally loaded in controlled diabetic patients. Al-Amiri et al. found no significant difference in plaque and bleeding index between conventionally loaded and immediate loaded single implants inserted in controlled type II diabetic patients. The increased plaque and gingival scores in the flap group may be attributed to the flap surgery which discourage the patients to perform adequate cleaning to avoid pain in the surgical site. Even after healing the wound, they develop the habit of avoiding cleaning to avoid discomfort. The increased plaque scores could be responsible for increased gingival scores in the flap group due to peri-implant mucosal inflammation.

Pocket depth significantly increased with advance of time in both groups. This may reflect the increased marginal bone loss combined with gingival overgrowth that occurred in both groups. In agreement with this observation, Agrawal et al. reported significant increase in peri-implant probing depth in both flap and flapless groups when implants are conventionally loaded in controlled diabetic patients. In contrast, Kapur et al. showed no significant change in pocket depth with passage.
of time around implants supporting mandibular overdentures in patients with controlled diabetes. The flap group recorded significant higher pocket depth than flapless group at all observation times. The increased pocket depth with flap group could be attributed to peri-implant gingival enlargement caused by flap reflection, re-adaptation and suturing the flap over the abutments together with increased peri-implant bone loss in the flap group compared to flapless group 36. On the other hand, the flapless approach provides gap free connection with optimum mucosal barrier free of bacterial accumulation that may protect soft tissue and allow establishment of a tissue collar overlapping the bone implant interface with decreased pocket depth 37. The increased pocket depth with computer guided flapless approach concurred with the findings of another study.38

Keratinized mucosal width significantly increased from baseline to 3 months for flap group and insignificantly decreased for flapless group. Also flap group reported significant increase in the width of keratinized mucosa than flapless group. The increased keratinized mucosal width in the flap group could be attributed to the crestal incision and mucosal reflection and readaptation of the flap around the abutments to be in a more apical position. This brings the keratinized mucosa which present on the crest of the ridge to a more apical position similar to apical positioned flap which help in increase the width of keratinized mucosa39. The increase width of keratinized mucosa with flap surgery concurred with the results of other studies 18, 19, 40. The insignificant decrease in the width of keratinized mucosa for flapless group concurred with the results of Wang et al. 40 who reported limited reduction of keratinized mucosal width with stable keratinized mucosa when minimal invasive flapless surgical approach was used for single implant placement in non-diabetic patients.

For both groups, crestal bone loss significantly increased with time. This time dependent upon loss is usually occur as a result of one the healing, bone maturation and bone reaction to increased load41. Moreover, the immediate loading of the implants with acrylic prosthesis may increase implant micromotions and could be responsible for increased bone resorption in both groups42. It has been proposed that diabetes leads to decreased bone turnover, with reductions in both resorption and formation 43. However, the mean marginal bone loss after one year for both groups did not exceed 1.2mm which located within the normal limit of marginal bone loss reported to occur in the first year for non-diabetic patients 25. This suggests that influence of diabetes on bone and osteoblast function was shown to improve by good glycaemic control44. Furthermore, the high initial stability of the implants caused by increased bone density in mandibular bone together with splinting the implants with rigid fixed restoration could be responsible for maintaining the bone loss values within the normal limits. Another factor that could contribute to the high survival rate of the implants in both groups and the reduced bone loss is the status of opposing occlusion. The factors that opposing prosthesis was complete dentures in both groups may reduce the impact of occlusal forces to the mandibular fixed prosthesis and could contribute to reduced bone loss.

Flap group showed significant higher crestal bone loss than flapless group. The increased crestal bone loss in the flap group may be attributed to flap elevation, microbial contamination, mucosal stripping, and interferences with blood supply to mucoperiosteum caused by flap reflection compared to flap surgery45-47. The decreased blood supply to the mucoperiosteum may decrease oxygen tension and activities osteoclastic activity leading to increase the implant crestal bone loss45-47. In contrast, flapless surgery reduces surgical time, reduces the need of sutures, reduces postoperative swelling and discomfort. On the other hand, the flapless procedure reduced surgical time, have no sutures, minor or no swelling, with minimal postoperative discomfort48. Moreover, the mucoperiosteum is not disturbed which may enhance bone remodeling,
with slight changes in marginal bone levels compared with flap surgery. In agreement with our finding, systematic review reported that flapless cortical better short term clinical and radiographic outcomes compared to flap surgery for non-diabetic patients. Conversely, another study found no difference in marginal bone loss between flap and flapless surgical approach for conventionally loaded implants in controlled diabetic patients. The difference in the results would be attributed to the different loading protocol used in each study.

The limitations of the present study include the small patient sample size and short follow-up period. Randomized controlled trials with large patient sample and the longer follow up period are still needed to confirm the findings of this study. Also studying different opposing occlusion such as natural teeth or fixed partial dentures still need to be investigated.

CONCLUSION

Within the limitation of this short-term randomized trial, computer guided flapless surgical approach may be recommended for implants placed in edentulous type 2 controlled diabetic patients and immediately loaded with mandibular fixed prosthesis as it was associated with favorable peri-implant soft tissue response and crestal bone loss compared to implants placed with flap surgery after one year.

REFERENCES


