Short implants (<8mm) versus longer implants (≥8mm) with lateral sinus floor augmentation in posterior atrophic maxilla: A meta-analysis of RCT’s in humans

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Abstract
Background: One of the greatest challenges that dentists face today is to rehabilitate severe atrophied alveolar ridges in partially and completely edentulous patients with implants. Despite the high survival rate of implants placed next to sinus elevation, this technique presents complications that can be avoided by placing short implants, an option that also presents high survival rates.

For this reason, the aim of this study is to compare the survival rate, marginal bone loss and complications associated with short implants (<8 mm) versus longer implants (≥8mm) placed with lateral sinus floor elevation in posterior atrophic maxillae.

Material and Methods: A literature search was conducted by two independent reviewers in the PubMed/Medline (National Library of Medicine, Washington, DC) electronic database for articles published from January 2007 to July 2018. Seven qualified articles were selected for the meta-analysis.

Results: The test for overall effect did not find statistical significance in the survival rates, overall complications, intra-operative complications, post-operative complications and prosthetic complications. However, the test showed statistically significant differences in biological complications in favor of standard implants, and marginal bone loss between control and test groups in favor of short implants (<8mm) was found.
Conclusions: Within the limitations of the present study, prosthetic rehabilitations with short implants (<8mm) in posterior maxilla is a reliable treatment option as an alternative to lateral wall sinus floor augmentation.

**Key words:** Short implant, lateral sinus floor augmentation, Randomized controlled trial, Survival rate, Complications, Marginal bone loss.

**Introduction**

One of the greatest challenges that dentists face today is to rehabilitate severe atrophied alveolar ridges in partially and completely edentulous patients with implants. Following tooth loss, jaws undergo vertical collapse due to increased osteoclast activity, which takes place in response to the absence of functional load transmission to the alveolar bone. Bone resorption is aggravated by the physiological process of sinus pneumatization especially in the maxillary posterior area (1). Therefore, bone quantity and quality is often insufficient for the ideal three-dimensional (3D) implant positioning. Several bone augmentation techniques have been proposed to overcome these problems. Among these, Sinus floor elevation is considered to be the most reliable surgical technique for increasing bone height in the posterior maxilla (2). Two sinus floor elevation techniques have been described by Wang et al. [2008] (3): lateral approach (LSFE), when the residual bone volume is less than ≤5 mm, or crestal (CSFE) approach when residual bone height is more than 6 mm. Both techniques have reported Research reports high survival rates, 100% after 5-year follow-up (4) and 97% after a 10-year follow-up (5) and success rates 98% after a 3-year follow-up (6).

Due to the high percentage of anatomical variations among patients (7) and the sensitivity of the technique, these procedures are not exempt from complications: Schneiderian membrane perforation, sinusitis, nasal bleeding, hematomas, post-operative pain, dehiscence, graft failure, or migration of the implant into the sinus cavity are common complications associated with sinus floor elevation surgery (8,9). For this reason, several alternatives have been proposed to avoid sinus lifting, such as tilted implants or short implants (10). Recently, implants as short as 8 mm have been considered as standard implants in several published articles (11). Short implants are slowly being accepted by patients and clinicians because they are associated with a less invasive procedure, leading to a smaller scale intervention, shorter intra-operative time, less morbidity, and lower treatment cost (12). Traditionally, short implants have been related to lower survival rates and unpredictable outcomes. But more recently, technical and manufacturing developments have improved implant surfaces and connections and nowadays short implants have a failure rate of under 4% for ≤8mm implants, a failure rate similar to longer implants (13).

The aim of the present study was to compare the survival rate, marginal bone loss and complications associated with short implants (<8 mm) versus longer implants (≥8mm) placed with lateral sinus floor elevation in posterior atrophic maxillae.

**Material and Methods**

- **Study Registration**

This systematic review and meta-analysis has been registered in the Prospero Database (International prospective register of systematic reviews - https://www.crd.york.ac.uk/prospero/) under the title: “Short implants (<8mm) versus longer implants (≥8mm) with lateral sinus floor augmentation in posterior atrophic maxilla: A meta-analysis of RCT’s in humans” (ID:92413).

- **Focused Question**

Following Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines (14), a specific answerable question was formulated according to PICO(S) recommendations (Participants, Interventions, Control, Outcomes, Study): (P) Participants: Patients who received at least one dental implant in the posterior area of the upper maxilla. (I) Type of intervention: at least one short dental implant placement (<8mm) in the posterior area of the upper maxilla without lateral sinus floor elevation procedure. (C) Control intervention: at least one long dental implant (≥8mm) placed simultaneously or deferred with sinus membrane elevation via lateral sinus floor elevation procedure. (O) Outcome measures: implant survival rate, intra- and post-operative surgical complications, biological and prosthetic post-operative complications, marginal bone loss (MBL). (S) Study type: randomized controlled clinical trials (RCTs).

- **Search Strategy**

A literature search was conducted by two independent reviewers (LN and AA) in the MEDLINE, EMBASE, Cochrane Central Register of Controlled Trials, and Cochrane Oral Health Group Trials Register databases. The research included references up to July 2018, using different combinations and Boolean Operators (AND, OR, NOT) with the following search terms/key words: “short implants”, “longer implant”, “standard implant”, “bone augmentation”, “sinus lift”, “randomized control trial”, “atrophy maxilla”, “posterior maxilla”.

Following the electronic search, a further manual search
was performed in the websites of the leading scientific journals on dentistry and implant dentistry. Crossed-references were screened to identify other potentially relevant articles.

- Eligibility criteria
Studies were deemed eligible if they met the following criteria: 1) Human subjects with posterior maxilla atrophy; 2) Randomized Clinical Trials (RCT); 3) the presence of a study group (receiving one or more short implant (<8 mm) and a control group (receiving long implants [≥8mm] simultaneously or deferred with lateral sinus floor elevation); 4) studies with a minimum follow-up period of >12 months after prosthesis placement; 5) results providing data on survival rates, complications, and marginal bone loss; 6) articles published in English. Exclusion criteria comprised of: 1) animal studies; 2) human studies with less than 15 subjects with posterior maxilla atrophy; 3) studies with a follow-up of <12 months after prosthetic loading; 4) prospective cohort studies, case reports, case series, retrospective studies, systematic reviews; and 5) articles that failed to provide sufficient information.

- Data extraction
The following information was extracted from the publications included for analysis: 1) author and year of publication; 2) duration of follow-up; 3) patient and implant sample; 4) systemic, periodontal, and smoking status; 5) time of loading; 6) implant location; 7) setting and funding; 8) preoperative preparation; 9) treatment control group; 10) treatment study group; 11) residual bone height; 12) post-surgical instructions; 13) augmentation technique; 14) survival rate; 15) intra/post-operative and biological/prosthetic complications; 16) marginal bone loss (MBL); and 17) study conclusions as reported by the authors.

Two reviewers (LN and AA) carried out the selection process, screening the articles’ titles and abstracts. The full texts of all studies of possible relevance were then obtained, and eligibility assessment and data extraction were performed independently in an un-blinded standardized manner by the two authors; any disagreement between the reviewers was resolved through discussion. When the reviewers did not agree, a third reviewer (SO) analyzed the text to decide whether the article should be included or excluded.

- Quality Assessment
The reviewers A.A and N.L assessed the quality of each study independently. Disagreements on validity assessment were resolved by consensus and discussion; when consensus could not be reached, a third reviewer was consulted (JG). The methodological quality of the RCT’s were assessed using the Cochrane Risk of Bias Tool for Randomized Controlled Trials (Table 1).

- Statistical Analysis
Statistical analysis was performed using R Project software (The R Foundation, Bell Laboratories, formerly AT&T, now Lucent Technologies by John Chambers and Colleagues). The Chi² test was used to evaluate heterogeneity across the studies; subgroup analysis was performed when heterogeneity was significant (p-value <0.05) and the I² statistic expressed the percentage of heterogeneity, with 25% corresponding to low heterogeneity, 50% to moderate, and 75% to high. A test of overall effects was used to evaluate significance between the groups; a p-value of less than 0.05 was considered statistically significant. A forest plot was drawn to represent estimates of relative effect, expressed as risk ratio (RR) with a 95% confidence interval (CI).

Results
- Study selection
Search results based on the PRISMA guidelines are depicted in Fig. 1. The initial search identified 482 titles, 476 PubMed Embase Database and 6 additional records identified through hand-searching. After elimination by screening all titles and abstracts, twenty-one studies were left for full-text assessment. After full-text screening, 14 articles were excluded due to failed to meet the inclusion criteria (Table 1) (15-29), leaving a total of 8 RTCs (30-37) for inclusion in the meta-analysis (Table 2).

Fig. 1: PRISMA flowchart of the screening process in different databases.
Table 1: Excluded Studies.

| Reason for exclusion | Number | Studies                                                                 |
|----------------------|--------|-------------------------------------------------------------------------|
| Short implants ≥8mm  | 4      | Canizzaro et al. 2009; (15) Canizzaro et al. 2013; (16) Esposito et al. 2015; (17) Taschieri; (18) |
| Mean follow-up < 1 year | 4    | Felice et al. 2009; (19) Esposito et al. 2012; (20) Esposito et al. 2015; (21) Esposito et al. 2016 (22) |
| More recent follow-up papers available | 2 | Pistilli et al. 2013; (23) Thoma et al. 2015 (24) |
| Data for maxillary implants could not be separated from mandibular implants | 2 | Esposito et al. 2011; (25) Felice et al. 2016 (26) |
| OSFE instead of LSFE | 2 | Felice et al. 2015 (27); Zhang et al. 2017 (28); Yu et al. 2017 (29); |
| Total                | 14     |                                                                         |

Note. OSFE: Osteotome sinus floor elevation. LSFE: Lateral sinus floor elevation.

Table 2: General overview of the studies included for analysis.

| Study | Study design, follow-up times | Mean Age, Patients and Implants (N) | Systemic, periodontal status, smoking habits | Time of loading in relation to implant placement | Location | Site, setting and funding |
|-------|-------------------------------|------------------------------------|---------------------------------------------|-------------------------------------------------|----------|--------------------------|
| Bolle et al. 2018. (30) | RCT 1 year post-loading | Mean age: CG 56.4 (36-71); SG 60.75 (25-77) Patients n= 40 (CG 20; SG 20) Implants n= 78 (CG 41; SG 37) | Systemic and periodontal health; non-smokers, moderate smokers (<10 cig/day), or heavy smokers (>10 cig/day) | 4 months with provisional prosthesis, and after another 4 months with definitive prosthesis | Post. Max | Italy, University Hospital, private centers; Public Hospital; partially supported by Global D (Brignais, France) |
| Gastaldi et al. 2018. (31) | RCT 1 year post-loading | Mean age: CG 58.5 (45-75); SG 61.1 (45-70) Patients n= 40 (CG 20; SG 20) Implants n= 73 (CG 37; SG 36) | Systemic and periodontal health; non-smokers, moderate smokers (<10 cig/day), or heavy smokers (>10 cig/day) | 4 months with provisional prosthesis, and after another 4 months with definitive prosthesis | Post. Max | Italy, University Hospital, private centers; Public Hospital: partially supported by MegaGen Implant (South Korea) |
| Hadzik et al. 2018. (32) | RCT After surgery, 6 months and 12 months | Mean age: 45.5 y Patients n= 29 (CG 15; SG 14) Implants n= 29 (CG 15; SG 14) | ≤25 plaque index, no smokers, no bruxers | 36 months since implant placement, 30 months since loading | Post. Max | Lithuania University; no funding support reported |
| Thoma et al. 2018. (33) | RCT 1, 3, 5 years post-loading | Mean age 20-75 years Patients n=90 (CG 46; SG 44) Implants n= 124 (CG 64; SG 60) | Systemic and periodontal health; non-smokers, moderate smokers (<10 cig/day), or heavy smokers (>10 cig/day) | 6 months | Post. Max | Austria, Switzerland, Poland, Spain, USA, private centers; fully funded by Dentsply Implants (Mölndal, Sweden) |
| Bechara et al. 2017. (34) | RCT 1, 3 years post-loading | Mean age: 48.1±15.1 years Patients n= 53 (CG 20; SG 33) Implants n= 90 (CG 45; SG 45) | Systemic and periodontal health; non-smokers, moderate smokers (<10 cig/day), or heavy smokers (>10 cig/day). Bruxism was an exclusion criteria | 4 months | Post. Max | Italy, private centers, Public Hospital; partially supported by Tecnoss®, (Giaveno, Italy) and Southern Implants |
| Esposito et al. 2014. (35) | RCT 1, 3 years post-loading | Mean age: 56 (45-70) Patients n= 15 Implants n= 72 (CG 38; SG 34) | Systemic and periodontal health; non-smokers, moderate smokers (<10 cig/day). | 4 months with provisional prosthesis, and after another 4 months with definitive prosthesis | Post. Max | Italy, private centers, Public Hospital; partially supported by Dentsply Implants (Brignais, France) |
| Guljé et al. 2014. (36) | RCT 1 year | Mean age: 50 ± 10.1 (SG), 48 ± 8.9 (CG) Patients n= 41 (CG 20, SG 20) Implants n= 72 (CG 20; SG 20) | Systemic and periodontal health; non-smokers, moderate smokers (<10 cig/day), Controlled DM, no corticosteroids, no chemotherapy in the previous 5 years, no radiation, no systemic or local disease that compromise the treatment | 12 months with definitive prosthesis | Post. Max | Twentyone patients were treated in the centre in Apeldoorn and 20 patients were treated in the centre in Groningen |
| Pistilli et al. 2013. (37) | RCT 12 months post-loading | Mean age: 57.6 (45-80) Patients n= 20 Implants n= 83 (CG 44; SG 39) | Systemic and periodontal health; non-smokers, moderate smokers (<10 cig/day), or heavy smokers (>10 cig/day) | 4 months with provisional prosthesis, and after another 4 months with definitive prosthesis | Post. Max | Italy, University Hospital, private centers; Public Hospital; partially supported by Tecnoss®, (Giaveno, Italy) and Southern Implants |
The selection of RCTs included a total of 328 patients (although Esposito et al. (35) and Pistilli et al. (37) did not report the number of patients in each group), with a total of 621 implants (296 allocated to study group, and 316 to control group). Six studies only investigated partially edentulous patients (30-33,35,37); only two studies employed a split-mouth design (34,35), while the rest had parallel treatment arms (Table 3).

- Quality assessment
- Table 4 summarizes the results of bias risk assessment in the included RCTs and The Cochrane Risk of Bias Tool for Randomized Controlled Trials criteria.

### Table 3: General characteristics of the intervention and results.

| Study                        | Follow-up | Preoperative preparation | CG LSFE / implant length | SG short implant length | RBH (mm) | Bone graft / Membrane (CG) | Post-surgical treatment / Post-op instructions | Survival rate (%) | Complications | MBL (mm) | Authors conclusion |
|------------------------------|-----------|---------------------------|---------------------------|-------------------------|----------|---------------------------|-------------------------------------------------|------------------|---------------|----------|-------------------|
| Bolle et al. 2018. (30)      | 1 year    | CBCT, OHI, prophylactic ATB 1h before, CHX 0.2% 1 minute | Simultaneous/ >10 mm | 4 mm | 4-5 mm | Particulated porcine bone / resorbable collagen membrane | Amoxicillin 1gr twice/day / 7 days, NSAID (Ibufrofen 400 mg) 2 to 4 times/day, soft diet, not allowed to wear removable prosthesis up to 1 month | CG= 14 | Intra-operative= 3 | Post-operative= 11 | Biological= 9; Prosthetic= 2 | CG= 0.72±0.25 | SG= 0.63±0.15 | One year after loading, 4.0 mm long implants achieved similar results to longer implants in augmented jaws, but fewer complications |
| Gastaldi et al. 2018. (31)  | 3 years   | CBCT, OHI, prophylactic ATB 1h before, CHX 0.2% 1 minute | Simultaneous/ >10 mm | 5 mm | 4-6 mm | Particulated porcine bone / resorbable collagen membrane | Amoxicillin 1gr twice/day / 7 days, NSAID (Ibufrofen 400 mg) 2 to 4 times/day, CHX gel, soft diet, avoid brushing and trauma at surgical site. | CG= 6 | Intra-operative= 5 | Post-operative= 1 | Biological= 0; Prosthetic= 1 | CG= 1.79±0.59 | SG= 1.34±0.35 | Three years after loading 5.0 mm × 5.0 mm implants achieved similar results to longer implants with GBR |
| Hadzik et al. 2018. (32)    | 3 years   | OHI, Prophylactic ATB | Simultaneous 11 and 13 mm | 6mm | 6 mm height 6-7mm width | Xenogenic Bone Graft | 0.12% chlorhexidine solution, ATB and Analgesics (non specified) | CG= 100% | SG = 100% | - | CG= 0.34 ±0.24mm | SG= 0.22±0.46mm | Short implants can be successfully used to support single crowns in the lateral part of the maxilla |
| Thomas et al. 2018. (33)    | 5 years   | Prophylactic ATB 1h before, CHX 0.2% 1 minute | Simultaneous 11-15 mm | 6 mm | 5-7 mm | Particulated bovine bone mixed with autogenous bone chips / resorbable collagen membrane | NP | CG=100% | SG=98.5% | CG=24 | Intra-operative= 1 | Post-operative= 23 | Biological= 9; Prosthetic= 14 | SG= 0.46±0.1 SG= 0.54±0.87 | Short implants (6 mm) for single-tooth restorations in the posterior maxilla as a viable solution versus longer implants in combination with sinus lift. |
| Study            | Duration | CBCT, OPG, Prophylactic ATB before | Sinus lift before | Implant Length | Implant Material | Implant Comp. | OHI, Antibiotics | CG Intra-op. | CG Post-op. | SG Intra-op. | SG Post-op. | Notes |
|------------------|----------|-----------------------------------|-------------------|---------------|-----------------|--------------|----------------|---------------|--------------|---------------|--------------|-------|
| Becham et al. 2017 | 3 years  | CBCT, OPG, Prophylactic ATB 1h before, CHX 0.2% 1 minute | Simultaneous>10 mm | 6 mm ≥4 mm Collagenated porcine particulate bone graft/Pericardium porcine resorbable collagen membrane | Amoxicillin 500mg plus clavulanic acid, NSAID (Ibuprofen 600mg), CHX 0.2% for 2 weeks, not allowed to wear removable dentures up to 1 month postoperative | CG=95.6% SG=100% | CG=19 Intra-operative= 3 Post-operative=17 (Biological=17; Prosthetic=0) | CG=0.27±(0.232-0.313) SG=0.20±(0.166-0.236) | At three years short (6-mm) dental implants and longer (≥20-mm) dental implants in combination with sinus floor elevation provided good results. However, with short implants, the treatment was faster and less expensive. |
| Espósito et al. 2014 | 3 years  | CBCT, OHI, Prophylactic ATB 1h before, CHX 0.2% 1 minute | Simultaneous>10 mm | 5 mm 4-6 mm Particulated bovine bone/resorbable collagen membrane | Amoxicillin 1gr twice/day/7days, NSAID (Ibuprofen 400mg) 2 to 4 times/day, CHX gel, soft diet. | CG=97.36% SG=91.17% | CG=1 Intra-operative= 1 Post-operative= 0 (Biological=1; Prosthetic=0) | CG=1.54±0.35 SG=1.02±0.47 | Both techniques provided acceptable results up to 3 years after loading. However, with 5-mm implants treatment was faster and cheaper. |
| Guljé et al. 2014 | 1 year   | CBCT, OPG, Prophylactic ATB 1 hour before, | Simultaneous 11mm | 6 mm 6-8 mm Autologous bone + Particulated bovine bone + collagen membrane | Chlorhexidine mouthrinse | CG=100% SG=100% | CG=0 Intra-operative= 0 Post-operative= 0 (Biological=0; Prosthetic=0) | CG=0,1±0,3mm SG=0,1±0,2mm | 6-mm implants and 11-mm implants combined with sinus floor elevation surgery are equally successful to support a single crown in the resorbed posterior maxilla after 1-year follow-up. |
| Pistilli et al. 2013 | 1 year   | CBCT, OHI, Prophylactic ATB 1h before, CHX 0.2% 1 minute | Simultaneous>10 mm | 6 mm 5-7 mm Particulated porcine bone/resorbable collagen membrane | Amoxicillin 1gr twice/day/7days, NSAID (Ibuprofen400mg) 2 to 4 times/day, CHX 0.2% for 2 week, soft diet for 1 week, to avoid brushing and trauma at surgical site | CG=100% SG=100% | CG=4 Intra-operative= 4 Post-operative= 0 (Biological=0; Prosthetic=0) | CG=1.09±0.05 SG=1.02±0.06 | 6 mm-long implants achieved similar results to longer implants placed in augmented bone. |

Note. CBCT: Cone Beam Computed Tomography. OHI: Oral Hygiene Instruction. LSFE: lateral sinus floor elevation. NSAID: Non-Steroidal Anti-Inflammatory Drugs. CG: control group (longer implants [≥8mm] simultaneously or deferred lateral sinus floor elevation). SG: study group (short implants<8 mm)
- Survival rate
A random-effects model was used to assess the survival rate of implants; statistically significant heterogeneity was not found among the publications ($I^2=0\%; p=0.81$). The test for overall effect showed no statistically significant differences in the survival rate of short implants (<8mm) compared to longer implants (≥8mm) with lateral sinus floor elevation (Risk Ratio [RR] of 0.98; 95% CI: [0.42-2.83]; $p=0.8$) (Fig. 2).

- Marginal Bone Loss
The random-effect model showed highly significant heterogeneity between the studies ($I^2=97.9\%; p=0.00$). The overall effect test showed statistically significant differences in marginal bone loss between control and study groups ($p=0.026$). A RR of 0.86; 95% CI: [0.75-0.98] in favor of short implants (<8mm) with lateral sinus floor elevation is significantly higher than patients receiving longer implants (≥8mm) (Fig. 2). This finding implies that the risk of marginal bone loss in patients receiving longer implants (≥8mm) with lateral sinus floor elevation is significantly higher than patients receiving short implants (<8mm). However, these results should be treated with caution due to the different follow-up periods among the studies analyzed.

- Complications
The test for overall effect did not find statistical significance (RR of 0.60; 95% CI: [0.25-1.47]; $p=0.262$). For this variable the random-effects model showed a statistically significant heterogeneity between the studies ($I^2=60.2\%; p=0.03$) (Fig. 2). So, complications were divided into four groups: (3a) intra-operative complications; (3b) post-operative complications; (3c) biological complications; and (3d) prosthetic complications.

Intra-operative complications: The Chi$^2$ test showed homogeneity between the studies ($I^2=22.9\% p=0.33$); and the overall effect test found no statistically significant differences between control and study groups (RR of 0.51; 95% CI: [0.16-1.63]; $p=0.258$), in relation to intra-operative complications (Fig. 3).

Post-operative complications: The Chi$^2$ test showed homogeneity between the studies ($I^2=36.1\% p=0.15$); and the overall effect test found no statistically significant differences between control and study groups (RR of 0.76; 95% CI: [0.33-1.74]; $p=0.517$), in relation to post-operative complications (Fig. 3).

Biological complications: The Chi$^2$ test did not find statistically significant heterogeneity between the studies ($I^2=0.0\% p=0.43$). The overall effect test demonstrated that there were more biological complications in the control group (RR of 0.46; 95% CI: [0.22-0.95]; $p=0.037$) (Fig. 3).

Prosthetic complications The Chi$^2$ test demonstrated homogeneity between the studies ($I^2= 0.0\% p=1.00$); and the overall effect test didn’t find statistically significant differences between the control and study groups ($p=0.110$). A high number of studies did not suffered any prosthetic complications, either for short implant groups or longer implant groups. A RR of 1.52; 95% CI: [0.91, 2.54] favored the control group slightly (Fig. 3).

Table 4: Bias risk assessment for the included RCTs using The Cochrane Risk of Bias Tool for Randomized Controlled Trials.

| Study           | Random sequence generation | Allocation concealment | Blinding of participants and personnel | Blinding of outcome assessment | Incomplete outcome data addresses | Selective reporting | Other bias | Overall risk of bias |
|-----------------|---------------------------|------------------------|---------------------------------------|-------------------------------|----------------------------------|---------------------|------------|----------------------|
| Boile et al. 2018 (3) | Low | Low | High | Low | Low | Low | Low | Moderate |
| Gastaldi et al. 2018 (31) | Low | Low | High | Low | Low | Low | Low | Moderate |
| Hadzik et al. 2018 (32) | Low | Low | Low | NP | Low | Low | Low | Low |
| Thoma et al. 2018 (33) | Low | Low | Low | NP | Low | Low | Low | Low |
| Bechara et al. 2017 (34) | Low | Low | NP | Low | Low | Low | Low | Low |
| Espósito et al. 2014 (35) | Low | High | NP | Low | Low | Low | Low | Moderate |
| Guljé et al. 2014 (36) | Low | Low | NP | NP | Low | Low | Low | Low |
| Pistilli et al. 2013 (37) | Low | High | High | Low | Low | Low | Low | High |
Fig. 2: Forest plot for the event: (a) “implant survival rate; (b) “marginal bone loss””; (c) “complications”.

Survival RR (a)

Marginal Bone Loss (b)

Complications (c)
Fig. 3: Forest plot for the event: (a) "intra-operative complications"; (b) "post-operative complications"; (c) "biological complications"; (d) "prosthetic complications".
Discussion

Short implants are considered a reliable and predictable alternative to bone augmentation procedures (7,11,13), reducing the rate of complications, intra-operative time, patient morbidity, and treatment costs (12). The fourth European Association for Osseointegration (EAO) consensus conference (31) reported a survival rate of 99.0% for short implants (<8 mm) after 16-18 months follow-up, considering their use a routine treatment. The present systematic review and meta-analysis defined the term ‘short’ to describe implants of less than 8 mm (<8 mm) in length in accordance with the definition proposed recently by Plonka et al. [2018] (11).

- Survival rate
The results of meta-analysis did not find statistically significant differences in survival rates between short implants (<8mm) and longer implants (≥8mm) with lateral sinus floor elevation. None of the RCTs analyzed reported statistically significant differences between control and test groups. Similar results were reported by Hadzik et al. (32), Guljé et al. (36) and Pistilli et al. (37), who obtained 100% survival rates for both control and test implants. However, these results should be treated with caution because of the small numbers of failed implants in both groups and the short follow-up periods.

Short implants might be expected to suffer more failures than long implants after loading because of their bio-mechanical disadvantages. However, the results of the RCTs in this review did not demonstrate this effect. The high survival rate of short implants could be attributed to improved implant surfaces and connections. Traditionally, machined surface implants with external connections were used, but the development of internal connections and rough surfaces have increased the implants surface area, favoring bone-to-implant contact, reduced treatment time, implant diameter and length, so that they now produce similar or even better results in comparison with machined implants (38).

The length of the implants included in the study groups ranged between 4 mm and <8 mm. Anitura et al. (38) obtained a 98.2% survival rate for 114 extra-short implants (<6.5 mm) after a follow-up period of 26 months. Recently, Srinivasan et al. (39), 690 6-mm short implants were assessed, obtaining a cumulative survival rate of 93.7% for maxillary implants and 98.6% for maxillary and mandibular implants together.

Furthermore, in a retrospective study published by Tetsch et al. (11) showed implants of ≥10 mm with lateral sinus floor elevation had 98.3% implant survival rate after 15.5 years of follow-up.

- Complications
In reference to complications, two studies (30,34) reported statistically significant differences in favor of short implants, although the random-effects model showed statistically significant heterogeneity between studies. For this reason, complications were divided into four groups: 1) intra-operative complications; 2) post-operative complications: 2a) biological complications; and 2b) prosthetic complications. When complications were divided into subgroups, only Bechara et al. (34) reported a significantly higher number of post-operative biological complications in the control group. The complications associated with longer implants with lateral maxillary sinus augmentation were, in order of frequency: pain/swelling > sinus membrane perforation > nasal bleeding and post-operative headache > intra-operative bleeding > infection of the grafting material > migration of the implant into sinus maxillaris.

In the group of short implants (<8 mm) the most frequent complications were: sinus membrane perforation > nasal bleeding > migration of the implant into sinus maxillaris. The most common prosthetic complication in both groups was screw loosening/fracture. In brief, incidence was slightly higher in study groups (short implants) although the difference was not statistically significant. Only a few studies reported prosthetic complications, comparing study groups with control groups, but the higher number of implants in the control groups suggest that longer implants (>8mm) with lateral sinus floor elevation suffered fewer prosthetic complications.

- Marginal Bone Loss (MBL)
In the present study, MBL in patients receiving longer implants (≥8mm) with lateral sinus floor elevation was statistically higher compared with patients who received short implants (<8mm). This results can be justified by the article of Galindo-Moreno et al. (40), evaluated the MBL of implants placed in native bone or in grafted sinus lift in the maxilla. Concluded that “implants placed in sites that received maxillary sinus augmentation exhibited more marginal bone loss than implants placed in pristine bone, although marginal bone loss mainly occurred during the first 12 months after functional loading”. In the RCT by Bechara et al. (34), the study group included short implants (6 mm in length) placed in healed sites and post-extraction sockets; however, no statistically significant differences in MBL between the two groups were found at either 1- or 3-year follow-ups.

Due to the heterogeneity of the publications reviewed and the lack of information this systematic review suffered some limitations. Three out eight reviewed publications were considered with high risk of bias what might affect the obtained results. Furthermore, it was not possible to draw any definitive conclusions regarding the success rate, impact of implant diameter, implant design, and the type of prosthetic restoration on the variables investigated. Little information was available in the studies reviewed regarding the type of prosthetic reconstruction (single unit or multiple units).
number of prosthetic reconstructions, and the number of implants per prosthetic unit. Moreover, the outcomes of the present review should be interpreted with caution given the small sample size and short follow-up times of the studies analyzed. In addition, five (30,33-35,37) out the 8 papers analyzed were published by the same research team.

Conclusions
Within the limitations of the present systematic review, prosthetic rehabilitations with short implants (<8mm) in the maxillary posterior areas are a predictable treatment option as an alternative to sinus floor elevation. No statistically significant differences in the survival rate and complications were found between short (<8mm) and longer implants (≥8mm) with lateral sinus floor elevation. Nevertheless, longer implants (≥8mm) in combination with lateral sinus elevation presented significantly greater marginal bone loss.

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Conflict of interest
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