Efficacy of bone-substitute materials use in immediate dental implant placement: A systematic review and meta-analysis

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Abstract
Objective: To assess the efficacy of using a bone substitute material (BSM) in the fixture–socket gap in patients undergoing tooth extraction and immediate implant placement.

Materials and methods: MEDLINE, EMBASE, and CENTRAL databases were searched for randomized controlled trials (RCTs). RCTs were screened for eligibility, and data were extracted by two authors independently. Risk of bias (ROB) was assessed using Cochrane’s ROB tool 2.0. Primary outcomes were implant failure, overall complications, and soft-tissue esthetics. Secondary outcomes were vertical buccal bone resorption, vertical interproximal bone resorption, horizontal buccal bone resorption, and mid-buccal mucosal recession. Meta-analysis was performed using random-effects model with generic inverse variance weighing. GRADE was used to grade the certainty of the evidence.

Results: After screening 19,544 potentially eligible references, 20 RCTs were included in this review, with a total of 848 patients (916 sites). Most included RCTs were deemed of some concerns (53%) or at low (38%) risk of bias, except for overall complications (high ROB). Implant failure did not differ significantly RR = 0.92 (confidence intervals [CI] 0.34 to 2.46) between using a BSM compared with not using a...
BSM (NoBSM). BSM use resulted in less horizontal buccal bone resorption (MD = −0.52 mm [95% CI −0.74 to −0.30]), a higher esthetic score (MD = 1.49 [95% CI 0.46 to 2.53]), but also more complications (RR = 3.50 [95% CI 1.11 to 11.1]) compared with NoBSM. Too few trials compared types of BSMs against each other to allow for pooled analyses. The certainty of the evidence was considered moderate for all outcomes except implant failure (low), overall complications (very low), and vertical interproximal bone resorption (very low).

**Conclusion:** BSM use during immediate implant placement reduces horizontal buccal bone resorption and improves the periimplant soft-tissue esthetics. Although BSM use increases the risk of predominantly minor complications.

**KEYWORDS**
bone regeneration, bone substitutes, dental implant, tooth extraction, tooth socket

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1 | **INTRODUCTION**

Tooth extraction marks the initiation of the socket healing process, which consists of three main stages, namely inflammatory, proliferative, and bone (re)modeling. Consequently, the bundle bone resorbs and the alveolar ridge tends to return to its pre-eruption form, thus resulting in reduction of the alveolar ridge dimensions. After atraumatic tooth extraction, immediate implant placement (IIP) is a common procedure with an approximate success rate of 95%. Moreover, IIP reduces treatment duration, and omits the need for another implant placement surgery when compared with delayed implant placement.

The difference in the circumference between the extracted tooth and the implant creates a gap; the “fixture-socket gap.” Several studies suggested that the use of a bone-substitute material (BSM) in the fixture-socket gap preserves socket volume, minimizes socket remodeling, and supports de-novo bone formation. This might improve the peri-implant bone regeneration process.

Various types of BSMs can be used during IIP, such as autograft, allograft, xenograft, or alloplast. Each of these BSMs has merits and drawbacks based on their properties. The type of BSM used can affect the dimensional change of the socket, percentage of regenerated vital bone, and the amount of connective tissue present in the site.

Trials assessing the efficacy of using a BSM in the fixture-socket gap during IIP are often limited in sample size, tend to suffer from heterogeneous reporting, and frequently have conflicting results. An overview, critical appraisal, and pooled analysis of studies looking at the efficacy of using BSMs for IIP is still lacking.

This systematic review aims to assess the efficacy of using a BSM in the fixture-socket gap compared with not using any grafting material (NoBSM) in patients undergoing tooth extraction and IIP. In addition, we aimed to evaluate whether efficacy was different between the different types of BSMs that were used.

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2 | **MATERIAL AND METHODS**

2.1 | Reporting

The methodology of this systematic review followed the Cochrane handbook for systematic reviews of interventions. Reporting was in accordance with the PRISMA checklist. The protocol of this systematic review was registered in the PROSPERO database (CRD42020164451).

2.2 | Eligibility criteria

RCTs assessing the efficacy of BSMs in adult participants (≥18 years old) undergoing tooth extraction and IIP with a minimum follow-up of
4 months were eligible for inclusion. Trials that only used the patient’s blood extracts, or tooth graft to fill the fixture–socket gap were excluded. Additionally, trials that used soft-tissue augmentation as the primary intervention were excluded.

2.3 | Information sources and search strategy

Searches were conducted up to September 15, 2020 in MEDLINE (via PubMed), EMBASE and the Cochrane’s Central Register of Controlled Trials (CENTRAL), and Google Scholar. The search strategy for MEDLINE is presented in (Table S1). Additionally, all online issues of the following specific journals were screened by two authors (JZ and NY): Journal of Clinical Periodontology, Journal of Periodontology, Journal of Dental Research, Journal of Clinical Implant Dentistry and Related Research, the International Journal of Oral and Maxillofacial Implants, the International Journal of Periodontics and Restorative Dentistry, and the European Journal of Oral Implantology. One author (JZ) performed snowballing by searching the reference lists of systematic reviews on similar topics and searching for relevant trials protocols on the Pan-African (https://pactr.samrc.ac.za) and national Institutes of Health’s (https://clinicaltrials.gov) clinical-trials registries.

All references were imported in Endnote X9 software,19 in which records were screened and duplicates were removed. Two authors (JZ and NY) first screened titles and abstracts of articles independently and subsequently assessed full-texts of the articles for inclusion in the review. Conflicts were resolved through discussion, and in case of disagreement, a third author (AE) was the referee for the final verdict. Data extraction was performed independently by two authors (JZ and NY) and discrepancies were resolved through discussion. The authors used a customized table that was first piloted then amended as necessary. When efficacy outcomes were not available in the full text or supporting information files, JN and NY attempted to contact the corresponding author to acquire the necessary missing information.

2.4 | Outcomes

Implant failure was defined as clinically detectable implant mobility.20 Any complications reported by the authors of the original article were included under the overall complications as efficacy outcome. Peri-implant soft-tissue esthetics was assessed clinically through the pink esthetic score (PES).21 The PES ranges from 0 to 14 points, with the minimum clinically acceptable value ≥ 8 points.22,23 Vertical bone resorption outcome was stratified into vertical buccal and vertical interproximal bone resorption. Horizontal bone resorption outcome was stratified into horizontal buccal and overall horizontal bone resorption. All bone resorption outcomes were measured either clinically or radiographically. Mid-buccal mucosal recession was measured clinically, and defined as any apical migration of the mid-buccal implant mucosa from baseline. When multiple follow-up durations were reported, the longest was used in meta-analysis. When an RCT used multiple BSM arms, we combined groups that were similar (e.g., multiple types of BSMs).16

2.5 | Risk of bias

Two authors (JZ and AE) independently assessed risk of bias (ROB) using the Cochrane risk of bias tool version 2.0 for RCTs.24 All conflicts were resolved through discussion. The Robvis web application was used to produce and visualize relevant plots.25 Authors JN and AE conducted the Grading of Recommendations, Assessment, Development and Evaluations (GRADE) independently via the GRADEpro GDT web application and resolved conflicts through discussion.26,27

2.6 | Data analysis

Meta-analysis was performed for all outcomes for which quantitative analysis was feasible, using review manager (Revman) 5.4 software.28 Summary estimates and 95% confidence intervals (CI) were generated using the generic inverse variance method in a random-effects model. Risk ratios (RR) and mean differences (MD) were used for dichotomous (implant failure, overall complications) and continuous outcomes (soft-tissue esthetics, vertical and horizontal bone resorption and mucosal recession), respectively. Heterogeneity was evaluated using the P-value of the $\chi^2$ test, $I^2$, Tau$^2$, and visual inspection of overlap between confidence intervals.26,29 When substantial ($I^2 = 50\%-90\%$) heterogeneity was present, pooled estimates were not calculated. If measures of dispersion (e.g., standard deviation) were not reported, values were imputed using mean values obtained from other trials reporting the measure of dispersion for that outcome.16 Descriptive summary of individual trials was provided when summary estimates could not be calculated.

Subgroup analysis was performed to identify potential effect modification when at least two trials were present per subgroup.16 Subgroups that were analyzed included the duration of the follow-up (short-term [≤ 1 year] vs long-term [>1-year]), use of a barrier (membrane vs no membrane), surgical approach (flap/no flap), and healing protocol (submerged/non-submerged). Trials reporting change from baseline and trials reporting only follow-up measurements were pooled together for the respective outcome.16 Sensitivity analysis was performed for trials with imputed values, and for trials at high risk of bias.

3 | RESULTS

3.1 | Study selection

The electronic search (Figure 1) retrieved 19 544 references after removal of duplicates. After title and abstract screening, 85 references remained for full-text evaluation, of which 20 were deemed eligible for inclusion (Table S2). Meta-analysis was possible for 17 trials while the other three were discussed narratively.
Characteristics of included studies

Characteristics of the included trials are presented in (Table 1 and Table S3). The 17 trials comparing BSM to NoBSM included 743 randomized participants (804 implants). Longest reported follow-up ranged from 4 months from placement to 48 months from prosthetic loading, and the predominant site of implant placement was the anterior maxilla.

Eleven trials used a xenograft,30,32,33,35,37-40,42,44,46 two used an alloplast,34,43 one used an autograft,31 one an allograft,36 and one
| Author (year) | Country (setting) | MF ratio | Mean age in years ± SD or (range) | Longest follow-up in months (reference) | Sample size (patients randomized) | Number of implants | Maxilla or Mandible (anterior or posterior) | Intervention | Co-intervention | Control | Outcomes used in the meta-analysis |
|--------------|-----------------|----------|-----------------------------------|------------------------------------------|-----------------------------------|-------------------|------------------------------------------|--------------|----------------|-----------|----------------------------------|
| Bittner et al. | USA (educational) | 9:23 | 52.3 ± 4 | 12 (placement) | 16 | 16 | 16 | 16 | maxilla (anterior) | Xenograft (Bio-oss collagen) | None | None | Mid-buccal mucosal recession |
| Chen et al. | Australia (private practice) | 15:12 | 41.15 ± 1.2 | 24 (loading) | 27a | 35a | 27a | 35a | maxilla (anterior) | Autograft | Resorbable membrane (Resolut) | Resorbable membrane (Resolut) | Non-resorbable membrane (Gore-tex) | Implant failure |
| Chen et al. | Australia (private practice) | 7:13 | 46.8 ± 10.8 | 36 (loading) | 20a | 10 | 20a | 10 | maxilla (anterior) | Xenograft (Bio-oss) | Collagen membrane (BioGuide) | Collagen membrane (BioGuide) | Connective-tissue graft | Complications |
| Cornelini et al. | Italy (educational) | 9:11 | 45 years (21–61) | 6 (placement) | 10 | 10 | 10 | 10 | Both (anterior) | Xenograft (Bio-oss) | Collagen membrane (BioGuide) | Collagen membrane (BioGuide) | Connective-tissue graft | Vertical interproximal bone resorption |
| Daif | Egypt (educational) | 10:18 | 34 (22–48) | 6 (loading) | 14 | 14 | 14 | 14 | mandible (anterior) | Alloplast (β-TCP) | None | None | N/A |
| Angelis | Italy (private practice) | 21:19 | 46.4 (20–77) | 47.3 (24–75) | 12 (loading) | 40 | 40 | 40 | 40 | Both (Both) | Xenograft (Endobon) | Collagen membrane (OsseoGuard) | Collagen membrane (OsseoGuard) | Soft-tissue esthetics | Implant failure |
| Gher et al. | USA (educational) | 15:5 | 58 (26–80) | 6 (placement) | 20 | 20 | 22 | 21 | Both (Both) | Allograft (DFDBA (Calibone)) | Non-resorbable membrane (Gore-tex) | Non-resorbable membrane (Gore-tex) | Vertical buccal bone resorption |
| Girlanda et al. | Brazil (educational) | 4:18 | (21–58) | 6 (placement) | 11 | 11 | 11 | 11 | Maxilla (anterior) | Xenograft (Bio-oss collagen) | None | None | Mid-buccal mucosal recession |
| Grassi et al. | Italy (educational) | 6:9 | 47.3 ± 8.7 | 42.6 ± 10.7 | 6 (placement) | 15 | 15 | 14 | 14 | Maxilla (anterior) | Xenograft (Bio-oss) | None | None | Vertical buccal bone resorption |
| Jacobs et al. | USA (educational) | 8:11 | 68 | 53 ± 20 | 10 (placement) | 19 | 14 | 19 | 14 | Maxilla (anterior) | Xenograft (Bio-oss) | Collagen dressing (Collagen plug) | None | Soft-tissue esthetics |

(Continues)
| Author (year)      | Country (setting) | M:F ratio | Mean age in years ± SD or (range) | Longest follow-up in months (reference) | Sample size (patients randomized) | Number of implants | Maxilla or Mandible (anterior or posterior) | Intervention | Co-intervention | Control | Outcomes used in the meta-analysis |
|------------------|------------------|-----------|----------------------------------|----------------------------------------|----------------------------------|-------------------|------------------------------------------|--------------|----------------|---------|-------------------------------|
| Mastrangelo et al.40 | Italy (private practice) | 31:20 | 44 ± 6.7 | 36 (loading) | 51 | 51 | 64 | 51 | Maxilla (anterior) | Xenograft (Bio-oss) | Collagen membrane (Osteobial) | None | Soft-tissue esthetics |
| Mohamed et al.41 | Egypt (educational) | 2:8 | 33 ± 5 | 6 (placement) | 10a | 5 | 14b | 7 | maxilla (anterior) | Allograft (Puros) | Alloplast (Bio-resorp) | None | Vertical interproximal bone resorption |
| Paknejad et al.42 | Iran (educational) | 3:17 | 38.8 (37-57) | 4–6 (placement) | 20 | 14 | 13 | maxilla (anterior) | Xenograft (Compact bone) | None | None | Vertical buccal bone resorption |
| Prosper et al.43 | Italy (educational) | 39:44 | 46.2 ± 14.3 | 48 (placement) | 83 | 56 | 55 | Both (posterior) | Alloplast (Biosite) | None | Resorbable membrane (Osteoques) | Implant failure |
| Sanz et al.44 | Italy, Spain, UK (educational) | 22:21 | 19:24 | ? | 4 (placement) | 45b | 46b | 43 | 43 | Maxilla (anterior) | Xenograft (Bio-oss collagen) | None | None | Implant failure |
| Spinato et al.45 | Italy (private practice) | 11:30 | 42.5 (22–70) | 32 (placement) | 41 | 22 | 23 | Maxilla (anterior) | Xenograft | Allograft | Autograft | Combination | None | Complications |
| Yuenyongaram et al.46 | USA (educational) | 4:6 | 2.8 | 65.4 (41-83) | 12 (placement) | 10 | 10 | 10 | 10 | Maxilla (anterior) | Xenograft (Bio-oss) | None | None | Implant failure |

*Combined groups.*

*aOne patient had a failure and four lost to the follow up.*

? = not mentioned.
used a combination. For one trial, we combined both the alloplast and allograft BSM groups into one intervention group. As an adjunct to the BSM, six trials used a barrier membrane, one a collagen plug, and another a connective-tissue graft. Five trials used a barrier membrane in the control group.

Four trials with a total of 105 participants (112 implants) compared different types of BSMs with each other. Two trials compared allograft to alloplast, one autograft to allograft, and one alloplast to autograft. The trials did not use barrier membranes, though one trial did use platelet-rich fibrin (PRF) to seal the gap between the implant and the socket. The longest follow-up ranged from 6 to 44 months from placement.

3.3 | Risk of bias assessment

For the outcome implant failure (Figure S1), two trials were at low risk of bias, while the other four were of some concerns. For overall complications as outcome, one trial was at low risk of bias, with the remaining three trials at a high risk of bias (Figure appx1). This was mainly due to deviation from the intended protocol, or only reporting complications narratively rather than quantitatively. Two out of the three trials that evaluated soft-tissue esthetics (Figure appx3) were at low risk, while the third had some concerns.

For the outcome vertical buccal bone resorption (Figure appx6), five trials were judged to have some concerns, and two as low ROB. Two trials were judged to have a low ROB with regard to vertical interproximal resorption (Figure appx8), with the other three at some concerns. For horizontal buccal bone resorption, two trials were at low ROB, and two trials had some concerns (Figure appx10). One trial reported the overall horizontal bone resorption, which was at low ROB. Two out of five trials assessing mid-buccal mucosal recession (Figure appx12) outcome were at low ROB while three were of some concerns.

Of the trials comparing different types of BSMs, three trials were at a high ROB for vertical interproximal bone resorption. One trial was of some concerns for vertical buccal bone resorption. One trial evaluated implant failure and crestal bone resorption. This trial was judged at high ROB and of some concerns, for these outcomes, respectively.

3.4 | Synthesis of results

3.4.1 | Implant failure

Fifteen trials assessed implant failure, but no trial reported time to implant failure. Only six of these trials reported implant failure as a dichotomous (yes/no) outcome, with at least one event observed. The pooled RR for implant failure was 0.92 (95% CI 0.34 to 2.46) (Figure 2) with no detected heterogeneity ($I^2 = 0$). No significant difference for the sub-groups based on duration of follow-up, surgical approach, or use of a barrier was observed. Only one trial involved in the second question contributed a single failure event in the allograft group with a statistically nonsignificant result (RR = 0.33 [95% CI 0.01 to 7.81]).

3.4.2 | Overall complications

Twelve trials reported complications, but only four reported one or more events. The pooled RR was 3.50 (95% CI 1.11 to 11.08) with no detected heterogeneity ($I^2 = 0$), indicating that use of a BSM in patients undergoing IIP increased the risk of complications (Figure 3). The 12 complications in the BSM group, and the three complications in the NoBSM group, were predominantly minor (93.4%). Complications in the BSM group included three events of abscess formation, and two events of pain at prosthetic loading. Other complications consisted of peri-implant infection, persistent inflammation of the buccal mucosa, peri-implant mucositis, small lesions in the peri-implant mucosa, post-operative pain, cover screw loosening, and loosening of the provisional abutment. Complications observed in the NoBSM group consisted of de-cementation of the final prosthesis, post-operative pain, and cover screw loosening. Sensitivity analysis for trials with low ROB for overall complications resulted in only one trial. Though the point estimate of the pooled analysis and this trial were similar, it was not statistically significant (RR = 3.00 [0.64 to 13.98]) (Figure appx22).

3.4.3 | Soft-tissue esthetics

Three trials reported soft-tissue esthetics using the PES. The mean PES was 1.49 (95% CI 0.46 to 2.53) points higher in the BSM

| Study or Subgroup | BSM Events | Total | NoBSM Events | Total | Weight | IV, | Risk Ratio Random, 95% CI | Risk Ratio Random, 95% CI |
|-------------------|-----------|-------|--------------|-------|--------|----|------------------------|------------------------|
| Chen 2005         | 2         | 27    | 0            | 35    | 10.9%  | 6.43 [0.32, 128.60]     |                        |
| De Angelis 2011   | 2         | 40    | 4            | 44    | 39.0%  | 0.40 [0.08, 1.94]       |                        |
| Mastrangelo 2018  | 1         | 64    | 1            | 51    | 12.9%  | 0.80 [0.05, 12.43]      |                        |
| Prosper 2003      | 1         | 56    | 2            | 55    | 17.3%  | 0.49 [0.05, 5.26]       |                        |
| Sanz 2016         | 1         | 45    | 0            | 46    | 9.7%   | 3.07 [0.13, 73.32]      |                        |
| Yuenyongorarn 2020| 1         | 10    | 0            | 10    | 10.2%  | 3.00 [0.14, 65.90]      |                        |
| Total (95% CI)    | 242       | 237   | 100.0%       | 0.92  [0.34, 2.46] | 8 | 8 | Heterogeneity: $\chi^2 = 0.00$; $\chi^2 = 4.08$, df = 5 (P = 0.54); $I^2 = 0$% | Test for overall effect: $Z = 0.17$ (P = 0.86) |

**FIGURE 2** Forest-plot of “implant failure” using a bone-substitute material (BSM) vs no filling material (NoBSM)
group compared with not using a BSM (Figure 4). Only xenografts were used in the trials and there was moderate heterogeneity ($I^2 = 57\%$).

For trials reporting long-term follow-up ($\geq 1$ year), the mean difference was $1.82$ (95% CI $1.09$ to $2.55$) with no detected heterogeneity.

### 3.4.4 Vertical buccal bone resorption

Thirteen trials reported vertical bone resorption, of which seven$^{32,36,38,41,42,44}$ reported vertical buccal resorption. Six trials$^{32,36,38,41,42,44}$ reported the change from baseline and one trial$^9$ reported only the follow-up measurements. The pooled mean difference in vertical buccal bone resorption was $-0.11$ mm (95% CI $-0.33$ to $0.10$) using a BSM compared with control (NoBSM) (Figure 5), with limited heterogeneity ($I^2 = 11\%$). Sub-group analysis of trials that used membranes either in the BSM group only$^{32}$ or in both groups$^{36}$ reported a significantly lower mean resorption of $-1.5$ mm (95% CI $-2.66$ to $-0.34$) in the BSM group (Figure 6). One trial$^{4}$ compared vertical buccal resorption for different types of BSMs, though did not find a statistically significant difference.
3.4.5 Vertical interproximal bone resorption

Of the 13 trials looking at vertical bone resorption, five reported interproximal resorption (Figure 7). Four of these reported change from baseline, while the fifth only reported the follow-up measurement. Two trials did not report standard deviations for vertical interproximal bone resorption, hence these values were imputed accordingly. One trial measured the vertical interproximal bone resorption, but did not provide the mean or standard deviation and was excluded from the analysis. Vertical interproximal bone resorption was lower in the BSM group than the NoBSM group, though not statistically significant (−0.05 mm [95% CI −0.16 to 0.06]). Combining trials reporting change from baseline and trials reporting only follow-up measurements, resulted in substantial heterogeneity ($I^2 = 94.2$) and as such no pooled estimate was calculated. Three trials compared different types of BSMs, and all reported statistically non-significant differences regarding vertical interproximal bone resorption.

3.4.6 Horizontal buccal bone resorption

Five trials reported horizontal bone resorption, of which four reported horizontal buccal resorption. Xenograft was the only type of BSM used in these trials. One trial did not report standard deviations for horizontal buccal bone resorption, hence these values were imputed accordingly. One trial did not provide the mean or standard deviation and was excluded from the analysis. Significantly less horizontal buccal bone resorption was observed in the BSM group (−0.52 mm [95% CI −0.74 to −0.30]), with no heterogeneity ($I^2 = 0$) between trials (Figure 8). Both subgroup analyses (Figures appx20 and appx21) based on
and healing protocol, did not show a significant difference regarding horizontal buccal bone resorption. Only one trial\(^37\) reported the bucco-palatal horizontal dimensions through the ridge width at 1 mm apical to the crest. A lower mean horizontal buccal bone resorption of \(0.22 \text{ mm (95\% CI 0.19 to 0.25)}\) was observed in the BSM group compared with the NoBSM group. One trial\(^49\) compared the mean horizontal buccal bone resorption of allograft (0.72 mm ± 1.46) to autograft (0.08 mm ± 1.95). This trial reported significantly \((P = 0.026)\) less resorption when using an allograft.

### 3.4.7 | Mid-buccal mucosal recession

Five trials\(^30,37,39,45,46\) reported the amount of mid-buccal mucosal recession as a continuous variable. One trial\(^32\) reported recession as present or absent with no threshold mentioned, and thus was excluded from the analysis. Pooled analysis of the five other trials showed a non-significant increase in mid-buccal mucosal recession of 0.02 mm (95\% CI −0.23 to 0.28) in the BSM group, with no detectable heterogeneity \(I^2 = 0\%\); Figure 9). There was no difference between the three trials\(^30,45,46\) reporting change from baseline (0.02 mm [95\% CI −0.25 to 0.30]), and the other two trials\(^37,39\) reporting only follow-up measurements (0.02 mm [95\% CI −0.68 to 0.71]).

### 3.4.8 | Grade

Grading of evidence and reasons for downgrading are presented in Table S4. A moderate certainty of evidence was judged for soft-tissue esthetics, vertical buccal bone resorption, horizontal buccal bone resorption, and mid-buccal mucosal recession outcomes. Implant failure was judged to be at low certainty of evidence judged by the wide confidence intervals and small sample size. Overall complications and vertical interproximal bone resorption were judged to be at very low certainty of evidence. The reasons for this downgrading were wide confidence intervals, small sample size, and heterogeneity.

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**FIGURE 8**  Forest-plot of “horizontal buccal bone resorption” (in mm) using a bone-substitute material (BSM) vs no filling material (NoBSM)

**FIGURE 9**  Forest-plot of “mid-buccal mucosal recession” (in mm) using a bone-substitute material (BSM) vs no filling material (NoBSM)
size, few complications events, and substantial heterogeneity ($I^2 = 94\%$) for the vertical interproximal bone resorption. Moreover, the majority of the involved trials in these two outcomes were of some concerns or at high ROB.

4 | DISCUSSION

In this systematic review, we have evaluated the evidence on the efficacy of using a BSM in the fixture-socket gap in patients undergoing tooth extraction and IIP. In view of the wide confidence intervals, BSM may increase or decrease the risk of implant failure (low certainty). Overall complications were more frequent (very low certainty) though these were predominantly minor. Esthetic scores were significantly higher (moderate certainty) in patients undergoing IIP with BSM.

The results should be interpreted with the considerations that a significant number of trials had at least some concerns for ROB for the relevant outcomes. As a consequence, most recommendations were based on a moderate certainty of evidence except for implant failure (low), complications (very low) and vertical inter-proximal bone resorption (very low). The certainty of the evidence was downgraded because of imprecision, ROB, and inconsistency (Table S4).

Finding no significant differences for implant failure could be explained by the very high success rates of IIP in both review arms, and low absolute risk of developing (serious) complications which might cause implant failure. No trials reported the time to implant failure as an outcome. Although there does not appear to be a clear reason, one possible explanation could be unfamiliarity in the dentistry field with more complex statistical methods, such as survival analysis. Not using survival analysis methods, and therefore, not taking selective censoring into account may result in biased estimates of implant failure.

We found more complications when using a BSM compared with not using one. However, the complications were predominantly minor, mainly peri-implant mucositis, post-operative pain, prosthetic complications, and only one case of peri-implantitis. The higher risk of complications may be due to the use of regenerative materials and the associated technique-sensitive approach. Poor reporting combined with the use of ambiguous definition of what constitute as complications could explain the limited number of complication events observed in included RCTs. The effect of BSM use on soft-tissue esthetics could be explained by the incorporation of the BSM particles in the soft-tissue, and the less horizontal resorption of the supporting bone. This might result in thicker and more supported soft tissues with improved long-term soft-tissue esthetics.

Our review did not demonstrate a mitigating effect of a BSM on vertical buccal bone resorption. The loss of the bundle bone during extraction, in addition to the amount of cancellous component at the alveolar crest of thin labial plates, seems to outweigh the potential positive effect of BSM on the amount of vertical buccal bone resorption. Moreover, the trials’ inclusion criteria that favor IIP such as the presence of an intact post-extraction labial plate, minimized the expected resorption in both arms. Another explanation could be the limited diagnostic performance of the Cone-beam computed tomography (CBCT) in measuring the thin peri-implant bone. Furthermore, the slow resorption rates of the radio-opaque graft materials, could decrease the validity of radiographic and clinical measurements of the vertical buccal bone resorption in the BSM group.

An effect in favor or against the use of a BSM could not be demonstrated for interproximal bone resorption. This finding could be explained by the dependency of the interproximal bone height mainly on the attachment level on the neighboring natural teeth. Also, the benefit of using a BSM on the mid-buccal mucosal recession could not be demonstrated nor refuted. The intactness of the labial plate and the non-submerged healing protocol in the five trials reporting mid-buccal mucosal recession could help understanding this finding.

To our knowledge, there is no clear histological explanation for the significantly less horizontal buccal resorption when using a BSM. All of the trials involved in this outcome used a xenograft, which has a slow resorption rate. Therefore, the hard-tissue structure in the fixture-socket gap could be vital bone, non-resorbed BSM particles in a connective-tissue matrix, or a combination of both. A radiographic, histologic, or clinical distinction should be made to identify the nature of the peri-implant hard-tissue structure, and its implications. After tooth extraction, the horizontal resorption is the most pronounced, compared with the vertical resorption. Mitigating horizontal bone resorption in sites with thin buccal plate (≤2 mm) would provide the patients with many benefits; especially in the esthetic zone. These benefits include better and stable peri-implant soft-tissue esthetics, in addition to improved implant function, and hygiene. Thin biotype and inadequate oral hygiene are risk factors for developing peri-implantitis and subsequent reduction in the quality of life.

This review systematically assessed the efficacy and strength of evidence of BSMs on a wide range of clinical outcomes assessed in RCTs published in both generic and dentistry-specific databases. Both quantitative and qualitative assessments of various clinically relevant outcomes were performed. It also complied to the guidelines provided by international literature regarding reporting, risk of bias assessment, and evaluation of the strength of evidence.

There are some limitations to fully appreciate the findings in this review. No trials reported the time to implant failure; consequently, implant failure was analyzed as a dichotomous outcome. Only a limited number of trials reporting implant failure as the primary outcome were found. These trials had a small sample size, and as a result, the precision of our pooled estimate is limited. The inconsistent methods of reporting and heterogeneity in the measurements and definitions used for the various outcomes posed a challenge, resulting in some concessions (e.g., combining outcome categories). Sub-groups based on type of BSM used, labial plate thickness, or fixture-socket dimensions could not be performed because only one trial was present for some of these subgroups. Moreover, due to the paucity of trials we could not compare the efficacy of different types of BSMs to each other regarding implant failure, overall complications, and soft-tissue esthetics.

Five systematic reviews on the topic of this article have been published. Each of these reviews included both observational and
experimental study designs. Two reviews\textsuperscript{12,15} could not provide a conclusion in favor or against the use of a BSM due to insufficient evidence. Another review\textsuperscript{12} did not provide a quantitative analysis because of high heterogeneity of the included studies. However, this review concluded that the use of guided bone regeneration techniques preserves the peri-implant hard and soft tissues. One review\textsuperscript{23} studied the broad outcome “crestal bone loss”; and reported a non-significant difference with substantial heterogeneity ($I^2 = 59.6\%$). Our findings are in accordance with a systematic review by Lee and colleagues\textsuperscript{14} who reported a MD of 0.01 (95% CI –0.05 to 0.08), and a weighted mean difference (WMD) of 0.84 mm (95% CI 0.53 to 1.14) for implant failure, and the change in horizontal bone dimensions, respectively. However, in this review the definition of the outcome assessing changes in ridge width was not clear. Moreover, this review included only seven trials, one of which been retracted due to the absence of the ethical approval to conduct the trial.\textsuperscript{74}

5 | CONCLUSIONS

1. Compared with no BSM, a beneficial or harmful effect of BSM on implant failure could not be demonstrated nor refuted (low level of certainty).
2. BSM use results in an increased risk of minor complications (very low level of certainty), reduces the amount of horizontal buccal resorption (moderate level of certainty), and improves long-term peri-implant soft-tissue esthetics (moderate level of certainty) compared with no BSM. Therefore, BSM use is recommended in the esthetic zone and sites with thin buccal plate, after discussing the benefits and drawbacks of using a BSM with the patient.
3. A difference in the amount of vertical buccal or interproximal bone resorption (moderate level of certainty), or the amount of mid-buccal mucosal recession (moderate level of certainty) between BSM use and no BSM use could not be demonstrated nor refuted.

6 | IMPLICATIONS FOR FUTURE RESEARCH

1. RCTs of sufficient sample size and consistent reporting of the outcomes.
2. RCTs assessing the added value of using a membrane with a BSM in IIP.
3. RCTs comparing the efficacy of various BSMs to xenografts in IIP.
4. Studies assessing the validity and reliability of the instruments used to measure peri-implant bone dimensions and mucosal recession.

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CONFLICT OF INTEREST

The authors declare no conflicts of interest.

AUTHOR CONTRIBUTION

John Zaki: Conceived the idea; authored the manuscript, screened the titles, selected the trials and extracted the data; assessed the risk of bias, and graded the evidence; designed the analysis plan. Rob J.P.M Scholten: Conceived the idea; designed the analysis plan; authored the manuscript; revised the manuscript. Nernim Yussif: Conceived the idea; screened the titles, selected the trials and extracted the data. Ahmed El-Khadem: Assessed the risk of bias, and graded the evidence. Kevin Jenniskens: Authored the manuscript; revised the manuscript.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available in the supplementary material of this article and from the corresponding author upon reasonable request.

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SUPPORTING INFORMATION
Additional supporting information may be found online in the Supporting Information section at the end of this article.

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