Minimaterized Electromagnetic Device Abutment Improves Stability of the Dental Implants

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Dental implant longevity is largely determined by osseointegration. Factors pertinent to the success and reliability of dental implants include primary stability, bone to implant contact formation, quality and quantity of residual bone. Ideal implant materials should display advantageous biocompatibility and mechanical characteristics.1,2 Currently, implants constructed of titanium are ubiquitous in dentistry secondary to their favorable characteristics including properties that allow the formation of an osseous integration of the implant. However, implants currently on the market customarily require a 2 to 6-month period without loading of the device to allow adequate time for osseointegration. Current literature demonstrates that attenuating the unloading time postimplantation increases failure rate by 2 or 3 times, especially in unsplinted dental devices.3–5 Osseointegration or union of alveolar bone and dental implant is one of the primary goals of dental implantation. Thus, multiple therapies targeting enhancing osseointegration or reducing time to achieve osseointegration have been proposed.

Pulsed electromagnetic field (PEMF) is one modality commonly used to stimulate bone generation throughout various clinical settings including orthopedic surgery. Pulsed electromagnetic field has been shown to primarily effect vascular generation, formation, and neovascularization.6,7 In dentistry PEMF stimulation may be a useful tool to encourage bone formation, ingrowth of bone on dental implants, and increased bone stock. This may help decrease time to osseointegration and allow patients to return to normal loaded eating activities sooner (Table 1).

A recently published study by Barak et al8 reported that in rabbits PEMF devices stimulated early osseointegration and ingrowth of bone onto dental implants by more than 3 times. It was therefore hypothesized that dental implant devices locally generating PEMF stimulation would significantly stimulate bone growth and increase osseointegration around the implant PEMF devices themselves.

This study was designed to retrospectively assess the effects of the miniaturized electromagnetic device (MED) on the implants stability for the first time in human subjects, in a prospective case controlled series (Fig. 1).

METHODS

Subjects

The local institutional review board approved this retrospective cohort study. Medical records and radiographic images of patients who presented to our center between 2014 and 2016 were reviewed. Twelve partially edentulous patients (7 females) in the mandible or the maxilla with a buccolingual ridge width of at least 6 mm and with opposing dentition of natural teeth or tooth-
implant-supported fixed reconstructions were assessed. Twelve Magdent healing caps (MED) and 16 regular control healing caps were inserted.

The mean age at implant surgery was 49 years (range 34–69). Prior to inclusion in the study, all patients were examined according to a standardized protocol with clinical and radiographic examination.

The following inclusion criteria were used:
- controlled oral hygiene
- absence of any lesions in the oral cavity
- sufficient residual bone volume to receive implants of at least 3.7 mm in diameter and 11.5 mm in length.

### Surgical Procedures

Surgical procedures were performed in outpatient clinics under local anesthesia. Implants (Alpha-Bio Tec Ltd, Petah Tikva, Israel) were placed according to standard protocols. To achieve the highest standardization of the initial bone-level situation, each implant was placed at the bone level. Antibiotics will be prescribed according to the standard practice at the clinic. After implantation, a MED was applied to the implant. Electromagnetic stimulation was administered continuously for 24 hours a day using a MED healing cap (Magdent Ltd, Tel Aviv, Israel). Identical devices and the same follow-up protocol were used for both control and PEMF cases.

### Postoperative Procedures

Patients were educated in postoperative dental hygiene and care procedures. They followed standard instructions for implant patients, including instructions not to brush in the treated area and to rinse twice per day for 1 minute with 0.1% to 0.2% chlorhexidine digluconate. Sutures were removed 2 weeks after surgery. The extent of healing and any local inflammation of the soft tissue around the study implant were assessed.

### Resonance Frequency Analysis

During healing, resonance frequency analysis (RFA) were performed, starting with an assessment immediately following implant installation (Week 0) and then at following weeks: mandible—after 2w and 4w, maxilla: 2, 4, and 8w, postoperatively. The wireless Ostell device was used for this study (Ostell Mentor, Integration diagnostics AB, Sävedalen, Sweden). Resonance frequency analysis measurements were carried out in 2 perpendicular directions (mesio-distal [M-D] and oro-facial [O-F]), twice in each direction. Resonance frequency analysis was performed at implant placement and abutment connection and an implant stability quotient value was given for each implant.

### Statistical Analyses

Descriptive statistics were calculated with measures of central tendency (mean and median), measure of dispersion (standard deviation), and 95% confidence interval. Data from 28 implants were analyzed using SPSS21.0 software (SPSS Inc, Chicago, IL), applying the Wilcoxon test according to the nature of the data. The significance level was established at $P < 0.05$.

### RESULTS

Twenty-eight dental implants were included in the current study. Following implantation, the patients reported no or only minor discomfort at the surgical sites.

No statistical significant difference in stability was found between the mandibular implants at the first 15 days. Maxillary implants stability was significantly higher with MED healing cups compared with controls at the day 15 postimplantation (66.2 vs 62.1, $P = 0.0008$). Resonance frequency analysis test performed at day 30 postimplantation demonstrated a significantly higher stability results in MED as compared with the control 73.5 ± 3.2 versus 66.7 ± 4.8 at mandible and 74 ± 1.7 versus 65 ± 2.3 at maxilla. At the day 50 postimplantation, RFA test revealed markedly higher stability of the maxillary implants with MED active healing caps compared with nonactive 75.4 ± 5.1 versus 68.5 ± 8.5, respectively.

### DISCUSSION

The principal outcomes of this study demonstrated improved MED implant at 30 and 50 days postimplantation. Moreover, maxillary implants stability was significantly higher in MED healing cups than in control cups at the early 15 day postimplantation time point.

Establishing implant stability and long-term maintenance of implant stability are imperative for clinical success of dental implants. Contact area between bone stock and implant, implantation technique, and bone quality are all factors influencing osseointegration and successful implantation outcomes. Initial mechanical implant stability is largely mediated by contact area, and thus friction or macro-retentions between the implant and its insertion...
site. Later biological stability is a product of the boney ingrowth and tissue integration which occurs after allowing ample time for healing.8

Dental implant failures are often classified as early failures and late failures. Early failures occur at abutment connection surgery, whereas late failures occur secondary to occlusal loading after healing. This classification and knowledge of the time course for healing suggests that early and late failures are associated with different mechanisms. Early failure is the result of inadequate bone to implant contact area. Bone healing and ingrowth in these cases may be impaired.3 Other factors that have the potential to influence bone healing include poor oral hygiene, risky behaviors, and systemic diseases. Additionally, radiotherapy, surgical conditions, and medication usage may all play a role in the ultimate clinical outcome of dental implants.

To achieve successful dental implant therapy achieving osseointegration in the shortest time possible is imperative.16 Targeted treatments have been proposed to improve and accelerate osseointegration at the dental implant–bone interface. Low-level laser therapy is one technique studied in animals which was shown to increase bone–implant contact on a microscopic level.20,22 Furthermore, low-level laser therapy was shown to enhance the association between native bone and prosthetic implant18 increases the percentage of calcium and phosphorus in local bone stock12 and increases production of OPG, RANKL, and RANK in the local environment.13 These results are supported by recent literature of dental implants in a rabbit model.17 However, there are no clinical studies reporting statistically significant macroscopic results.16,17

Research has shown that electromagnetic stimulation promotes osteogenic activity and multiple studies have demonstrated the effectiveness of externally applied electromagnetic stimulation in various clinical settings including dentistry.18 These studies have demonstrated reduced time to osseointegration surrounding dental implants when external source of electromagnetic stimulation is applied. Recent work using a rabbit model has shown that PEMF devices stimulate early bone formation around dental implants and produce higher peri-implant BIC as well as increased bone mass with as little as 2 weeks of therapy. This suggests accelerated osseointegration with PEMF stimulation in the setting of dental implants.8

This study has several limitations including the small sample size. Furthermore, Ostell was employed as the sole instrument used for functional assessment, using other instruments could have strengthened the results.

This is the first clinical study examining the effects of dental implant devices with internal PEMF stimulators. This device produces an electromagnetic field around the implant similar to that of external devices. The advantage of this device is that the effective electromagnetic field is directed exclusively around the dental implant and there is no need to use an external PEMF source. This allows for continuous activation of the PEMF device for 24 hours a day with the goal of achieving improved implant stability in the early postimplantation stage. Moreover, patient compliance with external hardware does not interfere with the treatment. Further randomized controlled trials in larger cohorts assessing the clinical outcomes after implantation of the MED device are warranted.

REFERENCES