Clinical Evaluation of the Effect of Implant-Supported and Implant Retained Distal Extension Removable Partial Dentures on the Supporting Structures

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Abstract

**AIM:** The objective of the study is to evaluate and compare the effect of implant-supported versus implant-retained removable partial denture (RPD) restoring Kennedy's class I cases on the supporting structures by measuring modified gingival index, probing depth for both abutment teeth and implant, as well as measuring marginal bone loss for implants.

**SUBJECTS AND METHODS:** Twelve patients; who had Kennedy class I mandibular partially edentulous ridge extending distal to the first or second premolar, all patients have full dentition in the maxillary arch or restored by fixed restoration, all patients were selected from the outpatient clinic, Prosthodontics Department, Faculty of Dentistry, Minia University. The patients were classified into two equal groups; according to the implant superstructures either dome-shaped abutment or ball and socket attachment. Each patient of both groups had two implants in the second molar position (one in each side) and received RPD of the same design. Patients were followed up for 1 year clinically and radiographically. Group I: six patients received RPD supported by dome-shaped short abutments. Group II six patients received RPD retained by ball and socket attachments.

**RESULTS:** Results revealed that there was no statistically significant difference between the two groups regarding marginal bone loss for implants.

**CONCLUSION:** The use of dome-shaped abutment or ball and O-ring attachment have the same effect on gingival index and probing depth around natural abutments and implants in Kennedy class I mandibular situations, The use of dome-shaped abutment produce less marginal bone loss than ball abutment.

Introduction

Removable dentures remain an essential prosthetic consideration in many conditions of oral rehabilitation, especially when restoring edentulous spaces posterior to the anterior remaining teeth [1].

Kennedy class I presents challenges for clinicians, as these dentures require support from the teeth, the mucosa and the underlying residual alveolar ridges. In particular, the distal extension removable partial denture (RPD) is subjected to vertical, horizontal and torsional forces that may have adverse effects during functional and para-functional activities [2].

The problems could be attributed to the absence of the posterior abutment [3]. Since the difference in displacement between the mucosa and the periodontal ligament of last standing abutment was estimated to be up to 25 times [4]. Consequently, when functional pressure is applied to the distal extension base RPD, the resultant forces are extremely damaging to the abutment teeth and must be controlled if clinical treatment is to be successful [5].

The use of distal implants to support and retain RPDs has been reported in the literature to minimize dislodgement, improve aesthetics and mastication as well as increase patient satisfaction in cost-effective manner [6], [7], [8].

The use of dental implant as a distal abutment can convert a distal extension RPD from a tooth and tissue-supported prosthesis to a tooth and implant-supported and retained prosthesis. A posteriorly placed implant provides a definite stop and stability and eliminates the problems often associated with a tooth and tissue-supported distal extension RPD [9], [10].

Dental implants reported in many studies a high success rate when they are combined with a partial denture in free end cases, especially in the mandible which can be a suitable treatment option because the pressure applied to tooth abutments can be decreased [11].
An implant-retained RPD provides patients with unique service and benefits not possible with more conventional treatment options [12]. It has been noted that the placement of dental implants in an edentulous space provides the biologic advantage of reducing bone resorption, also provides distinct biomechanical advantage in reducing the effect of the reciprocal arm of a conventional RPD and improving the fulcrum line position, as well as offering superior retention and elimination of unaesthetic clasps in the aesthetic zone. Adding flange on such prosthesis also allows for restoring facial esthetics and extraoral soft tissue support lost from advanced ridge resorption [13].

Combination of natural tooth and implant-supported RPD was reported. This design of connection allows stress control on the fixtures and natural abutments, provides strength, esthetics, fulfilling patients’ desire and increasing the long-term prognosis for the remaining teeth. The only disadvantages are that it is “removable” and some patients may not tolerate this type of prosthesis and the cost factor [14].

Several authors have been debated to connect natural teeth and implants together in a fixed partial denture [15], [16], [17], [18], fears of possible negative effects resulting from splinting natural teeth with osseointegrated implants have always played a superior role in not accepting this option as a mode of treatment and the majority of implant dentists worldwide avoid splinting the two in a fixed prosthesis [19].

Using ball attachment and O-ring to retain mandibular RPD on a bilateral single molar implants in cases of Kennedy class I cases helped to support and retain the mandibular RPD and present a cost-effective treatment [10].

The dome-type attachments have the function to allow displacement or rotation of the overdenture during function [20].

The aim of the study was to compare between two types of implant superstructures, namely dome and ball attachments regarding their effect on the longevity of both implants and natural abutments in cases restored by RPDs.

Subjects and Methods

The study and sample size

The study was a comparative study. Superstructures were randomly distributed for patients in both groups. Sample size was calculated to be 12 implants in each group.

Patient selection

Twelve patients with partially edentulous lower arch Kennedy’s class I were selected from the outpatient clinic, Prosthodontics Department, Faculty of Dentistry, Minia University. All the patients had their first or second premolar as posterior abutments, had full dentition in the maxillary arch, free from any systemic disorder that may affect dental implant rigid fixation to the bone and with average ridge form and height to receive implant of (diameter 3.7 mm and length 8.5 mm). The patients were informed by all procedures of our study. Only motivated patients who showed cooperation participated in the study and an informed consent were assigned, also the approval of Research Ethics Committee of the Faculty of Dentistry Minia University was obtained.

Clinical procedures

Stage 1: Construction of acrylic partial denture for the lower arch.

Each patient had received lower acrylic partial denture with conventional method.

Stage 2: Fabrication of surgical guide and radiographic examination.

A customized surgical guide was fabricated using CAD/CAM technology through the data obtained from the cone-beam computed tomography (CBCT), captured images by CBCT were imported into viewing software then sent for fabrication of the guide.

Stage 3: Surgical procedure

The surgical procedures were performed in one step under aseptic conditions. The implants were located at the second molar site in the edentulous area of the mandible. All patients were anaesthetized by local nerve block and infiltration at the site of the surgical field. The incision was made at the crest of the ridge, crestal flap was achieved by a sharp scalpel number 15 blades. The scalpel was pressed firmly to bone and the incision was made once for clean-cut or by tissue punch. Pilot drill was pointed down through the hole reaching down to the alveolar bone and punching it to make a point that acts as a guide for drilling. The sterilized surgical stent was placed securely in the oral cavity with its hole corresponding to the planned implant position. Drilling was done through the stent’s hole with light intermittent finger pressure using sterile saline solution irrigation. Drilling was performed starting with the pilot drill (2.3 mm) in diameter then intermediate drill (2.8 mm D) was used and driven to the full depth of the planned implant, and finally with (3.5 mm D). The paralleling rod was inserted into the drill hole to make sure that the implant was in its right position. The implant was removed from the sterile pack with the fixture mount and was inserted to the osteotomy till the implant collar by handpiece then manual using ratchet wrench. A surgical cover screw corresponding to the...
The mouth preparations were made as the following:

- Mesial occlusal rest seat for the 1st or 2nd premolar (the main abutment) adjacent to the edentulous area
- Canine rest or distal occlusal rest seat adjacent to the main abutment

Both groups were included for constructing the conventional Cobalt-Chromium RPD after the healing period with the same design RPD with the conventional way. Maxillary and mandibular preliminary impressions, final impression, duplication of the master cast was performed to obtain a refractory model for waxing up the partial overdenture framework., try in of metal framework (Figures 3 and 4) Jaw relation registration, partial overdenture try in was made with normal acrylic teeth, flasking, finishing and polishing then insertion.

- Mesial occlusal rest for the 1st or 2nd premolar (the main abutment) adjacent to the edentulous area

Mouth preparations

All patients had to complete phase I therapy including supra and subgingival scaling, root planning and curettage. Proper oral hygiene instructions including the appropriate brushing technique and interdental cleaning procedures were implemented. This phase also included minor occlusal adjustments when needed, occlusal analysis and correction for occlusal reconstruction were made either by selective grinding, obturation of carious lesions, crowning or removal of overhanging margins and uncovering the implant by removing the tissue above it using tissue punch. The surgical cover screw was removed using the screwdriver and the implant abutment was screwed into the implant. Inside the patient mouth for Group I they received dome-shaped abutment, while Group II received ball abutment (Figures 1 and 2).

Stage 4: Prosthetic procedure

The diameter of the implant was placed and tightened into position with hand screwdriver. A hand debridement and irrigation of the surgical site was carried out. The flap was repositioned around the implant and sutured by interrupted sutures using 3-0 silk sutures. Surgical technique was repeated for the other side either by surgical flap or using tissue punch.
pick up technique using monomer free- cold cure acrylic resin material (Figure 6) Partial overdenture was inserted into the patient’s mouth and was checked for retention, stability and support. Instructions were given to the patient about how to use and clean the partial denture.

**Clinical evaluation**

Each case was evaluated clinically at the time of denture insertion, 3, 6 and 12 months later. Clinical evaluation included assessment modified gingival index, probing depth around the main abutment and implant abutments.

**Modified gingival index**

Modified Gingival index was measured for main abutments and implant in Groups I and II (Table 1 and 2), according to Lobene and Weatherford (1986) [21] for implants in Group I and II. The circumference of the gingival margin was divided into buccal, mesiobuccal, distobuccal, and lingual [22], [23]. Each of them was scored from 0 to 4 according to the following:

| Score | Description |
|-------|-------------|
| 0     | Absence of inflammation, normal gingival. |
| 1     | Mild inflammation, slight color change, little change in texture of any portion of, but not entire, marginal, or papillary gingival unit (localized). |
| 2     | Mild inflammation, slight color change, little change in texture involving the entire or papillary gingival unit (generalized). |
| 3     | Moderate inflammation, redness, and edema. |
| 4     | Severe inflammation, marked redness, edema, ulceration, and spontaneous bleeding. |

**Probing depth**

It refers to the distance from the gingival margin to the bottom of clinical pocket according to Glavind and Loe 1967 [24]. Moreover, gingival sulcus depth around implants [25].

Probing depth was evaluated using pressure-sensitive color coded plastic probe* with perceptible click and divided into 3/5/7/10 mm.

a. Periodontal probing depth for the main abutment (Figure 7 and Table 3).

b. Peri-implant probing depth around implant (Figure 8 and Table 4).

The measurements were taken from each of the four axial surfaces of main abutments buccal, mesiobuccal, distobuccal and lingual, for Groups I and II. Peri-implant probing depths were measured according to Atassi [17] at four surfaces of implants:

**Radiographic evaluation of marginal bone loss**

For both groups, marginal bone level around the implants was examined using CBCT. Marginal bone level
was measured using OnDemand3D Application software (Soredex-Scannora® 3D). The distance from the marginal bone to the apex of the implant was calculated in millimetres using straight line tool of the system. The mesial and distal bone heights were measured on the sagittal view screen, while the buccal and lingual bone heights were measured on the coronal view screen using the linear assessment OnDemand3D software. The mean value of readings was taken, tabulated and statistically analyzed (Table 5).

Results and Statistical Analysis

Data were fed to the computer using IBM SPSS software package version 24.0. Qualitative data were described using number and percentage. Comparison between different groups regarding categorical variables was tested using Chi-square test.

Quantitative data were described using mean and standard deviation for normally distributed data while abnormally distributed data were expressed using median, minimum and maximum. For normally distributed data, comparison between two independent population was done using independent t-test while more than two population were analyzed F-test (ANOVA) to be used. The results of this study were represented by tables. The significant level was set at P ≤ 0.05.

The type of that study is randomized controlled trial with calculated samples. Randomization in this study the investigators randomly distribute the patients with the two different types of abutments before any surgical procedures.

Discussion

Selection of implant-retained overdenture treatment for the posterior edentulous mandible can provide both the patient and clinician with several advantages [26]. Mandibular implant-retained overdenture treatment has significantly increased the scores for retention and stability of the denture, masticatory function and general denture satisfaction. Furthermore, it may have favorable psychological and social effects on the patient [27]. Furthermore, the pressure applied to tooth abutments can be decreased as such; however, implant-abutment tooth attachment type is a matter of discussion [11].

In rigid attachments, higher stress accumulation can cause higher bone loss and increase the frequency

| Table 1: Modified gingival index for abutment teeth in Group I (dome-shaped abutment) and II (O-ring attachment) at different period of follow-up |
|---|---|---|---|---|---|
| Group | Insertion | 3 months | 6 months | 12 months |
| | No % | No % | No % | No % | No % |
| Group I | | | | | |
| Grade 0 | 12 | 100.0 | 10 | 100.0 | 12 | 100.0 |
| Grade 1 | 0 | 0.0 | 2 | 16.7 | 0 | 0.0 |
| Grade 2 | 0 | 0.0 | 0 | 0.0 | 0 | 0.0 |
| Grade 3 | 0 | 0.0 | 0 | 0.0 | 0 | 0.0 |
| Grade 4 | 0 | 0.0 | 0 | 0.0 | 0 | 0.0 |
| P1 | 0.365 N.S. | 1.00 N.S. | 1.00 N.S. |
| Group II | | | | | |
| Grade 0 | 12 | 100.0 | 8 | 66.7 | 8 | 66.7 |
| Grade 1 | 0 | 0.0 | 4 | 33.3 | 2 | 16.7 |
| Grade 2 | 0 | 0.0 | 0 | 0.0 | 2 | 16.7 |
| Grade 3 | 0 | 0.0 | 0 | 0.0 | 0 | 0.0 |
| Grade 4 | 0 | 0.0 | 0 | 0.0 | 0 | 0.0 |
| P1 | 0.042* | 0.040* | 0.365 N.S. |
| P2 | 1.00 N.S. | 0.452 N.S. | 0.40 N.S. | 0.365 N.S. |

Results: P1 comparison between time of insertion with other time of follow up in the same group. P2 comparison between the two groups at the same time. *Significant difference. N.S.: Not significant. p value was calculated by using the Chi-square test.

Table 2: Modified Gingival index in Group I (dome-shaped abutment) and II (O-ring attachment) for implant at different period of follow up

| Group | Insertion | 3 months | 6 months | 12 months |
|---|---|---|---|---|
| | No % | No % | No % | No % |
| Group I | | | | |
| Grade 0 | 10 | 83.3 | 10 | 83.3 | 12 | 100.0 |
| Grade 1 | 2 | 16.7 | 0 | 0.0 | 0 | 0.0 |
| Grade 2 | 0 | 0.0 | 2 | 16.7 | 0 | 0.0 |
| Grade 3 | 0 | 0.0 | 0 | 0.0 | 0 | 0.0 |
| Grade 4 | 0 | 0.0 | 0 | 0.0 | 0 | 0.0 |
| P1 | 0.108 N.S. | 0.365 N.S. | 1.00 N.S. |
| Group II | | | | |
| Grade 0 | 10 | 83.3 | 10 | 83.3 | 10 | 83.3 |
| Grade 1 | 2 | 16.7 | 2 | 16.7 | 0 | 0.0 |
| Grade 2 | 0 | 0.0 | 0 | 0.0 | 2 | 16.7 |
| Grade 3 | 0 | 0.0 | 0 | 0.0 | 0 | 0.0 |
| Grade 4 | 0 | 0.0 | 0 | 0.0 | 0 | 0.0 |
| P1 | 0.108 N.S. | 0.365 N.S. | 1.00 N.S. |
| P2 | 0.365 N.S. | 1.00 N.S. | 0.365 N.S. |

Results: P1 comparison between time of insertion with other time of follow up in the same group. P2 comparison between the two groups at the same time. *Significant difference. N.S.: Not significant. p value was calculated by using the Chi-square test.

Table 3: Probing depth in Group I and II at different periods of follow-up

| Abutment teeth | Insertion | 3 months | 6 months | 12 months |
|---|---|---|---|---|
| Group I | | | | |
| Rang | 1.5–2.3 | 1.8–2.5 | 2.2–2.5 | 2.6–3.5 |
| Mean | 1.78 | 2.09 | 2.35 | 2.94 |
| SD | 0.32 | 0.03 | 0.16 | 0.39 |
| P1 | 0.006 | 0.031* | 0.022* |
| Group II | | | | |
| Rang | 1.5–2.3 | 2.1–2.5 | 2.1–2.8 | 2.1–3 |
| Mean | 2.02 | 2.41 | 2.41 | 2.77 |
| SD | 0.26 | 0.14 | 0.21 | 0.27 |
| P1 | 0.113 | 0.044* | 0.027* |
| P2 | 0.254 | 0.075 | 0.311 |

Results: P1 comparison between time of insertion with other time of follow up in the same group. P2 comparison between the two groups at the same time. *Significant difference. N.S.: Not significant. p value was calculated by using ANOVA test.
of prosthetic complications such as fatigue fracture and screw loosening compared to resilient attachments. In non-rigid attachments may cause dental intrusion. Intrusion of tooth abutment decreases support and increases cantilever stresses in implants and the supporting bone [28].

However, in the attachment of natural teeth and implant by a RPD, intrusion of natural teeth does not seem to be problematic because permanent attachment of the prosthesis to tooth and implant does not exist [29].

Implant used in this study was placed into the second molar area, to avoid the posterior rotation of the partial overdenture on the implant’s abutment as a fulcrum [30], [31].

Two-stage surgery for the implant placement primarily have been considered by many authors for two reasons; to prevent early failure due to loading and to prevent infection [32], [33]. The number of stages for implant placement depends on the primary stability of the implant and the quality of the bone.

Choosing dome-shaped abutment was to reduce the load on the implant by permitting slight rottational movements [34]. O-ring ball abutment is an excellent method for increasing the retention and stability of such dentures and has several advantages, including ease of use, hygiene, and maintenance, and low cost [35].

Success criteria of osseointegration included: no clinical mobility; no peri-implant radiolucency or infection; no complaints of pain, neuropathy, or paresthesia; and crestal bone loss that does not exceed 1.5 mm at the end of the 1st year [36].

Accordingly, patients were scheduled for follow-up at partial denture insertion after 3, 6, and 12 clinically compare modified gingival index, probing depth around the abutment teeth and implant, as well as evaluation of marginal bone loss around the implant radiographically by using CBCT.

Measurements of periodontal probing depths around main abutments and peri-implants probing depths were made using the pressure-sensitive probe with a perceptible (click), to eliminate excessive forces generated by using conventional types [25], [37].

The results of the clinical evaluation of the present study of modified gingival index for abutment teeth demonstrated a non-significant change in all follow-up periods in group I. Despite continuous patient motivation for oral hygiene measures for all patients in both groups, Group II showed significant decrease in the percentages from grade 0 after 3 and 6 months this could be due to careless handling for oral hygiene measure in this group. These results agree with result conducted by Augustin et al., that found there are significant differences between study and control groups regarding gingival index, they attributed the cause due to poor oral care [38]. Continuous and routine follow-up of the implant patient with periodic assessment of plaque and calculus for both groups, after 12 months there was non-significant change from time of insertion.

The comparison between the results of modified gingival index for implant between the two groups showed a non-significant difference at any given period of follow-up. This result was in agreement with Alam-Elddeen [39], who concluded that regardless the type of the retainer used with natural abutments, gingival index had no significant differences effect on implants and natural abutment teeth.

### Table 4: Peri-implant probing depth in Group I and II at different periods of follow-up

| Abutment teeth | Insertion | 3 months | 6 months | 12 months |
|---------------|-----------|----------|----------|----------|
| Group I       | Rang      | Mean     | SD       | P1       | Group II   | Rang      | Mean     | SD       | P1       |
|               | 1.5–2.4   | 1.79     | 0.27     | 0.107    | 0.038*    | 0.034*    |
|               | 1.8–2.5   | 2.03     | 0.29     | 0.18     | 0.045     |
|               | 2.2–2.5   | 2.40     | 0.18     | 0.038    | 0.045*    |
|               | 2.1–3.5   | 2.51     | 0.18     | 0.045    |
|               | 0.211     | 0.26     | 0.18     | 0.038    |
|               | 0.65      | 0.26     | 0.18     | 0.038    |
|               | 2.1–2.6   | 2.47     | 0.26     | 0.038    |
|               | 2.5–3.1   | 2.88     | 0.26     | 0.038    |
|               | 1.5–2.4   | 2.08     | 0.21     | 0.096    |
|               | 2.32      | 0.16     | 0.072    |
|               | 2.47      | 0.26     | 0.038    |
|               | 0.32      | 0.26     |
|               | 2.1–2.8   | 2.47     | 0.26     |
|               | 0.32      | 0.26     |
|               | 0.107     | 0.072    |
|               | 0.038*    | 0.072    |
|               | 0.038*    |

*P1 comparison between time of insertion with other time of follow-up in the same group. P2 comparison between the two groups at the same time. p was significant if <0.05. * Significant difference. N.S.: Not significant. P1 value was calculated using ANOVA test. P2 value was calculated by using student t-test.

Furthermore, this result was in agreement with Saleh [40], who study the effect of two different clasps, Aker clasp and reverse Aker clasp on tissue health around abutment teeth and implants assisting mandibular bilateral removable partial overdenture, the results were no statistically significant differences between results in regard oral tissue health around the abutment teeth and implants located in the first molar area.

Results of clinical evaluation of probing depth of abutment teeth for both groups, there was insignificant increase in probing depth after 3 months while there was significant increase in 6 and 12 months follow-up periods, there was no significant difference between the two groups at different periods of follow-up.

A significant increase in the measured probing depth of abutments in both groups overdentures was detected throughout the study period, that was shown., this change is considered within the normal reported range of change in probing depth (0.5–3 mm) [41].

These results agreed with Brill et al that stated that increase in probing depth could be correlated
to alteration in the oral ecology and to the stresses produced by introducing partial overdentures to the oral cavity [42].

Results of clinical evaluation of peri-implant Probing depth for Group I showed that there was insignificant increase in peri-implant probing depth after 3 months, while there was significant increase after 6 and 12 months in comparison to time of insertion. For Group II there was insignificant increase after 3 and 6 months, while there was significant increase after 12 months.

While comparing the results of both groups together there was insignificant difference regarding peri-implant probing depth at different period of follow up. Group I showed that peri-implant probing depth was less insignificantly than Group II. This may be due to difficulty to obtain proper oral hygiene and more plaque accumulation in ball abutment (Group II) than dome-shaped abutment in Group I. Several studies have indicated that successful implants allow probe penetration of approximately 3 mm (Nishimura et al.; 1997) [43].

Authors concluded that increased pocket depth could be correlated with a higher degree of inflammation of the peri-implant mucosa [44] but not necessarily with peri-implant bone loss [45].

Results of radiographic evaluation of marginal bone loss for both groups showed there was a significant increase in all periods of follow-up, while when comparing the two groups together there were significant differences after 3 and 6 months, where Group II implant-retained group with ball abutment showed significant increase in mean marginal bone loss than Group I the supported group with dome-shaped abutment this may be related to the presence of space between the components of the resilient ball attachment, which may permit free vertical rotation of the overdenture during function with concentration of diverse forces on the residual ridge and the implant. This may be also attributed to that the presence of effective vertical implant support that may decrease the rotation potential of denture base during functional loading [46], [47], [12], our results agreed with Abdou ELsyad who stated that implant-supported partial overdentures appear to be associated with reduced posterior mandibular ridge resorption when compared to implant-retained partial overdentures [48].

The marginal bone loss was statistically significant, after 12 months in both groups and insignificant between both groups. Mitrani et al., suggested that any mechanical wear may occur at the interface between the implant and the denture base will allow the opportunity for the rotation potential to occur, consequently, the implant overloading may occur during function this will lead to marginal bone resorption in both groups [8], [49].

Conclusion

Within the limitations of this study, the following conclusions could be drawn:

- The use of dome-shaped abutment or ball and O-ring attachment have similar effects on natural abutments and implants in Kennedy class I mandibular situations
- There are no significant differences between the two groups on, the modified gingival index, or the increase of probing depth around the natural abutments or dental implants in such cases
- Significant increase in marginal bone loss in group II implant-retained with ball abutment than Group I implant-supported with dome-shaped abutment after 3 and 6 months of denture insertion

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