Abstract: A method for enhancing the binding of growth factors and/or cells to an allograft structure by applying an effective quantity of a coating material to the surface of the allograft structure, producing a thin coated allograft structure, administering to the thin coated allograft structure a growth factor, cells or a combination thereof, and implanting the thin coated allograft structure into a host bone.
SURFACE TREATMENTS OF AN ALLOGRAFT TO IMPROVE BINDING OF GROWTH FACTORS AND CELLS

FIELD OF THE INVENTION

The present invention relates to methods for improving the binding of growth factors to an allograft by treating the surface of allograft with a thin coating, administering the growth factors and implanting into a host bone.

BACKGROUND OF THE INVENTION

Allografts are used in repair of bone structures damaged by disease, trauma and surgery. Inadequate amounts of available autografts and the limited size and shape of a person's own bone makes allografts to be commonly used in reconstructive surgery. Using allograft tissue eliminates the need for a second operative site to remove autograft bone or tendon, reduces the risk of infection, and safeguards against temporary pain and loss of function at or near the secondary site. Moreover, allograft bone is a reasonable graft substitute for autologous bone and is readily available from cadavers. Allograft bone is essentially a load-bearing matrix comprised of cross-linked collagen, hydroxyapatite, and osteoinductive Bone Morphogenetic Proteins (BMP). Human allograft tissue widely used in orthopaedic surgery is strong, integrates with the recipient host bone, and can be shaped either by the surgeon to fit the specific defect or shaped commercially by a manufacturing process.

Allograft bone is available in two basic forms: cancellous and cortical. Cortical bone is a highly dense structure comprised of triple helix strands of collagen fiber reinforced with hydroxyapatite. The hydroxyapatite component is responsible for the high compressive strength and stiffness of bone while the collagen fiber component contributes to its elastic nature, as well as torsional, shear, and tensile strength. Cortical bone is the main load-bearing component of long bones in the human body.

Many devices of varying shapes and forms can be manufactured from cortical allograft tissue. Surgical implants such as pins, rods, screws, anchors, plates, and intervertebral spacers have all been made and used successfully in human surgery.

Even though allograft has certain advantages over the other treatments, one of the main drawbacks of the allograft treatment is that the in-growth of the host bone into the
grafted bone may take longer than in an autograft. As a result, allograft treatment may be less effective than the autograft. Attempts have been made to overcome these drawbacks by modifying the allograft surface. For example, U.S. Patent No. 6,511,509 discloses a textured graft, wherein the texturing comprises a plurality of closely spaced continuous or discrete protrusions.

U.S. Patent No. 6,458,168 teaches a graft comprising a combination of two cortical bone portions and a cancellous bone portion located between the cortical bone portions. According to the disclosure, the portions of the composite graft are held together by means other than adhesive and are not demineralized.

U.S. Patent No. 6,899,107 discloses a graft coated with a biopolymer seeded with periosteal cells harvested from either the graft recipient or from an allogenic or a xenogenic source.

U.S. Patent Application Publication No. 20040228899 teaches the use of bone grafts, including allografts, characterized by tartrate-resistant acid phosphatase (TRAP) adsorbed to a porous hydroxyapatite substratum.


U.S. Patent No. 6,294,041 discloses chemical linkages of the surface of collagen to osteoimplants. U.S. Patent No. 6,752,834 discloses a multilayer membrane containing predominantly collagen II for reconstruction of bone or cartilage tissue.

Despite the advances recently made in the art, new methods promoting in-growth of the host bone into the allograft are needed to better utilize the advantages of allograft treatment.

SUMMARY OF THE INVENTION

The present invention provides a method of enhancing the binding of growth factors and cells to an allograft structure comprising: applying an effective quantity of a
coating material to at least a portion of the surface of the allograft structure; producing a thin coated allograft structure; administering to the thin coated allograft structure a growth factor, cells or a combination thereof; and implanting the thin coated allograft structure into a host bone.

The coating material modifies the allograft surface to bind growth factors, or cells, such as, for example, cultured cells, or a combination of growth factors and cells. A person of ordinary skill in the art will appreciate that the invention is not limited to growth factors only.

In another aspect of the invention provides for a method for enhancing in-growth of a host bone to an allograft structure comprising applying an effective amount of a coating material to at least a portion of the surface of the allograft structure; producing a thin coated allograft structure; administering to the thin coated allograft structure a growth factor, a cell culture or a combination thereof; binding of growth factors and cell cultures to the thin coated allograft structure; and implanting the thin coated allograft structure into a host bone.

DETAILED DESCRIPTION OF THE INVENTION

To aid in the understanding of the invention, the following non-limiting definitions are provided:

The term "allograft" refers to a graft of tissue obtained from a donor of the same species as, but with a different genetic make-up from, the recipient, as a tissue transplant between two humans. Allograft is generally referred to as an implant.

The term "autologous" refers to being derived or transferred from the same individual's body, such as for example an autologous bone marrow transplant.

The term "morbidity" refers to the frequency of the appearance of complications following a surgical procedure or other treatment.

The term "osteochondral" refers to the ability to stimulate the proliferation and differentiation of pluripotent mesenchymal stem cells (MSCs). In endochondral bone formation, stem cells differentiate into chondroblasts and chondrocytes, laying down a cartilaginous ECM, which subsequently calcifies and is remodeled into lamellar bone. In intramembranous bone formation, the stem cells differentiate directly into osteoblasts,
which form bone through direct mechanisms. Osteoinduction can be stimulated by osteogenic growth factors, although some ECM proteins can also drive progenitor cells toward the osteogenic phenotype.

The term "osteoconduction" refers to the ability to stimulate the attachment, migration, and distribution of vascular and osteogenic cells within the graft material. The physical characteristics that affect the graft's osteoconductive activity include porosity, pore size, and three-dimensional architecture. In addition, direct biochemical interactions between matrix proteins and cell surface receptors play a major role in the host's response to the graft material.

The term "osteogenic" refers to the ability of a graft material to produce bone independently. To have direct osteogenic activity, the graft must contain cellular components that directly induce bone formation. For example, a collagen matrix seeded with activated MSCs would have the potential to induce bone formation directly, without recruitment and activation of host MSC populations. Because many osteoconductive scaffolds also have the ability to bind and deliver bioactive growth factors, their osteoinductive potential will be greatly enhanced.

The term "patient" refers to a biological system to which a treatment can be administered. A biological system can include, for example, an individual cell, a set of cells (e.g., a cell culture), an organ, or a tissue. Additionally, the term "patient" can refer to animals, including, without limitation, humans.

The term "treating" or "treatment" of a disease refers to executing a protocol, which may include administering one or more drugs to a patient (human or otherwise), in an effort to alleviate signs or symptoms of the disease. Alleviation can occur prior to signs or symptoms of the disease appearing, as well as after their appearance. Thus, "treating" or "treatment" includes "preventing" or "prevention" of disease. In addition, "treating" or "treatment" does not require complete alleviation of signs or symptoms, does not require a cure, and specifically includes protocols which have only a marginal effect on the patient.

The term "xenograft" refers to tissue or organs from an individual of one species transplanted into or grafted onto an organism of another species, genus, or family.
The term "pit" refers to a defined space formed beneath the surface of the allograft and can be used interchangeably with the terms "depression", "cavity", "indentation", "hollow", or "hole".

The term "plug" refers to the material that covers the pit and is complementary to the shape of the pit. A "plug" may be porous or solid.

The term "coating" refers to a layer of material that is sprayed, dipped or otherwise applied on to the surface of the allograft. A "thin coating" refers to a coating layer having a thickness of at least one molecular layer to several micrometers.

Aspects of the present invention provide for a method for enhancing binding of growth factors to an allograft and a method for enhancing in-growth of host bone. Applicants have found that coating an allograft surface, administering a cell culture, a growth factor or a combination thereof and implanting the thin coated allograft structure resulted in increased binding of the growth factors to the allograft.

The coating could be applied to an allograft comprising a smooth or porous surface. Such porosity could be cancellous in structure or introduced by external means, such as adding pits or surface irregularities for the coating to mechanically lock on to.

Suitable coating materials in the formation of the thin coating include gelatin, collagen, hyaluronic acid, glycosaminoglycans, glycerol, calcium phosphate, calcium sulfate, glycerol, demineralized bone matrix, mineral components from bone, organic components, organic components from bone or other tissues, chitin, starch, cellulose, carboxy methyl cellulose, alginate, heparin, and synthetic and naturally degradable polymers. Coatings that are compatible with surface allografts are preferred. Further, coating materials that have been used in the processing of demineralized bone matrix products are preferred.

It may be desirable to administer to said implanted, coated allograft structure a growth factor, cells or a combination thereof all of which are capable of binding to said coated allograft structure. Suitable growth factors are, for example, BMP-2, rhBMP-2, BMP-4, rhBMP-4, BMP-6, rhBMP-6, BMP-7[OP-1], rhBMP-7, GDF-5, Statin, LIM mineralization protein, NeI-I protein, platelet derived growth factor (PDGF), transforming growth factor D (TGF-D), insulin-related growth factor-I (IGF-I), insulin-related growth
factor-II (IGF-II), fibroblast growth factor (FGF), beta-2-microglobulin (BDGF II), and rhGDF-5.

Cells may also be used instead of or in addition to growth factors, such as growth factors. Non-limiting examples of suitable cell types include mesenchymal stems cells, pluripotent stem cells, embryonic stem cells, osteoprogenitor cells, osteoblasts and osteoclasts. Growth factors may be added at time of surgery.

According to one embodiment of the invention, the allograft structure is selected from the group consisting of cortical bone, cancellous bone, subchondral bone and any combination of the various bone tissue types. According to another embodiment, the allograft structure comprises a composite bone which includes a bone powder, a polymer and a demineralized bone.

Depending upon the condition of the patient, new bone in-growth is accomplished by one or more mechanisms such as osteogenesis, osteoconduction and osteoinduction. It can be appreciated that the needs of a child are different from an aging patient afflicted with osteoporosis. Accordingly, there is no "one size fits all" approach towards optimizing the healing conditions in a patient.

In one embodiment, the surface of the allograft is coated in such a way as to adsorb and/or bind growth factors, other proteins and cells affecting osteogenesis, osteoconduction and osteoinduction.

In another embodiment, bone structures include but are not limited to cortical bone, cancellous bone, subchondral bone, or any combination of the various bone tissue types.

In another embodiment, the allograft surface coated in such a way that the original chemical forces naturally present are altered so that the implant attracts and binds proteins, such as, for example, growth factors and cells, including cells from cell cultures.

In another embodiment the adsorption occurs through the intramolecular or intermolecular attractions between atoms are formed through weak chemical forces, which include hydrogen bonds, van der Waals forces, ionic bonds and hydrophobic interactions. These weak forces create bonds that are constantly forming and breaking at physiological temperature and are readily reversible under physiological conditions. The transient bonds between metabolites and macromolecules, and hormones and receptors, and all the other
cellular moieties necessary for life are required for biomolecular interactions since rigid, static bonds will inhibit, if not paralyze, cellular activities.

In one embodiment, pits formed or occurring as indentations on the surface are beneficial for host bone and allograft contact, and since the growth agents are mechanically held in place creating enhanced binding. The pits can also retain growth agents and other bioactive agents.

In an embodiment the shape of the pits are selected from the group consisting of irregular, regular, wedge, cylinder, ellipse, curved linear, square, pyramidal and combinations thereof.

In one embodiment the growth factor and/or the cells may be administered to the coated allograft both before and after implanting the coated allograft into the host bone. An example of a suitable allograft structure is a cortical allograft structure such as a allograft structure in any size and shape. Another non-limiting example of the allograft structure is a bone composite. According to one embodiment of the invention, the bone composite comprises a bone powder, a polymer and a demineralized bone. In different embodiments of the invention, bone powder content ranged from about 5% to about 90% w/w, polymer content content ranged from about 5% to about 90% w/w, and demineralized bone particles content comprised the reminder of the composition. Preferably, the demineralized bone particles comprise from about 20% to about 40% w/w while the polymer and the bone powder comprise each from about 20% to about 60% w/w of the composition.

In embodiment, the allograft surface may be coated in a targeted manner to produce an appropriately charged allograft structure. The charged surface may be the entire allograft structure or to selected portions thereof. Generally, the growth factor or cell culture is applied within minutes, for example from about 1 to about 120 minutes before implantation into the patient.

Growth factors suitable for use in the practice of the invention include but are not limited to bone morphogenic proteins, for example, BMP-2, rhBMP-2, BMP-4, rhBMP-4, BMP-6, rhBMP-6, BMP-7(OP-1), rhBMP-7, GDF-5, and rhGDF-5, as disclosed, for example, in the U.S. Patent Nos. 4,877,864; 5,013,649; 5,661,007; 5,688,678; 6,177,406; 6,432,919; 6,534,268, and 6,858,431, and in Wozney, J. M., et al. (1988) Science,
Bone morphogenic proteins have been shown to be excellent at growing bone and there are several products being tested. Extensive animal testing has already been undertaken, and human trials are finished and in process for these products. rhBMP-2 delivered on an absorbable collagen sponge (INFUSE® Bone Graft, Medtronic Sofamor Danek, Memphis, TN) has been used inside titanium fusion cages and resulted in fusion in 11 out of 11 patients in a pilot study and 99% of over 250 patients in a pivotal study. In July, 2002 INFUSE® Bone Graft received FDA approval for use in certain types of spine fusion. A pilot study with BMP-2 delivered on a ceramic carrier was recently published and reported a 100% successful posterolateral fusion rate. BMP-7 (OP-I) has reported 50-70% successful posterolateral lumbar fusion results in human studies to date. On May 4, 2004, INFUSE® Bone Graft was approved for acute, open fractures of the tibial shaft (Bosse et al. NEJM 347(24): 1924-1931, 2002; Govender et al. JBJS 84(12): 2123-2134, 2002). Studies with these and other BMP’s are underway. However, it is important to note that use of BMP’s may add cost to an already very expensive operation.

Additionally, suitable growth factors include, without limitation, Statin, NeI-I protein, LIM mineralization protein, platelet derived growth factor (PDGF), transforming growth factor D (TGF-D), insulin-related growth factor-I (IGF-I), insulin-related growth factor-II (IGF-II), fibroblast growth factor (FGF), beta-2-microglobulin (BDGF II), as disclosed in the U.S. Patent No. 6,630,153.

In yet another embodiment, the allograft structure is treated with a coating that is positively-charged such that cells, in particular cell cultures having a negative surface charge bind to the positively-charged allograft structure. Examples of cells which are suitable for use in the practice of the invention include but are not limited to mesenchymal stem cells, pluripotent stem cells, embryonic stem cells, osteoprogentior cells and osteoblasts.

In one embodiment of the present invention the allograft mechanically locks with the host bone by means of surface imperfections and/or created pits in the surface. The pits have at least one plug inserted such that load bearing capacity is maintained. The mechanical locking of the pits and the coating promotes binding of the growth agents and enhanced in-growth of the allograft.
In one embodiment the pits have a size which are described in previously filed patent application serial number 11/158,924, filed on June 22, 2005, which is incorporated herein by reference.

EXAMPLES

EXAMPLE 1: BONE CONSTRUCT PREPARATION

Ovine cortical bone cylinders (4 mm diameter and 5 mm height) were machined from cadaver tibias and metatarsals. After machining, the Surface demineralized bone sample group(s) were first immersed in 0.6 N HCl (EMD Chemicals, Inc., Gibbstown, NJ) for 30 minutes with constant agitation and washed with water.

All bone constructs were washed in 0.5% (w/w) SDS (Bio-Rad Laboratories, Hercules, CA) / 0.5% (v/v) Triton X-100 (Sigma-Aldrich Co., St. Louis, MO) for 120 minutes under vacuum and constant agitation, and washed with water. These constructs were placed into Chex-all II Instant Sealing Sterilization Pouches (Propper Manufacturing Co., Inc., Long Island City, NY) and freeze dried (Freeze Dry System, Labconco Corporation, Kansas City, MO) for 48 hours. All bone cylinders were gamma irradiated (Nuteck Corporation, Hayward, CA) to simulate the terminal sterilization treatment of allograft bone commonly utilized at tissue banking facilities.

Oxygen vacuum plasma treatments were used to create negative charge on the surfaces of bone plugs. The treatment was carried out aseptically in a bell jar reactor. Bone plugs were placed upright on a stainless steel tray on the powered electrode. The treatment chamber was pumped down to a base pressure of 7 mTorr as measured with a Baratron sensor placed between the reactor and the vacuum pump. The pressure was controlled with a throttle valve placed between the pressure sensor and the pump. Oxygen gas was metered into the system with mass flow controllers at a flow rate of 10 seem to an operating pressure of 600 mTorr. The reactor was allowed to equilibrate for 10 minutes. 300 Watts of Radio Frequency (RF, 13.56 MHz) power delivered through an arrangement of one powered and two grounded planar electrodes was applied to the system for 2 minutes. Applied and reflected power was balanced using a matching network. With the RF power off, the chamber was then brought up to atmospheric pressure. The bone plugs were turned upside down so that the surface that had been in contact with the tray would
be exposed during a second plasma treatment. Finished samples were removed from the plasma reactor and placed in sterile packages.

**EXAMPLE 2: OVINE CORTICAL DEFECT MODEL**

Twenty (20) skeletally mature adult domestic sheep were assigned to one group corresponding to an implantation period of eight weeks post-operative. Animals were initially screened to exclude acute and chronic medical conditions, including Q-fever and Johne's disease, during a one-week quarantine period prior to surgery. Specific attention was paid to selecting animals of uniform size and weight to limit the variability of loading.

Phenylbutazone (1 g p.o.) and Cefazolin sodium were administered approximately 20 to 30 minutes prior to anesthesia induction. Induction of anesthesia was administered by intramuscular (IM) injection of examine (11 mg/kg) and xylazine (2 mg/kg). Following induction, anesthesia was maintained by endotracheal tube delivered isoflurane. The right hind leg was shaved and prepped with povidone-iodine solution, and draped in a sterile fashion.

A lateral approach to expose the right tibia and fused 3rd and 4th metatarsals was performed by blunt dissection. Four 4 mm diameter holes were drilled in each bone for a total of 8 implants per animal. The defect was irrigated with saline to remove bone particles or fragments prior to inserting the appropriate plug into the hole, flush with the host bone. A marking screw was inserted near the 2 defects at each end of the bone.

Placement verification for post-mortem analyses was made by measuring the distance between the defect and the screw, and noted on the animal's surgical sheet. The subcutaneous (SC) layer was closed with running suture, and the skin closed with staples.

All groups (Untreated allograft, surface demineralized bone, straight pits (1mm diameter x 1 mm deep), undercut pits (1 mm deep with a surface diameter of 1 mm and an interior diameter of 1.5 mm), and cortical bone with a negative surface charge) were all examined *in vivo* with and without a biologically active compound. All groups without a bioactive compound (n=13) were inserted directly into the defect without hydration of the bone plugs. The groups receiving rhBMP-2 (n=13) had 500 DI of 0.43 mg/mL rhBMP-2 dripped onto the construct, and occasionally rolled in the resulting pool of rhBMP-2.

After the 15-minute soak, the appropriate cortical allograft/rhBMP-2 construct was inserted into the defect. One group of straight pits (n=13) received 0.1 g of sheep DBM
mixed with 20 mL warm saline (37°C). The resulting paste was smeared across the surface of the straight pit construct prior to insertion into the defect.

Following the procedure, a Fentanyl patch was applied, and an additional dose of Cefazolin sodium was administered. Post-operative radiographs were obtained to obtain baseline densities within the defect and to verify placement.

The animals were not immobilized following surgery, and supplied chow and water *ad lib*. Animals were kept in recovery cages for several hours post-operatively after which they were transferred to standard cages so that motion was limited. After ten days, the animals were transferred to the off-site housing facility and allowed unrestricted motion in a naturalistic environment.

All animals were sacrificed eight weeks post-operatively using an intravenous barbiturate overdose. The overlying soft tissues will be sharply dissected from the defect site, the tibias and metatarsals examined for any gross deformities, and the operative section of the bones retrieved. Tibia and metatarsal bones from euthanized sheep were labeled and transported from necropsy to the Orthopaedic Bioengineering Lab (OBRL). The defects were identified by the intra-operative marking screw. Defects and surrounding bone were dissected using an Exakt Bone Saw (Exakt Technologies, Oklahoma City, OK). For defects undergoing biomechanical testing, 2 cm of host bone was retained; for histological specimens, 1 cm of bone surrounding the defect was retained.

**EXAMPLE 3: Biomechanical testing**

Biomechanical Specimens were tested on the day of euthanasia. Specimens were placed on a custom fixture allowing orientation of the defect (allograft plug) to be perpendicular to the direction of load application. The testing fixture contained a support plate that supported the host bone surrounding the allograft plug. The clearance of the hole in the support jig was 0.7 mm (diameter of support plate hole = 5.0 mm + 1.4 mm = 6.4 mm). A cylindrical pin with a flat loading surface (3.5 mm diameter) was used to push out the allograft plug. Using a servo-hydraulic testing system (MTS Bionix 858, Eden Prairie, MN), the pin applied a load to the allograft construct at a displacement rate of 2 mm/min with load and displacement data acquired at 100 Hz. Once the break load
was reached, the test was stopped. Peak load was identified as the highest load prior to a significant drop (maximum force).

After the allograft construct was pushed out, the empty allograft plug hole was bisected with the Exakt Saw. Cortical bone thickness at the hole was measured with digital calipers. The engineering analyses of the biomechanical data were:
1) Ultimate Force  
Maximum force

\[ \text{Shear Strength} = \frac{F}{\pi \cdot D \cdot H} \]

wherein

2) Shear Strength

\( F = \text{Ultimate force} \)
\( D = \text{Outer diameter of cylindrical implant (4 mm in all cases)} \)
\( H = \text{Average transcortical bone interface thickness} \)

\[ \text{Shear Modulus} = \frac{F(\ln R_2 - \ln R_i)}{2 \cdot U \cdot d \cdot H} \]

wherein

3) Shear Modulus

\( F = \text{Ultimate force} \)
\( R_2 = \text{Radius of defect} \)
\( R_i = \text{Radius of implant} \)
\( d = \text{Displacement at ultimate force} \)

\( H = \text{Transcortical interface length (as measured by digital calipers)} \)
Using the combined data from two studies, the effects of allograft treatment on biomechanical properties (ultimate load, shear strength, and shear modulus) were determined using a one-way ANOVA. Effects of allograft treatment on histomorphometric parameters were analyzed for significance comparing only two groups at a time using the Parametric Unpaired t-test (two-tailed, P-value with 95% Confidence Intervals).

Ultimate load at failing was significantly greater for the Straight pits/rhBMP-2 group compared to the Negative surface charge, Untreated allograft, Negative surface charge/rhBMP-2, and Undercut pit groups (p<0.05). However, the importance of this biomechanical parameter is questionable and potentially misleading, as ultimate load measures alone can not adequately describe the mechanical integrity of the bone-graft interface. Mechanical integration or resistance to push out is better represented by shear strength and shear modulus measures since the contact area between the plug and the host cortical bone is both considered in the calculation as well as used to normalize these values, thus allowing direct comparison to the values to be more indicative of the true effect of the treatment groups. The shear strength for the Surface demineralized/rhBMP-2 and Straight pits/rhBMP-2 were approximately 30% and 25% higher than that for the Undercut pits, Untreated allograft/rhBMP-2, Undercut pits/rhBMP-2, Negative surface charge, and Negative surface charge/rhBMP-2 (p<0.05) respectfully. Additionally, the shear strength was shown to be statistically greater for the Straight pits treatments compared to the Negative surface charge and Negative surface charge/rhBMP-2 treatments (p<0.05). A 50% improvement was seen in the shear modulus for the Surface demineralized/rhBMP-2 treatment group compared to Negative surface charge, Untreated allograft/rhBMP-2, Undercut pits/rhBMP-2, and Negative surface charge/rhBMP-2 (p<0.05). Many of the treatments were equivalent to Surface demineralized/rhBMP-2 including Straight pits/rhBMP-2, Straight pits, SDM, Untreated allograft, and Straight pits/DBM. However, Surface demineralized/rhBMP-2 consistently produced better interface mechanical properties than the other treatments.

The analysis also indicated trends showing that certain allograft treatments have adverse affects on the biomechanical properties. For the three biomechanical measured, the Undercut pit constructs had performance values below that of the Straight pit constructs. This is interesting because these treatments are similar except that the
Undercut pits treatment had been undercut at a depth to a diameter of 1.5 mm while the Straight pits had not. The biomechanical data implies the undercutting process has adverse effects. Further inspection also indicated a negative surface charge decreased the biomechanical performance of the allograft constructs. The shear strength and modulus data showed that the Negative surface charge and the Negative surface charge/rhBMP-2 treatments were statistically less than the top ranking constructs, as well as Untreated allograft.

**EXAMPLE 4: Histological analysis**

The trimmed samples were fixed in 70% ethyl alcohol (ETOH) for 1 week. The specimens were dehydrated in graded solutions of ETOH (70%, 95%, and 100%) over the course of approximately 3 weeks with increasing concentrations of Technovit 7000 (embedding resin). The final solution contained 100% of the embedding resin which was polymerized using light activation. An average of 10 sections (7 Dm thick) of each specimen was taken in the sagittal plane to include the implant and the adjacent bone. The sections were cut from the specimen block along the longitudinal axis of the defect using an Exakt diamond blade bone saw (Exakt Technologies, Oklahoma, OK). All sections were ground flat using an Exakt microgrinder to 10-20 µm thickness. Sections were made at equal intervals. The sections were stained with a modified Van Gieson bone stain. Histological images were acquired using an Image Pro Imaging system (Media Cybernetics, Silver Spring, MD) and a Nikon E800 microscope (AG Heinze, Lake Forest, CA), Spot digital camera (Diagnostic Instruments, Sterling, Heights, MI), and a pentium IBM-based IBM compatible computer with expanded memory capabilities (Dell Computer Corp., Round Rock, TX). Histomorphometric parameters measured included: Defect Area (mm²), Bone Area within Defect Area (mm²), Percent Bone Area within Defect (%), Graft Area within Defect Area (mm²), and Percent Graft Area within Defect (%). Qualitative assessment of bone morphology and cellularity were made including: lamellar vs. woven bone, cellularity, inflammatory cells, and bone integration with graft material.

Graft resorption was increased by the addition of rhBMP-2, with the effects of this growth factor more evident at the endosteal region. Bone formation was also improved by the addition of rhBMP-2.
With the exception of the negative surface charge and Undercut pits/rhBMP-2 groups, all treatments had better de novo bone formation in the defect than Untreated allograft: Straight pits/rhBMP-2, Straight pits/DBM (p<0.001) > Untreated allograft/rhBMP-2, Surface demineralized/rhBMP-2, Straight pits, Negative surface charge/rhBMP-2, and Xenograft (p<0.01) > Undercut pits (p<0.05). Untreated allograft/rhBMP-2 was significantly better than Negative surface charge (p<0.05). Surface demineralized/rhBMP-2 was better than Undercut pits/rhBMP-2 (p<0.05) and Negative surface charge (p<0.01). Straight pits was better than Undercut pits/rhBMP-2 (p<0.05) and Negative surface charge (p<0.05). Straight pits/rhBMP-2 is better than Undercut pits (p<0.01), Undercut pits/rhBMP-2 (p<0.01), Xenograft (p<0.01), and Negative surface charge (p<0.001). Straight pits/DBM had significance over Undercut pits (p<0.05), Undercut pits/rhBMP-2 (p<0.05), Xenograft (p<0.05), and Negative surface charge (p<0.001). Negative surface charge/rhBMP-2 was significant over Undercut pits (p<0.05), Undercut pits/rhBMP-2 (p<0.05), Xenograft (p<0.05), and Negative surface charge (p<0.01). Xenograft was better than Negative surface charge (p<0.01).

The remaining histomorphometric analysis is presented in the groups in which they were originally analyzed: Group I (Untreated allograft, Surface demineralized, Surface demineralized/rhBMP-2, Straight pits, Straight pits/rhBMP-2, Straight pits/DBM, and Xenograft) and Group II (Untreated allograft/rhBMP-2, Undercut pits, Undercut pits/rhBMP-2, Negative surface charge, and Negative surface charge/rhBMP-2. In group I, significantly more graft remained in the defect for the Untreated allograft and Xenograft groups (p<0.005). The percent ο de novo bone in the periosteal callus was significantly lower in the Untreated allograft group when compared to the other treatment groups (p<0.05). In group II, graft resorption within the defect was significantly improved for the Negative surface charge/rhBMP-2 group compared to the Negative surface charge group (p<0.05). There was greater graft resorption within the endosteal callus for all three treatments enhanced with rhBMP-2 compared to the two groups not exposed to the morphogen (p<0.05), except for Undercut pits which was not different than Undercut pits/rhBMP-2. All three rhBMP-2 treatment groups showed better de novo bone formation at the periosteal surface than the Negative surface charge treatment group (p<0.05). Negative surface charge/rhBMP-2 and Untreated allograft/rhBMP-2 had more de novo bone formation at the endosteal callus than the remaining treatment groups (p<0.05).
Histomorphometric results indicate that the addition of rhBMP-2 is responsible for an increase in bone *de novo* bone formation and a decrease on the amount of implanted allograft. In essence, treatments enhanced with growth factor stimulated new bone formation and also stimulated graft resorption. Osteoblastic stimulation was expected as a consequence of the addition of rhBMP-2 to the allograft plugs. However, osteoclasts, the cells responsible for bone and graft resorption, were also responsive to rhBMP-2.

Synopsis of histopathology is presented in Tables 1 and 2. An inflammatory reaction was only observed in the bovine xenograft sections. All treatments showed good incorporation of graft with host bone. Endosteal and periosteal graft incorporation was also observed for all treatments.

Allografts enhanced with rhBMP-2 showed better resorption than allograft alone, and rhBMP-2 seemed to aid osteoblasts activity. The results of the individual surface treatments were variable. Untreated allograft/rhBMP-2 showed the highest scores of bone remodeling, with the highest percentage of lamellar bone with allograft. Straight pits/DBM had the best cellular activity characterized by osteoclastic resorption of the graft, while straight pits/rhBMP-2 had the best osteoblastic new bone formation. Straight pits/rhBMP-2 showed the most consistent remodeling and allograft incorporation with large portions of the allograft plug remodeled. There was a more intense callus formation observed on the periosteal surface for undercut pits/rhBMP-2. Negative surface charge had the lowest scores for graft integration with host bone, and there was some fibrous tissue within the defect observed. Negative surface charge/rhBMP-2 showed the highest extension of graft resorption.
<table>
<thead>
<tr>
<th>Treatment</th>
<th>Present</th>
<th>Cells</th>
<th>Resorpt.</th>
<th>Surface</th>
<th>Oelasts</th>
<th>Oblast</th>
<th>Remodelli ng</th>
<th>Present in</th>
<th>Allo plug in</th>
<th>Allo plug desc.</th>
<th>Callus</th>
<th>Size</th>
<th>Size Removal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allograft</td>
<td>y</td>
<td>0.0</td>
<td>0.3</td>
<td>0.50</td>
<td>0.83</td>
<td>0.3</td>
<td>y,n</td>
<td>3</td>
<td>e,</td>
<td>o,p</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Allograft/ rhBM</td>
<td>y</td>
<td>0.0</td>
<td>2.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>n</td>
<td>0</td>
<td>e</td>
<td>p</td>
<td>e,</td>
<td>1.6</td>
<td>7</td>
</tr>
<tr>
<td>Straigt Pits</td>
<td>y</td>
<td>0.0</td>
<td>1.0</td>
<td>1.3</td>
<td>1.17</td>
<td>0.0</td>
<td>c,e</td>
<td>p</td>
<td>p</td>
<td>e</td>
<td>p,</td>
<td>0.0</td>
<td>1.0</td>
</tr>
<tr>
<td>Straigt Pits/ rhBM</td>
<td>y</td>
<td>0.0</td>
<td>2.0</td>
<td>3.0</td>
<td>2.0</td>
<td>0.0</td>
<td>c</td>
<td>p</td>
<td>p</td>
<td>b</td>
<td>p,</td>
<td>0.0</td>
<td>1.0</td>
</tr>
<tr>
<td>Straigt Pits/D BM</td>
<td>y</td>
<td>0.0</td>
<td>1.0</td>
<td>1.6</td>
<td>1.17</td>
<td>0.0</td>
<td>c,e</td>
<td>b,e</td>
<td>e</td>
<td>p</td>
<td>p,</td>
<td>0.0</td>
<td>1.0</td>
</tr>
<tr>
<td>Undercut Pits</td>
<td>y</td>
<td>0.0</td>
<td>0.0</td>
<td>0.9</td>
<td>1.5</td>
<td>0.0</td>
<td>0</td>
<td>e</td>
<td>e</td>
<td>p</td>
<td>p,</td>
<td>1.5</td>
<td>0</td>
</tr>
<tr>
<td>Undercut Pits/ rhBM</td>
<td>y</td>
<td>0.0</td>
<td>2.0</td>
<td>1.2</td>
<td>1.8</td>
<td>0.0</td>
<td>e</td>
<td>b</td>
<td>p</td>
<td>8</td>
<td>80</td>
<td>0</td>
<td>1.8</td>
</tr>
</tbody>
</table>
Histopathological analyses also revealed that the addition of rhBMP-2 stimulates *de novo* bone formation, bone remodeling, allograft incorporation and cellular activity.

More exuberant callus tissue was also observed for the treatments enhanced with the rhBMP-2. Additionally, the histology showed that callus formation was more substantial on the periosteal surface. This was probably due to the graft usually being inserted all the
way into the medullary cavity; thus, the periosteal surface of the graft was often leveled with host bone, while the endosteal surface was frequently protruding into the medullary cavity.

Table 2. Summary Of Histopathology Analysis For Each Treatment

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Untreated allograft</td>
<td>Minimal incorporation mostly originating from the graft-host interface. Minimal remodeling. Graft appears to be inert.</td>
</tr>
<tr>
<td>Untreated allograft/rhBMP-2</td>
<td>Allograft included in all sections. No inflammatory reaction present. Graft resorbed to some extent. Moderate cellular activity was accompanied by excellent new bone remodeling. Periosteal callus was more evident compared to endosteal callus. Woven and lamellar bone was similarly observed in the healing callus.</td>
</tr>
<tr>
<td>Surface demineralized</td>
<td>Variable response. Some plugs appear to have extensive remodeling and plug resorption; whereas, others have activity limited to the surface of the allograft. In general, active bone formation and remodeling was observed in all sections.</td>
</tr>
<tr>
<td>Surface demineralized/rhBMP-2</td>
<td>Variable response. Half of the sections show great resorption of the graft and the other half show minimal allograft resorption. All plugs show excellent integration and active remodeling.</td>
</tr>
<tr>
<td>Straight pits</td>
<td>Variable response. Some plugs appear to have extensive remodeling; whereas, others have activity limited to the surface of the pits. In general, active bone formation and remodeling occurred at the pits.</td>
</tr>
<tr>
<td>Straight pits/rhBMP-2</td>
<td>Great remodeling and plug incorporation. Osteoblast actively was impressive. Large portion of the plug remodeled.</td>
</tr>
<tr>
<td>Straight pits/DBM</td>
<td>Response was variable. Some specimens had great plug resorption and remodeling; whereas, others were well integrated, especially in the pits, but not good resorption. In general, good remodeling and activity.</td>
</tr>
<tr>
<td>Undercut pits</td>
<td>Allograft present in all specimens. Very little bone activity observed. Inflammatory cells were not detected. Endosteal and</td>
</tr>
<tr>
<td>Undercut pits/rhBMP-2</td>
<td>Periosteal callus present in 2 of 6 specimens. Minimal graft resorption.</td>
</tr>
<tr>
<td>----------------------</td>
<td>-------------------------------------------------------------------</td>
</tr>
<tr>
<td>Negative Surface Charge</td>
<td>Allograft was present in all specimens. No inflammatory reaction observed. Some graft resorption detected. Good osteoblastic and osteoclastic activity accompanied by moderate new bone remodeling. Periosteal callus more evident than endosteal callus formation, consisting, of mainly, lamellar bone.</td>
</tr>
<tr>
<td>Negative Surface Charge/rhBMP-2</td>
<td>Allograft present in all specimens. Minimal bone remodeling with presence of endosteal or periosteal callus in 2 of 6 specimens. Graft resorption was observed to some extent. Fibrous tissue was present in 2 of 6 specimens compromising host-graft integration. No inflammatory reaction detected</td>
</tr>
<tr>
<td>Bovine Xenograft</td>
<td>Inflammatory cells present. Most integration coming from the host bone. Remodeling evident, yet limited in scope.</td>
</tr>
</tbody>
</table>

Undercut pits/rhBMP-2 specimens were frequently graded as having excellent callus formation with intense cellular activity, and very good callus maturity. Negative surface charge/rhBMP-2 specimens demonstrated the best graft resorption, and also established adequate callus maturity. Undercut pits or negative surface charged specimens without rhBMP-2 had inferior bone healing, with minimal cellular activity, graft resorption and callus formation.

All publications cited in the specification, both patent publications and non-patent publications, are indicative of the level of skill of those skilled in the art to which this invention pertains. All these publications are herein fully incorporated by reference to the same extent as if each individual publication were specifically and individually indicated as being incorporated by reference.
Although the invention herein has been described with reference to particular embodiments, it is to be understood that these embodiments are merely illustrative of the principles and applications of the present invention. It is therefore to be understood that numerous modifications may be made to the illustrative embodiments and that other arrangements may be devised without departing from the spirit and scope of the present invention as defined by the following claims.
WE CLAIM:

1. A method for enhancing the binding of growth factors and cells to an allograft structure comprising:
   applying an effective amount of a coating material to at least a portion of the surface of the allograft structure;
   producing a thin coated allograft structure;
   administering to the thin coated allograft structure a growth factor, cells or a combination thereof; and
   implanting the thin coated allograft structure into a host bone.

2. The method of claim 1, wherein the allograft structure has surface irregularities which have been added to allow the coating material to physically adhere to the allograft surface.

3. The method of claim 2, wherein the allograft structure has surface irregularities which have been added to allow the coating material to mechanically lock to the allograft surface.

4. The method of claim 2, wherein the surface irregularities which have been added are pits.

5. The method of claim 3, wherein the pits have a shape selected from a member of the group consisting of irregular, regular, wedge, cylinder, ellipse, curved linear, square, pyramidal and combinations thereof.

6. The method of claim 1, wherein the allograft structure is selected from the group consisting of cortical bone, cancellous bone, composite bone, subchondral bone and any combination of the various bone tissue types.

7. The method of claim 1, wherein the growth factor, the cells or a combination thereof is administered to the allograft structure prior to the implantation of the thin coated allograft structure into the host bone.
8. The method of claim 1, wherein the growth factor is selected from the group consisting of BMP-2, rhBMP-2, BMP-4, rhBMP-4, BMP-6, rhBMP-6, BMP-7[OP-1], rhBMP-7, GDF-5, Statin, NeI-I protein, LIM mineralization protein, platelet derived growth factor (PDGF), transforming growth factor D (TGF-D), insulin-related growth factor-I (IGF-I), insulin-related growth factor-II (IGF-II), fibroblast growth factor (FGF), beta-2-microglobulin (BDGF II), rhGDF-5, and tartrate-resistant acid phosphatase.

9. The method of claim 1, wherein the cell culture is selected from the group consisting of mesenchymal stem cells, periosteal cells, pluripotent stem cells, embryonic stem cells, osteoprogenitor cells, osteoblasts, osteoclasts, bone marrow-derived cell lines, and any combination thereof.

10. The method of claim 1, wherein the growth factor is rhBMP-2.

11. The method of claim 1, wherein the allograft structure is an acortical allograft.

12. The method of Claim 1, wherein the thin coating has a thickness of at least one layer of molecules.

13. The method of claim 1, wherein the allograft structure comprises a composite bone.

14. The method of claim 7, wherein the composite bone comprises a bone powder, a polymer and demineralized bone particles.

15. A method for enhancing in-growth of a host bone to an allograft structure comprising:

   applying an effective amount of a coating material to at least a portion of the surface of the allograft structure;

   producing a thin coated allograft structure;

   administering to the thin coated allograft structure a growth factor, cells or a combination thereof;

   binding of growth factors and cell cultures to the thin coated allograft structure; and

   implanting the thin coated allograft structure into a host bone.