Minimally Invasive Alveolar Ridge Augmentation Procedure (Tunneling Technique) Using rhPDGF-BB in Combination with Three Matrices: A Case Series

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This study investigated a minimally invasive surgical procedure for alveolar ridge augmentation that combined recombinant human platelet-derived growth factor BB (rhPDGF-BB) and three different matrices. The minimally invasive tunneling ridge augmentation procedure was applied to 12 patients randomized into three groups: rhPDGF-BB (0.3 mg/mL) was combined with freeze-dried bone allograft (FDBA; group A), anorganic bovine bone graft (ABBG; group B), or anorganic bovine bone graft/mineralized collagen bone substitute (ABBG/MCBS; group C). Computed tomography (CT) scans were obtained presurgically and prior to 14-week re-entry surgery. Clinical reentry revealed adequate bone volume to place implants in all patients in groups A and B and two of four patients in group C. Trephine core biopsies were obtained and evaluated by microCT, backscatter scanning electron microscopy (BE-SEM), and light microscopy. New bone formation was consistently observed with BE-SEM and histologic analysis for group A and B specimens. Newly formed woven and lamellar bone were in close contact with graft particles. The ABBG/MCBS specimens (group C) had more variable results, with fibrous encapsulation of graft particles and limited histologic evidence of new bone formation. Within the limits of this study, the FDBA and ABBG carriers appear to be appropriate scaffolds to deliver rhPDGF-BB for ridge augmentation via minimally invasive surgical techniques. (Int J Periodontics Restorative Dent 2009;29:371–383.)

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This surgical approach appears to reduce postoperative discomfort and swelling. A number of variations on the subperiosteal tunneling technique have been attempted and reported, with particulate hydroxyapatite and particulate human mineralized bone allograft used to enhance thin alveolar ridges. Decortication of the recipient site and the use of a resorbable collagen membrane in conjunction with anorganic bovine bone graft have been reported. Recombinant human platelet-derived growth factor BB (rhPDGF-BB) is a natural biologic molecule that mediates and regulates key cellular events such as cell proliferation, chemotaxis, and matrix synthesis by binding to specific cell-surface receptors. The wound-healing response is activated when PDGF is released locally during clotting by platelets at sites of soft and hard tissue injury. Once released from the platelets, PDGF binds to cell surface receptors to promote rapid cellular migration (chemotaxis) and proliferation (mitogenesis). It is proangiogenic in that it acts in synergy with endogenous vascular epithelial growth factor to stimulate neovascularization at the defect site.

Preclinical and clinical studies, including a large-scale randomized controlled trial, have demonstrated the clinical safety and efficacy of rhPDGF-BB–mediated therapy in both periodontal regeneration and implant site development. These studies of ridge augmentation have provided evidence of the periosteal influence on rhPDGF-BB–mediated osteogenesis.

The present study evaluated a minimally invasive ridge augmentation procedure (tunneling technique) that used rhPDGF-BB in combination with three particulate scaffolds, including freeze-dried bone allograft (FDBA), anorganic bovine bone graft (ABBG), and mineralized collagen bone substitute (MCBS).

**Method and materials**

This investigation was designed and implemented as a single-center, prospective, open-label clinical study. Subjects were selected for enrollment from the population of patients requesting dental implant placement in the maxillary anterior region and presenting with a ridge that was inadequate in width to place 4-mm-diameter dental implants. To be eligible, prospective subjects had to be between the ages of 18 and 70 years and willing and able to follow the study protocols. Subjects signed informed consent documents according to the Declaration of Helsinki. In addition, subjects were excluded from entry into the study if (1) they presented with systemic disorders that would prevent them from undergoing surgery or (2) they were current smokers (within 6 months of entry into the study). Pregnant women were also excluded from the study.

At the screening visit, the eligibility criteria were assessed with reviews of dental and medical histories, along with extraoral and intraoral examinations. Once the eligibility criteria were met, the patient was scheduled for clinical photographs, periapical radiographs, dental computed tomography (CT) scan, and dental prophylaxis, and was scheduled for the surgery.

Alveolar ridge augmentation was performed in the maxillary anterior region with local anesthesia (2% lidocaine with 1:100,000 epinephrine). A minimally invasive tunneling ridge augmentation procedure using two short vertical releasing incisions adjacent to the site of augmentation was performed. The buccal tissue was carefully dissected and lifted from the alveolar crest with a periosteal elevator to form a subperiosteal pouch. Decortication of the buccal plate was performed manually with a custom-fabricated decorticator to stimulate bleeding into the defect and allow for access to progenitor cells and angiogenesis.

Prior to grafting, the 12 subjects were randomized into three experimental groups (Table 1).

- **Group A:** FDBA (The University of Miami Tissue Bank) hydrated with rhPDGF-BB (Gem21S, Osteohealth)
- **Group B:** ABBG (Bio-Oss, Osteohealth) hydrated with rhPDGF-BB
- **Group C:** ABBG/MCBS (Bio-Oss Collagen, Osteohealth) hydrated with rhPDGF-BB

In groups A and B, 1 g of collagen (Hemostop, Technew) was mixed with graft material to enhance its handling characteristics.

The grafts were delivered to the deficient area in a syringe (3 mL, Embraplast) (Fig 1). Following its insertion, the graft was packed and condensed until stable. Primary wound coaption was achieved with simple
interrupted sutures (nylon sutures, Ethicon). Subjects were instructed to take oral antibiotics (amoxicillin 500 mg three times daily for 7 days) and ibuprofen (800 mg four times daily for 3 days), use a mouth rinse (0.12% chlorhexidine two times a day), and avoid brushing of the treated site for 2 weeks. They were also provided instructions on when to return for follow-up visits, as indicated in the protocol. Oral hygiene instruction was reviewed at each visit. The 3-month visit included periapical radiographs, a CT scan, and dental prophylaxis.

Surgical reentry of the treatment site was performed 3.5 months (14 weeks) after augmentation surgery under local anesthesia (2% lidocaine with 1:100,000 epinephrine) with the elevation of a full-thickness mucoperiosteal flap. Trephine drills were used to obtain core biopsies (minimum 2 mm wide by 7 mm deep) from the center of the treated area as well as the lateral surface (horizontal core). These non-decalcified bone cores were scanned, and the data were quantified using three-dimensional (3D) microCT (µCT 40, Scanco Medical). Dental implants were placed in the augmented sites when primary stability could be achieved.

### Table 1

<table>
<thead>
<tr>
<th>Group/ Subject</th>
<th>Age</th>
<th>Gender</th>
<th>Site(s)*</th>
<th>Graft (g)</th>
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<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>301</td>
<td>45</td>
<td>F</td>
<td>12,11,21,22</td>
<td>2 g FDBA</td>
</tr>
<tr>
<td>305</td>
<td>51</td>
<td>F</td>
<td>11,21,22</td>
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<tr>
<td>310</td>
<td>56</td>
<td>F</td>
<td>12,11,21,22</td>
<td>3.5 g FDBA</td>
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<tr>
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<td></td>
<td></td>
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</tr>
<tr>
<td>302</td>
<td>37</td>
<td>M</td>
<td>12,11,21,22</td>
<td>3 g ABB</td>
</tr>
<tr>
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<td>M</td>
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</tr>
<tr>
<td>309</td>
<td>40</td>
<td>F</td>
<td>12,11,21,22</td>
<td>3 g ABB</td>
</tr>
<tr>
<td>Group C</td>
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</tr>
<tr>
<td>304</td>
<td>31</td>
<td>F</td>
<td>12,11</td>
<td>2 g ABB/1 g MCBS</td>
</tr>
<tr>
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<td>M</td>
<td>14–11</td>
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<td>2 g ABB/0.25 g MCBS</td>
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<td>2 g ABB/0.25 g MCBS</td>
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</table>

Two milliliters of rhPDGF-BB were used in each patient in combination with the selected graft material.
*FDI tooth-numbering system used.
The primary outcome variable was bone quality and quantity assessed by light microscopy, histomorphometry, and microCT and backscatter SEM. The secondary outcome variables included bone volume, bone quality, soft tissue healing, and incidence of unanticipated adverse healing events.

Results

Twelve patients (four men with a mean age of 37.35 years and eight women with a mean age of 42.25 years) were enrolled based on the inclusion criteria and treated with the tunneling technique for ridge augmentation. There were no serious adverse events during the course of the study.

Clinical Results

Early wound healing demonstrated typical postsurgical sequelae, such as localized facial edema and gingival/mucosal erythema. Two patients in group B and one patient in group C showed fenestration of soft tissues at the 7-day postoperative visit. All of the sites subsequently healed, with complete wound closure by the 28-day postoperative visit. The fenestrations occurred at the buccal aspect of the grafted sites, where the continuity of the mucosa had been compromised by instrumentation during elevation or condensing of the graft particulate.

Group A

Clinical examination of the group A patients at the 14-week postoperative visit revealed enhancement of ridge width in all four patients, as confirmed by the 12-week CT scan. The newly grafted bone was more radiopaque than the native bone. However, it was obvious that the ridge shape lacked uniformity, with significant irregularity of the surface contour.

Surgical reentry demonstrated well-incorporated bone particulate presenting increased ridge volume with minor surface irregularities and partial undercuts in the apical third of the ridge (Fig 2). The regenerated bone appeared to be firm on the surface, but variable bone density was noticed at the time of implant site preparation. All planned implants were placed.

Group B

Clinical examination of the group B patients at the 14-week postoperative visit revealed significant enhancement of the ridge width in all four patients. The CT images evidenced well-incorporated graft material. Flap elevation and removal of soft tissues revealed enhanced edentulous ridge form, with clinical quality consistent with new bone (Fig 3). The augmented sites allowed for implant placement in all patients.

Group C

Clinical examination of the group C patients at the 14-week postoperative visit revealed moderate enhancement of the ridge width in all four patients. The graft particulate was embedded in the soft tissue flap and the lack of incorporation noted on the CT scan (Fig 4).

Fig 2 (left) Well-incorporated bone particulate was noted in group A sites. HC = horizontal core; VC = vertical core.

Fig 3 (right) Visual identification of the graft particles in group B sites. HC = horizontal core; VC = vertical core.
Two of the four patients showed no increase in ridge width, prohibiting implant placement.

### Quantitative CT scan evaluation

Postsurgical CT scans indicated that there was a volumetric increase at the defect sites in all three groups (Fig 5). Measurements were performed by an independent (blinded) radiologist to compare presurgical to postsurgical defect dimensions, evaluating for a width that would be sufficient for implant placement. The height of the ridge extending to a 6-mm width was evaluated (Table 2). Eleven of the 12 CT scans, including 20 of the 23 potential implant sites, were available for evaluation. The other three readings were discarded owing to poor CT scan technique.

### Table 2: Preoperative and postoperative defect dimensions, as derived from CT scans

<table>
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<th>Subject</th>
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*Width at 6 mm.

\( ^\dagger \)FDI tooth-numbering system used.
Figs 6a and 6b  MicroCT representation of sample from patient 310 (group A). Red = host bone; white = FDBA particles.

Figs 7a and 7b  MicroCT representation of sample from patient 302 (group B). Red = host bone; white = ABBG particles.

Figs 8a and 8b  MicroCT representation of sample from patient 306 (group C). Red = host bone; white = ABBG/MCBS particles.
MicroCT and histomorphometric analyses

The microCT evaluation could not differentiate between local bone, newly formed bone, and graft material because the grey levels were too similar (Figs 6 to 8). Thus, only the total amount of mineralized tissue per core was evaluated by microCT. The mean percentage (± SD) of mineralized tissue in group A was 34.6% ± 8.7%; in group B it was 38.2% ± 8.7%; and in group C it was 52.9% ± 12.9%.

MicroCT evidenced intense osteogenesis with extensive new bone formation. It was possible to isolate remaining matrix. The graft particulate was apparent to varying degrees in all of the samples.

BE-SEM and histologic evaluation

Group A

BE-SEM and histologic evaluations showed FDBA chips and newly formed cancellous bone surrounding them and between them, indicating osteoconductive bone formation (Fig 9a). Bone formation was ongoing at the time of biopsy sampling (Figs 9b to 9d).
Group B
Group B specimens showed the presence of ABBG surrounded by new bone. BE-SEM grey levels indicated the degree of mineralization (Fig 10a). Light microscopy demonstrated intensive new bone formation with varying degrees of maturation, including newly formed woven bone and lamellar bone forming bridges between ABBG particles (Figs 10b to 10d). Some specimens demonstrated new bone formation within the ABBG particles.

Group C
Group C results varied with location of the bone core. Multiple cores demonstrated MCBS particles surrounded by connective tissue (Fig 11). Newly formed bone was observed in one specimen. Other specimens showed solid bone with limited remnants of MCBS particles.
Implant placement and success

Twenty implants were placed into the trephined biopsy sites at the 14-week postoperative visit (seven implants in group A, eight implants in group B, and five implants in group C). Implant placement was not possible in two patients in group C owing to immature bone quality and/or limited bone volume. Three implants in group A had to be removed because of mobility and buccal dehiscence at 6 months (one implant) and 8 months (two implants). The implants were removed prior to beginning the restorative phase of treatment. The patients used provisional removable partial denture appliances prior to restoration. The remaining implants were clinically and radiographically stable at the 12-month evaluation.

Discussion

There is a desire for minimally invasive surgery that offers a predictable and simple method to augment localized edentulous ridge defects. The present authors used a tunneling technique to obtain access to deficient ridges (maxillary anterior region) with minimally invasive incisions and minimal tissue elevation. This technique reduces the possibility of postoperative soft tissue loss or deformity.

All patients enrolled in the study presented with ridge defects that prohibited implant placement. They were segregated into three different treatment groups based on combinations of scaffolds and rhPDGF-BB. All eight patients in groups A (FDBA and rhPDGF-BB) and B (ABBG and rhPDGF-BB) demonstrated consistent increases in ridge dimension and graft maturity, and implant placement proceeded in all potential implant sites.

The concept of stabilization and intimate adaptation of a membrane to the surrounding bone to create a secluded space and to prevent the ingrowth of competing nonosteogenic cells into the defect area has been challenged recently by the results that have been obtained with rhPDGF-BB therapy. Preclinical and clinical studies have supported the use of xenografts without a membrane for significant ridge augmentation in both horizontal and vertical defects. It is thought that a membrane may inhibit progenitor cell migration and angiogenesis by presenting a physical barrier to chemotaxis. To achieve graft stability without a membrane, there was a need for a large volume of the graft material in each treatment site (2 to 3 g), compared to that reported by Hasson (1 to 1.5 g).

New bone formation was consistently observed with BE-SEM and histologic analysis for groups A and B.

Fig 11. Light microscopic view of the fibrous encapsulation of the mineralized collagen bone substitute (MCBS) particles. CT = connective tissue.
specimens. Newly formed woven and lamellar bone were consistently observed in close contact with graft particles in groups A and B specimens. The FDBA and ABBG particulate grafts appear to act as sufficient scaffolds to deliver the rhPDGF-BB signaling molecules to the recipient site and allow for normal new bone growth to enhance the edentulous ridge. In contrast, the MCBS specimens (group C) had more variable results, with fibrous encapsulation of graft particles and limited histologic evidence of new bone formation.

On reentry, the augmented ridges presented with variable contours and undercuts beyond the new bone. It appears that the use of a membrane might be reconsidered for the benefit of improving graft containment and the contour of the final grafted sites with this technique.

Qualitative assessment of the sites via CT scan revealed findings consistent with the clinical results but lacked the capability to discern radiolucent encapsulation of opaque graft materials. Therefore, in this study CT scans were able to measure bone augmentation quantitatively, but their ability to assess bone quality was limited. It appears that a combination of qualitative and quantitative assessment is necessary when using opaque grafting materials.

The time point of reentry for this study was only 14 weeks postoperative. This is an early time point compared with other clinical studies of particulate bone grafting. Nevins et al reentered extraction sites grafted with MCBS and rhPDGF-BB at 16 weeks and reported mature bone healing. Fiorellini et al demonstrated mature healing of extraction sites treated with recombinant human bone morphogenetic protein-2 after 16 weeks. The present study demonstrated clinically immature tissue in some sites upon reentry. The histologic results demonstrated ongoing bone formation and remodeling, and it is possible that a longer healing period would enhance implant site development.

The limitations of the present study were the small sample size for each group and a lack of control groups (grafting group without the rhPDGF-BB or grafting group with the use of membrane). However, several previously reported studies have already investigated these control groups and have obtained similar results. Future studies are needed with larger sample sizes to verify the scientific value of these promising preliminary results.

**Conclusion**

Within the limits of this study, the freeze-dried bone allograft and anorganic bovine bone graft carriers appeared to be appropriate scaffolds to deliver recombinant human platelet-derived growth factor BB for ridge augmentation using minimally invasive surgical techniques.

**Acknowledgment**

This study was supported by a grant from Osteohealth.
References


