Ridge Preservation and Augmentation Using Regenerative Materials to Enhance Implant Predictability and Esthetics

Abstract

Dentistry has entered an era in which patients no longer need to accept an edentulous or partially edentulous condition, or one in which their candidacy for tooth structure replacement (ie, implants with subsequent restoration) must be dismissed because of insufficient alveolar bone volume, height, or width. The supporting bone can be preserved at the time of tooth extraction, or augmented at the time of case presentation, using a variety of available regenerative materials. Among them are mineralized human allograft bone and collagen membranes that can be placed in combination with specific growth factor complexes and implant designs. This article reviews the challenges associated with adequately preserving or augmenting the alveolar bone after tooth extraction or loss and before implant placement. The research and benefits to support using allogenic bone graft and membrane materials for such procedures are detailed, and 3 clinical cases are presented to demonstrate the clinically successful incorporation of these materials with the host tissues.

When dental treatment involves restoration of an extracted tooth, one that is missing because of trauma, or one missing congenitally, preservation and/or augmentation of the alveolar ridge dimensions is of paramount importance. The bone volume in the maxillary and mandibular alveolar ridges in buccolingual and apicocoronal directions influences a variety of factors related to oral health and potential restorative treatment, including the location and position of implants when placed, their subsequent success or failure, and the esthetics of the definitive restorations. Long-term esthetic success requires vital alveolar bone to support healthy keratinized tissue and dictate the soft-tissue profile, which follows the underlying bone contour. Alveolar bone loss can occur after tooth extraction and/or trauma. Atrophy of the alveolar bone also may occur in areas of congenitally missing teeth, and adverse alveolar bone conditions

Learning Objectives

After reading this article, the reader should be able to:

• explain why preservation and augmentation of the alveolar bone at the time of tooth extraction is significant to subsequent implant placement and restoration processes.

• discuss some of the bone graft materials and techniques in use today to facilitate bone preservation and augmentation.

• describe the clinical and patient benefits of using allograft materials for bone preservation and augmentation before implant placement.

• explain the manner in which bone graft, barrier membranes, biologic modifiers, and other state-of-the-art technologies can be used in conjunction with one another to facilitate predictable treatment outcomes.

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also can result from advanced periodontal disease and
failed endodontic therapy.4 When the resorptive patterns
of the alveolar bone progress horizontally and vertically, a
more lingually positioned ridge can result.2 It has been
suggested that if the alveolar ridge is not preserved at the
time of tooth extraction or tooth loss, alveolar ridge
height and width—as well as position—may be lost, par-
ticularly in the area of the buccal (ie, facial) plate.2 Major
changes have been demonstrated to occur in extraction
sites during the first year after tooth extraction,1 and losses
of between 3 mm and 6 mm horizontally and 1 mm and
2 mm vertically have been reported.3,5,6,7,8

Preserving the alveolar ridges at the time of tooth
extraction helps to minimize difficulties during subse-
tquent implant placement.2 A site planned for implant
placement requires sufficient bone height and width to
ensure stability and ideal soft-tissue contours.3 The mini-
imum amount of bone recommended on the buccal or lin-
gual aspects is 1 mm, with at least 2 mm of facial bone
required for implants in the esthetic zone.3,10

The most predictable way to maintain the width,
height, and position of the alveolar ridges is preservation at
the time of tooth extraction.2,5,6 When there is insufficient
existing bone volume (eg, resulting from atrophy and con-
genitally missing dentition), the ridge can be augmented
using appropriate surgical techniques, including guided
bone regeneration with barrier membranes, bone grafts, and
biologic modifiers.11-13 The materials and surgical tech-
iques in use today facilitate bone preservation and augmen-
tation before implant placement and enable clinicians to more pre-
dictably ensure the functional and esthetic outcomes of the
implants and subsequent restorations that are delivered.1-3

Bone grafting can be completed using a variety of ma-
teials, such as autografts (ie, the patient’s own bone, which
requires multiple and potentially painful procedures), allo-
grafts (ie, human bone, not from the patient), and xenografts (ie, animal bone) or alloplasts (ie, synthetic
bone).3,5 Autografts are osteogenic and contain bone-form-
ing cells; allografts can be osteoconductive and osteoinduc-
tive, attracting cells that are bone-forming; and xenografts
and alloplasts are osteoconductive, acting as “scaffolds” for
other cells that are bone-forming.5 Allografts may be dem-
ineralized or mineralized freeze-dried bone allograft materi-
als. Mineralized bone allografts preserve the calcium and
phosphate salts and, therefore, resorb slower than deminer-
alized freeze-dried bone allografts.1,3

One human mineralized cancellous particulate bone
allograft (Puros®) is slowly resorbed and replaced with the
patient’s natural bone.13 This is because the material pre-
serves the natural collagen matrix, trabecular pattern, and
porosity of human cancellous bone. Histological analysis
following placement after tooth extraction demonstrates
the efficacy of this material.14,15

The most predictable way to maintain the width, height,
and position of the alveolar ridges is preservation at the
time of tooth extraction.

When properly prepared and delivered according to a
standardized surgical technique that has been evaluated in
the literature,16 a novel, sterilized, antigen-inactivated min-
eralized block allograft (Puros Block Allograft®) demon-
strates a high degree of predictability. Technique requisites
include proper contouring and shaping to fit the defect
site, screw fixation to prevent block allograft rotation, con-
touring of the block allograft to prevent soft-tissue compli-
cations, and coverage of the graft site with a barrier mem-
brane that may or may not have been saturated in platelet-
rich plasma.16 Additionally, soft-tissue closure must be
accomplished without tension.16

When placed in atrophic human maxillae, this miner-
alized bone allograft has been shown to form new bone,
and all subsequently placed Tapered Screw-Vent implants achieved osseointegration.4,15 What’s more, research also
determined that resorption and replacement by new bone
occurred more rapidly with the regenerative, mineralized
bone allograft that was tested.15

In one report of 4 cases, it was found that the allogenic
bone block that was used (Puros J-Block®) was an effective

a Zimmer Dental, Carlsbad, CA; www.zimmerdental.com
alternative to harvesting and grafting autogenous bone for implant site development—one that clinically generates new bone fill for implant placement. Using mineralized, cortico-cancellous bone allograft can eliminate the additional surgical procedure necessary for harvesting an autograft; however, it requires precise shaping and contouring to adapt the allograft to the defect site. This aspect of the technique sensitivity associated with the preparation and placement of block allografts may be mitigated using state-of-the-art imaging and modeling technology.

Specifically, data from computed tomography (CT) scans can be used to fabricate accurate 3-D stereolithic models of a patient's jaw that can be presurgically fitted with the planned allograft and sterilized for later transfer to the mouth for use as a template. Using 3-D CT imaging, the clinician can critique the allograft from different perspectives, without such intraoral challenges as bleeding, flaps, and restricted access. As a result, an additional advantage is the reduced risk of complications or failure from poorly contoured allografts. Further, the time required for the grafting procedure can be reduced by eliminating the need to prepare the block allograft at the time of the surgical appointment.

When implants ultimately are to be placed, clinicians can insert them with confidence that they will function successfully long-term in ridges that have been preserved and/or augmented with regenerative materials. One investigation showed that bone regenerated using a membrane technique reacts to implant placement like nonregenerated bone, achieving successful tissue integration with functional ankylosis. In a later study that analyzed 526 implants placed and loaded in regenerated bone, the researchers found that the regenerated bone reacted to implant placement in a clinically similar manner as native bone.

The barrier membranes mentioned earlier may be resorbable (eg, collagen), nonabsorbable (eg, expanded polytetrafluoroethylene), or acellular dermal matrices that serve as barriers and can be used when the buccal wall is deficient to protect the bone graft after placement.
Biologic modifiers include preparations rich in growth factors harvested from the patient’s own blood. They are used to form a fibrin matrix that can further protect the graft and induce cell proliferation.24,25 All of these materials or some combination thereof have demonstrated clinical efficacy for site preservation, augmentation, and implant integration at the affected site.4,15,17,26 The 3 cases described here highlight the use of specific regenerative materials and techniques to preserve the alveolar ridge after tooth extraction and/or augment the bone to accommodate anticipated implant and restorative requirements.

Case No. 1 (Dr. Keith): Single-Tooth Extraction Site Precipitated By Endodontic Failure

A 63-year-old woman, a nonsmoker, presented with a dental history of chronic decay and periodontal disease. Her medical history included allergy to penicillin, as well as use of calcium and bupropion at the time of presentation. During the examination, the endodontic failure of tooth No. 13 was revealed, necessitating its extraction. The extraction of tooth No. 13 resulted in a large apical defect at the site, in addition to significant destruction of the labial plate (Figure 1A). Mineralized particulate bone (Puros) was placed in the defect (Figure 1B) and covered using a regenerative collagen matrix membrane (Puros Pericardium) (Figure 1C). The particulate bone/membrane complex was sutured using Vicryl 4/0 sutures, leaving incomplete soft-tissue coverage of the membrane.3,23 However, 2 weeks after surgery, the absorbable membrane remained intact (Figure 1D), serving as a barrier during this critical period of wound healing. Three months after surgery, adequate areas of keratinized tissue had been achieved as a result of using the Pericardium membrane (Figure 1E). For this case, the patient ultimately received an implant and esthetic single-unit restoration and, 2 years after surgery, excellent soft-tissue contours (Figure 1F) and implant osseointegration had been realized, as seen in a 2-year postoperative radiograph (Figure 1G).

Case No. 2 (Dr. Keith): Bone Atrophy and Congenitally Missing Lateral Incisor

An 18-year-old woman presented with a chief complaint regarding the congenitally missing tooth No. 10 (Figure 2A). Her medical history was negative, she was a nonsmoker, and she had undergone prior orthodontic therapy. The patient was treatment planned to receive an implant-supported crown restoration after augmentation of the severely atrophied alveolar bone in the area of missing tooth No. 10 using an allograft bone block graft (Puros) and regenerative collagen membrane (BioMend Membrane).16 The allograft bone block was placed and after 5 months of healing, demonstra-
ed the desired cortical/trabecular patterns necessary for dimensional ridge augmentation (Figure 2B).

Alternatively, the case could have been treatment planned for ridge augmentation and implant placement using a 3-D software program, CT scan, and a stereolithic model, onto which the bone graft material would be prefit (See Box).17-19,26

The preoperative panoramic radiograph identified the root positions of teeth Nos. 9 and 11, as well as their mesiodistal spatial relationship. This mesiodistal space was compared with the contralateral maxillary right lateral incisor (Figure 2C) for reference. Once placed, the bone block allograft was covered by the BioMend membrane (Figure 2D), and its placement confirmed radiographically.

At 5 months, the block allograft was surgically exposed (Figure 2E) and the titanium screws were removed. A 3.3-mm Tapered Screw Vent implant was surgically placed in the No. 10 site (Figure 2F), and its placement/position confirmed radiographically (Figure 2G). A provisional crown restoration was placed at this visit. Upon completion of the definitive crown restoration 3 months later, the temporary was removed and the final implant-supported crown restoration was placed (Figure 2H).

Case No. 3 (Dr. Salama): Sudden Edentulous Segment/Fixed Partial Denture Fracture

A 52-year-old woman presented after her 4-unit fixed partial denture in the maxillary left quadrant fractured off, leaving teeth Nos. 11 and 12 (ie, abutments) fractured at the gumline and hopeless (Figure 3A). As a result, the patient was now edentulous in that area. The teeth were extracted atraumatically (Figure 3B) and the sockets planned for implant placement.

Using 3-D computer software (SimPlantb), implant placement was preplanned and a surgical guide fabricated (SurgiGuideb) based on the predetermined appropri-
ate location for implant insertion to accommodate future restorative needs, not on the location of current bone volume. This guide was prefabricated with die-cut holes to indicate the precise location for the drill/implants during surgery, as determined by the 3-D software.

Three implants (3.7-mm Tapered Screw Vent implants\textsuperscript{a}) were placed into the edentulous ridge and extraction sockets (ie, Nos. 11, 12, and 13 sites) (Figure 3C). Healing abutments (eg, 3 mm) were placed on the implants in the areas of Nos. 11 and 12, and a cover screw was placed in the area of No. 13.

However, it can be appreciated that the buccal plate in front of these implants was very thin. To prevent further resorption of this bone, the inside and outside of the socket were grafted with a 50:50 mixture of cancellous particulate bone allograft (Puros) and bovine bone (Bio-Oss) with PRGF.\textsuperscript{13,27-30} (Figure 3D). The cancellous bone allograft mixture was combined with a preparation rich in growth factor (PRGF).\textsuperscript{24,25} The bone graft complex was covered with fibrin, over which a regenerative collagen membrane (Puros Pericardium Allograft Membrane) was placed to protect the bone graft, as well as add thickness.
to the tissue and ridge (Figure 3E), and a tension-free closure was achieved. Note that the autologous fibrin demonstrates an elastic consistency that functions as an ideal biomaterial membrane for sealing postextraction wounds and inducing cell proliferation.24,25 Four months after surgery, the ridge had been successfully and significantly augmented both horizontally and vertically (Figure 3F).

At approximately 5 months after surgery, the surgical site was re-entered to expose the implants, at which time a significant increase in ridge dimensions was again observed. Specifically, the bone had been augmented buccally (ie, horizontally) up to between 4 mm and 5 mm, and also vertically above the level of the fixture heads on the implants at site Nos. 11 and 12 (Figure 3G). Additionally, remnants of the collagen membrane barrier were still in place and well incorporated with the surrounding natural bone and soft tissue to protect the bone graft (Figure 3H). This is significant because professionals who choose to use a collagen barrier require a material that is first accepted by the body, incorporated by the body, and not resorbed too quickly to protect the underlying graft.1,3,23

The extent of bone augmentation was verified radiographically (Figure 3I). To expose the implants to tissue, longer 5-mm healing abutments were placed (Figure 3J). At the time of writing, the patient was scheduled for routine impression taking and delivery of the final restoration in about 4 weeks.

Conclusion

Myriad regenerative products (eg, bone allografts, cancellous bone, bioactive modifiers, PRGF, fibrin membranes)12 in combination with careful treatment strategies that may, on occasion, involve 3-D imaging, CT scans, and surgical planning using stereolithic models and surgical guides, are helping to change the face of implant and restorative dentistry and contributing to more predictable long-term results. Given the propensity for the alveolar ridges to resorb after tooth extraction and/or as a result of atrophy, it behooves clinicians to understand the benefits and elements of socket and bone preservation/augmentation techniques and the materials that can be used for these purposes31 to enhance
the ultimate outcome of implant placement and subsequent restorative treatment.

Disclosure

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References

1. Bone volume in the alveolar ridges influences which of the following?
   a. esthetics of definitive restorations
   b. success or failure of implants
   c. location/position of implants
   d. all of the above

2. Alveolar bone loss can occur after:
   a. extraction only.
   b. trauma only.
   c. extraction or trauma.
   d. orthodontic treatment.

3. What may occur when the alveolar bone resorbs horizontally and vertically?
   a. A lingually positioned ridge may result.
   b. Bone volume may increase by 3.0 mm horizontally.
   c. Bone volume may increase by 2.0 mm vertically.
   d. No additional changes occur.

4. If the alveolar ridge is not preserved at the time of tooth extraction or tooth loss, alveolar ridge height and width may be lost, particularly:
   a. in the area of the buccal plate.
   b. along the maxillary ridge.
   c. along the mandibular ridge.
   d. along the midline.

5. In the first year after extraction, which of the following has been reported?
   a. bone loss between 1 mm and 2 mm horizontally
   b. bone loss between 3 mm and 6 mm horizontally
   c. bone loss between 2 mm and 3 mm vertically
   d. bone loss between 3 mm and 6 mm vertically

6. Which of the following is not required for implant placement?
   a. sufficient bone height
   b. sufficient bone width
   c. maximum bone on the buccal and lingual aspects of 1 mm
   d. at least 2 mm of facial bone in the esthetic area

7. Which of the following is the most predictable way to maintain width, height, and position of the alveolar ridge?
   a. ridge preservation at the time of tooth extraction
   b. bone regeneration with barrier membranes
   c. bone regeneration with bone grafts
   d. bone regeneration with biologic modifiers

8. Which category of bone graft materials contains human bone?
   a. autografts and allografts
   b. allografts and alloplasts
   c. alloplasts and autografts
   d. xenografts and alloplasts

9. Xenografts and alloplasts:
   a. are osteoinductive.
   b. are osteoconductive.
   c. can be demineralized freeze-dried bone.
   d. can be mineralized freeze-dried bone.

10. Because mineralized freeze-dried bone allografts contain calcium and phosphate salts they:
    a. resorb slower than demineralized freeze-dried allografts.
    b. resorb faster than demineralized freeze-dried allografts.
    c. resorb faster than xenografts.
    d. resorb faster than alloplasts.
11. Which of the following is not part of the requisite technique for placing a sterilized, mineralized block allograft?
   a. proper contouring and shaping of the block
   b. screw fixation to prevent rotation
   c. tight soft-tissue closure with visible tension
   d. coverage of the graft site with a barrier membrane

12. When placed in atrophic human maxillae, one mineralized bone allograft has been shown to:
   a. form new bone.
   b. achieve osseointegration of all placed tapered screw-vent implants.
   c. provide rapid resorption and replacement by new bone.
   d. all of the above

13. Cortico-cancellous bone allograft can eliminate which of the following?
   a. precise shaping and contouring to adapt the allografts to the defect site
   b. use of imaging and modeling technology
   c. surgery necessary for harvesting an autograft
   d. preparation and placement of block allografts

14. Using 3-D CT scanning eliminates which of the following?
   a. bleeding
   b. flaps
   c. restricted access
   d. all of the above

15. Using CT scans in the preparation of block allografts:
   a. adds time and stress to the procedure.
   b. limits the clinicians ability to critique the allograft before placement.
   c. reduces the risk of complications or failures from poorly contoured allografts.
   d. may result in increased failures.

16. One study showed that bone regenerated using a membrane technique reacts to implant placement like:
   a. nonregenerated bone.
   b. cortico-cancellous bone allograft.
   c. xenografts.
   d. allogenic bone.

17. Barrier membranes may be:
   a. resorbable.
   b. nonresorbable.
   c. acellular dermal matrices.
   d. all of the above

18. Which of the following is a resorbable barrier membrane?
   a. collagen
   b. expanded polytetrafluoroethylene
   c. PRGF
   d. cortico-cancellous bone

19. Biologic modifiers include preparations rich in growth factors harvested from?
   a. the patient's own blood
   b. animal bone
   c. human bone not from the patient
   d. synthetic materials

20. Biologic modifiers and barrier membranes or a combination thereof have demonstrated clinical efficacy for:
   a. site preparation.
   b. augmentation.
   c. implant integration.
   d. all of the above
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