Intraoperative complications and early implant failure after transcrestal sinus floor elevation with residual bone height ≤5 mm: A retrospective multicenter study

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Abstract

Objective: Clinical indications for maxillary sinus floor elevation with transcrestal techniques have increased in recent years even in sites with minimal residual bone height (RBH). Nevertheless, limited information is currently available on incidence of intraoperative complications and early implant failure in these cases.

Material and Methods: This retrospective multicenter study was performed on anonymized clinical and radiographic records of patients who underwent transcrestal sinus floor elevation in seven clinical centers. Influence of different factors related to patient, and sinus anatomy and surgical technique on the incidence of intraoperative complications and early implant failure rate after transcrestal sinus lift were investigated.

Results: A total of 430 patients treated with transcrestal sinus floor elevation for single-implant insertion in sites with RBH ≤5 mm were included in the final analysis. After 1 year of loading, 418 implants of 430 were satisfactorily in function. Early implant failure was recorded in 12 cases (2.8%); results were significantly associated with the presence of large sinus cavities and with the occurrence of membrane perforation.

The following adverse events were recorded: membrane perforation (7.2%), acute sinusitis (0.9%), implant displacement into the sinus cavity (0.7%), oro-antral fistula (0.2%), and benign paroxysmal positional vertigo (0.5% of osteotome cases). A strong direct correlation between sinus membrane perforation and bucco-palatal sinus width (p = .000) was demonstrated.

Conclusions: Early implant failure after transcrestal sinus elevation showed significant direct correlation with bucco-palatal maxillary sinus width and the presence of membrane perforation. Sinus membrane perforation was strongly associated with bucco-palatal sinus width (extremely low perforation rate in narrow and much higher incidence in wide sinuses).
1 | INTRODUCTION

The anatomical configuration of the edentulous posterior upper jaw is often characterized by severe bone shrinking resulting from the combination of both teeth extraction and sinus pneumatization. A widespread surgical option for treating these vertical bone deficiencies is sinus floor elevation with lateral or transcrestal approach (Boyne & James, 1980; Tatum, 1986). The criteria for choosing between these two techniques have been based on residual bone height (RBH), calculated as the distance from sinus floor to the bony crest. The Sinus Consensus Conference held in Boston (1996) and subsequent classifications suggested the transcrestal approach with RBH of 6–7 to 9 mm and lateral window sinus augmentation in the presence of 5 mm or less of bone below the sinus floor (Jensen et al., 1998; Wang & Katranji, 2008).

However, it should be considered that maxillary sinus involvement in the therapeutic plan increases morbidity, costs, and operative risk. A recent retrospective study conducted on 3900 patients who had oral and periodontal surgeries between 1990 and 2018 at the University of Michigan School of Dentistry indicated that lateral sinus floor elevation (together with surgical extraction of impacted teeth) is the procedure associated with more frequent and severe complications compared with other oral, periodontal, and implant surgeries (Askar et al., 2019).

In the last decade, minimally invasive options for the rehabilitation of the atrophic posterior maxilla have been extensively investigated. Recent evidence demonstrated as short and ultrashort implants in the atrophic posterior maxilla represent a rapid and predictable treatment alternative, both for splinted and single-unit implant-supported prostheses. Short implants showed a similar medium-term survival rate, lower morbidity and incidence of complications, better peri-implant marginal bone stability, and reduced treatment time and cost when compared to longer implants placed in augmented sinuses (Al-Moraissi et al., 2019; Ravidà et al., 2019; Ravidà et al., 2021; Yan et al., 2019).

Furthermore, clinical indications for maxillary sinus floor elevation with transcrestal approach have greatly increased in recent years (Block, 2016; Stacchi et al., 2020). Several studies explored the possibility to perform one-stage or two-stage transcrestal sinus augmentation even in sites with residual bone height ≤5 mm, resulting in minimal invasivity and excellent implant survival rate (Bernardello et al., 2011; Gonzalez et al., 2014; Lin et al., 2020; Lombardi et al., 2017; Sisti et al., 2012; Sonoda et al., 2020; Stacchi et al., 2018; Toffler, 2004). Clinical and histologic studies showed that the transcrestal technique is more predictable in terms of endo-sinus new bone formation in narrow than in wide sinuses, irrespective of crestal bone height (Lombardi et al., 2017; Spinato et al., 2015; Stacchi et al., 2018). Therefore, the distance between buccal and palatal sinus bone walls is fundamental when selecting the best surgical option, coupling minimal invasiveness with high predictability of clinical outcomes. A width between buccal and palatal bone walls of 12 mm represents the threshold dividing narrow (≤12 mm) and wide sinuses (>12 mm) (Lombardi et al., 2017; Spinato et al., 2015; Stacchi et al., 2018; Zheng et al., 2016).

Nevertheless, limited information is currently available on the potential influence of anatomical and surgical variables on incidence of intraoperative complications and early implant failure after transcrestal sinus floor elevation with minimal residual bone height.

The present retrospective study aimed to analyze the possible influence of different factors (related to patient, sinus anatomy, and surgical technique) on early implant failure and intra- and post-operative complications in patients requiring sinus floor elevation with transcrestal approach in the presence of RBH ≤5 mm.

2 | MATERIAL AND METHODS

2.1 | Patient recruitment

The present retrospective multicenter study was conducted on anonymized clinical and radiographic records of patients who underwent transcrestal sinus floor elevation between 2000 and 2020 in seven private clinical centers in Italy (C.S., Gorizia; F.B., Terranegra di Legnago (VR); S.S., Sassuolo (MO); T.L., Cassano allo Ionio (CS); R.M., Arco (TN); M.P., Torino; and L.C., Roma). In the informed consent prior to sinus surgery, patients were informed that their clinical and radiographic data could have been used anonymously for research purposes. This study has been reported according to the Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) guidelines (von Elm et al., 2014). The study protocol was approved by the relevant Ethical Committee (Comitato Etico di Ateneo, University of Trieste, nr. 114-31/05/2021).

2.2 | Inclusion and exclusion criteria

All patients (>18 years old) treated in the aforementioned centers with unilateral transcrestal sinus floor elevation were considered eligible for inclusion.

Inclusion criteria were the following: (1) immediate post-operative periapical radiograph; (2) periapical radiograph after 6 months of healing; (3) well-documented medical charts reporting details of the surgical procedure and notes in relation to intra- and post-operative complications; and (4) minimum follow-up of 12 months after prosthetic crown delivery. Exclusion criteria included: (1) incomplete or low-quality clinical and radiographic documentation; (2) pre-operative residual bone height of the alveolar crest >5 mm at the planned augmentation site; and (3) no antibiotic therapy prescribed after sinus surgery.

2.3 | Data collection

Medical records were collected by one trained examiner per center (C.S., F.B., S.S., T.L., R.M., M.P., and L.C.). In order to standardize data collection and study variables assessment, examiners participated in
a calibration meeting prior to the beginning of the study. Data were recorded in a specific case report form.

The following patient-level information was collected:

- age;
- gender;
- systemic diseases;
- medications;
- smoking habits;
- history of periodontal disease.

The following sinus- and implant-level information was collected:

- bone height (BH) at the augmentation site before sinus floor elevation (mm);
- sinus floor inclination at the augmentation site (flat / sloped);
- presence of Underwood septa at the augmentation site;
- bucco-palatal sinus width (SW) at the augmentation site (mm), if pre-operative cone beam computed tomography (CBCT) was available;
- crestal antrostomy technique (rotary instruments / osteotomes / ultrasonic inserts);
- type of grafting material (allograft / xenograft / alloplastic);
- graft characteristics (≤1 mm granules / >1 mm granules / sponge / injectable gel or paste);
- timing of implant placement (simultaneous/staged);
- implant surface treatment (minimally rough / moderately rough / rough) (Albrektsson & Wennerberg, 2004).

The occurrence of the following intra- and post-operative complications was recorded:

- sinus membrane perforation;
- benign paroxysmal positional vertigo (BPPV);
- oro-antral fistula;
- acute sinusitis;
- implant displacement into the sinus cavity;
- early implant failure (implant lost before loading or within the first year of prosthetic function).

2.4 | Radiographic measurements

Periapical radiographs were taken using the long-cone paralleling technique with a Rinn film holder. No attempt was made for further standardization. Pre-operative BH was measured on a 30-inch led-backlit color diagnostic display using a specific software (ImageJ 1.48a, National Institutes of Health, Bethesda, USA). Pre-operative BH was calculated on each periapical radiograph as the linear measurement of the distance between the most coronal point of the alveolar crest at implant site and the sinus floor.

When available, CBCT scan was used to evaluate sinus width (SW) at the augmentation site, defined as the distance between buccal and palatal wall measured at 10-mm height, comprising the alveolar crest at implant site and the sinus floor.

2.5 | Predictor and outcome variables

The present retrospective study aimed to test the possible influence of different factors (related to patient, sinus anatomy, and surgical technique) on early implant failure and intra- and post-operative complications.

2.6 | Primary outcome measure

- early implant failure.

The predictor variables tested for the primary outcome were the following:

- patient level (age, gender, smoking habits, and history of periodontitis);
- sinus level (BH before augmentation, SW, sinus floor inclination, and presence of Underwood septa);
- surgical variables (surgical technique, implant surface treatment, timing of implant placement, and Schneiderian membrane perforation during surgery).

Secondary outcome measures:

- Schneiderian membrane perforation during surgery;
- any complications or adverse events.

The predictor variables tested for the secondary outcomes were the following:

- patient level (age, gender, smoking habits, and history of periodontitis);
- sinus level (BH before augmentation, SW, sinus floor inclination, and the presence of Underwood septa);
- surgical variables (surgical technique, implant surface treatment, and timing of implant placement).

2.7 | Statistical analysis

All statistical analyses have been performed by using the software Stata 16.0 (StataCorp LLC, College Station, USA). Descriptive
statistics has been performed by calculating frequencies for dichotomous data and means with standard deviations for continuous variables. Univariate logistic regression analysis was first performed to select factors associated with the presence of the primary and secondary outcomes of the present study. Subsequently, the predictor variables which resulted significant at the univariate analysis were inserted in a stepwise multivariate logistic regression model, setting a $p$-value of .157 in the stepwise backward model as suggested by Heinze and Dunkler (2017).

3 | RESULTS

3.1 | Clinical outcomes

A total of 783 patients treated with transcrestal sinus floor elevation and single-implant insertion matched inclusion criteria and were further screened. Three-hundred and thirty-six patients were excluded: 28 had incomplete clinical or radiographic documentation, 2 patients did not undergo post-operative antibiotic therapy, and 306 presented pre-operative residual bone height >5 mm. A total of 447 patients were included in the preliminary analysis. Surgery and prosthesis were performed in seven clinical centers (F.B. = 108; L.C. = 20; T.L. = 41; R.M. = 96; M.P. = 68; S.S. = 63; and C.S. = 51), with a minimum follow-up of 1 year after prosthetic loading.

Transcrestal sinus access was performed with subtractive techniques [specific burs (Cosci & Luccioli, 2000) or ultrasonic inserts (Kim et al., 2014; Sentineri & Dagnino, 2011): 53.2% and 3.8% of the subjects, respectively] or by bone compaction [osteotomes (Franceschetti et al., 2014; Summers, 1994): 43.0% of the subjects] and sinus membrane elevation was obtained by incremental grafting material insertion. As only 3.8% of the subjects ($n = 17$) had the procedure performed with ultrasonic inserts, it was decided to exclude them from the logistic regression analysis to avoid possible skewing of results. Finally, 430 patients (207 males and 223 females; age range: 30–84 years; mean: $58.3 \pm 11.5$ years) were included in the final analysis. Selection process has been summarized in Figure 1. Demographic data of patients included in the study have been presented in Table 1, while maxillary sinus

![Figure 1: Patient selection process flowchart](image)
characteristics and details of surgical technique have been listed in Table 2.

After 1 year of prosthetic loading, 418 of 430 implants were satisfactorily in function (97.2%). Early implant failure occurred in 12 cases (2.8%): stepwise multivariate logistic regression analysis demonstrated a significant correlation with the presence of large sinus cavities [OR = 8.50; 95%CI (1.02–70.42); p = .047] and with the occurrence of sinus membrane perforation [OR = 4.21; (95%CI: 1.10–16.05); p = .035] (Table 3).

3.2 | Radiographic outcomes

BH at the augmentation site before sinus floor elevation ranged from 1.0 to 5.0 mm (mean 4.0 ± 0.9 mm). BH at the augmented site 6 months after transcrestal sinus augmentation ranged from 1.0 to 19.0 mm (mean 9.4 ± 2.8 mm), with a mean vertical bone gain of 5.4 ± 2.9 mm (range 0 to 18.0 mm).

SW at the augmentation site, measured on CBCT at 10-mm level comprising the residual alveolar crest, was ≤12 mm (narrow sinus) in 176 patients and >12 mm (wide sinus) in 161 patients. It was not possible to classify sinus width in 93 patients, due to CBCT Unavailability.

ICC score for radiographic measurements (>0.92) resulted in an excellent intraexaminer repeatability: mean difference in BH and SW was 0.12 and 0.17 mm, respectively.

3.3 | Intra- and post-operative complications

The most frequent adverse event was Schneiderian membrane perforation (n = 31; incidence 7.2%) (Figure 2): 2 perforations occurred in 176 narrow sinus cavities (incidence 1.1%), 26 perforations in 161 wide sinus cavities (incidence 16.1%), and 3 perforations in sinuses with unknown bucco-palatal width. When perforation was detected after crestal osteotomy, collagen sponges or resorbable membranes were inserted prior to implant placement, without the use of bone substitutes. Other complications comprised of acute sinusitis (n = 4; 0.9%), implant displacement into the sinus cavity (n = 3; 0.7%), and oro-antral fistula (n = 1; 0.2%). Benign paroxysmal positional vertigo was recorded in one case, following the use of osteotomes (0.5% of osteotome cases).

Univariate logistic regression analysis showed a strong direct correlation between sinus membrane perforation and bucco-palatal

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**Table 1** Demographic characteristics of the included patients. Data are expressed as mean ± standard deviation

| Gender   | 207 males (48.1%) | 223 females (51.9%) |
|----------|-------------------|---------------------|
| Age      | 58.3 ± 11.5 years - range 30–84 years |

| Smoking Status | No (%) | Yes (%) |
|----------------|--------|--------|
| History of Periodontitis | 317 (73.7%) | 113 (26.3%) |
| Systemic Disease | 268 (62.3%) | 162 (37.7%) |
| Medication | 366 (85.1%) | 64 (14.9%) |
| Medication | 372 (86.5%) | 58 (13.5%) |

**Table 2** Surgical site and surgical intervention characteristics. BH: bone height

**Surgical site (n = 430)**
- Sinus Floor Shape
  - Flat: 319 (74.2%)
  - Sloped: 111 (25.8%)
- Sinus Width
  - Narrow ≤12 mm: 176 (40.9%)
  - Wide >12 mm: 161 (37.4%)
  - Not Available: 93 (21.7%)
- Underwood Septa
  - Absent: 383 (89.1%)
  - Present: 47 (10.9%)
- Pre-operative BH mean 4.0 ± 0.9 mm range 1.0–5.0 mm
- BH after 6 months mean 9.4 ± 2.8 mm range 1.0–19.0 mm

**Surgical intervention (n = 430)**
- Technique
  - Burs: 238 (55.3%)
  - Osteotomes: 192 (44.7%)
- Graft Type
  - Xenograft: 289 (67.2%)
  - Allograft: 68 (15.8%)
  - Collagen: 61 (14.2%)
  - Synthetic: 7 (1.6%)
  - No graft: 5 (1.2%)
- Graft Formulation
  - Granules ≤1 mm: 221 (51.4%)
  - Granules >1 mm: 112 (26.0%)
  - Sponge: 61 (14.2%)
  - Gel: 31 (7.2%)
  - No graft: 5 (1.2%)
- Implant Placement
  - Simultaneous: 385 (89.5%)
  - Staged: 45 (10.5%)
- Implant Surface
  - Moderately Rough: 414 (96.3%)
  - Rough: 16 (3.7%)
Multivariate analysis was not performed as no other predictor variable reached statistical significance. It was not possible to perform a statistical analysis for the post-operative complications due to their extremely low numerosity.

### DISCUSSION

In the present study, bucco-palatal maxillary sinus width seems to play a crucial role in the success of implant rehabilitations: wide sinus cavities (>12 mm at 10-mm height, comprising the alveolar crest) resulted significantly associated with early implant failure. A wide sinus cavity has been shown by numerous studies to represent a biologically unfavorable environment for new bone formation after both lateral and crestal sinus lift (Avila et al., 2010;
TABLE 4 Univariate analysis for the outcome “Membrane Perforation”

| Number of implants = 430 | Univariate Analysis |  |  |  |
|-------------------------|---------------------|---|---|---|
| Membrane perforation    | OR [95% CI]         | p-value |
| Age                     | 0.99 [0.96–1.03]    | .711 |
| Gender                  |                     |     |
| Female                  | 1                   |     |
| Male                    | 1.59 [0.74–3.41]    | .235 |
| Periodontal Disease     | 1.19 [0.55–2.57]    | .612 |
| Smoking                 |                     |     |
| No                      | 1                   |     |
| Yes                     | 1.54 [0.69–3.43]    | .287 |
| Underwood septa         |                     |     |
| Absent                  | 1                   |     |
| Present                 | 1.73 [0.63–4.78]    | .287 |
| Sinus Floor Shape       |                     |     |
| Flat                    | 1                   |     |
| Sloped                  | 0.78 [0.32–1.88]    | .584 |
| Sinus Width             |                     |     |
| Narrow                  | 1                   |     |
| Wide                    | 15.47 [3.59–56.59]  | .000* |
| Residual Bone Height    | 0.76 [0.51–1.13]    | .176 |
| Technique               |                     |     |
| Burs                    | 1                   |     |
| Osteotomes              | 1.33 [0.63–2.82]    | .460 |
| Implant Placement       |                     |     |
| Simultaneous            | 1                   |     |
| Staged                  | 1.88 [0.68–6.13]    | .224 |

Note: OR, Odds Ratio.*p-value <.05

FIGURE 3 The force required for membrane detachment is directly correlated with the surface of the elevated area. Transcrestal techniques allow adequate membrane elevation in narrow sinuses (a), where elevated surface is smaller than in wide ones. In wide sinuses (b), the force required to properly elevate the membrane could exceed its deformation capacity, resulting in increased perforation risk.
Main limitations of the present study are inherent to the retrospective study design as well as to the sample characteristics: as they were used records not designed for the study, some available data may be of poor quality and/or potential confounding factors may not have been properly controlled.

Moreover, Schneiderian membrane perforation during transcervestral sinus floor elevation may not have been always recognized and their final number could have been underestimated (Garbacea et al., 2012). Finally, possible heterogeneity in clinical practice among the seven clinical centers could be an additional confounding factor in interpreting the results of the present study.

ACKNOWLEDGMENTS

The authors would like to thank Gaia Olga Guercio for the drawings included in the present article. Open access funding provided by Universitat Bern.

AUTHOR CONTRIBUTIONS

Claudio Stacchi: Conceptualization (equal); formal analysis (equal); methodology (equal); project administration (equal); supervision (equal); validation (equal). Fabio Bernardello: Investigation (equal); resources (equal); validation (equal); writing – review and editing (equal). Sergio Spinato: Data curation (equal); investigation (equal); writing – review and editing (equal). Rossano Mura: Data curation (equal); investigation (equal); writing – review and editing (equal). Michele Perelli: Investigation (equal); writing – review and editing (equal). Giuseppe Troiano: statistical analysis (equal); review and editing (equal). Luigi Canullo: conceptualization (equal); methodology (equal); project administration (equal); investigation (equal); writing – review and editing (equal).

DATA AVAILABILITY STATEMENT

Data available on request from the authors

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How to cite this article: Stacchi, C., Bernardello, F., Spinato, S., Mura, R., Perelli, M., Lombardi, T., Troiano, G., & Canullo, L. (2022). Intraoperative complications and early implant failure after transcrestal sinus floor elevation with residual bone height ≤5 mm: A retrospective multicenter study. Clinical Oral Implants Research, 33, 783–791. https://doi.org/10.1111/clr.13959