Periodontal Repair in Dogs: Evaluation of a Bioabsorbable Space-Providing Macro-Porous Membrane with Recombinant Human Bone Morphogenetic Protein-2

Ulf M.E. Wikesjö,*† Won Hee Lim,† Robert C. Thomson,‡ Alonzo D. Cook,‡ John M. Wozney,§ and W. Ross Hardwick‡

Background: Recombinant human bone morphogenetic protein-2 (rhBMP-2) technologies have been shown to significantly support alveolar bone formation. Biomaterial limitations, however, have restricted the biologic potential for onlay indications. The objective of this study was to evaluate regeneration of alveolar bone and periodontal attachment, and biomaterials reaction following surgical implantation of a space-providing, bioabsorbable, macroporous, polyglycolic acid-trimethylene carbonate (PGA-TMC) membrane combined with a rhBMP-2 construct in a discriminating onlay defect model.

Methods: Routine supraalveolar periodontal defects were created at the mandibular premolar teeth in 9 beagle dogs. Contralateral jaw quadrants in subsequent animals were randomly assigned to receive the dome-shaped PGA-TMC (100 to 120 µm pores) membrane with rhBMP-2 (0.2 mg/mL) in a bioresorbable hyaluronan (Hy) carrier or the PGA-TMC membrane with Hy alone (control). The gingival flaps were advanced to submerge the membranes and teeth and sutured. Animals were euthanized at 8 and 24 weeks postsurgery for histologic observations.

Results: Jaw quadrants receiving the PGA-TMC membrane alone experienced exposures at various time points throughout the study. Jaw quadrants receiving the PGA-TMC/rhBMP-2 combination remained intact, although one site experienced a late minor exposure. Newly formed alveolar bone approached and became incorporated into the macroporous PGA-TMC membrane in sites receiving rhBMP-2. The PGA-TMC biomaterial was occasionally associated with a limited inflammatory reaction. Residual PGA-TMC could not be observed at 24 weeks postsurgery. Residual Hy could not be observed at any time interval. Regeneration of alveolar bone height (means ± SD) was significantly increased in sites receiving the PGA-TMC/rhBMP-2 combination compared to control (3.8 ± 1.3 versus 0.7 ± 0.5 mm at 8 weeks and 4.6 ± 0.8 versus 2.1 ± 0.4 mm at 24 weeks; P < 0.05). Limited cementum regeneration was observed for PGA-TMC/rhBMP-2 and PGA-TMC control sites. Ankylosis compromised regeneration in sites receiving PGA-TMC/rhBMP-2.

Conclusions: The bioabsorbable, space-providing, macroporous PGA-TMC membrane appears to be a compatible biomaterial for bone augmentation procedures. rhBMP-2 significantly enhances alveolar bone augmentation and soft tissue healing when combined with the PGA-TMC membrane. J Periodontol 2003; 74:635-647.

KEY WORDS
Alveolar ridge augmentation; animal studies; bone regeneration; membranes, barrier; membranes, bioabsorbable; periodontal attachment; wound healing.
Design criteria for guided tissue/bone regeneration (GTR/GBR) devices include biocompatibility, cell/tissue occlusion, space maintenance, tissue integration, and ease of use. Studies have shown that regeneration of alveolar bone is critically dependent on space provision by the tissue occlusive GTR or GBR devices (references 3 through 7 and unpublished data). Limited or no regeneration of alveolar bone is observed in supraalveolar periodontal defects and in alveolar ridge defects following gingival flap surgery without a space providing GTR/GBR device, when the device inadvertently had collapsed or had been compressed into the defect, or when the space underneath the membrane during surgery had been filled with slowly resorbing or non-resorbable biomaterial in an attempt to enhance space provision (references 3 through 7 and unpublished data).

Karaki et al. evaluated the influence of space provision without connective tissue occlusion on osteogenesis. Contralateral horizontal periodontal defects were surgically created between the mandibular premolar teeth in dogs. A tissue-expanding gold mesh was applied on one side while the contralateral side served as surgical control. Callus formation was enhanced in defects treated with the gold mesh compared to the surgical control. Evidently, osteogenesis in a periodontal environment may proceed in presence of space provision without strict provisions for gingival tissue occlusion. A concept of space provision without tissue occlusion for alveolar regeneration emerges from this study. This concept is supported by observations from a more recent study reporting the influence of biomaterials characteristics on bone healing in the craniofacial skeleton; i.e., the influence of expanded polytetrafluoroethylene (ePTFE) barrier porosity on osteogenesis. In a rat calvaria model, ePTFE barrier devices with a porosity of 20 to 25 and 100 µm increased the rate of osteogenesis compared to less porous devices (<8 µm). The authors proposed that increased permeability allowing transposition of extracellular matrix and cellular elements favored osteogenesis.

Most recently a non-resorbable, macroporous (300 µm pores, 0.8 mm apart) ePTFE membrane was used in a proof-of-principle study designed to test the concept of tissue occlusivity for GTR. It was shown that tissue occlusivity was not a critical requirement for GTR. Clinically relevant amounts of alveolar bone and cementum regeneration with functionally inserting fibers were demonstrated following installation of tissue occlusive and non-occlusive space providing macroporous ePTFE devices. It was also shown that the macroporous ePTFE device was considerably more clinically effective than the occlusive device. None of the sites receiving the macroporous device experienced wound failure in contrast to 50% of the sites receiving the occlusive ePTFE device. Wound failure, membrane exposure, contamination, and infection are not uncommon following use of tissue occlusive GTR devices in the clinic and commonly compromise clinical outcomes following GTR (references 3 through 5, 10 through 13, and unpublished data). Taken together, these observations suggest that macroporous devices may prove clinically beneficial in GTR.

In clinical practice, non-resorbable devices require surgical removal upon completion of regenerative treatment, which causes an added inconvenience to the patient and clinician. Macroporous devices for GTR must use bioabsorbable biomaterials because the porous nature of the device results in considerable tissue integration, which may make surgical removal of a non-resorbable device complicated and result in considerable morbidity. To circumvent the need for surgical removal, resorbable biomaterials for GTR have been suggested, evaluated, and manufactured for clinical use. Natural products like collagen-based devices; dura mater; cargile membrane; oxidized cellulose; laminar bone; and synthetic bioabsorbable devices based on polyactic acid, polyglycolic acid, and their copolymers have been developed and marketed for the treatment of patients. While bioabsorbable devices do not require a second surgery, they commonly present limitations including space provision, early/late resorption, and adverse inflammatory reactions in their resorption process resulting in fragmentation and associated foreign body reactions. Recently, Tatakis and Trombelli reported abscess formation including foamy macrophages in over 50% of a group of patients treated with GTR using a DL-polylactic acid based device. Thus, it appears critical that biomaterials developed for clinical use receive considerable scrutiny in relevant, discriminating preclinical models prior to clinical release.

Other studies have evaluated bone morphogenetic proteins (BMPs) as a candidate therapy for alveolar augmentation and periodontal regeneration. rhBMP-2 technologies have been shown to significantly increase bone regeneration in craniofacial applications (references 16 through 21 and unpublished data). However, limitations of rhBMP-2 technologies have been reported relative to biomaterials evaluated as carriers. In particular, carrier limitations have restricted the biologic potential of rhBMP-2 for indications where soft tissue compressive forces may limit the space for bone formation. The objective of this study was to evaluate a space-providing, macroporous, bioabsorbable polyglycolic acid-trimethylene carbonate membrane intended for GTR and to evaluate this bioabsorbable device in presence of rhBMP-2 induced bone in a discriminating preclinical onlay defect model.

MATERIALS AND METHODS

Animals
Nine male beagle dogs (age 18 to 24 months, weight approximately 15 kg) exhibiting intact mandibular pre-
molar dentition without crowding or evidence of periodontal disease, obtained from an approved dealer, were used. Animal selection and management, surgery protocol, and periodontal defect preparation followed routines approved by the Animal Care and Use Committee, W.L. Gore & Associates, Inc., Flagstaff, Arizona. The animals were identified by an implanted microchip. The animals had access to standard laboratory diet and water until the beginning of the study. Oral prophylaxis was performed within 2 weeks prior to the experimental surgeries.

**Macroporous GTR Device**

A bioabsorbable, space-providing, macroporous, polyglycolic acid-trimethylene carbonate membrane (PGA-TMC) was used (Fig. 1). The devices incorporated a proprietary fibrous non-woven construction designed to integrate with the host tissue during healing. In addition, 100 to 120 µm through-and-through holes were laser-etched in the devices to allow fibrovascular penetration into the space created by the device when implanted. Device configuration was custom-made to adapt specifically to the experimental model and to be sufficiently rigid to resist collapse from overlying tissue pressure. This macroporous PGA-TMC material has been shown to be biocompatible.26 Experimental studies have shown membrane configurations based on this biomaterial to support periodontal regeneration and alveolar bone augmentation.14

**rhBMP-2/Hyaluronan Construct**

rhBMP-2 (purified rhBMP-2 >98%) was formulated in storage-stable buffer at a concentration of 0.2 mg/ml surgical implant volume. A bioabsorbable non-woven hyaluronic acid ester (Hy; non-woven felt) was used for drug delivery in jaw quadrants receiving rhBMP-2 (Fig. 1). Four strips of the Hy biomaterial, each measuring 28 × 10 × 1.5 mm, were cut from a larger piece of sterile material. Using a sterile syringe, 2.0 ml of 0.2 mg/mL rhBMP-2 solution was withdrawn and uniformly dispensed over the surface of the 4 Hy strips. Following a minimum 10-minute binding period, the rhBMP-2 soak-loaded Hy strips were layered within the defect space, the interdental space, and in each furcation.

**Surgical Protocol**

Food was withheld the night before surgical procedures. The animals were premedicated with atropine (0.02 mg/kg IM), buprenorphine (0.04 mg/kg IM), and flunixin meglumine (0.1 mg/kg IV). A prophylactic antibiotic (cefazolin; 22 mg/kg IV) was administered. General anesthesia was induced with diazepam (0.2 mg/kg IV) and ketamine (6 mg/kg IV). An endotracheal tube was placed and the animals were maintained on isoflurane gas (1% to 2%) in 100% oxygen using positive pressure ventilation. A sterile catheter was placed and the animals received a slow constant rate infusion of lactated Ringer’s solution (10 to 20 ml/kg/hour IV) to maintain hydration while anesthetized. Routine dental infiltration anesthesia with epi-nephrine was administered at the surgical sites.

The maxillary first, second, and third premolar teeth were surgically extracted. The maxillary fourth premolars were reduced in height to alleviate potential trauma from the maxillary teeth to the experimental mandibular sites postsurgery and exposed pulpal tissues were sealed.**

Supraalveolar, critical size, periodontal defects were created around the third and fourth mandibular premolar teeth in right and left jaw quadrants in each animal (Fig. 1). Briefly, buccal and lingual mucoperiosteal flaps were reflected following buccal and lingual mucoperiosteal flaps were reflected following buccal and lingual sulcular incisions from the

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**Figure 1.**

Mandibular jaw quadrant with surgically induced, critical size, discriminating supraalveolar periodontal defect (A). The defect is implanted with rhBMP-2/Hy (B) and a macroporous PGA-TMC membrane is placed over the teeth and the rhBMP-2/Hy construct and secured with stainless steel tacks (C). The mucogingival flaps are then advanced to cover the teeth and implanted biomaterials, and sutured (D).
canine tooth to the second molar. The first and second premolar teeth were extracted, and the first molar teeth were amputated to the level of the surgically reduced alveolar crest. Alveolar bone was removed around the circumference of the remaining premolar teeth using chisels and water-cooled rotating burs. The root surfaces were instrumented with curets, chisels, and water-cooled rotating diamonds to remove the cementum. The crowns of the teeth were reduced to approximately 2 mm coronal to the cemento-enamel junction (CEJ) and the cut surfaces smoothed. Exposed pulpal tissues were sealed. Clinical defect height, from the CEJ to the reduced alveolar crest, was 6 mm as measured with a periodontal probe.

**Wound Management**

The precut rhBMP-2/Hy constructs were surgically implanted in one jaw quadrant in each animal. Contralateral jaw quadrants received Hy soak-loaded with buffer (2 ml) without rhBMP-2 (split-mouth design).

Experimental conditions were alternated between left and right jaw quadrants in subsequent animals. Dome-shaped PGA-TMC membranes were subsequently installed over the rhBMP-2/Hy and Hy control constructs (Fig. 1). The membranes were fixed to the reduced alveolar bone with medical grade stainless steel tacks‡‡ designed for these applications. Following placement of the membranes, periostea were fenestrated at the base of the flaps to allow tension-free flap apposition. The flaps were advanced, flap margins being adapted 3 to 4 mm coronal to the PGA-TMC membranes and sutured.†† Intrasurgery photographs were taken prior to and immediately after placement of the barrier membrane and following wound closure.

**Post-surgery Protocol**

Animals were fed a canned soft dog food diet the first 14 days postsurgery. Thereafter, the animals received a standard laboratory diet soaked in warm water until thoroughly soft. The animals received postoperative analgesia (buprenorphine; 0.04 mg/kg IM QID) for 2 days postsurgery. A broad-spectrum antibiotic (enrofloxacin; 2.5 mg/kg, IM, bid) was used for postsurgery infection control for 14 days. Plaque control was maintained by twice daily topical application of a 2% solution (enrofloxacin) until gingival suture removal, there-after, once daily (Monday through Friday) until completion of the study.

Radiographs were obtained immediately postsurgery, and at 4, 8, 12, 16, and 24 weeks postsurgery. Gingival sutures were removed under sedation at approximately 10 days postsurgery. Observations of experimental sites with regards to gingival health, maintenance of suture line closure, edema, and evidence of tissue necrosis or infection were made daily until suture removal, and at least twice weekly thereafter and recorded. The animals were scheduled for euthanasia at 8 and 24 weeks postsurgery. Following euthanasia, teeth with surrounding soft and hard tissues were removed en bloc. Membranes were not removed during the healing interval unless indicated as a consequence of wound failure.

**Histological Processing and Evaluation**

The tissue blocks were fixed in 10% buffered formalin for 3 to 5 days. Following fixation and complete decalcification with 20% EDTA, specimens were washed in running tap water and subsequently processed using an automatic tissue processor.§§ Standard methods were followed and specimens were infiltrated and embedded in methylmethacrylate. Specimens were allowed to polymerize for 3 to 5 days at room temperature. Five µm sections¶¶ were taken 100 µm apart. All sections were stained with a modified Goldner's trichrome stain.

The most central stained section of each root of the third and fourth premolar tooth was identified by the size of the root canal. This section and the adjacent stained step-serial sections were subjected to histometric analysis. Thus, 3 subsequent step-serial sections, encompassing approximately 0.2 mm of the mid-portion of the mesial and the distal root for each premolar tooth, were used for analysis. Analysis was performed using incandescent and polarized light microscopy, a microscope digital camera system, and a PC-based image analysis system by one calibrated investigator masked to the specific experimental conditions. The following measurements were recorded for the buccal and the lingual tooth surfaces for each section. Defect height: distance between apical extension of the root planing and the CEJ. Junctional epithelium: distance between gingival margin and apical termination of the junctional epithelium. Connective tissue repair: distance between apical termination of junctional epithelium and apical extension of the root planing. Cementum regeneration (height): distance between apical extension of the root planing and the coronal extension of a continuous layer of new cementum or cementum-like deposit on the planed root. Bone regeneration (height): distance between the apical extension of root planing and the coronal extension of regenerated alveolar bone along the planed root. Root resorption: combined linear heights of distinct resorption lacunae on the planed root. Ankylosis: combined linear heights of ankylosis union between the regenerated alveolar bone and the planed root.

†† FRIOS Augmentation System, Friatec, Mannheim, Germany.
‡‡ Xttrium Laboratories, Inc., Chicago, IL.
§§ Tissue-Tek, Sakura, Torrance, CA.
¶¶ Reichert Jung Polycut, Leica, Deerfield, IL.
## Media Cybernetic, Silver Spring, MD.
Data Analysis
Summary statistics (means ± SD) based on animal means for the experimental conditions were calculated using the selected step serial sections. Differences between experimental conditions were analyzed using paired t-tests.

RESULTS
Clinical Observations
One animal was euthanized 12 days postsurgery due to bilateral wound failure and subsequent infection of the implanted sites. A second animal was euthanized at 6 weeks due to a chronic fistula in the control site. Four animals were euthanized at 8 weeks, and 3 animals were euthanized at 24 weeks postsurgery. For the histologic and histometric analysis, data from the animals euthanized at 6 and 8 weeks postsurgery were pooled. Jaw quadrants receiving the PGA-TMC control experienced membrane exposure at various time-points. Healing was generally uneventful in sites receiving the PGA-TMC/rhBMP-2 combination (Fig. 2). All sites remained intact; however, one site experienced late exposure.

Radiographic Observations
Radiographic observations at 4 weeks postsurgery presented no appreciable bone formation in both PGA-TMC control and PGA-TMC/rhBMP-2 treated sites. Radiographic observations at 8 and 24 weeks postsurgery showed significant bone formation encompassing the top surface of teeth in sites receiving the PGA-TMC/rhBMP-2 combination (Fig. 3). The newly formed bone also filled the furcation areas. Two animals exhibited radiographic evidence of seromas in sites treated with PGA-TMC/rhBMP-2 at 8 weeks postsurgery that partially resolved at 24 weeks postsurgery. Jaw quadrants receiving the PGA-TMC control exhibited bone fill approximating 10% to 50% and 30% to 70% of the defect height at 8 and 24 weeks, respectively.

Histologic Observations
Limited alveolar bone regeneration was observed in all jaw quadrants receiving the PGA-TMC control at 8 weeks postsurgery (Fig. 4). In contrast, sites receiving the PGA-TMC/rhBMP-2 combination exhibited substantial bone regeneration encompassing approximately 70% of the defect height (Fig. 5). The newly formed bone was predominantly a mixture of lamellar and woven bone. Fatty marrow and fibrovascular tissue was also observed in sites receiving PGA-TMC/rhBMP-2.

The newly formed bone defined a periodontal ligament space. New cementum approximating 10% and 40% of the defect height was observed in the sites receiving the PGA-TMC control and PGA-TMC/rhBMP-2, respectively. The newly formed periodontal attach-
ment included a combination of parallel and functionally oriented fibers. Functionally oriented fibers were a predominant finding in sites receiving the PGA-TMC treatment without rhBMP-2.

Ankylosis was a common finding in the CEJ area in sites receiving PGA-TMC/rhBMP-2. Root resorption of surface erosion character was observed in almost all sites. Undermining root resorption was observed without predilection to the applied treatment. Seroma formation was not observed in the 8-week specimens.

There were no traces of the Hy biomaterial in any site at 8 weeks postsurgery. In contrast, the PGA-TMC membrane remained intact in all sites. The host tissue response to the PGA-TMC material appeared relatively benign and was composed largely of fibrovascular elements in direct contact with the fibers comprising the material structure. When present, inflammatory elements were composed of non-foamy macrophages or multinucleated cells in direct contact with the material. Inflammatory infiltrates were localized to the material surface and were not diffuse in the surrounding tissue. Bone formation frequently approximated and incorporated the membrane structure in sites receiving PGA-TMC/rhBMP-2. In one site, bone formation was also observed immediately outside the boundaries of the membrane.

Bone formation approximated 90% and 40% of the defect height in sites receiving PGA-TMC/rhBMP-2 and the PGA-TMC membrane without rhBMP-2, respectively, at 24 weeks postsurgery (Figs. 6 and 7). Newly formed bone represented a mixture of lamellar and woven bone. Fatty marrow was the predominant observation in sites receiving rhBMP-2 and fibrovascular tissue in sites receiving the PGA-TMC device without rhBMP-2.

Cementum regeneration approximated one-third of the defect height following both treatments. Jaw quadrants implanted with PGA-TMC/rhBMP-2 exhibited parallel-oriented fibers approximating the root surface (not shown). In contrast, sites implanted with PGA-TMC alone exhibited functionally oriented fibers inserting into the newly formed cementum (Fig. 6).

Ankylosis including the CEJ area and other aspects of the root surface was observed in sites receiving PGA-TMC/rhBMP-2 (Fig. 7). In contrast, sites without rhBMP-2 exhibited limited, if any, ankylosis. Root resorption of surface erosion character was a common finding following both protocols. Seroma formation was observed in one jaw quadrant receiving PGA-TMC/rhBMP-2 (not shown). There were no traces of residual PGA-TMC biomaterial in any site at the 24-week observation interval.

**Histometric Analysis**

The histometric analysis is presented in Tables 1 through 4. Regeneration of alveolar bone was significantly enhanced in sites receiving the PGA-TMC/rhBMP-2 combination compared to control (3.8 ± 1.3 versus 0.7 ± 0.5 mm at 8 weeks and 4.6 ± 0.8 versus 2.1 ± 0.4 mm at 24 weeks, respectively; *P* <0.05).
Cementum regeneration was significantly increased in sites receiving PGA-TMC/rhBMP-2 compared to those receiving the PGA-TMC control at 8 weeks (2.1 ± 1.0 versus 0.7 ± 0.4 mm, respectively; \(P<0.01\)). Similar amounts of cementum regeneration were observed for the PGA-TMC/rhBMP-2 and the PGA-TMC control treatment at 24 weeks (1.3 ± 0.5 versus 1.7 ± 1.0 mm, respectively; \(P>0.05\)). Functionally oriented fibers were rare observations in sites receiving PGA-TMC/rhBMP-2. In contrast, sites implanted with the macroporous PGA-TMC device characteristically showed functionally oriented fibers.

Ankylosis was significantly increased in sites receiving PGA-TMC/rhBMP-2 compared to sites receiving the PGA-TMC control. Root resorption of surface erosion character was observed in all animals.

**DISCUSSION**

The objective of this study was to evaluate regeneration of alveolar bone and periodontal attachment and to evaluate the biomaterial reaction following surgical implantation of a space-providing, bioabsorbable, macroporous PGA-TMC membrane combined with a rhBMP-2 construct in a discriminating onlay defect model. Contralateral critical size, suprabalveolar periodontal defects in 9 beagle dogs received the dome-shaped PGA-TMC membrane with or without rhBMP-2/Hy. Animals were euthanized at 8 and 24 weeks postsurgery for histologic and histometric analysis.

The PGA-TMC membrane appeared to be suitable biomaterial for bone augmentation procedures. At 8 weeks, the PGA-TMC membrane remained apparently intact and bone formation frequently approximated and incorporated into its micro- and macroporous structure in sites receiving rhBMP-2. The PGA-TMC membrane appeared completely resorbed at 24 weeks. Bone regeneration, including ankylosis, was significantly increased in sites receiving rhBMP-2 compared to controls. Similar and limited amounts of cementum regeneration were observed. There were notable differences in the nature of the periodontal attachment between rhBMP-2 and control sites. Functionally oriented fibers were commonly observed in control sites, however, they were a rare observation in rhBMP-2 sites.

There was a significant difference in coronal regrowth of alveolar bone in defect sites receiving PGA-TMC/rhBMP-2 compared to controls at both 8 (67% versus 13% of the defect height) and 24 (85% versus 38% of the defect height) weeks postsurgery. Similar observations were reported by Wikesjö et al.\(^3\) in this animal model using non-resorbable GTR membranes in conjunction with rhBMP-2. Regeneration of alveolar...
bone averaged 4.8 mm (100% of defect height) in sites receiving a macroporous ePTFE membrane combined with rhBMP-2/ACS compared to 2.0 mm (43% of defect height) using the macroporous ePTFE membrane and buffer/ACS following an 8-week healing interval. Alveolar bone regeneration following surgical implantation of rhBMP-2/ACS without provisions for GTR approximated 75% of the defect height following an 8-week interval in a previous study using this animal model. Bone regeneration in surgical controls without rhBMP-2 averaged 0% of the defect height. Collectively, these data suggest that rhBMP-2 has a unique potential to induce clinically relevant regeneration of alveolar bone.

Cementum regeneration averaged 37% and 13% of the defect height, respectively, in sites receiving PGA-TMC/rhBMP-2 and the PGA-TMC control at 8 weeks postsurgery. At 24 weeks, the corresponding values were 24% and 31%. Functionally oriented fibers, commonly observed in the PGA-TMC control, were rare observations in sites receiving rhBMP-2. Similar observations were made in our previous study evaluating the macroporous ePTFE membrane with or without rhBMP-2/ACS following an 8-week healing interval. Cementum regeneration approximated 33% and 22% of the defect height, respectively, in sites receiving the macroporous ePTFE membrane with or without rhBMP-2/ACS. Additional studies have reported cementum regeneration averaging less than 10% of the defect height in sites receiving rhBMP-2/ACS in this animal model also using an 8-week healing interval. Cementum regeneration in control sites averaged 0% of the defect height. Cementum regeneration has also been evaluated in the critical size supraalveolar periodontal defect model following guided tissue regeneration. Sigurdsson et al. reported cementum regeneration including functionally oriented fibers averaging 41% of the defect height in sites receiving an occlusive ePTFE membrane at 8 weeks postsurgery. Cementum regeneration in surgical controls averaged 1% of the defect height. Recently
cementum regeneration was evaluated following surgical installation of space-providing occlusive and macroporous ePTFE membranes using an 8-week healing interval. Cementum regeneration including functional oriented fibers averaged 94% and 50% of the defect height in sites receiving occlusive and macroporous membranes, respectively. All together these observations suggest that rhBMP-2 may not play a decisive role in the regeneration of the periodontal attachment.

Ankylosis appears to be a considerable short-coming of the rhBMP-2/Hy technology for periodontal indications. Ankylosis commonly compromises healing following application of other BMP technologies as well. Wikesjö et al. [8,30,31] observed that cellular cementum extending from the apical extension of the defect often merged with ankylosic bone in supraalveolar periodontal defects implanted with rhBMP-2/ACS. Other studies using rhBMP-2 [16,17,22,32-35] or rhOP-1/BMP-7 [36,37] in various carrier systems provide evidence of ankylosis in a variety of periodontal defects in rodent, canine, and non-human primate models. Ankylosis has not been a complication following surgical implantation of BMPs in the absence of extensive bone regeneration. [36,38,39] In context, root resorption of surface erosion character as observed herein is a common finding following reconstructive periodontal surgery and is thus not unique to BMP technologies. [40]

Various biomaterials have been evaluated as candidate carriers for BMPs. These include collagen, decalcified bone matrix, hyaluronic, hydroxypatite, calcium phosphates, a hydroxypatite-collagen composite, various poly (α-hydroxy acids), and titanium. [41] This study used a bioabsorbable non-woven hyaluronic acid ester non-woven felt to deliver rhBMP-2 to periodontal sites. In a previous study, Hunt et al., [21] using a large saddle-type alveolar ridge defect in the dog, showed that a Hy sponge supported significant bone induction by rhBMP-2 and that the Hy sponge alone possessed no apparent osteoconductive potential. Similar observations were made in the current study. The Hy non-woven felt was easy to prepare and exhibited favorable clinical characteristics upon soak

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Defect Height</th>
<th>Connective Tissue Repair</th>
<th>Junctional Epithelium</th>
<th>Cementum Height</th>
<th>Bone Height</th>
<th>Ankylosis</th>
<th>Root Resorption</th>
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<tr>
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<td>5.7 ± 0.1</td>
<td>0.0 ± 0.0</td>
<td>2.1 ± 1.0</td>
<td>3.8 ± 1.3</td>
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Figure 7.
Photomicrographs of supraalveolar defect receiving the PGA-TMC/rhBMP-2 combination at 24 weeks postsurgery. Overview (A; original magnification 2.5× Goldner’s trichrome) and higher magnification of the lingual (B and C; 8×) aspect of the defect. Continuous cementum regeneration may be observed extending from the apical aspect of the defect (B) through the coronal extension of new bone forming an ankylosic union with the root (C). There is limited, if any, appreciable evidence of formation of a periodontal ligament. Fatty marrow is observed in immediate contact with the newly formed cementum. There are no apparent residues of the Hy and PGA-TMC biomaterials.
Table 2.
Frequency of Teeth Exhibiting Junctional Epithelium, Functionally Oriented Fibers, Ankylosis, Root Resorption, or Seroma Formation Following 8-Week Observation Interval

<table>
<thead>
<tr>
<th>Animal</th>
<th>PGA-TMC/rhBMP-2</th>
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<tr>
<td></td>
<td>Junctional Epithelium</td>
<td>Functionally Oriented Fibers</td>
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<tr>
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<tr>
<td>%</td>
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<td>30</td>
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</table>

Table 3.
Comparisons between Jaw Quadrants Receiving the PGA-TMC/rhBMP-2 Combination Versus the PGA-TMC Membrane without rhBMP-2 Following 24-Week Observation Interval (N = 3; means ± SD in mm)

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Defect Height</th>
<th>Connective Tissue Repair</th>
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<th>Ankylosis</th>
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<td>PGA-TMC</td>
<td>5.5 ± 0.2</td>
<td>4.9 ± 0.6</td>
<td>0.3 ± 0.3</td>
<td>1.7 ± 1.0</td>
<td>2.1 ± 0.4</td>
<td>0.1 ± 0.1</td>
<td>0.7 ± 0.6</td>
</tr>
<tr>
<td>P value</td>
<td>0.5960</td>
<td>0.2278</td>
<td>0.1976</td>
<td>0.6377</td>
<td>0.0280</td>
<td>0.0840</td>
<td>0.2825</td>
</tr>
</tbody>
</table>

Table 4.
Frequency of Teeth Exhibiting Junctional Epithelium, Functionally Oriented Fibers, Ankylosis, Root Resorption, or Seroma Formation Following 24-Week Observation Interval

<table>
<thead>
<tr>
<th>Animal</th>
<th>PGA-TMC/rhBMP-2</th>
<th>PGA-TMC</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Junctional Epithelium</td>
<td>Functionally Oriented Fibers</td>
</tr>
<tr>
<td>6</td>
<td>0/2</td>
<td>0/2</td>
</tr>
<tr>
<td>7</td>
<td>0/2</td>
<td>0/2</td>
</tr>
<tr>
<td>8</td>
<td>0/2</td>
<td>0/2</td>
</tr>
<tr>
<td>Group</td>
<td>0/6</td>
<td>0/6</td>
</tr>
<tr>
<td>%</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>
loading. The rhBMP-2/Hy construct maintained its volume following application to the large supraalveolar periodontal defects. The clinical and radiographic examinations showed that rhBMP-2/Hy implanted sites restored tissue contour and readily filled with bone. The radiographic and histometric examination suggested that rhBMP-2/Hy implanted sites exhibited bone density comparable to that of the immediate resident bone without obvious residues of the Hy biomaterial. These properties all point to Hy biomaterials as candidate carriers for rhBMP-2.

When rhBMP-2/ACS constructs have been applied as onlays in alveolar defects, bone formation has been limited probably due to compression of the construct by forces from or transmitted through the gingival flaps. Previous studies have used occlusive ePTFE membranes in conjunction with implantation of rhBMP-2 constructs; however, occlusive membranes have been shown to impair the bone inducing capacity of rhBMP-2. Proof-of-principle studies have evaluated the efficacy of space-providing macroporous ePTFE membranes to support bone formation by rhBMP-2/ACS (references 30, 31, 42, and unpublished data). The macroporous ePTFE technology significantly increased bone formation by rhBMP-2/ACS when used as an onlay for alveolar augmentation. The bioabsorbable, space-providing, macroporous PGA-TMC membrane used in our study was developed to advance this novel treatment concept towards clinical application. The PGA-TMC macroporous biomaterial remained intact at 8 weeks postsurgery. Bone formation frequently approximated and incorporated into the membrane structure in sites receiving rhBMP-2. A limited inflammatory reaction was occasionally observed associated with the PGA-TMC biomaterial. These observations suggest a role for the bioabsorbable, space-providing, macroporous PGA-TMC membrane as a biocompatible device to support and delineate BMP induced bone formation.

In summary, the space-providing, macroporous PGA-TMC membrane is resorbed within 24 weeks and associated with only a minimal inflammatory reaction, bone may form up to and within the membrane; the PGA-TMC membrane provides for bone and cementum regeneration with a functionally oriented periodontal ligament (guided tissue regeneration); substantially increased bone formation occurs following implantation of rhBMP-2/Hy, geometry of bone formation being controlled by the PGA-TMC membrane; cementum regeneration with a functionally oriented periodontal ligament is rare in sites receiving rhBMP-2/Hy; ankylosis is common following rhBMP-2/Hy induced bone formation; and limited root resorption mainly of surface erosion character appears common following use of the PGA-TMC membrane with or without rhBMP-2/Hy.

CONCLUSIONS

1) The PGA-TMC membrane appears a biocompatible, space-providing device suitable for GTR and for defining rhBMP-2 induced bone formation and 2) rhBMP-2 significantly enhances alveolar bone augmentation and soft tissue healing when combined with the PGA-TMC membrane.

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REFERENCES


42. Wikesjö UME, Qahash M, Thomson RC, et al. Space providing ePTFE devices define rhBMP-2 induced alveolar augmentation at dental implants. *Clin Implant Dent Relat Res* 2002; accepted for publication.

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