Single-crown restorations in premolar–molar regions: short (≤ 6.5) vs longer implants: retrospective cohort study

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Abstract

Purpose: To compare the survival, changes in marginal bone level and prosthetic complications rate of short (≤ 6.5 mm) and longer implants (≥ 7.5) supporting a single-crown restoration in the maxillary/mandibular premolar or molar region.

Methods: This cohort study was conducted following the STROBE statement recommendations for observational studies. Clinical outcomes of 88 short implants in 78 patients and 88 long implants in 88 patients were examined. All the implants had been placed by the same surgeon and restored following the same prosthetic concept; using a transepithelial abutment (intermediate abutment) and a screw retained restoration.

Results: All the implants were in function after the follow-up period since insertion (median: 31 months; range 11 to 84 for SiG vs median: 35 months; range: 6–117 for CG; p = 0.139). No statistical differences (p = 0.342) were observed related to prosthetic complications (screw loosening 2/88 vs 5/88 CG, ceramic chipping 1/88 vs 0/88, temporary crown resin chipping 1/88 vs 0/88 for SiG and CG, respectively) or related to marginal bone level (Mesial or Distal MBL ≥ 2 mm in 1/88 implants for SiG vs 3/88 for CG; p = 0.312).

Conclusions: Within the limitations of this study, no survival differences have been observed between short implants and longer implants in single-crown restorations in posterior maxilla/mandible.

Keywords: Dental implants, Short implants, Marginal bone loss, Crown-to-implant ratio

Introduction

Tooth loss leads to changes in the integrity of the alveolar bone and soft tissues [1, 2]. The healing of extraction sockets leads to histological and dimensional changes in the remaining alveolar ridge [1–4]. Progressive atrophy of the alveolar process begins at this moment both horizontally and vertically. Resorption of the alveolar ridge occurs due to a combination of different factors, such as loss of the periodontal ligament (and lack of the vascularization of the bundle bone), loss of function (and the stimulus on the bone), fractures in the alveolar wall during extraction or the subsequent occurrence of infectious processes [1–4].

The physiological post-extraction resorption (RRR, Residual Ridge Reduction) [5] can hinder the implant rehabilitation of missing teeth. It not only affects the available volume, but also the density of the remaining bone [5]. These changes could hamper the subsequent placement of an implant in an optimal position or affect the esthetic results. [6]

Clinicians are often faced with the challenge of treating patients with severe vertical bone atrophy. Different treatment options allow the use of standard implants in the posterior region where the nerve canal or the
maxillary sinus limit the residual bone height, such as guided bone regeneration (GBR), maxillary sinus grafting, inlay or onlay bone graft, distraction osteogenesis, nerve lateralization or others [7, 8]. These techniques require greater knowledge and surgical skills on the part of the professional and potentially increase the complication rate [9–12]. Recent advances in implant design and sizing (short, extra-short implants) have provided new solutions and alternatives or have allowed optimization of existing ones [13–15]. In addition to facilitate the procedures, the use of short implants reduces the risk of reaching anatomical structures at the time of drilling, minimizes the number of surgeries, reduces the time and cost of treatment, and saves the need for bone augmentation. [7, 16–18]

In cases of vertical atrophy, the prosthetic space is larger thus increasing the crown-to-implant ratio. Former recommendations about the ideal proportions seem outdated in sight of the diverse clinical and biomechanical studies demonstrating the safety and good clinical performance of short and extra-short implants [19, 20]. Crown-to-implant ratios ranging from 0.9 to 2.2 did not influence the occurrence of technical or biological complications [21]. Indeed, it has been claimed that within the range of 0.6 to 2.36, the higher the crown-to-implant ratio, the less the peri-implant marginal bone loss (MBL) [20]. As the length of the implant is reduced, it has been suggested to increase the diameter to enhance the bone–implant contact and optimize the distribution of stress in the bone, particularly in cases of low bone density. [22]

Short implants are not limited to cases of limited available bone. Currently, short implants can be preferred to maintain as much pristine bone as possible, even when standard implants could be housed. [23, 24]

Some authors reported that short implants could have lower survival rates than standard implants [25, 26] but recent systematic reviews have shown that short implants had a better or equal performance compared with standard [27–29] and did not seem to have a significant influence on marginal bone loss [30]. Several systematic reviews and meta-analysis have been conducted to clarify the controversies on the clinical performance of short implants [25, 31–34] but their results should be individually interpreted with caution to assess the eventual presence of uncontrollable confounding factors in the included studies [31] (as studies including both splinted and non-splinted restorations, implants placed in both grafted or pristine bone, different implant designs and surfaces or different types of restorative design).

There is also another controversy regarding the classification of short implants that could result in a misinterpretation of the results. While some authors considered short implants those under 10 mm [35, 36], others considered a length under 8 mm [37] and more recently, others under 6.5 mm [38] or 6 mm. [39, 40] Recent evidence from clinical trials has shown the need for more studies and longer periods of follow-up before the recommendation of short implants to support single-crown restorations [19, 40]. The objective of this study has been the comparison between short implants and longer implants in terms of implant survival, marginal bone remodeling and prosthetic complications of single-crown restorations.

Materials and methods
Study design
The present unicentric observational retrospective study was conducted following the STROBE statement recommendations for observational studies and in compliance to the principles of the Declaration of Helsinki on clinical research involving human subjects. Before starting, the permission of the ethics committee was obtained from the Basque drug research committee.

Sample size estimation
A study of a continuous response variable of matched pairs of study subjects was planned. The sample size was estimated based on previous bone loss data at 12-month follow-up indicating that the difference in response of matched pairs was normally distributed with a standard deviation of 0.4320 [41]. If the true difference in marginal bone loss at 12 months of matched pairs was 0.13, 88 pairs of implants should be necessary to reject the null hypothesis that this difference in response is zero with a probability (power) of 0.8. The probability of type I error associated with this test of the null hypothesis was 0.05.

Patients
Data were retrospectively collected from 88 short (≤ 6.5 mm; 78 patients) and 88 longer implants (≥ 7.5 mm; 88 patients) randomly selected from a cohort of 16,780 implants placed from 2012 to 2019 at the same center (Eduardo Anitua Clinic, Vitoria, Spain). After sample size estimation, this cohort was divided in two groups (short implant group; SIG and control group; CG) and simple random sampling was conducted using the SPSS software, (SPSS for Windows, Version 15.0. Chicago, SPSS Inc) to select 88 implants form each group. The inclusion criteria were:

- Implants placed both in maxilla or mandible
- Implants supporting a single-crown screw-retained, restored using a transepithelial (intermediate abutment).
- Patients over 18 years
To address sources of bias, all patients included in this study had been previously treated by the same team, using the same implant system (UnicCa®, BTI Biotechnology, Vitoria, Spain) and the same surgical and prosthetic protocols. All the treatments were performed following the usual clinical practice of the participating center for the insertion and subsequent loading of short and standard implants in the mandible and/or maxilla.

Data collection methods
The outcomes measured were survival (presence of the implant at the last visit), MBL and prosthetic (technical) complications. The bone level assessment was performed vertically measuring the distance from the bone crest to the first bone-implant contact both mesially and distally. Measurements to estimate MBL were performed at loading time and at the time of the last available radiograph using the Sidexis software (Dentsply Sirona; York, US) and the length of the implant was used as calibrator. Among the technical complications screw loosening/break and ceramic/resin chipping were considered.

Other clinically relevant variables recorded were implant diameter, location, insertion torque, bone type, sex and age, residual bone height, type of antagonist teeth and the need for additional surgical techniques. Follow-up time was calculated since implant insertion (until last recall) and implant loading (until last recall).

The crown-to-implant ratio was determined by dividing the length of the crown together with the transepithelial (intermediate) abutment by the length of the implant. Residual bone height was measured from the bone ridge crest to the maxillary sinus/nerve canal at the implant position, using the radiography obtained previously to the surgery.

Information about smoking habit, alcohol intake, diabetes or hypertension was also retrieved from medical records. Bone type quality [42] was rated with the aid of computer software (BTI Scan, BTI Biotechnology, Vitoria, Spain).

Statistical analysis
A statistical analysis was performed using specialized software (SPSS for Windows, Version 15.0. Chicago, SPSS Inc). Categorical variables were expressed in absolute and relative frequencies. Continuous variables were expressed as median and range. Before statistical analysis, the normal distribution of the continuous variables was evaluated using the Saphiro–Wilks normality test. Statistical differences between categorical variables were performed by the Chi-square test, and statistical differences between dichotomous and continuous categorical variables were performed with the Mann–Whitney test. The effect of the crown to implant ratio on the marginal bone loss was assessed by linear regression analysis. The statistical significance was set at $p < 0.05$.

Results
The study included 176 dental implants placed in 166 patients that complied with the inclusion/exclusion criteria. The Short implant Group (SiG) was composed by 88 short implants ($\leq 6.5$ mm) placed in 78 patients (53 females; 35 males) and the Control Group (CG) was composed by 88 implants ($\geq 7.5$ mm) placed in 88 patients (52 females; 36 males). Further demographic data are presented in Table 1.

From the SiG, 71 implants were 6.5 mm-length and 17, 5.5 mm-length. Attending to their location, 4 corresponded to Upper Premolars (UP), 3 to Lower Premolars (LP), 52 to Upper Molars (UM) and 29 to Lower Molars (LM). Figures 1,2. From the CG, 69 were 7.5-length, 17 8.5 mm-length and 2 were 10 mm-length, and corresponded to 23 UP, 11 LP, 20 UM and 34 LM. Diameter of the implants is presented in Fig. 3. The diameter of SiG implants was higher ($p < 0.001$).

Residual bone height was higher ($p < 0.001$) in CG (12.6 vs 7.7 mm) and crown-to-implant ratio was higher ($p < 0.001$) in SiG (1.7 vs 1.3 mm). Figure 4. Conversely, there were no statistical differences between both groups regarding insertion torque, bone type or antagonist type. Further information is available in Table 2.

At the time of implant placement, no differences could be observed attending to the number of implants placed equicrestal (−0.5 to 0.5 mm), subcrestal > 0.5 mm or supracrestal < −0.5 mm in mesial, but a higher proportion of short implants ($p \leq 0.001$) were placed equicrestal.

At the time of loading and last radiography, no differences in bone level were observed between both groups in mesial. CG implants were more subcrestal in distal at loading time than SiG implants ($p \leq 0.001$). In addition, during the last radiography, a slight difference ($p < 0.05$) in distal bone level was observed in favor to CG implants. Moreover, the crown to implant ratio did not significantly affect the marginal bone loss ($p = 0.781$).

| Table 1 Demographic data       | SiG  | CG  | $p$-value |
|--------------------------------|------|-----|-----------|
| Number of patients ($n = 176$) | 78   | 88  | NA        |
| Number of implants ($n = 186$) | 88   | 88  | NA        |
| Age (years; median (range))    | 56 (20 to 78) | 53 (18 to 76) | 0.159$^a$ |
| Sex (females (males))          | 52 (36) | 53 (35) | 0.878$^b$ |
| Smokers                        | 7    | 8   | 0.787$^b$ |

$^a$ Mann–Whitney test

$^b$ Chi-square test
Fig. 1  Left: 6.5 length implant placed in #3.7 position. Center: 2 years later; single screw-retained crown over 4 mm. straight transepithelial (intermediate abutment). Right: 6 years after implant placement. No marginal bone loss observed after 6-year follow-up.

Fig. 2  Left: 8.5 length implant placed in #1.4 position. Center: 2 years later; single screw-retained crown over 2 mm. Straight transepithelial (intermediate abutment). Right: 6 years after implant placement. No marginal bone changes after 6-year follow-up.

Fig. 3  Implant diameter. SiG (short implant group), CG (control group)
The follow-up of dental implants, since insertion had a median of 31 months (range 11 to 84) for SiG and 35 months for CG (range 6–117).

All the implants from both group were in function at the last recall (100% survival) and the number of prosthetic complication did not statistically differ (3 events for SiG vs. 6 for CG). Attending to the Health Scale for Dental Implants [43], Success rate for SIG group was 100% and 98.9% for CG. No statistical differences in Marginal Bone Loss (MBL) were observed either in

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**Table 2** Surgical data, crown-to-implant ratio, and follow-up data

| Group                        | p-value |
|------------------------------|---------|
| **Residual bone height (mm; median (range))** |         |
| Experimental                 | 7.7 (4.2 to 17.6) | 12.6 (7.7 to 20.4) | 0.000<sup>a</sup> |
| Control                      |         |
| **Insertion torque (Ncm; median (range))** |         |
| Experimental                 | 42.5 (5.0 to 70.0) | 35.0 (5.0 to 70.0) | 0.188<sup>a</sup> |
| Control                      |         |
| **Bone type (number of implants)** |     |
| Type I                       | 4       | 4 | 0.234<sup>b</sup> |
| Type II                      | 44      | 49 |
| Type III                     | 33      | 22 |
| Type IV                      | 7       | 13 |
| **Antagonist (number of implants)** |         |
| Tooth                        | 63      | 66 | 0.609<sup>b</sup> |
| Implant                      | 25      | 22 |
| **Crown to implant ratio (median (range))** |         |
| Experimental                 | 1.7 (1.1 to 2.5) | 1.3 (0.9 to 2.1) | 0.000<sup>a</sup> |
| Control                      |         |
| **Immediate loading (number of implants)** |         |
| Experimental                 | 49      | 49 | 1.000<sup>b</sup> |
| Control                      |         |
| **Follow-up since insertion (months; median (range))** |         |
| Experimental                 | 31 (11 to 84) | 35 (6 to 117) | 0.139<sup>a</sup> |
| Control                      |         |
| **Follow-up since loading (months; median (range))** |         |
| Experimental                 | 27 (4 to 84) | 31 (6 to 67) | 0.249<sup>a</sup> |
| Control                      |         |
| **Follow-up of marginal bone level (months; median (range))** |         |
| Experimental                 | 24 (3 to 92) | 30 (1 to 96) | 0.095<sup>a</sup> |
| Control                      |         |

<sup>a</sup> Mann–Whitney test

<sup>b</sup> Chi square test
mesial or distal between both groups after the follow-up period. There were no differences in the number of implant areas (mesial/distal) showing MBL ≥ 2 mm (SiG 1/166; 6/166 CG). Additional information about the clinical performance is available in Table 3.

Discussion
The results of the present retrospective cohort study showed no differences in survival, changes in marginal bone levels or prosthetic complications between SiG and CG implants supporting a single-crown restoration over a transepithelial (intermediate abutment) in posterior maxilla or mandible. These results are in line with those of the systematic review and meta-analysis published by Tolentino da Rosa de Souza et al. [7] Short implants in posterior single crown had similar survival rates, low MBL and low prosthetic and surgical complications rate for a 1-year follow-up time. On the contrary Xu et al. [41] on their meta-analysis stated that the survival rate of short implants in the maxilla may be lower than that of long implants, while in the mandible both type of implants showed similar survival rate. Furthermore, short implants have been associated with lower MBL and biological complications but higher technical complications. It Worth mentioning that short and longer implant definition was different in these studies [7, 41]. Moreover, most of authors selected a unique length in the SiG and a unique length in the CG instead of defining a limit length between both groups. The type of included articles (only RCTs [41] or CCTs and RCTs [7]), the ratio of maxillary/mandibular implants, the proportion of implants placed with additional techniques, or the follow-up period could also explain the differences between both meta-analysis.

Systematic reviews comparing Short and Standard implants including both splinted and non-splinted implants [31, 44, 45] should be interpreted with caution as the biomechanical performance of splinted implants is substantially different from non-splinted ones. In multiple restorations splinted implants can disperse the stress on each single implant thereby reducing the implant overload and the possible incidence of mechanical complications [7, 46, 47]. In relation to the biomechanical performance of short implants supporting single crowns, similar outcomes have been reported when the implant is placed in the most distal position in the arch or between adjacent teeth or other implants. [48]

Many articles have been published comparing the performance of short implants in native bone vs longer implants along with sinus grafting in the maxilla. A recent umbrella review of meta-analysis from Vetromilla et al. [28] concluded that short implants showed fewer biological complication rates, reduced cost, and an overall similar satisfaction rate of the patients.

In the present study, the crown-to-implant ratio of all the implants in the CG where within the ranges that have been stated not to negatively influence the performance of implants; from 0.9 to 2.2 [21] or 0.6 to 2.36 [20]. From the SiG, only 2/88 implants slightly exceeded this second

| Technical complications (number of implants) | Short Implants | Standard | p value |
|---------------------------------------------|---------------|---------|--------|
| None                                        | 85            | 82      | 0.342b |
| Screw loosening                             | 2             | 5       |        |
| Resin chipping                              | 1             | 0       |        |
| Ceramic chipping                            | 0             | 1       |        |
| Mesial MB level (mm; median (range))         |               |         |        |
| Loading                                     | 0.7 (−0.7 to 2.5) | 0.7 (−0.4 to 2.3) | 0.173a |
| Last visit                                  | 0.8 (−1.2 to 2.9) | 0.6 (−3.3 to 2.9) | 0.262a |
| Distal MB level (mm; median (range))         |               |         |        |
| Loading                                     | 0.3 (−2.5 to 2.3) | 0.6 (−1.3 to 2.3) | 0.001a |
| Last visit                                  | 0.4 (−2.1 to 2.1) | 0.5 (−3.8 to 2.4) | 0.018a |
| Change in MB level (mm; median (range))      |               |         |        |
| Mesial                                      | 0 (−1.5 to 1.3) | 0.1 (−4.0 to 1.4) | 0.322a |
| Distal                                      | 0.1 (−2.2 to 1.0) | 0.1 (−4.0 to 1.0) | 0.758b |
| Change in MB level ≥ 2 mm                   |               |         |        |
| Mesial                                      | 0             | 1       | 0.316b |
| Distal                                      | 1             | 3       | 0.312b |
| Change in MB level ≥ 3 mm                   |               |         |        |
| Mesial                                      | 0             | 1       | 0.316b |
| Distal                                      | 0             | 1       | 0.316b |

Marginal bone (MB) level (−): below the most coronal part of the implant shoulder; (+): above the most coronal part of the implant shoulder

a Mann–Whitney test
b Chi square test
previously published range and 5/88 the first one. The differences between SiG and CG in the crown-to-implant in the present study were lower than in other studies [20, 21] probably because the length of the CG implants was lower.

The use of transepithelial abutments in all the implants probably contributed to achieve a low number of technical complications in both groups and helped to dissipate implant loads thus positively influencing in the biological performance too [22, 49, 50]. This total homogeneity in the type of prosthetic restoration and in the type of implant surface together with a well-balanced sampling in relation to patients’ sex and age, number of smokers, bone type, insertion torque or type of antagonist, are relevant strengths of the present study. The unicentric design of the study ensured that all implants were placed and prosthetically rehabilitated by the same team, following the same protocol. This could have helped to reduce bias but at the same time the lack of data from other centers could be at the same time considered a limitation of the study. Other limitations of the study were the limited follow-up time or the reduced ability to control confounding factors of the retrospective designs. New long-term prospective studies are recommended to confirm these results.

Conclusions
Within the limitations of this study, short (≤ 6.5 mm) and standard (≥ 7.5 mm) implants supporting a single-crown restoration over a transepithelial (intermediate abutment) in the posterior maxilla/mandible show similar clinical performance (survival, MBL, prosthetic complications).

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Author contributions
All authors have made substantial contributions to the conception or design of the work or the acquisition, analysis, or interpretation of data for the work, AND drafting the work or revising it critically for important intellectual content; AND have given final approval of the version to be published; AND agree to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. All authors read and approved the final manuscript.

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Availability of data and materials
The data sets used and/or analysed during the current study are available from the corresponding author on reasonable request.

Declarations
Ethics approval and consent to participate
The study protocol was approved by the ethical committee of the University Hospital of Araba (FIBEA-02-ER/22/Extracortos).

Consent for publication
Not applicable.

Competing interests
EA is the Scientific Director and the president of BTI Biotechnology Institute, a dental implant company that investigates in the fields of oral implantology and PRGF-Endorex technology, and the president of Eduardo Anitua Foundation. MHA is a researcher (employee) at BTI Biotechnology Institute. AE is a researcher at the Eduardo Anitua Foundation.

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