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Use of plasma rich fibrin and Bio-OSS/SINT-Oss for implant-prosthetic rehabilitation in maxillary atrophy with sinus pathology: A 48-month follow-up

F Inchingolo,1 A Ballini,2 SA Mura,2 D Farronato,3 N Cirulli,2 F Pettini,1 E Gheno,1 D Vermesan,4 P Pederzoli,1 G Resta,2 M Caprio,5 F Muollo,2 G Marinelli,2 AD Inchingolo,2 G Malcangi,2 S Cantore,2 M Del Corso,2 M Debededitiss,2 AM Inchingolo,6 M Serafini,2 S Diteodoro,2 F Schinco,2 R Cagiano,5 D De Vito,7 R Cortelazzi1 and G Dipalma2

Abstract
The maxillary sinus floor elevation procedure has gained popularity with predictable results, and is a safe, acceptable technique for bone augmentation, providing a base for dental implant treatment. Faint radiopaque lesions at the base of the maxillary sinus are frequent diagnoses on radiographs and must be identified during dental implant planning. The use of autografts, xenografts, allografts, and alloplasts or a combination between them has been demonstrated to be effective for increasing bone height and bone volume in maxillary sinus. The objective of this study was to evaluate the outcome of subjects with considerable sinus membrane pathology (test group) undergoing maxillary sinus floor augmentation using PRF as a filling material, in association with the Bio-Oss and Sint-Oss and simultaneous implant placement in a one-stage surgical procedure. All patients reported no pain to percussion, no sign of tissue suffering to the soft peri-implant tissues, the presence of an optimal primary stability of the inserted implants, and the increase in the peri-implant bone density. No complications were encountered during follow-up periods in these patients, including no negative evolution in the sinusitis and all implants are functioning successfully. In conclusion, the use of PRF and Piezosurgery reduced the healing time, favoring optimal bone regeneration and allowing sinus membrane integrity to be maintained during surgical procedures, according to evidence-based dentistry.

Keywords
Bio-Oss, chronic rhinosinusitis, plasma rich fibrin, Sint-Oss, sinus lift

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Introduction

Dental implants have revolutionized dentistry. After almost 50 years, the placement of a titanium screw with an attached tooth has now grown to be society’s standard for permanently replacing missing or severely damaged teeth.\(^1\)

The enthusiastic intention to treat edentulous maxillary areas with endosseous implants has often clashed with the assumption that the loss of teeth is reflected in progressive bone reabsorption.\(^2\)

This bone deficiency may result from an excessive pneumatization/inflammation of the maxillary sinus, from an important atrophy of the alveolar crest referable to dental extractions and/or periodontal diseases, from both of these causes.\(^3\)

Inflammation of the maxillary sinus, such as rhinitis and sinusitis usually co-exist and are concurrent in most individuals; thus, the correct terminology is now rhinosinusitis.\(^4\) Because dental implantation and sinus augmentation have been widely performed in recent years, one of their possible complications, maxillary sinusitis, has become a major concern for both dentists and otolaryngologists.\(^4\)\(^–\)\(^6\)

A widely used technique for augmenting bone tissue volume in the maxilla is maxillary sinus lifting, in which, by means of osteotomy of the anterior maxillary sinus wall, the sinus membrane is exposed, and must be divulsed to obtain a receptor site for the bone tissue.\(^7\)

One of the most common accidents in this technique is perforation of the sinus membrane while performing osteotomy which, in some cases may make a further surgical procedure unfeasible.\(^8\)

Vercellotti et al.\(^9\) was a pioneer when he presented the use of piezoelectric ultrasound as a new alternative in performing osteotomies in maxillary sinus lifting surgeries. Piezosurgery appears to be an extremely advanced and conservative tool when compared with the existent methods for the treatment of bone and soft tissues.\(^10\) The osteotomy is performed by means of pendular vibratory movements, with nanometric variations of amplitude, by the use of various points.\(^11\)

Nowadays, international consensus or scientific literature about the possibility to perform oral rehabilitation and maxillary sinus grafting in the case of pre-existing sinus membrane pathology are not present.\(^\square\) Bone density and local inflammatory processes influence the operative protocol and the choice of the type of implant used in order to replace the lost teeth\(^12\) and maxillary bone losses often require additional surgical procedures.\(^9\)

Cone beam computerized tomography (CBCT) favors the obtaining of images with lower radiation doses and reduced cost, due to its technology capable of producing images with submillimetric isotropic spatial resolution.\(^13\) and for that, it is indicated especially for the dentomaxillofacial region.\(^14\) A grafting material takes the role of substitute of the insufficient bone tissue if it meets biocompatibility criteria, if it has an optimal response to biomechanical stress and a great capacity to replace the functions of synthesis/reshaping of the bone structure, essential for a correct turnover and for a good functionality of the tissue.\(^15\) In recent years, PRF concentrates have been widely used as a supplement to tissue regeneration procedures.\(^16\)\(^–\)\(^17\)

For these reasons, the aim of the present study was to evaluate the outcome of subjects with considerable sinus membrane pathology undergoing maxillary sinus floor augmentation using Platelet Rich Fibrin (PRF®) as a filling material, in association with the deproteinized bovine bone (Bio-Oss®) and beta tricalcium phosphate (Sint-Oss®) and simultaneous implant placement.

Materials and methods

Patients

Fifty-six patients (32 men, 24 women) in the age range of 55–73 years (mean age, 64 years), who received a dental implant between June 2009 and June 2010, were selected from multicentric private practice dental cabinets in Bari, Barletta, Carate Brianza, Milan, and Chieti.

In order to minimize the operator’s bias, surgical procedures were performed by only two operators, maintaining the same equipment in all performed surgical procedures.

The present study followed the rules established in the 1975 Declaration of Helsinki, then revised in 1983. Before any treatments, the patients were properly informed about the advantages and disadvantages of all the planned procedures, first verbally and then with a written form for the informed consent.

A total of 175 implants were placed in these patients. Patients were included in the study according to the following criteria: (1) patients having multiple consecutively edentulous space both in maxillary and mandibular region and with compromised dental elements needing multiple extractions; (2) adequate bone quantity and quality at the implant site; (3) patients well motivated for...
implant therapy and maintaining good oral hygiene. Patients who were either non-compliant with appointments or had acute sinusitis were excluded from the study prior the surgical procedure.

Radiographic examination

An X-ray orthopantomogram was performed before the surgical procedure (Figure 1).

Patients were evaluated for sinus pathology for a period of 6–48 months after bone transplantation and implant insertion using preoperative and postoperative panoramic radiological imaging and CBCT scan for the purposes of diagnosing their sinus pathology (Figure 2).

Furthermore, CT is employed in the diagnosis and follow-up of patients who are candidates for, or have been submitted to grafting techniques, which allows bone areas to be evaluated by means of measurements and different densities with great precision.13,14

Imaging was employed to monitor the progress of all patients.

The above CBCT scan examinations have shown that 26 patients of Group A (Control Group) were not showing alterations of the anatomy and physiology of the maxillary sinus, such as thickening of the Schneider membrane or radiographic opacification which could suggest the presence of an inflammatory process.

In the other 30 patients of Group B (Test Group), on the basis of the radiographic findings, it was possible to hypothesize the presence of opacification of the maxillary sinus.

All patients were questioned for complaints and symptoms of sinusitis preoperatively, and any positive findings were assessed by an otolaryngology consultation prior to the surgical procedure.

Surgery preparation

All the patients underwent a careful hematologic and cardiologic evaluation (blood routine, cardiological examination, ECG) in specialist wards to optimize and customize the prophylaxis and the preoperative pharmacologic therapy (details shown in surgical procedure).

The bone loss was assessed by means of radiological investigations and CBCT; the choice of implants was case-specific, according to the rehabilitative necessity (Stone® -I.D.I.-EVOLUTION, Monza, Italy).

Before each surgical treatment, a full mouth disinfection protocol18 consisting of scrupulous oral hygiene was practiced by supragingival and subgingival scaling with adjuncts of antimicrobial (chlorhexidine, CURASEPT 0.2%, Curadent, Italy). Moreover, before surgery, the authors decided to administer per os amoxicillin/clavulanic acid (NEODUPLAMOX, Procter & Gamble, Italy, of 2 g, 1 h before surgery and 1 g 6 h after surgery).

Preparation of PRF®

As reported from previous experience,19 the protocol is performed in accordance with Choukroun’s procedure, introduced with the European Directive no. 2004/23/CE of 31 March 2004.

The patients were subjected, before surgery, to complete hemogram analysis. Before starting the surgical procedure, 5 mL of the patient’s venous
blood was drawn from the antecubital vein in a sterile 10 mL glass test tube (free from anticoagulant) and centrifuged at 3000 rpm for 10 min (Centrifugette model 4206, ALC International).

The upper straw-colored layer was then removed and the middle fraction was collected, 2 mm below lower dividing line, which was the PRF.

This PRF was used in two ways: a part of it was included in a sterile cup and mixed with Bio-Oss®/Sint-Oss®, until a homogeneous material is obtained; the other part was divided into two sterile compresses, one of which was modeled as a resistant fibrin membrane, transferable to the Schneiderian membrane, and the other was transferred to the material placed before closing the wound.

**Surgical procedure**

The authors opted for the surgical procedure of one-stage sinus lift (implant insertion occurred concurrently with sinus lift, and the healing and integration time was 6–9 months), except for eight selected cases in which the height of the residual bone was lower than 5 mm.

After local and regional anesthesia administration (20 mg/mL mepivacaine chloride CARBOPLYINA-Dentsply, Italy) containing adrenaline as a vasoconstrictor (1:100,000), all patients underwent a lateral window approach, in accordance with Tatum’s technique.19

The crestal mucosa was incised and two release incisions were also made. The muco-periosteal flap was then turned over in order to expose the bone surface beneath.

The osteotomy to delimit the bone window was performed by Surgybone® Piezosurgery (Silfradent Co., Sofia, Italy) (Figure 3) and the Valsalva maneuver was performed to be sure of the membrane integrity.10

The sinus membrane was carefully elevated from the sinus floor and medial sinus wall; radiographic depth gauge was used to confirm that the depth was correct.

PRF was used in combination with autogenous bone, an organic bone material, and organic bone substitutes (Bio-Oss® and Sint-Oss®).

This mixture was inserted in the surgical alveolus by means of a syringe or an amalgam carrier (Kavo Kerr Group, Italy), and pushed upwards with the osteotome, in order to lift the Schneiderian membrane and keep clear the first part of the implant bed (Figure 4), and at the end of this stage, two membranes were placed.

They were obtained from the flattening of the remaining PRF between two sterile gauzes and then placed in order to close the access area to the sinus site.

At this phase, implants were inserted, and the length was in the range of 2–4 mm in excess, compared to the height of the available bone.

Lastly, the bone window was closed to avoid traumatisms, the flap was replaced and then the mucosal areas were sutured through a resorbable material.19

During implant insertion, which had to be very slow, the filling material displaced the Schneiderian membrane and was placed around the implant, thus avoiding the collapse of the membrane on its surface.

At the end of surgery, an endoral periapical radiography with a positioner was performed to assess the implant position compared to the maxillary sinus and the implants did not make contact with the sinus membrane in any of the patients.

In the eight patients in which the height of the residual bone was lower than 5 mm, the fixture was
inserted in a later stage (4 months after the grafting procedure), in order to allow the stabilization and integration of the grafted material, and then the restoration of an adequate bone quantity to ensure the predictability and stability of implant rehabilitation.

Patients were given postoperative instructions, and prescribed antibiotics (amoxicillin/clavulanic acid 1000 mg every 12 h, for 7 days; NEODUPLAMOX, Procter & Gamble, Italy), analgesics/anti-inflammatory drugs (intramuscular injection of ARTROSILENE, Dompè SpA, Italy; ketoprofen 160 mg, every 12 h for 5 days), and mouthwash with clorexidine (CURASEPT 0.12%, Curadent, Italy, every 12 h for 14 days).

Patients in Group B also received instructions for aerosol therapy consisting of 2 mL of nebulized corticosteroid (hydrocortisone and diprofillin, CORT-INAL, Teofarma SRL, Italy) and 500 mg/4 mL of antibiotic (Thiamphenicol glycinate acetyl-cysteinate; FLUIMUCIL ANT. IN. TOP, Zambon, Italy), twice per day for 10 days before and 10 days after surgery.

Follow-up

Synthetic absorbable sterile surgical suture stitches (Vicryl® sizes 3.0–4.0) were removed after 7 days.

After implant placement, clinical parameters were recorded and evaluated at baseline, 3, 6, 9, and 12 months. The clinical parameters recorded were the modified plaque index,20 modified gingival index,21 probing depth around the implant at four sites using the Tissue pressure sensitive (TPS) probe,22 implant mobility according to clinical implant mobility scale,23 and the absence or presence of any infection around the implant.

X-ray orthopantomogram and a second-level radiographic examinations (Denta-Scan) were performed at 15, 60, 90, 120, and 180 days and 6, 12, 24, 36, and 48 months after surgery (Figures 5 and 6).

At the 1-year follow-up, patients were shown auxiliary cleaning methods given that they had been instructed on how to brush in order to improve oral hygiene. [AQ: 3]

Results

A total of 175 dental implants were inserted in 56 patients (32 men, 24 women) between 2009 and 2010.

All procedures were successfully concluded without significant procedural or postprocedural complications or implant failure. Healing, in general, was uneventful with minimal discomfort to all the patients. All the sites maintained excellent peri-implant soft and hard tissue conditions.

In 30 patients with a pre-existing radiographic opacification which could suggest the presence of an inflammatory process (54% of both groups), the sinus pathology regressed or remained unchanged in 18 patients (32% in Group B), while in two patients in Group B the sinus membrane pathology was limited to evaluation by periapical X-rays and in six patients (11% in Group B) it completely disappeared as shown in CBCT scan examinations.

All the patients receiving the procedure reported a subjective improvement of nasal breathing with increased perception of the air passageway.
From a clinical point of view, the authors noticed peri-implant tissue healing, with a successful implant-prosthetic rehabilitation. All patients reported no pain to percussion, no sign of tissue suffering to the soft peri-implant tissues, and the presence of an optimal primary stability of the inserted implants, as confirmed by the fulfillment of Albrektsson’s implant success criteria.\(^\text{24}\)

The radiological examination, performed after 6 months, revealed the presence of newly-formed bone tissue, well amalgamated with the residual bone, and also showed an average increase in the peri-implant bone density of 31\% (Figures 5 and 6).

In the apical region, the authors also noticed close contact of implants with the newly-formed bone.

One year after loading there were no dropouts and no failure of the definitive prosthesis occurred.

**Discussion**

The basis of reduction of the volume of the maxillary sinus is multifactorial and includes congenital and acquired variants; among the latter are neoplastic, traumatic, iatrogenic, inflammatory, and systemic causes.\(^\text{4,25}\)

Maxillary sinus grafting is a predictable and reliable procedure that has been routinely performed for more than 30 years.\(^\text{5}\)

The complication rate is low, but some cases may require additional surgery, and the outcome of oral rehabilitation may be affected.\(^\text{6}\)

The use of Piezosurgery involves considerable advantages for the oral surgeon: an osteotomy with micrometric cut for complete surgical accuracy and a high intraoperative sensibility; a site-specific selective cut to minimize the damage to the soft tissues, which is an important quality to preserve the sinus membrane while creating the trans-cortical access window in sinus lift; and an excellent intraoperative visibility thanks to a bloodless surgical field, ensured by the effect of cavitation.

Both the measurements previously performed on radiograms and the tactile sensation of contact with the cortical surface allowed the experienced clinician to appreciate the contiguity with the sinus floor.

In the present study, even in cases where the surgery was considered risky and not suitable for the presence of a clear pathological state of departure that could affect the final result and exposure to a higher risk of complications, there was full success in centering the objective of regenerative bone therapy and the authors also performed the collateral objective of no negative evolution in the sinusitis.

For this purpose, the choice of the ‘filling’ material represents an essential aspect for a complete and real achievement of the above-mentioned objectives.

In fact, in recent years, various techniques have been proposed and several types of materials have been assessed (autologous and eterologous); these were critically reviewed by means of clinical studies and histological investigations.\(^\text{26}\)

The continuous pursuit of the ideal filling material has often focused the attention on the adoption and marketing of several bone substitutes, such as autologous bone, demineralized and freeze-dried bone, hydroxyapatite, and different combinations of these materials, often with satisfactory results in terms of biocompatibility, induction of bone formation, and of course, implant stability.\(^\text{26}\)

As a supplement to procedures of tissue regeneration, a platelet concentrate called PRF was tested for the first time in France by Choukroun et al. and introduced with the European Directive no. 2004/23/CE of 31 March 2004.\(^\text{27}\)

PRF belongs to a new generation of platelet concentrates; unlike Platelet Rich Plasma (PRP), PRF is obtained without the addition of anticoagulants such as heparin, EDTA, bovine thrombin, and so on.\(^\text{27}\)

During the production of PRF, apart from platelets, other cellular elements are activated, such as leukocytes, which release cytokines after the artificial hemostatic and inflammatory phenomenon induced by centrifugation.\(^\text{28}\)

There are, therefore, three pro-inflammatory cytokines (IL-1\(\beta\), IL-6, and TNF\(\alpha\)), an anti-inflammatory cytokine (IL-4), and an angiogenesis promoter (VEGF).\(^\text{27}\)

Hence, PRF is able to regulate inflammation and stimulate the immune process of chemotaxis.\(^\text{28}\)

PRF is an autologous graft material that eliminates any risk of disease transmission; besides, its gelatinous consistency enhances clotting and graft stability.\(^\text{19,28}\) However, it has an important defect: as it is a biomaterial directly obtained from the patient’s blood, its quantity is modest.

PRF has the special feature to undergo polymerization during centrifugation, naturally and slowly; active thrombin and fibrinogen concentrations are almost physiological, because the material does not require any addition of bovine thrombin.\(^\text{27,28}\)
As a result, the fibrin assumes a three-dimensional conformation, and the combination of fibrin monomers leads to the formation of a trimolecular structure, thus forming a soft and permeable mesh network and allowing rapid colonization of the scar cells.27 This natural material actually seems to accelerate the physiological wound healing; besides, in association with bone grafts, it seems to accelerate new bone formation.29

Most of the authors who used PRF in their dental practice affirm that there was no risk of infection, disease transmission, or side effects.26 In this regard, a careful revision of literature on the possible effects of the therapeutic use of growth factors (GFs) (including PRF), with relation to carcinogenesis, to their influence on tissues with epithelial dysplasia or oral carcinoma, and to their relation with growth and tumor infiltration is absolutely necessary.

Besides, it is essential to underline that little is known about the ideal concentration of each GF or the optimal dosage for each actual therapeutic situation, and it seems that concentrated growth factor (CGF) is similar to PRF and may be useful in the future to increase the success rate of bone grafting.29

In the present study, all patients participated until the end of the study with no clinical dropout reported; all patients showed good compliance in oral hygiene maintenance and followed other instructions and no complications were recorded in any of the cases.

Furthermore, the use of PRF® and Piezosurgery reduced the healing time, favoring optimal bone regeneration and allowing sinus membrane integrity to be maintained during surgical procedures, according to evidence-based dentistry.30

In conclusion, the authors conclude by stressing the need to continue this study on a wider cohort of patients, and to carry out large-scale analyses which could provide statistically significant criteria about the success of this procedure.

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