Effects of Corticobasal Implant Protrusion inside the Nasal and Maxillary Sinus

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Abstract

Background: Implant protrusion into the nasal and maxillary sinuses presents a challenge in cases of severely resorbed maxillae. Aim: The aim of this study was to evaluate the clinical and radiographic effects of BECES® implant penetration depth into the nasal and maxillary sinuses. Setting and Design: This was an observational study conducted in a tertiary institution. Materials and Methods: Forty-nine BECES® implants were inserted into the maxilla of patients who presented with severely resorbed ridges but no history of sinusitis. Forty-five implants protruded into the sinus cavities. Patients were examined clinically and radiographically at 1 week and 3, 6, 12, and 18 months after insertion. Maxillary sinus health, survival and success rates, and peri-implant health were assessed using the plaque index (PI), calculus index, modified gingival index (MGI), and probing pocket depth (PPD). Statistical Analysis: Wilcoxon signed-rank test and Mann–Whitney test were used in this study. Results: Four (8.16%) of the 45 implants that penetrated the cavities reached the sinus floor without disrupting the membrane; the penetration depth was ≥4 mm in 20 implants (44.44%) and <4 mm in 25 (55.56%). No patient showed clinical or radiographic signs of sinusitis during the observation period. There were significant differences in the PI, MGI, and PPD values between baseline and the 18-month follow-up with no association with the penetration depth. All implants showed radiographically direct bone-to-implant contact. Where the implant tip barely reached the sinus floor, the membrane healed uneventfully while when deeply penetrating the sinus, the membrane healed around the implant but did not cover the tip. All prostheses and implants survived during the observation period. Conclusion: Penetration depth of polished implants with cortical engagement into the maxillary sinus or the floor of the nose does not negatively affect implant survival, the success rate of the treatment, nor peri-implant soft-tissue health. It also does not provoke the development of sinusitis.

Keywords: BECES® implant, cortical engagement of implants, corticobasal implant, maxillary sinus, nasal sinus

INTRODUCTION

Many methods have been described for the management of complete and partial edentulism. Recently, implant treatment has become the gold standard for replacing missing teeth if adequate bone is present.[1] Cases of severe alveolar ridge resorption remain a challenge in dental implantology.[1-3] The posterior maxilla often presents with limited bone height and poor bone quality, which jeopardizes the primary stability of the implant. A further problem is the complication of maxillary sinus “pneumatization.”[1-3] Conventional maxillofacial therapy has devised methods to increase the bone quantity in cases of severe bone loss, such as sinus lifting and sinus augmentation procedures with bone grafts harvested from the hip, tibia, ribs, or chin followed by insertion of the implant.[1,3] This procedure was first documented by Boyne and James in 1980[7] and subsequently

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by Tatum.[9] Summers[9] introduced the sinus floor elevation technique using an osteotome with a flapless crestal approach, possibly accompanied by simultaneous placement of an implant.

Although a bone augmentation procedure can increase the bone height and width, it has some disadvantages, including the need for two independent invasive surgical procedures, which not only prolongs the treatment significantly but also results in additional costs and risks as well as patient discomfort.[10,11]

Numerous postoperative complications of sinus lifting procedures have been reported, including wound dehiscence, acute/chronic sinusitis, mucocele formation, swelling, discharge of graft material into the sinus, and graft infection resulting in complete loss of the graft and continuing morbidity at the donor site.[10-15]

Furthermore, sinus lifting procedures have been associated in many studies with the risk of perforation of the sinus membrane. The literature reports a 44% incidence of sinus membrane perforation during implant osteotomies and/or implant placement.[15] Consequently, implants protrude into the nasal or maxillary sinuses. Reports on the effects of this protrusion on the health of the nasal and maxillary sinuses are mixed. Several groups have found no adverse effects[1,3,4,16-25] while others have reported mild adverse effects.[1,24]

An electronic and manual literature search by Ragucci et al.[3] reported after penetration of implants through the floor of the maxillary sinus a clinical complication rate of 14.8%, among which epistaxis being the most common side effect. They also found a radiographic complication rate of 3.4%, the most common being thickening of the Schneiderian membrane, with no significant difference according to the level of implant penetration. However, they documented a survival rate of about 95.6%, with no significant effect of level of implant penetration. They classified the implant penetration depths into two groups (>4 mm and ≤4 mm), and the results for both groups were the same.

In the past few years, several authors have recommended the use of Corticobasal® implants to achieve bicortical anchorage (e.g., crest of the ridge and maxillary sinus floor) and to ensure implant stability in patients with severe ridge resorption or compromised bone support.[26-36] Consequently, implants may have exposed to the nasal and maxillary sinuses [Figure 1] with possible challenging effects. Although many studies have reported that Corticobasal® (BECES®) implants have very favorable outcomes with limited and manageable complications,[27,29,34-36] there is still some doubt concerning the effect of their penetration into the nasal and maxillary sinuses. In 2018, Lazarov[37] published a prospective cohort study in which patients treated with Strategic Implant® technology who underwent placement of 375 BCS® implants in 105 maxillary sinuses were followed for 3–4.5 years. The maxillary sinuses were assessed preoperatively on panoramic views and clinical examination. Patients with symptoms of acute sinusitis were not treated until their symptoms had disappeared. Postoperative control panoramic views were taken and clinical examinations were performed in all patients at 3 years and 4.5 years. Only one sinus showed radiographic and clinical symptoms of a sinusitis. Lazarov concluded that polished implants placed in such a way to intentionally or unintentionally penetrate the maxillary sinus do not cause any sinus reaction, and they can remain stable for many years. Furthermore, all symptoms of sinusitis were transient and resolved completely after various treatments even though the implants were left in place. Little information is available concerning the peri-implant health of the mucosa around penetration areas in the maxillary and nasal sinuses.

The aim of the present study was to evaluate the effect of protrusion of a BECES® implant inside the nasal and maxillary sinuses and to investigate the effect of penetration depth on peri-implant health and the implant survival rate.

**Materials and Methods**

**Characteristics of the study sample**

The study was approved by the ethics committees of the hospital and the Ministry of Health, Khartoum, Sudan. All patients who underwent Corticobasal® implant treatment between 2015 and 2017 and presented with severe alveolar ridge resorption with possible protrusion of an implant inside the nasal or sinus cavity were asked to participate in the study. All patients provided informed consent for participation in the study and publication of the results after receiving a detailed explanation of the objectives of the study, the protocol, and the possible complications.

The study inclusion criteria were age 18–80 years and complete or partial edentulism. All patients had severe ridge resorption and requested implant treatment. The patients were questioned about their history and symptoms of maxillary sinusitis,[38] including nasal bleeding, congestion, or obstruction, nasal secretion, and pain or tenderness in the infraorbital region.

![Figure 1: A clinical photograph showing an implant that is potentially protruding into the nasal cavity using sinoscopy](image-url)
Patients with a history of sinusitis or bisphosphonate therapy were excluded.

**Surgical and prosthetic procedures**

Preoperative cone-beam computed tomography (CBCT) scans (Planmeca ProMax; Planmeca, Budapest, Hungary) were obtained for all patients. The scans were obtained using a standardized technique with the same scanner. All images were analyzed under standardized conditions. The implants were placed under local infiltration anesthesia with 2% lidocaine and epinephrine 1:100,000 after application of Betadine 5% at the implant sites. The implant osteotomy was performed using a drill with copious irrigation, followed by cortical penetration, administration of 2 ml of Betadine 5%, and placement of the implant. All implants (BECES® Brand, Manufacturer Simpladent GmbH, CH-8737, Gommiswald, Switzerland) were inserted using a standard one-stage surgical procedure. The length and width of the implant were selected according to the amount of available bone. High primary stability was achieved and tested using the reverse-torque technique at 35 Ncm. Amoxicillin and clavulanate potassium (Meganox, 1 mg; Hikma Pharmaceuticals, Amman, Jordan), diclofenac potassium (Rapidus, 50 mg; Tabuk Pharmaceutical Manufacturing Co., Tabuk, Kingdom of Saudi Arabia), and xylometazoline (Otrivine, adult nasal drops; GlaxoSmithKline, Brentford, UK) were prescribed for all patients. The penetration depth of the implant was measured on CBCT scans obtained postoperatively using the machine’s software.

Impression copings were secured to the head of the implant, and an impression was taken using monophase vinyl polysiloxane (Ivoclar Vivadent AG, Schaan, Liechtenstein). The metal framework was constructed on the following day, and a heavy body silicone jaw relation was taken. This metal framework was used to provide implant splinting, to reduce load per unite area, and to ensure better force distribution that minimizes the possible causes of implant overloading. Either acrylic or composite veneer material was applied to the framework depending on the extent of loss of soft and hard tissues. One day later (within 72 h), the final prosthesis was inserted and cemented using Fuji cement (Fuji I Luting Cement, GC Corporation, Tokyo, Japan). Patients were scheduled for follow-up clinical and radiographic examinations at 1 week and 3, 6, 12, and 18 months.

**Follow-up evaluation**

**Sinuses**

Any signs and symptoms of sinusitis such as mucopurulent drainage, nasal obstruction, facial pain/pressure/fullness, headaches, and an impaired sense of smell were noted.[38]

**Implant success and survival rates**

In line with the literature, implant survival was defined as the presence of the implant in the mouth at the time of examination.[39] The implant success rate was assessed using the modified James–Misch implant quality of health scale[39] (I, success [optimum health]; II, survival [satisfactory health]; III, survival [compromised health]; and IV, failure [clinical or absolute failure]), the Albrektsson[40] criteria for implant success, which includes a bone loss of <1.5 mm in the 1st year and <0.2 mm yearly thereafter, and the presence of suppurative and peri-implant infection, continuous pain, mobility, or persistent peri-implant radiolucency.[40] The sample was categorized into two groups, i.e., those with an implant penetration depth ≥4 mm and those with a penetration depth <4 mm measured on CBCT.[3]

**Clinical examination**

**Peri-implant soft tissue**

The plaque index (PI) is a three-point scale devised by Mombelli et al.[41]: 0, plaque detected; 1, plaque detected by running a probe across the implant; 2, plaque visible to the naked eye; and 3, abundance of plaque). The modified gingival index (MGI) was determined according to the modified Loe and Silness index[42] (0, absence of inflammation [normal peri-implant mucosa]; 1, mild inflammation [slight change in color] and slight edema; 2, moderate inflammation [redness, edema, and glazing]; and 3, severe inflammation [marked redness and edema] and ulceration). The calculus index (CI) was reported based on its presence or absence of calculus around the implants both labially and lingually (1, presence and 0, absence). The CI was reported as a yes/no variable. The probing pocket depth (PPD) was measured in millimeters using a short-shank probe with gentle pressure from the mucosal margin to the bottom of the pocket in both a labial/buccal and palatal direction.[43] If complete peri-implant soft-tissue healing around the implants had been achieved, the pocket depth was reported as 0 mm; introduction of the probe was prohibited given that “consensus on probing around basal implants”[43] does not recommend probing as a routine step. However, these data should be assessed and reported for comparison with other implant studies. Patients were asked about their level of satisfaction, any complaints, and whether or not they would choose the same treatment modality again.

The implant penetration depth and the mean PI, MGI, and PPD scores were recorded at five follow-up visits. The mean was then categorized as: 0 = 0, 0.2–0.4 = 1, 0.6–1.0 = 2, and 2 = 3 for calculation of the associations between penetration depth and the index scores and PPD using contingency coefficient and Chi-square tests.

**Prostheses**

The prostheses were inspected for decementation, lip support, extent of visible smile, mobility, unnatural wear of the opposing dentition, and fracture of the veneer material.

**Radiographic evaluation**

Radiographic assessment was performed using CBCT to detect implant loss and/or fracture, to evaluate bone-to-implant contact, and to detect any changes in the nasal and maxillary sinuses at 1 week and 6, 12, and 18 months. The intrarater reliability was determined using repeated measurements at varying time intervals spanning several days and was 0.95, 1, 1, and 0.90 for the PI, CI, MGI, and PPD, respectively.
Statistical analysis
The Wilcoxon signed-rank test was used to compare the PI, modified MGI, and PPD between baseline and the 18-month follow-up visit. All data recording and statistical analyses were performed using the Statistical Package for the Social Sciences (SPSS) software (SPSS version 22; IBM Corp., New York, NY, USA). \( P < 0.05 \) was considered statistically significant. The 95% confidence interval was also calculated.

RESULTS
Sample distribution
Forty-nine implants were found to have penetrated into the nasal and maxillary sinus cavities. The mean patient age was 63.6 ± 14.22 years. The immediate (baseline) postoperative radiographs revealed that 4 (8.16%) of the 45 implants that had penetrated the nasal and sinus cavities had reached the sinus floor without disrupting the membrane; the penetration depth was ≥4 mm for 20 implants (44.44%) and <4 mm in 25 (55.56%).

Nasal and maxillary sinuses
Clinical findings
None of the patients showed any signs or symptoms of rhinitis or sinusitis during follow-up. All patients maintained good oral hygiene without pain or implant-related exudates.

Radiographic findings
Comparison of the radiographs obtained at 6, 12, and 18 months with those obtained at baseline did not reveal any inflammatory reactions at or around the implants, signs of osteolytic reactions, polyps or granulation around the implants inside the sinuses, lost implants, or implant fractures. All implants showed newly regenerated bone surrounding the threads, indicating an increase in the bone-to-implant contact despite the different penetration depths [Figures 2 and 3]. For those implants that merely penetrated or disrupted the sinus membrane, it was found that the membrane had healed and tended to cover the implant tips [Figure 2a and b]. When the implant penetrated the maxillary sinus deeply, there was still an increase in bone contact around the implant; in these cases, the membrane healed around the tip of the implant but did not cover it [Figure 3a and b].

Figure 2: Cone-beam computed tomography scans showing an implant that just barely made contact with the Schneiderian membrane. (a) Immediately after implant insertion. (b) At the 18-month follow-up visit. There are no inflammatory reactions at or around the implant, no signs of an osteolytic reaction, no polyps, and no granulation around the implant inside the sinuses. The implant shows an increase in bone–implant contact

Implant health and survival rate
According to the implant health assessment scale,[39] all patients presented with optimum health at the 18-month follow-up, with no pain or tenderness on function, no (zero) mobility, not more than 2 mm of radiographic bone loss from baseline, and no history of exudates. Therefore, the implant survival and success rates were both 100%.[39]

Peri-implant soft tissue
There was a statistically significant difference in the PI score between baseline and the 18-month follow-up \([ P = 0.038, \text{ Wilcoxon signed-rank test} \); Table 1]. Despite the reported increase in the PI readings in some patients, the score ranged between 0 and 2 [Table 1]. The CI did not change over time in any case \([ \text{mean} 0.00 \pm 0.00; \text{ Table 1} \). There was also a significant decrease in the MGI score \([ P = 0.025, \text{ Wilcoxon signed-rank test} \). The mean was 0.22 (range, 0–1). There was a significant decrease in the PPD \([ P = 0.003, \text{ Wilcoxon signed-rank test} \) with a range of readings from 0 mm to 3 mm. Clinical examination revealed that the soft tissue healed very rapidly even in sites where implants were inserted into fresh extraction sockets and a complete soft-tissue collar present around the vertical shaft of the implants.

Association between implant penetration depth and plaque index, modified gingival index, and probing pocket depth values
There was no significant difference in the PI, MGI, or PPD value according to implant penetration depth \([ P = 0.066, \ P = 0.283, \ P = 0.550, \text{ respectively, Mann–Whitney test}; \text{ Table 2} \). Moreover, there was no association between implant penetration depth (≥4 mm or <4 mm) and the PI, MGI, and PPD values \([ P = 0.116, P = 0.262, \ P = 0.480, \text{ respectively}; \text{ Tables 3-5} \). There was an apparent increase in the peri-implant bone level exhibited around the vertical shafts and horizontal plates in all implants [Figures 2 and 3].

Figure 3: A cone-beam computed tomography scan showing an implant that penetrated the Schneiderian membrane and beyond, deep into the maxillary sinus. (a) Immediately after implant insertion. (b) At the 18-month follow-up visit. There are no inflammatory reactions at or around the implant, no signs of an osteolytic reaction, no polyps, and no granulation around the implant inside the sinuses. The implant shows an increase in bone–implant contact
Patient satisfaction

All patients were satisfied with their treatment, reported an improvement in their quality of life, and said they would choose the same treatment again.

Discussion

Resorption of the residual alveolar ridge following a tooth extraction, as well as pneumatization of the maxillary sinus, can compromise the placement of a dental implant in the maxillary posterior region. Several researchers[26,32,33] have recommended anchorage of bicortical crestal bone and the sinus floor implant to improve implant stability in patients with limited bone quality. Consequently, implants may protrude into the maxillary or nasal sinuses, resulting in perforation of the maxillary sinus membrane.

Although some studies have reported an incidence of sinusitis associated with implant penetration inside the maxillary sinus, other studies have reported no change in the maxillary sinus, and there are a number of reports of epistaxis as common clinical complications and thickening of the Schneiderian membrane as the main radiographic complication. Therefore, there was a need for a clinical study with a longer observation period to investigate these controversial results.

According to the literature,[1,44] sinusitis of dental origin accounts for about 10%–12% of all cases of maxillary sinusitis and is associated with many etiological factors, including dental infections, such as periapical granuloma, periapical abscess, periodontal infection, inflammatory cyst, oroantral fistula, a foreign body introduced inside the sinus (such as filling material, root fragment, broken instrument, and large odontogenic cyst),[45-47] and a migrated dental implant.[5,48-51] In the present study, although some of the implants investigated showed penetration of >4 mm, there was no clinical or radiographic evidence of adverse effects in the nasal or maxillary sinuses, such as nasal congestion, sinusitis, polyp formation, or mucocele. This finding is in accordance with previous reports[1,4,16-23] that revealed no association between implant protrusion into the maxillary sinus per se with the incidence of maxillary sinusitis. This observation could be attributed to the smooth surface of the implant and the small insertion tip that limits the interruption of the blood supply, resulting in the rapid healing of peri-implant soft tissue.[27,28,32] Petruson[49] reported normal mucosal conditions with no signs of increased secretion or infection around implants in which the tip penetrated the nasal or maxillary sinus even though the surface of the implants used in their study was rough.[49]

Variable implant penetration levels were noted in this study; in the case of minor penetration, the membrane healed and covered the tip of the implant. This is confirmed the observations of Zhong et al.[4] who reported that a surgically disrupted membrane around the apical portion of an implant healed and covered the tips of the implants if the protruding depth was <2 mm. Jung et al.[17,18] found that the sinus mucosa covered implants penetrating the sinus floor by <2 mm in mongrel dogs. Computed tomography scans showed that implant protrusion of >4 mm in the maxillary sinus caused thickening of the sinus mucosa around the implants; however, these sinuses remained asymptomatic. In contrast, Nooh[24] reported that perforation of the membrane with dental implants may be associated with minor manageable complications, such as epistaxis and sinusitis.[24]

The high implant survival rate in this study was in accordance with that in the study by Ragucci et al.[3] who reported survival
rates of 99.5% and 98.5% with implants that have a penetration depth of ≤4 mm and >4 mm, respectively, with no significant difference between the two depths. Nooh\cite{24} mentioned an overall survival rate of about 98.4% after 1 year of follow-up. Seven patients showed mild epistaxis during the immediate postoperative period, and there was one case of sinusitis.

The mean PI reported in this study was in line with that reported by Mombelli et al.,\cite{52} who rated the amount of plaque accumulation around the surface of the submucosal aspect of the implant-abutment on a scale from 0 to 3. The use of acrylic resin as a veneering material to compensate for the severe loss of soft and hard tissues may have caused the increased plaque accumulation on the restorations. This finding confirms the observations of other investigators\cite{53-55} who reported that the micropores and structural inconsistencies of the acrylic surface supported adhesion of dental plaque and microorganisms.

The excellent results for the MGI are credited to the smooth polished surface that allows rapid attachment of soft tissue around the implants. The polished vertical implant slot prevents adherence of plaque and apical transmission of the bacterial load inside the basal aspect of the implant, far away from the site of bacterial infection (the oral cavity). Our observations confirmed the results of Hede et al.\cite{31,33} Lazarov,\cite{27} Ahmad et al.,\cite{28} and Singh et al.,\cite{29} who recommended this implant design for use at compromised implant sites, such as the maxillary sinus, the floor of the nose (where retrograde peri-implantitis could occur), or extraction sockets. These results are similar to the experimental and clinical results in maxillofacial traumatology and orthognathic surgery, in which penetration of smooth bone screws into the sinus and the nasal aspect of the floor of the nose did not result in any complications, infection, or bone loss.\cite{56,57}

The increased bone-to-implant contact found in this study is consistent with the finding of Khairnar and Gaur,\cite{20} who reported increased bone formation and excellent primary stability after indirect elevation of the nasal membrane with smooth polished implant. Zhong et al.\cite{14} found high removal torques and intimate bone-to-implant contact in bicortical implants in rabbits. Furthermore, Kim et al. reported a 100% implant survival rate with a significant increase in bone around the implants when the initial residual bone is <5 mm in height.

All patients in this study presented with severe ridge resorption that precluded rehabilitation with conventional implant-supported prostheses unless preceded by bone-grafting procedures. However, any removable prostheses made for such patients may be associated with compromised retention and stability due to the poor status of the denture-bearing area. Rehabilitation of these patients with fixed basal implant-supported prostheses greatly improved their satisfaction. This result is in line with other studies conducted with conventional dental implants.\cite{58-61}

**Conclusion**

Based on the clinical and radiographic evaluation of the patients in this study (within the limitations of a small sample size and short follow-up period), we can conclude that protrusion of a BECES® implant into the nasal or maxillary sinuses does not compromise the success or survival rate of the implant nor the health of the sinuses. Both the implants and the peri-implant soft tissues in this study were deemed to be in excellent health. Further research should be performed using larger samples and more extended follow-up periods.

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**Conflicts of interest**

There are no conflicts of interest.

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