Title: METHODS FOR TREATING ISCHEMIC TISSUE

Abstract: Compositions and methods for treating ischemic tissue using cultured three-dimensional tissue are provided herein.
METHODS FOR TREATING ISCHEMIC TISSUE

CROSS-REFERENCE TO RELATED APPLICATIONS

[0001] This application claims benefit under 35 U.S.C. § 119(e) to application Serial No. 60/691,731, entitled "Methods for Promoting Repair and Regeneration of Ischemic Tissues," filed June 17, 2005, and to application Serial No. 60/692,054, entitled "Methods and Compositions for Treating Congestive Heart Failure," filed June 17, 2005, the disclosures of which are incorporated herein by reference in their entirety.

BACKGROUND

[0002] Tissue damage and defects can be caused by many conditions, including, but not limited to, disease, surgery, environmental exposure, injury, and aging. Tissue damage can also be caused by, and can result in, ischemia, which is typically caused by an imbalance between oxygen supply and demand in the damaged tissue. Usually, the imbalance between oxygen supply and demand is due to a reduction or blockage in blood flow to the damaged tissue. For example, insufficient blood flow to the heart due to the narrowing or blockage of one or more coronary arteries can result in ischemia. The resulting ischemia can be temporary, in that the symptoms associated with ischemia can be reversed: in other instances, ischemia can become chronic as a result of prolonged reduction or blockage of blood flow to the damaged tissue.

[0003] Currently used clinical methods for improving blood flow in diseased or otherwise damaged tissues, such as the heart, can involve invasive surgical techniques such as coronary by-pass surgery, angioplasty, and endarterectomy. Such procedures involve high degrees of inherent risk both during and after surgery, and often only provide a temporary remedy to the underlying physiological changes associated with ischemia. Consequently, there is a need for additional treatments, especially those that can ameliorate and/or reverse the damage to ischemic tissue.
4. SUMMARY

[0004] The present disclosure relates to methods for promoting the healing of ischemic tissues and organs. In particular, the methods relate to the injection, implantation and/or attachment of a cultured three-dimensional tissue to prevent and/or reduce tissue thinning that is characteristic of the tissue remodeling observed in ischemic tissue, as well as promote endothelialization, tissue growth, vascularization and/or angiogenesis in ischemic tissues and organs.

[0005] In some embodiments, the methods described herein can be used to improve the performance of a heart clinically manifesting symptoms associated with the presence of ischemic tissue. For example, in some embodiments, the compositions and methods can be used to strengthen weakened heart muscle such that there is a demonstrable increase in pumping efficiency. Additionally, the compositions and methods described herein can be combined with conventional treatments, such as the administration of various pharmaceutical agents and surgical procedures, to treat individuals diagnosed with coronary disease, including coronary artery disease.

5. BRIEF DESCRIPTION OF THE FIGURES

[0006] FIG. 1 depicts histological evidence of new microvessel formation in canine dog hearts contacted with Anginera™ according to some of the embodiments described herein.

[0007] FIGS. 2A and 2B depict EDVI parameters during the 30 day ameroid period according to some of the embodiments described herein.

[0008] FIG. 3 depicts the cardiac output in the four treatment groups 30 days after placement of Anginera™ according to some of the embodiments described herein.

[0009] FIG. 4 depicts the cardiac output in the four treatment groups 90 days after placement of Anginera™ according to some of the embodiments described herein.

[0010] FIG. 5 depicts the left ventricular ejection fraction in the four treatment groups 30 days after placement of Anginera™.
FIG. 6 depicts the left ventricular ejection fraction in the four treatment groups 90 days after placement of Anginera™ according to some of the embodiments described herein.

FIG. 7 depicts the left ventricular end diastolic volume in the four treatment groups 30 days after placement of Anginera.

FIG. 8 depicts the left ventricular end diastolic volume in the four treatment groups 90 days after placement of Anginera™ according to some of the embodiments described herein.

FIG. 9 depicts the left ventricular systolic volume in the four treatment groups 30 days after placement of Anginera™.

FIG. 10 depicts the left ventricular systolic volume in the four treatment groups 90 days after placement of Anginera™ according to some of the embodiments described herein.

FIG. 11 depicts systolic wall thickening in the four treatment groups 30 days after placement of Anginera™ according to some of the embodiments described herein.

FIG. 12 depicts systolic wall thickening in the four treatment groups 90 days after placement of Anginera™ according to some of the embodiments described herein.

6. DETAILED DESCRIPTION

Disclosed herein are methods of treating ischemic tissue, comprising contacting a region of ischemic tissue with an amount of a cultured three-dimensional tissue effective to treat at least one clinical symptom or sign associated with the ischemic tissue. The cultured three-dimensional tissue comprises a variety of growth factors and/or Wnt proteins, both within and secreted by the cells of three-dimensional tissue that promote one or more biological processes that contribute to effective treatment, including but not limited to, prevention and/or reduction in tissue thinning, as is characteristic of the tissue remodeling observed in ischemic tissue, and/or promotion of endothelialization, tissue growth, vascularization and/or angiogenesis.

Biological properties that can be expressed by the three-dimensional tissue and/or secreted growth factors and/or Wnt proteins include, but are not limited to, prevention and/or
reduction of tissue thinning characteristic of the tissue remodeling observed in ischemic tissue, promotion of endothelialization, tissue growth, vascularization and/or angiogenesis.

[0020] The three-dimensional tissue can be used to treat ischemia in any tissue and/or organ. For example, the three-dimensional tissue can be used to treat patients presenting symptoms associated with heart disease, including but not limited to, coronary artery disease, silent ischemia, stable angina, unstable angina, acute myocardial infarction, and left ventricular dysfunction. Application of the three-dimensional tissue to an ischemic region in the heart of a patient diagnosed with heart disease promotes the healing of the ischemic tissue resulting in an overall improvement in the cardiac output of the treated heart.

Three Dimensional Tissue and Scaffolds

[0021] In various embodiments, the three-dimensional tissue capable of promoting healing of ischemic tissue can be obtained from various types of cells as discussed in more detail below. The three-dimensional tissue can be obtained commercially or generated de novo using the procedures described in U.S. Patent 6,372,494; 6,291,240; 6121,042; 6,022,743; 5,962,325; 5,858,721; 5,830,708; 5,785,964; 5,624,840; 5,512,475; 5,510,254; 5,478,739; 5,443,950; and 5,266,480; the disclosures of which are incorporated herein by reference in their entirety.

[0022] In some embodiments, the cultured three-dimensional tissue is obtained commercially from Smith & Nephew, London, United Kingdom. In particular, the product referred to as Dermagraft™, also referred to herein as Anginera™, can be obtained from Smith & Nephew.

[0023] Generally, the cultured cells are supported by a scaffold, also referred to herein as a scaffold, composed of a biocompatible, non-living material. The scaffold can be of any material and/or shape that: (a) allows cells to attach to it (or can be modified to allow cells to attach to it); and (b) allows cells to grow in more than one layer (i.e., form a three dimensional tissue).

[0024] In some embodiments, the biocompatible material is formed into a three-dimensional scaffold comprising interstitial spaces for attachment and growth of cells into a three dimensional tissue. The openings and/or interstitial spaces of the scaffold are of an appropriate size to allow the cells to stretch across the openings or spaces. Maintaining actively growing cells that are stretched across the scaffold appears to enhance production of the repertoire of growth factors responsible for the activities described herein. If the
openings are too small, the cells may rapidly achieve confluence but be unable to easily exit from the mesh. These trapped cells may exhibit contact inhibition and cease production of the growth factors described herein. If the openings are too large, the cells may be unable to stretch across the opening; which may decrease production of the growth factors described herein. When using a mesh type of scaffold, as exemplified herein, it has been found that openings at least about 140 μm, at least about 150 μm, at least about 160 μm, at least about 175 μm, at least about 185 μm, at least about 200 μm, at least about 210 μm, and at least about 220 μm work satisfactorily. However, depending upon the three-dimensional structure and intricacy of the scaffold, other sizes can work equally well. In fact, any shape or structure that allows the cells to stretch, replicate, and grow for a suitable length of time to elaborate the growth factors described herein can be used.

[0025] In some embodiments, the three dimensional scaffold can be formed from polymers or threads braided, woven, knitted or otherwise arranged to form a scaffold, such as a mesh or fabric. The materials can be formed by casting the material or fabrication into a foam, matrix, or sponge-like scaffold. In other embodiments, the three dimensional scaffold can be in the form of matted fibers made by pressing polymers or other fibers together to generate a material with interstitial spaces. The three dimensional scaffold can take any form or geometry for the growth of cells in culture as long as the resulting tissue expresses one or more of the tissue healing activities described herein. Descriptions of cell cultures using a three dimensional scaffold are described in U.S. Patent 6,372,494; 6,291,240; 6121,042; 6,022,743; 5,962,325; 5,858,721; 5,830,708; 5,785,964; 5,624,840; 5,512,475; 5,510,254; 5,478,739; 5,443,950; and 5,266,480; all publications incorporated herein by reference in their entireties.

[0026] A number of different materials can be used to form the scaffold. These materials can be non-polymeric and/or polymeric materials. Polymers when used can be any type of block polymers, co-block polymers (e.g., di, tri, etc.), linear or branched polymers, crosslinked or non-crosslinked. Non-limiting examples of materials for use as scaffolds or frameworks include, among others, glass fiber, polyethylene, polypropylene, polyamides (e.g., nylon), polyesters (e.g., Dacron), polystyrenes, polycrylates, polyvinyl compounds (e.g., polyvinylchloride; PVC), polycarbonates, polytetrafluoroethylenes (PTFE; TEFLOW), expanded PTFE (ePTFE), thermanox (TPX), nitrocellulose, polysaacharides (e.g., cellulos,
chitosan, agarose), polypeptides (e.g., silk, gelatin, collagen), polyglycolic acid (PGA), and dextran.

[0027] In some embodiments, the scaffold can be comprised of materials that degrade over time under the conditions of use, such as degradable materials. As used herein, a degradable material refers to a material that degrades or decomposes. In some embodiments, the degradable material is biodegradable, i.e., degrades through action of biological agents, either directly or indirectly. Non-limiting examples of degradable materials include, among others, poly(lactic-co-glycolic acid) (i.e., PLGA), trimethylene carbonate (TMC), co-polymers of TMC, PGA, and/or PLA, polyethylene terephthalate (PET), polycaprolactone, catgut suture material, collagen (e.g., equine collagen foam), polylactic acid (PLA), fibronectin matrix, or hyaluronic acid.

[0028] In embodiments in which the cultures are to be maintained for long periods of time, cryopreserved, and/or where additional structural integrity is desired, the three dimensional scaffold can comprise nondegradable materials. As used herein, a nondegradable material refers to a material that does not degrade or decompose significantly under the conditions in the culture medium. Exemplary nondegradable materials, include, but are not limited to, nylon, dacron, polystyrene, polyacrylates, polyvinyls, teflons, and cellulose. An exemplary nondegrading three dimensional scaffold comprises a nylon mesh, available under the tradename Nitex®, a nylon filtration mesh having an average pore size of 140 µm and an average nylon fiber diameter of 90 µm (#3-210/36, Tetko, Inc., N.Y.).

[0029] In other embodiments, the three dimensional scaffold can be a combination of degradable and non-degradable materials. The non-degradable material provides stability to the scaffold during culturing, while the degradable material allows interstitial spaces to form sufficient for formation of three-dimensional tissues that produce factors sufficient for promoting the healing of ischemic tissue. The degradable material can be coated onto the non-degradable material or woven, braided or formed into a mesh. Various combinations of degradable and non-degradable materials can be used. An exemplary combination is poly(ethylene terephthalate) (PET) fabrics coated with a thin degradable polymer film (poly[D-L-lactic-co-glycolic acid] PLGA).

[0030] In various embodiments, the scaffold material can be pre-treated prior to inoculation with cells to enhance cell attachment to the scaffold. For example, prior to inoculation with
cells, nylon screens can be treated with 0.1 M acetic acid, and incubated in polylysine, fetal bovine serum, and/or collagen to coat the nylon. In some embodiments, polystyrene can be analogously treated using sulfuric acid. In other embodiments, the growth of cells in the presence of the three-dimensional scaffold is further enhanced by adding to the scaffold, or coating with proteins (e.g., collagens, elastin fibers, reticular fibers), glycoproteins, glycosaminoglycans (e.g., heparan sulfate, chondroitin-4-sulfate, chondroitin-6-sulfate, dermatan sulfate, keratan sulfate, etc.), fibronectins, a cellular matrix, and/or other materials glycopolymer (poly[N-p-vinylbenzyl-D-lactoamide], PVLA) in order to improve cell attachment. Treatment of the scaffold or scaffold is useful to improve attachment of cells.

[0031] In other embodiments, the scaffold comprises particles so dimensioned such that cells cultured in presence of the particles elaborate the factors that promote healing of ischemic tissue. In some embodiments, the particles comprise microparticles, or other suitable particles, such as microcapsules and nanoparticles, which can be degradable or non-degradable (see, e.g., "Microencapsulates: Methods and Industrial Applications," in Drugs and Pharmaceutical Sciences, 1996, VoI 73, Benita, S. ed, Marcel Dekker Inc., New York). Generally, the microparticles have a particle size range of at least about 1 µm, at least about 10 µm, at least about 25 µm, at least about 50 µm, at least about 100 µm, at least about 200 µm, at least about 300 µm, at least about 400 µm, at least about 500 µm, at least about 600 µm, at least about 700 µm, at least about 800 µm, at least about 900 µm, at least about 1000 µm. Nanoparticles have a particle size range of at least about 10 nm, at least about 25 nm, at least about 50 nm, at least about 100 nm, at least about 200 nm, at least about 300 nm, at least about 400 nm, at least about 500 nm, at least about 600 nm, at least about 700 nm, at least about 800 nm, at least about 900 nm, at least about 1000 nm. The microparticles can be porous or nonporous. Various microparticle formulations can be used for preparing the three dimensional scaffold, including microparticles made from degradable or non-degradable materials used to form the mesh or woven polymers described above.

[0032] Exemplary non-degradable microparticles include, but are not limited to, polysulfones, poly (acrylonitrile-co-vinyl chloride), ethylene-vinyl acetate, hydroxyethylmethacrylate-methyl-methacrylate copolymers. Degradable microparticles include those made from fibrin, casein, serum albumin, collagen, gelatin, lecithin, chitosan, alginate or poly-amino acids such as poly-lysine. Degradable synthetic polymers polymers such as polylactide (PLA), polyglycolide (PGA), poly (lactide-co-glycolide) (PLGA), poly
(caprolactone), polydioxanone trimethylene carbonate, polyhydroxyalkonates (e.g., poly (y-hydroxybutyrate)), poly (Y-ethyl glutamate), poly (DTH iminocarbony (bisphenol A iminocarbonate), poly (ortho ester), and polycyanoacrylate.

[0033] Hydrogels can also be used to provide three-dimensional scaffolds. Generally, hydrogels are crosslinked, hydrophilic polymer networks. Non-limiting examples of polymers useful in hydrogel compositions include, among others, those formed from polymers of poly (lactide- co-glycolide), poly (N-isopropylacrylamide) ; poly (methacrylic acid-γ-polyethylene glycol); polyacrylic acid and poly (oxypropylene-co-oxyethylene) glycol; and natural compounds such as chondroitan sulfate, chitosan, gelatin, fibrinogen, or mixtures of synthetic and natural polymers, for example chitosan-poly (ethylene oxide). The polymers can be crosslinked reversibly or irreversibly to form gels sufficient for cells to attach and form a three dimensional tissue.

[0034] Various methods for making microparticles are well known in the art, including solvent removal process (see, e.g., US Patent No. 4,389,330); emulsification and evaporation (Maysinger et al., 1996, Exp. Neuro. 141: 47-56; Jeffrey et al., 1993, Pharm. Res. 10: 362-68), spray drying, and extrusion methods. Exemplary microparticles for preparing three dimensional scaffolds are described in US Publication 2003/021 1083 and US Patent Nos. 5,271,961; 5,413,797; 5,650,173; 5,654,008; 5,656,297; 5,114,855; 6,425,918; and 6,482,231, and the U.S. application entitled "Cultured Three Dimensional Tissues and Uses Thereof," filed concurrently herewith, the disclosures of which are incorporated herein by reference in their entireties.

[0035] It is to be understood that other materials in various geometric forms, other than those described above, can be used to generate a three dimensional tissue with the tissue healing characteristics described herein, and thus, the materials are not limited to the specific embodiments disclosed herein.

Cells and Culture Conditions

[0036] In some embodiments, the cultured three dimensional tissues can be made by inoculating the biocompatible materials comprising the three-dimensional scaffold with the appropriate cells and growing the cells under suitable conditions to promote production of a cultured three-dimensional tissue with one or more tissue healing properties. Cells can be
obtained directly from a donor, from cell cultures made from a donor, or from established cell culture lines. In some instances, cells can be obtained in quantity from any appropriate cadaver organ or fetal sources. In some embodiments, cells of the same species preferably matched at one or more MHC loci, are obtained by biopsy, either from the subject or a close relative, which are then grown to confluence in culture using standard conditions and used as needed. The characterization of the donor cells are made in reference to the subject being treated with the three-dimensional tissue.

[0037] In some embodiments, the cells are autologous, i.e., the cells are derived from the recipient. Because the three-dimensional tissue is derived from the recipient's own cells, the possibility of an immunological reaction that neutralizes the activity of the three-dimensional tissue is reduced. In these embodiments, cells are typically cultured to obtain a sufficient number to produce the three-dimensional tissue.

[0038] In other embodiments, the cells are obtained from a donor who is not the intended recipient of the culture medium. In some of these embodiments, the cells are syngeneic, derived from a donor who is genetically identical at all MHC loci. In other embodiments, the cells are allogeneic, derived from a donor differing at least one MHC locus from the intended recipient. When the cells are allogeneic, the cells can be from a single donor or comprise a mixture of cells from different donors who themselves are allogeneic to each other. In further embodiments, the cells comprise xenogenic, i.e., the are derived from a species that is different from the intended recipient.

[0039] In various embodiments herein, the cells inoculated onto the scaffold can be stromal cells comprising fibroblasts, with or without other cells, as further described below. In some embodiments, the cells are stromal cells, that are typically derived from connective tissue, including, but not limited to: (1) bone; (2) loose connective tissue, including collagen and elastin; (3) the fibrous connective tissue that forms ligaments and tendons, (4) cartilage; (5) the extracellular matrix of blood; (6) adipose tissue, which comprises adipocytes; and, (7) fibroblasts.

[0040] Stromal cells can be derived from various tissues or organs, such as skin, heart, blood vessels, skeletal muscle, liver, pancreas, brain, foreskin, which can be obtained by biopsy (where appropriate) or upon autopsy.
[0041] The fibroblasts can be from a fetal, neonatal, adult origin, or a combination thereof. In some embodiments, the stromal cells comprise fetal fibroblasts, which can support the growth of a variety of different cells and/or tissues. As used herein a fetal fibroblast refers to fibroblasts derived from fetal sources. As used herein neonatal fibroblast refers to fibroblasts derived from newborn sources. Under appropriate conditions, fibroblasts can give rise to other cells, such as bone cells, fat cells, and smooth muscle cells and other cells of mesodermal origin. In some embodiments, the fibroblasts comprise dermal fibroblasts. As used herein, dermal fibroblasts refers to fibroblasts derived from skin. Normal human dermal fibroblasts can be isolated from neonatal foreskin. These cells are typically cryopreserved at the end of the primary culture.

[0042] In other embodiments, the three-dimensional tissue can be made using stem and/or progenitor cells, either alone, or in combination with any of the cell types discussed herein. The term "stem cell" includes, but is not limited to, embryonic stem cells, hematopoietic stem cells, neuronal stem cells, and mesenchymal stem cells.

[0043] In some embodiments, a "specific" three-dimensional tissue can be prepared by inoculating the three-dimensional scaffold with cells derived from a particular organ, i.e., skin, heart, and/or from a particular individual who is later to receive the cells and/or tissues grown in culture in accordance with the methods described herein.

[0044] As discussed above, additional cells can be present in the culture with the stromal cells. Additional cell types include, but are not limited to, smooth muscle cells, cardiac muscle cells, endothelial cells and/or skeletal muscle cells. In some embodiments, fibroblasts, along with one or more other cell types, can be can be inoculated onto the three-dimensional scaffold. Examples of other cell types include, but are not limited to, such as cells found in loose connective tissue, endothelial cells, pericytes, macrophages, monocytes, adipocytes, skeletal muscle cells, smooth muscle cells, and cardiac muscle cells. These other cell types can readily be derived from appropriate tissues or organs such as skin, heart, and blood vessels, using methods known in the art such as those discussed above. In other embodiments, one or more other cell types, excluding fibroblasts, are inoculated onto the three-dimensional scaffold. In still other embodiments, the three-dimensional scaffolds are inoculated only with fibroblast cells.
[0045] Cells useful in the methods and compositions described herein can be readily isolated by disaggregating an appropriate organ or tissue. This can be readily accomplished using techniques known to those skilled in the art. For example, the tissue or organ can be disaggregated mechanically and/or treated with digestive enzymes and/or chelating agents that weaken the connections between neighboring cells making it possible to disperse the tissue into a suspension of individual cells without appreciable cell breakage. Enzymatic dissociation can be accomplished by mincing the tissue and treating the minced tissue with any of a number of digestive enzymes either alone or in combination. These include, but are not limited to, trypsin, chymotrypsin, collagenase, elastase, and/or hyaluronidase, DNase, pronase, and dispase. Mechanical disruption can be accomplished by a number of methods including, but not limited to, the use of grinders, blenders, sieves, homogenizers, pressure cells, or insonators to name but a few. For a review of tissue disaggregation techniques, see Freshney, *Culture of Animal Cells. A Manual of Basic Technique*, 2d Ed., A.R. Liss, Inc., New York, 1987, Ch. 9, pp. 107-126.

[0046] Once the tissue has been reduced to a suspension of individual cells, the suspension can be fractionated into subpopulations from which the fibroblasts and/or other stromal cells and/or other cell types can be obtained. This can be accomplished using standard techniques for cell separation including, but not limited to, cloning and selection of specific cell types, selective destruction of unwanted cells (negative selection), separation based upon differential cell agglutinability in the mixed population, freeze-thaw procedures, differential adherence properties of the cells in the mixed population, filtration, conventional and zonal centrifugation, centrifugal elutriation (counter-streaming centrifugation), unit gravity separation, countercurrent distribution, electrophoresis and fluorescence-activated cell sorting. For a review of clonal selection and cell separation techniques, see Freshney, *Culture of Animal Cells. A Manual of Basic Techniques*, 2d Ed., A.R. Liss, Inc., New York, 1987, Ch. 11 and 12, pp. 137-168.

[0047] Cells suitable for use in the methods and compositions described herein can be isolated, for example, as follows: fresh tissue samples are thoroughly washed and minced in Hanks balanced salt solution (HBSS) in order to remove serum. The minced tissue is incubated from 1-12 hours in a freshly prepared solution of a dissolving enzyme such as trypsin. After such incubation, the dissociated cells are suspended, pelleted by centrifugation and plated onto culture dishes. As stromal cells attach before other cells, appropriate stromal
cells can be selectively isolated and grown. The isolated stromal cells can be grown to confluence, lifted from the confluent culture and inoculated onto the three-dimensional scaffold (United States Patent No. 4,963,489; Naughton et al. 1987, J. Med. 18(3&4):219-250). Inoculation of the three-dimensional scaffold with a high concentration of cells, e.g., approximately $1 \times 10^6$ to $5 \times 10^7$ stromal cells/ml, can result in the establishment of a three-dimensional tissue in shorter periods of time.

[0048] In other embodiments, an engineered three-dimensional tissue prepared on a three-dimensional scaffold includes tissue-specific cells and produces naturally secreted growth factors and Wnt proteins that stimulate proliferation or differentiation of stem or progenitor cells into specific cell types or tissues. Moreover, the engineered three-dimensional tissue can be engineered to include stem and/or progenitor cells. Examples of stem and/or progenitor cells that can be stimulated by and/or included within the engineered three-dimensional tissue, include, but are not limited to, stromal cells, parenchymal cells, mesenchymal stem cells, liver reserve cells, neural stem cells, pancreatic stem cells and/or embryonic stem cells.

[0049] After inoculation of the scaffold with desired cell type(s), the scaffold can be incubated in an appropriate nutrient medium that supports the growth of the cells into a three-dimensional tissue. Many commercially available media, such as Dulbecco's Modified Eagles Medium (DMEM), RPMI 1640, Fisher's, and Iscove's, McCoy's, are suitable for use. The medium can be supplemented with additional salts, carbon sources, amino acids, serum and serum components, vitamins, minerals, reducing agents, buffering agents, lipids, nucleosides, antibiotics, attachment factors, and growth factors. Formulations for different types of culture media are described in reference works available to the skilled artisan (e.g., Methods for Preparation of Media, Supplements and Substrates for Serum Free Animal Cell Cultures, Alan R. Liss, New York (1984); Tissue Culture: Laboratory Procedures, John Wiley & Sons, Chichester, England (1996); Culture of Animal Cells, A Manual of Basic Techniques, 4th Ed., Wiley-Liss (2000)). Typically, the three-dimensional tissue is suspended in the medium during the incubation period in order to enhance tissue healing activity(ies), secretion of growth factors and/or Wnt proteins. In some embodiments, the culture can be "fed" periodically to remove spent media, depopulate released cells, and add fresh medium. During the incubation period, the cultured cells grow linearly along and envelop the filaments of the three-dimensional scaffold before beginning to grow into the openings of the scaffold.
Different proportions of various types of collagen deposited on the scaffold can affect the growth of the cells that come in contact with the three-dimensional tissue. The proportions of extracellular matrix (ECM) proteins deposited can be manipulated or enhanced by selecting fibroblasts which elaborate the appropriate collagen type. This can be accomplished using monoclonal antibodies of an appropriate isotype or subclass that is capable of activating complement, and which define particular collagen types. These antibodies and complement can be used to negatively select the fibroblasts which express the desired collagen type. Alternatively, the cells used to inoculate the framework can be a mixture of cells that synthesize the desired collagen types. The distribution and origin of different collagen types is shown in Table I.

Table I: Distribution and Origin of Different Collagen Types

<table>
<thead>
<tr>
<th>Collagen Type</th>
<th>Principle Tissue Distribution</th>
<th>Cells of Origin</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Loose and dense ordinary connective tissue; collagen fibers</td>
<td>Fibroblasts and reticular cells; smooth muscle cells</td>
</tr>
<tr>
<td></td>
<td>Fibrocartilage</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Bone</td>
<td>Osteoblast</td>
</tr>
<tr>
<td></td>
<td>Dentin</td>
<td>Odontoblasts</td>
</tr>
<tr>
<td>II</td>
<td>Hyaline and elastic cartilage</td>
<td>Chondrocytes</td>
</tr>
<tr>
<td></td>
<td>Vitreous body of the eye</td>
<td>Retinal cells</td>
</tr>
<tr>
<td>III</td>
<td>Loose connective tissue; reticular fibers</td>
<td>Fibroblasts and reticular cells</td>
</tr>
<tr>
<td></td>
<td>Papillary layer of dermis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Blood vessels</td>
<td>Smooth muscle cells; endothelial cells</td>
</tr>
<tr>
<td>IV</td>
<td>Basement membranes</td>
<td>Epithelial and endothelial cells</td>
</tr>
<tr>
<td></td>
<td>Lens capsule of the eye</td>
<td>Lens fiber</td>
</tr>
</tbody>
</table>
In various embodiments, the culture three-dimensional tissue has a characteristic repertoire of cellular products produced by the cells, such as growth factors. In some embodiments, the cultured three-dimensional tissues are characterized by the expression and/or secretion of the growth factors shown in Table II.

Table II: Three Dimensional Tissue Expressed Growth Factors

<table>
<thead>
<tr>
<th>Growth Factor</th>
<th>Expressed by Q-RT-PCR</th>
<th>Secreted Amount Determined by ELISA</th>
</tr>
</thead>
<tbody>
<tr>
<td>VEGF</td>
<td>8 x 10⁶ copies/ug RNA</td>
<td>700 pg/10⁶ cells/day</td>
</tr>
</tbody>
</table>
[0054] In some embodiments, the cultured three-dimensional tissue can be characterized by
the expression and/or secretion of connective tissue growth factor (CTGF). CTGF is a well-
known fibroblast mitogen and angiogenic factor that plays an important role in bone
formation, wound healing, and angiogenesis. See, e.g., Luo, Q., et al., 2004, *J. Biol. Chem.*, 15

<table>
<thead>
<tr>
<th>Gene</th>
<th>Expression/Secretion</th>
</tr>
</thead>
<tbody>
<tr>
<td>PDGF A chain</td>
<td>$6 \times 10^5$ copies/ug RNA</td>
</tr>
<tr>
<td>PDGF B chain</td>
<td>0</td>
</tr>
<tr>
<td>IGF-1</td>
<td>$5 \times 10^5$ copies/ugRNA</td>
</tr>
<tr>
<td>EGF</td>
<td>$3 \times 10^3$ copies/ug RNA</td>
</tr>
<tr>
<td>HBEGF</td>
<td>$2 \times 10^4$ copies/ug RNA</td>
</tr>
<tr>
<td>KGF</td>
<td>$7 \times 10^4$ copies/ug RNA</td>
</tr>
<tr>
<td>TGF-β1</td>
<td>$6 \times 10^6$ copies/ug RNA</td>
</tr>
<tr>
<td>TGF-β3</td>
<td>$1 \times 10^4$ copies/ug RNA</td>
</tr>
<tr>
<td>HGF</td>
<td>$2 \times 10^4$ copies/ug RNA</td>
</tr>
<tr>
<td>IL-1α</td>
<td>$1 \times 10^4$ copies/ug RNA</td>
</tr>
<tr>
<td>IL-1β</td>
<td>0</td>
</tr>
<tr>
<td>TNF-a</td>
<td>$1 \times 10^7$ copies/ug RNA</td>
</tr>
<tr>
<td>TNF-b</td>
<td>0</td>
</tr>
<tr>
<td>IL-6</td>
<td>$7 \times 10^6$ copies/ug RNA</td>
</tr>
<tr>
<td>IL-8</td>
<td>$1 \times 10^7$ copies/ug RNA</td>
</tr>
<tr>
<td>IL-12</td>
<td>0</td>
</tr>
<tr>
<td>IL-15</td>
<td>0</td>
</tr>
<tr>
<td>NGF</td>
<td>0</td>
</tr>
<tr>
<td>G-CSF</td>
<td>$1 \times 10^8$ copies/ug RNA</td>
</tr>
<tr>
<td>Angiopoietin</td>
<td>$1 \times 10^4$ copies/ug RNA</td>
</tr>
</tbody>
</table>

[0055] In addition to the above list of growth factors, the three dimensional tissue can also be characterized by the expression of Wnt proteins. "Wnt" or "Wnt protein" as used herein refers to a protein with one or more of the following functional activities: (1) binding to Wnt receptors, also referred to a Frizzled proteins, (2) modulating phosphorylation of Dishevelled protein and cellular localization of Axin (3) modulation of cellular β-catenin levels and corresponding signaling pathway, (4) modulation of TCF/LEF transcription factors, and (5) increasing intracellular calcium and activation of Ca^{2+} sensitive proteins (e.g., calmodulin dependent kinase). "Modulation" as used in the context of Wnt proteins refers to an increase or decrease in cellular levels, changes in intracellular distribution, and/or changes in functional (e.g., enzymatic) activity of the molecule modulated by Wnt.

[0056] Of relevance to the present disclosure are Wnt proteins expressed in mammals, such as rodents, felines, canines, ungulates, and primates. For instance, human Wnt proteins that have been identified share 27% to 83% amino-acid sequence identity. Additional structural characteristics of Wnt protein are a conserved pattern of about 23 or 24 cysteine residues, a hydrophobic signal sequence, and a conserved asparagine linked oligosaccharide modification sequence. Some Wnt proteins are also lipid modified, such as with a palmitoyl group (Wilkert et al., 2003, Nature 423(6938):448-52). Exemplary Wnt proteins and its corresponding genes expressed in mammals include, among others, Wnt1, Wnt2, Wnt2B, Wnt3, Wnt3A, Wnt4, Wnt4B, Wnt5A, Wnt5B, Wnt6, Wnt7A, Wnt7B, Wnt8A, Wnt8B, Wnt9A, Wnt9B, Wnt10A, Wnt11, and Wnt16. Other identified forms of Wnt, such as Wnt2, Wnt3, Wnt4, and Wnt15, appear to fall within the proteins described for Wnt 1-11 and 16. Protein and amino acid sequences of each of the mammalian Wnt proteins are available in databases such as SwissPro and Genbank (NCBI). See, also, U.S. Publication No. 2004/0248803 and U.S. application entitled, "Compositions and Methods Comprising Wnt Proteins to Promote Repair of Damaged Tissue," filed concurrently herewith, and U.S. application entitled, "Compositions and Methods for Promoting Hair Growth; the disclosures of which are incorporated herein by reference in their entireties.
[0057] Various techniques for the isolation and identification of Wnt proteins are known in the art. See, e.g., U.S. Publication No. 2004/0248803, the disclosure of which is incorporated herein by reference in its entirety.

[0058] In some embodiments, the Wnt proteins comprise at least Wnt5a, Wnt7a, and Wntll. As used herein, Wnt5a refers to a Wnt protein with the functional activities described above and sequence similarity to human Wnt protein with the amino acid sequence in NCBI Accession Nos. AAH74783 (gI:50959709) or AAA16842 (gI:348918) (see also, Danielson et al., 1995, J. Biol. Chem. 270(52):31225-34). Wnt7a refers to a Wnt protein with the functional properties of the Wnt proteins described above and sequence similarity to human Wnt protein with the amino acid sequence in NCBI Accession Nos. BAA82509 (gI:5509901); AAC51319.1 (GI:2105100); and 000755 (gI:2501663) (see also, Ikegawa et al., 1996, Cyto genetic Cell Genet. 74(1-2):149-52; Bui et al., 1997, Gene 189(1):25-9). Wntll refers to a Wnt protein with the functional activities described above and sequence similarity to human Wnt protein with the amino acid sequence in NCBI Accession Nos. BAB72099 (gI:17026012); CAA74159 (gI:3850708); and CAA73223.1 (gI:3850706) (see also, Kirikoshi et al., 2001, Int. J. Mol. Med. 8(6):651-6); Lako et al., 1998, Gene 219(1-2):101-10). As used herein in the context the specific Wnt proteins, "sequence similarity" refers to an amino acid sequence identity of at least about 80% or more, at least about 90% or more, at least about 95% or more, or at least about 98% or more when compared to the reference sequence. For instance, human Wnt7a displays about 97% amino acid sequence identity to murine Wnt7a while the amino acid sequence of human Wnt7a displays about 64% amino acid identity to human Wnt5a (Bui et al., supra).

[0059] The expression and/or secretion of various growth factors and/or Wnt proteins by the three dimensional can be modulated by incorporating cells that release different levels of the factors of interest. For example, vascular smooth muscle cells, are known to produce substantially more VEGF than human dermal fibroblasts. By utilizing vascular smooth muscle cells, instead of or in addition to fibroblasts, the expression and/or secretion of VEGF by the three dimensional tissue can be modulated.

Genetically Engineered Cells

[0060] Genetically engineered three-dimensional tissue can be prepared as described in U.S. Patent No. 5,785,964, the disclosure of which is incorporated herein by reference in its
entirety. A genetically-engineered tissue can serve as a gene delivery vehicle for sustained release of growth factors and/or Wnt proteins in vivo. For example, in certain embodiments, cells, such as stromal cells, can be engineered to express a gene product that is either exogenous or endogenous to the engineered cell. Stromal cells that can usefully be genetically engineered include, but are not limited to, fibroblasts (of fetal, neonatal, or adult origin), smooth muscle cells, cardiac muscle cells, stem or progenitor cells, and other cells found in loose connective tissue such as endothelial cells, macrophages, monocytes, adipocytes, pericytes, and reticular cells found in bone marrow. In various embodiments, stem or progenitor cells can be engineered to express an exogenous or endogenous gene product, and cultured on a three-dimensional scaffold, alone or in combination with stromal cells.

[0061] The cells and tissues can be engineered to express a desired gene product which can impart a wide variety of functions, including, but not limited to, enhanced function of the genetically engineered cells and tissues to promote tissue healing when implanted in vivo. The desired gene product can be a peptide or protein, such as an enzyme, hormone, cytokine, a regulatory protein, such as a transcription factor or DNA binding protein, a structural protein, such as a cell surface protein, or the desired gene product may be a nucleic acid such as a ribosome or antisense molecule. In some embodiments, the desired gene product is one or more Wnt proteins, which play a role in differentiation and proliferation of a variety of cells as described above (see, e.g., Miller, J.R., 2001, Genome Biology 3:3001.1-3001.15). For example, the recombinantly engineered cells can be made to express specific Wnt factors, including, but not limited to, Wnt5a, Wnt7a, and Wnt1.

[0062] In some embodiments, the desired gene products can provide enhanced properties to the genetically engineered cells, include but are not limited to, gene products which enhance cell growth, e.g., vascular endothelial growth factor (VEGF), hepatocyte growth factor (HGF), fibroblast growth factors (FGF), platelet derived growth factor (PDGF), epidermal growth factor (EGF), transforming growth factor (TGF), connective tissue growth factor (CTGF) and Wnt factors. In other embodiments, the cells and tissues can be genetically engineered to express desired gene products which result in cell immortalization, e.g., oncogenes or telomerese.

[0063] In other embodiments, the cells and tissues can be genetically engineered to express gene products which provide protective functions in vitro such as cyropreservation and anti-
desiccation properties, e.g., trehalose (U.S. Patent Nos. 4,891,319; 5,290,765; 5,693,788). The cells and tissues can also be engineered to express gene products which provide a protective function in vivo, such as those which would protect the cells from an inflammatory response and protect against rejection by the host's immune system, such as HLA epitopes, MHC alleles, immunoglobulin and receptor epitopes, epitopes of cellular adhesion molecules, cytokines and chemokines.

[0064] There are a number of ways that the desired gene products can be engineered to be expressed by the cells and tissues of the present invention. The desired gene products can be engineered to be expressed constitutively or in a tissue-specific or stimuli-specific manner. The nucleotide sequences encoding the desired gene products can be operably linked, e.g., to promoter elements which are constitutively active, tissue-specific, or induced upon presence of one or more specific stimulus.

[0065] In some embodiments, the nucleotide sequences encoding the engineered gene products are operably linked to regulatory promoter elements that are responsive to shear or radial stress. In these embodiments, the promoter element is activated by passing blood flow (shear), as well as by the radial stress that is induced as a result of the pulsatile flow of blood through the heart or vessel.

[0066] Examples of suitable regulatory promoter elements include, but are not limited to, tetracycline responsive elements, nicotine responsive elements, insulin responsive element, glucose responsive elements, interferon responsive elements, glucocorticoid responsive elements estrogen/progesterone responsive elements, retinoid acid responsive elements, viral transactivators, early or late promoter of SV40 adenovirus, the lac system, the tip system, the TAC system, the TRC system, the promoter for 3-phosphoglycerate and the promoters of acid phosphatase. In addition, artificial response elements can be constructed, comprising multimers of transcription factor binding sites and hormone-response elements similar to the molecular architecture of naturally-occurring promoters and enhancers (see, e.g., Herr and Clarke, 1986, J Cell 45(3): 461-70). Such artificial composite regulatory regions can be designed to respond to any desirable signal and be expressed in particular cell-types depending on the promoter/enhancer binding sites selected.

[0067] In some embodiments, the engineered three-dimensional tissue includes genetically engineered cells and produces naturally secreted factors that stimulate proliferation and
differentiation of stem cells and/or progenitor cells involved in the revascularization and healing of ischemic tissue.

Use of Cultured Three-Dimensional Tissues to Facilitate Healing of Ischemic Tissue

[0068] The three-dimensional tissues described herein find use in promoting the healing of ischemic tissue. The ability of the three-dimensional tissue to promote the healing of an ischemic tissue depends in part, on the severity of the ischemia. As will be appreciated by the skilled artisan, the severity of the ischemia depends, in part, on the length of time the tissue has been deprived of oxygen.

[0069] Without being bound by theory, application of the three-dimensional tissue to an ischemic tissue promotes various biological activities involved in the healing of ischemic tissue. Among such activities is the reduction or prevention of the remodeling of ischemic tissue. By "remodeling" herein is meant, the presence of one or more of the following: (1) a progressive thinning of the ischemic tissue, (2) a decrease in the number or blood vessels supplying the ischemic tissue, and/or (3) a blockage in one or more of the blood vessels supplying the ischemic tissue, and if the ischemic tissue comprises muscle tissue, (4) a decrease in the contractibility of the muscle tissue. Untreated, remodeling typically results in a weakening of the ischemic tissue such that it can no longer perform at the same level as the corresponding healthy tissue.

[0070] In some embodiments, the ischemic tissue includes cardiac muscle tissue. As illustrated in Example 1, application of one or more pieces of cultured three-dimensional tissue to ischemic regions of canine hearts improved ventricular performance and increased blood supply to the ischemic regions.

[0071] In some embodiments, the ischemic tissue includes skeletal muscle tissue, brain tissue e.g., affected by stroke or malformations of the arteries and veins covering the brain (i.e., AV malformations), kidney, liver, organs of the gastrointestinal tract, muscle tissue afflicted by atrophy, including neurologically based muscle atrophy and lung tissue. In further embodiments, the ischemic tissue is present in a mammal, such as a human.

[0072] In other embodiments, the ischemic tissue includes, but is not limited to, tissue wounds, such as skin ulcers and burns.
In other embodiments, the ischemic tissue does not include skin wounds, such as skin ulcers and burns.

In some embodiments, the ischemic tissue can be artificially created, i.e., can be created as a result of a surgical procedure.

In some embodiments, the ischemic tissue is heart tissue. Cardiovascular ischemia is generally a direct consequence of coronary artery disease, and is usually caused by rupture of an atherosclerotic plaque in a coronary artery, leading to formation of thrombus, which can occlude or obstruct a coronary artery, thereby depriving the downstream heart muscle of oxygen. Prolonged ischemia can lead to cell death or necrosis, and the region of dead tissue is commonly called an infarct.

Candidates for treatment by the methods described herein can be individuals who have been diagnosed with myocardial ischemia, but who have not been diagnosed with congestive heart failure. Diseases associated with myocardial ischemia include stable angina, unstable angina, and myocardial infarction. In some embodiments, candidates for the methods described herein will be patients with stable angina and reversible myocardial ischemia. Stable angina is characterized by constricting chest pain that occurs upon exertion or stress, and is relieved by rest or sublingual nitroglycerin. Coronary angiography of patients with stable angina usually reveals 50-70% obstruction of at least one coronary artery. Stable angina is usually diagnosed by the evaluation of clinical symptoms and ECG changes. Patients with stable angina may have transient ST segment abnormalities, but the sensitivity and specificity of these changes associated with stable angina are low.

In some embodiments, candidates for the methods described herein will be patients with unstable angina and reversible myocardial ischemia. Unstable angina is characterized by constricting chest pain at rest that is relieved by sublingual nitroglycerin. Anginal chest pain is usually relieved by sublingual nitroglycerin, and the pain usually subsides within 30 minutes. There are three classes of unstable angina severity: class I, characterized as new onset, severe, or accelerated angina; class II, subacute angina at rest characterized by increasing severity, duration, or requirement for nitroglycerin; and class III, characterized as acute angina at rest. Unstable angina represents the clinical state between stable angina and acute myocardial infarction (AMI) and is thought to be primarily due to the progression in the severity and extent of atherosclerosis, coronary artery spasm, or hemorrhage into non-
occluding plaques with subsequent thrombotic occlusion. Coronary angiography of patients
with unstable angina usually reveals 90% or greater obstruction of at least one coronary
artery, resulting in an inability of oxygen supply to meet even baseline myocardial oxygen
demand. Slow growth of stable atherosclerotic plaques or rupture of unstable atherosclerotic
plaques with subsequent thrombus formation can cause unstable angina. Both of these causes
result in critical narrowing of the coronary artery. Unstable angina is usually associated with
atherosclerotic plaque rupture, platelet activation, and thrombus formation. Unstable angina
is usually diagnosed by clinical symptoms, ECG changes, and changes in cardiac markers.

[0078] In some embodiments, candidates for the methods described herein will be patients
undergoing an acute myocardial infarction. Myocardial infarction is characterized by
constricting chest pain lasting longer than 30 minutes that can be accompanied by diagnostic
ECG Q waves. Most patients with AMI have coronary artery disease, and as many as 25% of
AMI cases are "silent" or asymptomatic infarctions. AMI is usually diagnosed by clinical
symptoms, ECG changes, and elevations of cardiac proteins, most notably cardiac troponin,
creatine kinase-MB and myoglobin.

[0079] In some embodiments, candidates for the methods described herein will be human
patients with left ventricular dysfunction and reversible myocardial ischemia that are
undergoing a coronary artery bypass graft (CABG) procedure, who have at least one graftable
 coronary vessel and at least one coronary vessel not amenable to bypass or percutaneous
coronary intervention.

[0080] As described in more detail below, one or more of the tissues comprising the wall of
the heart of an individual diagnosed with one of the disease states described above, can be
contacted with a cultured three-dimensional tissue, including the epicardium, the myocardium
and the endocardium.

Assays Useful for Determining Healing of Ischemic Tissue

[0081] In some embodiments, application of the cultured three-dimensional tissue to an
ischemic tissue increases the number of blood vessels present in the ischemic tissue, as
measured using laser Doppler imaging (see, e.g., Newton et al., 2002, J Foot Ankle Surg,
41(4):233-7). In some embodiments, the number of blood vessels increases 1%, 2%, 5%; in
other embodiments, the number of blood vessels increases 10%, 15%, 20%, even as much as
25%, 30%, 40%, 50%; in some embodiments, the number of blood vessels increase even more, with intermediate values permissible.

[0082] In some embodiments, application of the cultured three-dimensional tissue to an ischemic heart tissue increases the ejection fraction. In a healthy heart, the ejection fraction is about 65 to 95 percent. In a heart comprising ischemic tissue, the ejection fraction is, in some embodiments, about 20 - 40 percent. Accordingly, in some embodiments, treatment with the cultured three-dimensional tissue results in a 0.5 to 1 percent absolute improvement in the ejection fraction as compared to the ejection fraction prior to treatment. In other embodiments, treatment with the cultured three-dimensional tissue results in an absolute improvement in the ejection fraction more than 1 percent. In some embodiments, treatment results in an absolute improvement in the ejection fraction of 1.5%, 2%, 3%, 4%, 5%, 6%, 7%, 8%, even as much as 9% or 10%, as compared to the ejection fraction prior to treatment. For example, if the ejection fraction prior to treatment was 40%, then following treatment ejection fractions between 41% to 59% are observed in these embodiments. In still other embodiments, treatment with the cultured three-dimensional tissue results in an improvement in the ejection fraction greater than 10% as compared to the ejection fraction prior to treatment.

[0083] In some embodiments, application of the cultured three-dimensional tissue to an ischemic heart tissue increases one or more of cardiac output (CO), left ventricular end diastolic volume index (LVEDVI), left ventricular end systolic volume index (LVESVI), and systolic wall thickening (SWT). These parameters are measured by art-standard clinical procedures, including, for example, nuclear scans, such as radionuclide ventriculography (RIW) or multiple gated acquisition (MUGA), and X-rays.

[0084] In some embodiments, application of the cultured three-dimensional tissue to an ischemic heart tissue causes a demonstrable improvement in the blood level of one or more protein markers used clinically as indicia of heart injury, such as creatine kinase (CK), serum glutamic oxalacetic transaminase (SGOT), lactic dehydrogenase (LDH) (see, e.g., U.S. Publication 2005/0142613), troponin I and troponin T can be used to diagnose heart muscle injury (see, e.g., U.S. Publication 2005/0021234). In yet other embodiments, alterations affecting the N-terminus of albumin can be measured (see, e.g., U.S. Publications 2005/0142613, 2005/0021234, and 2005/0004485; the disclosures of which are incorporated herein by reference in their entireties).
The methods and compositions described herein can be used in combination with conventional treatments, such as the administration of various pharmaceutical agents and surgical procedures. For example, in some embodiments, the cultured three-dimensional tissue is administered with one or more of the medications used to treat heart failure. Medications suitable for use in the methods described herein include angiotensin-converting enzyme (ACE) inhibitors (e.g., enalapril (Vasotec), lisinopril (Prinivil, Zestril) and captopril (Capoten)), angiotensin II (A-II) receptor blockers (e.g., losartan (Cozaar) and valsartan (Diovan)), diuretics (e.g., bumetanide (Bumex), furosemide (Lasix, Fumide), and spironolactone (Aldactone)), digoxin (Lanoxin), beta blockers, and nesiritide (Natrecor) can be used.

In other embodiments, the cultured three-dimensional tissue can be administered during a surgical procedure, such as angioplasty, single CABG, and/or multiple CABG.

Additionally, the cultured three-dimensional tissue can be used with therapeutic devices used to treat heart disease including heart pumps, endovascular stents, endovascular stent grafts, left ventricular assist devices (LVADs), biventricular cardiac pacemakers, artificial hearts, and enhanced external counterpulsation (EECP).

Administration and Dosage of Cultured Three-Dimensional Tissue

A variety of methods can be used to attach and/or contact the cultured three-dimensional tissue to ischemic tissue. Suitable means for attachment include, but are not limited to, direct adherence between the three-dimensional tissue and the ischemic tissue, biological glue, synthetic glue, lasers, and hydrogel. A number of hemostatic agents and sealants are commercially available, including but not limited to, "SURGICAL" (oxidized cellulose), "ACTIFoAM" (collagen), "FIBRX" (light-activated fibrin sealant), "BOREAL" (fibrin sealant), "FIBROCAPS" (dry powder fibrin sealant), polysaccharide polymers p-Gl cNAc ("SYVEC" patch; Marine Polymer Technologies), Polymer 27CK (Protein Polymer Tech.). Medical devices and apparatus for preparing autologous fibrin sealants from 120ml of a patient's blood in the operating room in one and one-half hour are also known (e.g., Vivostat System).

In some embodiments, the cultured three-dimensional tissue is attached directly to the ischemic tissue via cellular attachment. For example, in some embodiments, the three-
dimensional tissue can be attached to one or more of the tissues of the heart, including the epicardium, myocardium and endocardium. When attaching a three-dimensional tissue to the heart epicardium or myocardium, typically the pericardium (i.e., the heart sac) is opened or pierced prior to attachment of the three-dimensional tissue. In other embodiments, for example when attaching a three-dimensional tissue to the endocardium, a catheter or similar device can be inserted into a ventricle of the heart and the three-dimensional tissue attached to the wall of the ventricle.

[0090] In some embodiments, a three-dimensional tissue can be attached to an ischemic tissue using a surgical glue. Surgical glues suitable for use in the methods and compositions described herein include biological glues, such as a fibrin glue. For a discussion of applications using fibrin glue compositions see, e.g., U.S. Patent Application Serial Number 10/851,938 and the various references disclosed therein; the disclosures of which is incorporated by reference herein in its entirety.

[0091] In some embodiments, a laser can be used to attach the three-dimensional tissue to an ischemic tissue. By way of example, a laser dye can be applied to the heart, the three-dimensional tissue, or both, and activated using a laser of the appropriate wavelength to adhere the cultured three-dimensional tissue to the heart. For a discussion of various applications using a laser see, e.g., U.S. Patent Application Serial Number 10/851,938, the disclosure of which is incorporated by reference herein in its entirety.

[0092] In some embodiments, a hydrogel can be used to attach the cultured three-dimensional tissue to an ischemic tissue. A number of natural and synthetic polymeric materials can be used to form hydrogel compositions. For example, polysaccharides, e.g., alginate, can be crosslinked with divalent cations, polyphosphazenes and polyacrylates ionically or by ultraviolet polymerization (see e.g., U.S. Pat. No. 5,709,854). Alternatively, a synthetic surgical glue such as 2-octyl cyanoacrylate (“DERMABOND”, Ethicon, Inc., Somerville, NJ) can be used to attach the three-dimensional tissue to an ischemic tissue.

[0093] In some embodiments, the cultured three-dimensional tissue can be attached to an ischemic tissue using one or more sutures as described in U.S. Patent Application Serial Number 10/851,938, the disclosure of which is incorporated by reference herein in its entirety. In other embodiments, the sutures can comprise cultured three-dimensional tissue as described in U.S. application no. ___________ entitled “Three Dimensional Tissues and
Uses Thereof," filed concurrently herewith; the disclosure of which is incorporated herein by
reference in its entirety.

[0094] The cultured three-dimensional tissue is used in an amount effective to promote tissue
healing and/or revascularize the ischemic tissue. The amount of the cultured three-
dimensional tissue administered, depends, in part, on the severity of the ischemic tissue,
whether the cultured three-dimensional tissue is used as an injectable composition (see, e.g.,
U.S. application no. __________, entitled, " Cultured Three Dimensional Tissues and Uses
Thereof," filed concurrently herewith; the disclosure of which is incorporated herein by
reference in its entirety), the concentration of the various growth factors and/or Wnt proteins
present, the number of viable cells comprising the cultured three-dimensional tissue, ease of
access to the ischemic tissue (e.g., is the ischemic tissue present on the surface of the skin or
present in an organ), and/or the tissue or organ being treated. Determination of an effective
dosages is well within the capabilities of the those skilled in the art. Suitable animal models,
such as the canine model described in Example 1, can be used for testing the efficacy of the
dosage on a particular tissue.

[0095] As used herein "dose" refers to the number of cohesive pieces of cultured three-
dimensional tissue applied to an ischemic tissue. A typical cohesive piece of cultured three-
dimensional tissue is approximately 35 cm². As will be appreciated by those skilled in the
art, the absolute dimensions of the cohesive piece can vary, as long it comprises a sufficient
number of cells to stimulate angiogenesis and/or promote healing of ischemic tissue. Thus,
cohesive pieces suitable for use in the methods described herein can range in size from 15
cm² to 50 cm².

[0096] The application of more than one cohesive piece of cultured three-dimensional tissue
can be used to increase the area of the ischemic tissue treatable by the methods described
herein. For example, in embodiments using a single piece of cohesive tissue, the treatable
area is approximately doubled in size. In embodiments using three cohesive pieces of
cultured three-dimensional tissue, the treatable area is approximately tripled in size. In
embodiments using four cohesive pieces of cultured three-dimensional tissue, the treatable
area is approximately quadrupled in size. In embodiments using five cohesive pieces of
cultured three-dimensional tissue, the treatable area is approximately five-fold, i.e. from 35
cm² to 175 cm².
[0097] In some embodiments, one cohesive piece of cultured three-dimensional tissue is attached to a region of an ischemic tissue.

[0098] In other embodiments, two cohesive pieces of cultured three-dimensional tissue are attached to a region of an ischemic tissue.

[0099] In other embodiments, three cohesive pieces of cultured three-dimensional tissue are attached to a region of an ischemic tissue.

[0100] In other embodiments, four, five, or more cohesive pieces of cultured three-dimensional tissue are attached to a region of an ischemic tissue.

[0101] In embodiments in which two or more cohesive pieces of cultured three-dimensional tissue are administered, the proximity of one piece to another can be adjusted, depending in part on, the severity of the ischemic tissue, the type of tissue being treated, and/or ease of access to the ischemic tissue. For example, in some embodiments, the cultured pieces of three-dimensional tissue can be located immediately adjacent to each other, such that one or more edges of one piece contact one or more edges of a second piece. In other embodiments, the pieces can be attached to the ischemic tissue such that the edges of one piece do not touch the edges of another piece. In these embodiments, the pieces can be separated from each other by an appropriate distance based on the anatomical and/or disease conditions presented by the patient. Determination of the proximity of one piece to another, is well within the capabilities of those skilled in the art, and may be tested using suitable animal models, such as the canine model described in Example 1.

[0102] In embodiments that comprise a plurality of pieces of cultured three-dimensional tissue, some, or all of the pieces can be attached to the area comprising the ischemic tissue. In other embodiments, one or more of the pieces can be attached to areas that do not comprise ischemic tissue. For example, in some embodiments, one piece can be attached to an area comprising ischemic tissue and a second piece can be attached to an adjacent area that does not comprise ischemic tissue. In these embodiments, the adjacent area can comprise damaged or defective tissue. "Damaged," or "defective" tissue as used herein refer to abnormal conditions in a tissue that can be caused by internal and/or external events, including, but not limited to, the event that initiated the ischemic tissue. Other events that can result in ischemic, damaged or defective tissue include disease, surgery, environmental exposure, injury, aging, and/or combinations thereof.
[0103] In embodiments that comprise a plurality of pieces of cultured three-dimensional tissue, the pieces can be simultaneously attached, or concurrently attached to an ischemic tissue.

[0104] In some embodiments, the pieces can be administered over time. The frequency and interval of administration depends, in part, on the severity of the ischemic tissue, whether the cultured three-dimensional tissue is used as an injectable composition (see, e.g., U.S. application no. __________, entitled, "Cultured Three Dimensional Tissues and Uses Thereof," filed concurrently herewith; the disclosure of which is incorporated herein by reference in its entirety), the concentration of the various growth factors and/or Wnt proteins present, the number of viable cells comprising the cultured three-dimensional tissue, ease of access to the ischemic tissue (e.g., is the ischemic tissue present on the surface of the skin or present in an organ), and/or the tissue or organ being treated. For example, if the ischemic tissue is present in a skin wound, two, three, four, five, six, seven, eight, or more applications of a cultured three-dimensional tissue can be applied in weekly or monthly intervals. Determination of the frequency of administration and the duration between successive applications is well within the capabilities of the those skilled in the art, and if desired can be tested using suitable animal models, such as the canine model described in Example 1.

[0105] In some embodiments, the cultured three-dimensional tissue is administered as an injectable composition as described in the U.S. application no. __________, entitled, "Cultured Three Dimensional Tissues and Uses Thereof," filed concurrently herewith. Guidance for the administration and effective dosage of injectable compositions for the treatment of ischemic tissue is provided in U.S. application no. __________, entitled, "Cultured Three Dimensional Tissues and Uses Thereof," filed concurrently herewith; the disclosure of which is incorporated herein by reference in its entirety.

[0106] Any literature and similar materials cited in this application, including but not limited to patents, patent applications, articles, books, and treatises, regardless of the format of such literature and similar materials, are expressly incorporated by reference in their entirety for any purpose. In the event that one or more of the incorporated literature and similar materials differs from or contradicts this application, including but not limited to defined terms, term usage, described techniques, or the like, this application controls.
[0107] All numerical ranges in this specification are intended to be inclusive of their upper and lower limits.

[0108] Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs. Unless mentioned otherwise the techniques employed or contemplated herein are standard methodologies well known to one of ordinary skill in the art. The materials, methods and examples are illustrative only and not limiting.

7. EXAMPLES

7.1 Treatment of Chronically Ischemic Tissue in a Dog Heart Study

a. Experimental Design

[0109] The three-dimensional cultured tissue, i.e., Anginera™ (also referred to herein as Dermagraft™), was manufactured by Smith & Nephew. Anginera™ is a sterile, cryopreserved, human fibroblast-based tissue generated by the culture of human neonatal dermal fibroblasts onto a bioabsorbable polyglactin mesh scaffold (Vicryl™). The process is carried out within a specialized growth container or bioreactor. Tissue growth is supported with cell medium that provides the required nutrients for cell proliferation. The closed bioreactor system used to manufacture Anginera™ maintains a controlled environment for the growth of a sterile, uniform and reproducible, viable human tissue.

[0110] The dermal fibroblasts used in the manufacture of Anginera™ were obtained from human neonatal foreskin tissue derived from routine circumcision procedures. Every lot of Anginera™ passes USP sterility tests before being released for use. It is cryopreserved at -75°C after harvest to provide an extended shelf life. Following thawing, about 60% of the cells retain viability and are capable of secreting growth factors, matrix proteins, and glycosaminoglycans.

[0111] A canine study was used to evaluate the safety and efficacy of Anginera™ for treating chronically ischemic heart tissue. Evaluation of the data from the canine study demonstrated Anginera™ to be safe at all dosing levels. The canine study was in compliance with the Food and Drug Administration Good Laboratory Practice Regulations (GLP) as set forth in Title 21 of the U.S. Code of Federal Regulations, Part 58.
Initially, chronic myocardial ischemia was induced in forty animals (four groups of five male and five female mongrel dogs) through the surgical placement of an ameroid constrictor on the ventral interventricular branch of the left anterior descending coronary artery (LAD). Approximately 30 days (± two days) following the surgical placement of an ameroid constrictor, the animals received one of four treatments: Group 1, sham surgical treatment; Group 2, surgical application of one unit of non-viable Anginera™; Group 3, surgical application of one unit of viable Anginera™; and, Group 4, surgical application of three units of viable Anginera™. Anginera™ used in this study was Dermagraft™ released by Smith & Nephew for clinical use. All investigators performing tests or analyzing data were blinded to the greatest extent possible as to the identity of an animal's treatment. Two animals per sex were necropsied on Day 30 (± one day), and three animals per sex from each treatment group were necropsied on Day 90 (± one day) (see Table 3).

Electrocardiograms and direct arterial pressure were continuously monitored during the surgical procedure. A left lateral thoracotomy was performed between the fourth and fifth ribs. Prior to heart isolation, lidocaine was given intravenously (2 mg/kg) and topically as needed to help control arrhythmias. The heart was isolated and a pericardial well was constructed. The ventral interventricular branch of the left anterior descending coronary artery (LAD) was identified and isolated for placement of an ameroid constrictor. The appropriately sized ameroid constrictor (Cardovascular Equipment Corporation, Wakefiled, Massachusetts, 2.0-3.0 mm) was placed around the proximal portion of the LAD. Any ventricular arrhythmias were treated using pharmacological agents, i.e., lidocaine, dexamethasone, bretyllium, as needed and as indicated. The study design is illustrated in Table 3.

Table 3: Canine Study Design

<table>
<thead>
<tr>
<th>Group Number</th>
<th>Number of Animals</th>
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Safety was assessed by evaluating clinical observations, physical and ophthalmic examinations, body weights, body temperatures, cardiac monitoring (including electrocardiography (ECG), arterial blood pressure, heart rate, and echocardiographic determination of left ventricular function), clinical pathology (including hematology, coagulation, serum chemistry, Troponin T, and urinalysis), anatomic pathology and histopathology of selected organs and tissues. Additional evaluation of the echocardiography data from all treatment groups at the Day 30 and Day 90 time points was performed. Finally, a separate analysis of heart histology was performed.

Echocardiograms were collected within four weeks prior to Day -30 (i.e., 30 days prior to surgery, surgery was done at Day 0), approximately eight days prior to Day 1, and approximately eight days prior to sacrifice/necropsy (Day 30 or 90). Trans-thoracic resting and stress echocardiography were performed using methods to standardize echocardiographic windows and views. Animals were manually restrained as much as possible and placed in right lateral recumbency (right side down). Echocardiographic evaluation was performed after the animals have achieved a stable heart rate followed by a second echocardiographic examination under dobutamine-induced increased heart rate. Dobutamine was administered intravenously starting at five micrograms/kg/min and titrated to a maximum infusion rate of 50 micrograms/kg/min to achieve 50% increase in heart rate (± 10%). Animal ID numbers, study dates, and views were annotated on the video recording of each study. All echo studies were recorded on videotape and images and loops were captured digitally and saved to optical disc. Short-axis images were recorded on both videotape and digitally on optical disc and included at least two cardiac cycles. Segmental contractility, measured as wall thickening (in centimeters), was quantified in the ischemic region and the control region of
the left ventricle. These measurements were performed in three cross sectional planes to include basal plane, mid papillary plane and a low-papillary plane. Left ventricular dimensional measurements were taken from 2 dimensional images. Two-chambered and four-chambered long axis images were recorded for the determination of left ventricular volumes, ejection fraction, and cardiac output. The mathematical model for this determination was the biplane, modified Simpson’s approximation. Electrocardiograms were recorded coordinate with the echocardiography.

[0117] Images saved to optical disc were stored on CDs in a DICOM image format for review in chronological order of the study by at least one board-certified veterinarian cardiologist (VetMed), blinded as to the identity of the samples.

[0118] Three measurements were performed on all echocardiographic data and reported as a mean of the three measurements.

[0119] Echocardiography was performed on all animals within four weeks prior to ameroid placement. Any animal identified with congenital heart disease or abnormal left ventricular function by echocardiogram was excluded from the study. Dobutamine stress echocardiography was performed to establish baseline comparisons.

[0120] Echocardiography was performed on all animals following surgery and placement of the ameroid constrictor on the LAD, and within approximately eight days prior to treatment application. Regional left ventricular wall motion was assessed under resting and dobutamine stress conditions. Any animals identified with transmural infarcts were excluded from the study.

[0121] Echocardiography was performed on all animals within approximately eight days of necropsy. Regional left ventricular wall motion was assessed under resting and dobutamine stress conditions. Global left ventricular function was assessed using a combination of left ventricular dimensional measurements, left ventricular volume determinations, ejection fraction, and cardiac output determinations.

[0122] A separate non-GLP echocardiography analysis was performed on the original echocardiography data to provide statistical comparisons of selected parameters. One-way analysis of variance (ANOVA) was used to determine a significant difference (p<0.05) between treatment groups. Comparisons were made between and within groups with specific
focus on parameter changes under resting conditions vs. dobutamine-stress conditions at both the 30 and 90 day time points. The parameters that were identified for comparison were:

1. Cardiac Output (CO): CO was expected to increase from resting to stress conditions. It is expected that diseased hearts would demonstrate a compromised ability to increase CO under dobutamine-stress conditions.

2. Left Ventricular Ejection Fraction (LVEF): LVEF was also expected to increase from resting to stress conditions. It is expected that diseased hearts would demonstrate a compromised ability to increase LVEF under dobutamine-stress conditions.

3. Left Ventricular End Diastolic Volume Index (LVEDVI): LVEDVI was expected to increase from resting to stress conditions. It is expected that diseased hearts would demonstrate greater increases in LVEDVI under dobutamine-stress conditions.

4. Left Ventricular End Systolic Volume Index (LVESVI): LVESVI was expected to decrease from resting to stress conditions. It is expected that diseased hearts would demonstrate a compromised ability to decrease LVESVI under dobutamine-stress conditions.

5. Systolic Wall Thickening (SWT): SWT values were expected to increase from resting to stress conditions. It is expected that diseased hearts would demonstrate a compromised ability to increase SWT values under dobutamine-stress conditions.

b. Results

[0123] None of the animals observed during the baseline pre-ameroid echo evaluation demonstrated significant left ventricular dysfunction or congenital heart disease. At baseline resting conditions all animals were evaluated to be within normal species ranges for hemodynamic values and wall dimensions. These data demonstrate that animals from all groups began the study with normal range values of left ventricular function. Furthermore, pre-treatment, post-ameroid left ventricular wall dimensions demonstrated a blunted response to dobutamine stress at the basilar (mitral valvular), high papillary, and low papillary levels in comparison to the pre-ameroid baseline assessment, demonstrating diminished wall function in the anterior and lateral wall of the left ventricle. These pre-treatment, post-ameroid echo observations are consistent with the ameroid experimental model that resulted in mild left ventricular dilation secondary to ventricular ischemia and demonstrate that the ameroid
canine model used in this study was successful at creating ventricular ischemia measurable by echocardiography.

[0124] No animal deaths were observed during the in-life phase of the study.

[0125] Clinical observations common to the surgical procedures associated with the exposure of the heart via thoracotomy (ameroid placement) or sternotomy (test article placement) were observed (e.g., swelling, erythema, open incisions, abrasions, etc.). The distribution and frequency of these clinical observations prior to Day 1 was similar between the final treatment groupings. Following the Day 1 surgical administration of treatment, clinical observations were similar between the four treatment groups with the exception that more animals in Groups 1, 2, and 4 (ischemia only, nonviable Anginera™ and three piece Anginera™ respectively), in comparison to Group 3 (single Anginera™) were observed with open surgical incisions. Most of the open incisions were observed following the sternotomy procedure (application of treatment), although one to two animals per group had open incisions observed following the thoracotomy procedure (application of ameroid occluder). All animals with open incisions were treated with antibiotics until the incisions were closed or had granulated and were dry.

[0126] Ophthalmologic examinations, physical examinations, body weights, body temperatures, hematology, coagulation, serum chemistry, Troponin T, urinalysis, surgical hemodynamic/cardiovascular monitoring, and weekly cardiovascular monitoring were all evaluated to be within normal species ranges and were not different between the four groups of animals. Collectively, these data demonstrate the safety of Anginera™ at all dosing levels within the parameters evaluated. Qualitative evaluation of the ECGs demonstrated normal cardiac rhythms for all but three animals (two Group 2 animals and one Group 3 animal). The arrhythmias or conduction disturbances observed in these three animals were evaluated to be either normal variants in dogs or a temporary residual effect of the surgical placement of the test article onto the myocardial surface.

[0127] Gross macroscopic pathology observations were limited to numerous myocardial adhesions (between the heart and the pericardium and the pericardium and the lungs or chest wall) and nodular lesions or discolorations in the myocardial tissue surrounding the ameroids. No differences were detected in the frequency or intensity of these observations amongst the four treatment groups of animals. These types of gross observations are consistent with the
surgical procedures utilized in this experimental protocol (i.e., thoracotomy and sternotomy). Microscopic pathology observations associated with the surgical placement of the test material included widespread fibrous thickening of the epicardium, correlating to the adhesions between the epicardial surface of the heart and the pericardium or lung, and limited serosanguineous exudates. These observations were noted in all animals in each of the four treatment groups and were felt to be related to the surgical procedures and not to specific treatment with the test article. Therefore, no safety concerns were evident from the histologic results, and the tissue responses observed were consistent with tissue injury attributed to the surgical procedures. Microscopic changes noted to be a result of the surgical placement of the ameroid occluder onto the coronary artery ranged from low-grade lymphoplasmacytic and histiocytic infiltrates, varying degrees of arterial intimal hyperplasia in the ameroided vessel, and areas of myocardial infarction. Transmural infarction was not observed in any of the tissue samples examined. Overall, no trends in the incidence or severity of infarction could be associated with specific treatment at the Day 30 or Day 90 evaluation time points.

[0128] In summary, evaluation of the primary safety endpoints (including hemodynamic, electrocardiographic, echocardiographic, and clinical and gross pathology observations) demonstrated the safety of Anginera™ at all dosing levels and at both time points.

[0129] Additional evaluations of heart histology were performed to identify evidence of new microvessel formation. These findings confirm previously reported and published findings of new microvessel formation with the presence of a mature microvasculature (arterioles, venules, and capillaries) (see FIG. 1).

[0130] Hematoxylin and eosin stained sections from the canine study were further analyzed to evaluate the cellular infiltrate in association with Anginera™ and the epicardial tissue. This analysis was performed on tissues that were in direct contact with the Anginera™ material. The following observations were made:

1. Scarring indicative of subendocardial ischemic damage was seen in all groups.

2. Group 1 (ischemia only) specimens showed minimal focal pericardial thickening without inflammation.
3. Group 2 (non-viable Anginera™) implants had diffuse mild and focally increased pericardial thickening with minimal inflammation and focal mesothelial proliferation.

4. Groups 3 (single dose Anginera™) and Group 4 (three pieces of Anginera™) had fibrous pericardial thickening with varying amounts of moderate, focal, multifocal or band-like inflammation between the patch and the epicardium, and focal foreign body reaction (most associated with sutures).

5. Less inflammation was seen at 90 than 30 days.

6. No definitive evidence of immunological reaction was seen.

7. In no case was there inflammation involving the myocardium.

8. Increased vasculature was seen focally in areas of pericardial inflammation.

[0131] These histopathological evaluations demonstrated no evidence of an immunologic reaction to Anginera™. There was a transient inflammatory response observed in all four treatment groups associated with the experimental conditions. In the viable Anginera™ groups there was evidence of a cellular response, which included an increase in microvasculature specific to the epicardium and pericardium. There was no evidence of a localized fibrosis, associated with the treatment, in the epicardium or myocardium that might lead to arrhythmias. The infiltrates had the morphologic appearance of macrophagic rather than lymphocytic cell types.

[0132] Prenecropsy echocardiographic assessment performed as part of the GLP study demonstrated dose-dependant decreases in left ventricular chamber volumes. Resting stroke volume and cardiac output indices were decreased in Group 3, but these mild decreases normalized in response to dobutamine infusion. Resting stroke volume and cardiac output indices decreased in Group 4 while decreases in left ventricular chamber volumes were marked compared with pretreatment values and was diminished over baseline values. These changes were more dramatic in Group 4 compared with Group 3. The response to dobutamine infusion in terms of percent difference in Group 4 was actually better than that seen in baseline values. Stroke volume and cardiac output indices did not return to normal baseline values, but were very close.
[0133] Group 3 animals (one unit dose Anginera™) at the 30-day prenecropsy time point had larger left ventricles than Group 3 animals at the 90-day prenecropsy time point or Group 4 animals (three units dose Anginera™) at either the 30 or 90-day prenecropsy time point. Group 4 animals had smaller left ventricles than Group 1, 2, or 3 animals. Compensatory mechanisms in and of themselves cause a decrease in left ventricular size (volume) as was seen in Group 1 untreated animals and Group 2 non-active Anginera™ treated animals. However, the fact that the left ventricular volumes were actually smaller in Group 4 animals than Group 1, 2, or 3 animals suggests a positive treatment effect. Decreases in left ventricular sizes/volumes are at least in part responsible for the decreases in stroke volume index (SVI) and cardiac output index (COI). These decreases returned both cardiac output index and stroke volume index to values similar to or better than normal baseline values that were also improved compared to the pre-treatment values. The most improved function compared with pre-treatment values was in Group 4 animals at the 90-day prenecropsy time point.

[0134] As part of the GLP study, segmental wall dimensions and segmental functional data suggested that application of treatment Groups 2, 3, and 4, increased wall dimensions where applied. It also suggested that in these regions there was a mild myocardial stiffening effect—evident in Group 2 dogs that received non-active test article alone. Data from this group also suggests that the non-active test article alone may cause an improvement in overall segmental function in adjacent segments. This may simply be a manifestation of compensatory responses in other segments. Group 3 animals demonstrated either mild increases in segmental function or no change over pretreatment values supporting the fact that Anginera™ was safe and at this dose mildly improved function in ischemic segments, but did not return segmental function to baseline normal values. Group 4 measurements at the basilar level demonstrated increased segmental function with return to close to baseline values and marked improvement over pretreatment values. This was not the case at the high papillary muscle level nor apical levels where segmental wall thickening was mildly depressed in response to dobutamine infusion in most segments. Segments that revealed mildly depressed segmental function had systolic wall dimensions that were increased significantly over either pretreatment values or normalized in response to treatment.

[0135] A separate non-GLP evaluation of left ventricular EDVI values was performed for two reasons. First, to specifically understand the changes in EDVI values following
treatment; and, second, to evaluate the 30 day ameroid period with respect to the canine model. Studies reported in the published literature have suggested that the canine is capable of significant collateralization of the coronary circulation. This can present limitations on the interpretation of functional data purposed to evaluate the benefit of a treatment. However, the canine model remains a well-established model within the published literature.

[0136] In light of these understandings of the canine model, EDVI parameters were evaluated in more detail. The parameters of EDVI during the 30 day ameroid period appear to suggest that animals treated with a single piece of Anginera™ (Group 3) and those treated with three pieces of Anginera™ (Group 4) may have had a more severe disease condition as suggested by EDVI values at the pre-treatment time point under dobutamine stress (FIG. 2). However, no statistically significant differences were seen in comparisons between these baseline EDVI values, possibly due to the large standard deviations and low sample size. Key to the X axis legend: Normal = pre-ameroid occlusion time point (-30 days); PreTx = ameroid occlusion, pre treatment time point (0 days); 30d PreNx = treatment after 30 days, prior to necropsy (30 days), and 90d PreNx = treatment after 90 days, prior to necropsy (90 days).

[0137] As previously described, a separate, secondary evaluation was performed on the original raw data that was collected in the primary echocardiography evaluation. The secondary evaluation focused on specific statistical comparisons of clinically relevant echocardiographic parameters. The general findings of the primary echo evaluation and the specific findings of the secondary echocardiography evaluation support of each other. In addition, in the secondary echocardiography evaluation, parameters of cardiac output (CO), left ventricular ejection fraction (LVEF), left ventricular end systolic volume index (LVESVI), and systolic wall thickening (SWT) support the conclusion that Anginera™ stimulates a positive biologic effect on chronically ischemic canine hearts. Furthermore, these data support the conclusion that treatment with viable Anginera™ improves ventricular performance and ventricular wall motion in chronically ischemic canine hearts after 30 days of treatment.

[0138] Following 30 days of treatment, dogs in the non-viable, single and multiple Anginera™ patch groups showed a significant (P<0.05) improvement in cardiac output with dobutamine (4273 ± 450, 4238 ± 268, and 4144 ± 236 ml/min, respectively) compared to their baseline, resting cardiac output. The sham surgical group did not significantly improve its CO with dobutamine infusion. However, at 90 days all dogs improved their CO with
dobutamine, including the sham operated animals (FIGS. 3 and 4). CO was expected to increase from resting to stress conditions. It is expected that diseased hearts would demonstrate a compromised ability to increase CO under dobutamine-stress conditions. These data suggest that dogs treated with non-viable, single, and multiple pieces of Anginera™ had a better CO response to dobutamine than the control sham group at 30 days. By 90 days, all groups performed statistically equivalent to each other.

[0139] LVEF demonstrated a similar stress response to dobutamine as CO at 30 and 90 days (FIGS. 5 and 6). Specifically after days of treatment, dogs in the non-viable, single and multiple Anginera™ patch groups showed a significant (P<0.05) improvement in LVEF with dobutamine compared to their baseline, resting LVEF. The sham surgical group did not significantly improve its LVEF with dobutamine infusion. However, at 90 days all dogs improved their LVEF with dobutamine, including the sham operated animals. LVEF was expected to increase from resting to stress conditions. It is expected that diseased hearts would demonstrate a compromised ability to increase LVEF under dobutamine-stress conditions. These data suggest that dogs treated with non-viable, single, and multiple pieces of Anginera™ had a better LVEF response to dobutamine than the control sham group at 30 days. By 90 days, all groups performed statistically equivalent to each other.

[0140] The LVEDV index was measured at rest and during stress in all groups at 30 and 90 days. At rest the LVEDV index was similar in all groups at 30 and 90 days. However, during stress at 90 days there is a significant (P<0.05) decrease in LVEDV index at the highest Anginera™ dose (Group 4) (FIGS. 7 and 8). LVEDVI was expected to increase from resting to stress conditions. It is expected that diseased hearts would demonstrate greater increases in LVEDVI under dobutamine-stress conditions. Therefore, the result of Group 4 animals at 90 days under dobutamine stress having significantly lower LVEDV index values suggests that the maximum treatment group (three pieces of Anginera™) provides additional benefit to the ischemic heart.

[0141] Consistent with the data from LVEF and CO, LVSV index values also significantly decreased with either viable or non-viable Anginera™ at stress compared to baseline at 30 days. At 90 days, there was also an improvement in the LVSV index with the sham surgery animals (FIGS. 9 and 10). LVESV index was expected to decrease from resting to stress conditions. It is expected that diseased hearts would demonstrate a compromised ability to decrease LVESVI under dobutamine-stress conditions. These data suggest that dogs treated
with non-viable, single, and multiple pieces of Anginera™ had a better LVESV index response to dobutamine than the control sham group at 30 days. By 90 days, all groups performed statistically equivalent to each other.

[0142] During the early ischemia period, dobutamine increased (P<0.05) SWT in all 4 randomized groups, however there appears to be a dose-dependent relationship since the most significant increase in SWT occurred in dogs that had the three patches of Anginera™ implanted (FIG. 11). At 30 and 90 days post treatment, within the chronic ischemia period, there is a gradual trend demonstrating increasing SWT in response to dobutamine over time, as plotted through a linear regression analysis of all data points (FIG. 12). This appears to occur in both untreated ischemic animals as well as those treated with Anginera™ patches. SWT values were expected to increase from resting to stress conditions. It is expected that diseased hearts would demonstrate a compromised ability to increase SWT values under dobutamine-stress conditions. These data suggest that during the early ischemia period (30 days after treatment), dobutamine increases (P<0.05) SWT in all 4 treatment groups; however, there appears to be a dose-dependent relationship since the most significant increase in SWT occurred in dogs that had the three patches of Anginera™ implanted.

[0143] The placement of either non-viable or viable Anginera™ patches, irrespective of the number of patches implanted resulted in an improved LV ejection fraction, increased cardiac output and reduced LV systolic volume index during stress with dobutamine at 30 days after induction of ischemia. In the chronic ischemia animals (group 1), this response was only seen at 90 days; at this time point the chronic ischemia animals were able to mount a response to dobutamine even though they had not received the Anginera™ treatment. This finding is congruent to the published literature where the canine model is described as a model that has an intrinsic ability for coronary collateralization.

[0144] In conclusion, the general findings of the primary echo evaluation as part of the GLP study and the specific findings of the separate non-GLP echocardiography analyses are in support of one another. In addition in the separate non-GLP echocardiography analyses, changes in CO, LVEF, LVESVI, and SWT support the conclusions that treatment with Anginera™ improves ventricular performance and ventricular wall motion in chronically ischemic canine hearts after 30 days of treatment.
CLAIMS

What is claimed is:

1. A method, comprising: contacting an ischemic tissue concurrently with at least a first cultured three-dimensional tissue and a second cultured three-dimensional tissue.

2. A method, comprising: contacting an ischemic tissue concurrently with at least a first cultured three-dimensional tissue, wherein the cultured three-dimensional tissue is in an amount sufficient to promote one or more biological activities associated with the healing of ischemic tissue.

3. The method of claim 2, in which the ischemic tissue is further contacted with at least a second cultured three-dimensional tissue.

4. The method of claim 1 or 2, wherein the cultured three-dimensional tissue is in an amount sufficient to reduce or prevent tissue remodeling associated with ischemia.

5. The method of claim 1 or 2, further comprising attaching the cultured three-dimensional tissue to the ischemic tissue using a degradable or non-degradable suture, a biologic glue, a synthetic glue, a laser dye, a hydrogel, or by cellular attachment.

6. The method of claim 1 or 2 in which the ischemic tissue is heart tissue.

7. The method of claim 6 in which the ischemia is reversible.

8. The method of claim 6 in which the ischemic tissue is epicardium.

9. The method of claim 6 in which the ischemic tissue is myocardium.

10. The method of claim 6 in which the ischemic tissue is endocardium.

11. The method of claim 6 in which the cultured three-dimensional tissue is in an amount sufficient to induce angiogenesis in the ischemic heart tissue.

12. The method of claim 6 in which the cultured three-dimensional tissue is in an amount sufficient to improve the ejection fraction of the treated heart.

13. The method of claim 6 in which the ischemic heart tissue is contacted with a first and at least second cultured three-dimensional tissues.

14. The method of claim 13 in which the ischemic heart tissue is contacted concurrently with said first and at least second cultured three-dimensional tissue.
15. The method of claim 6 further comprising attaching the cultured three-dimensional cells to the ischemic heart tissue using a degradable or non-degradable suture, a biologic glue, a synthetic glue, a laser dye, a hydrogel, or by cellular attachment.

16. A method of improving the ejection fraction of a diseased heart comprising contacting an ischemic region of the diseased heart with an effective amount of a cultured three-dimensional tissue.

17. A method of treating a patient suffering from coronary artery disease comprising contacting an ischemic region of the patient's heart with an effective amount of a cultured three-dimensional tissue.

18. A method of treating a patient suffering from left ventricular dysfunction and reversible myocardial ischemia comprising contacting an ischemic region of the patient's heart with an effective amount of a cultured three-dimensional tissue.

19. The method of claim 16, 17, or 18 in which the effective amount of the cultured three-dimensional tissue is sufficient to induce angiogenesis in the ischemic heart tissue.

20. The method of claim 16, 17, or 18 in which the effective amount of the cultured three-dimensional tissue is sufficient to improve the ejection fraction of the diseased heart.

21. The method of claim 16, 17, or 18 in which the ischemic heart tissue is contacted with a first and at least a second cultured three-dimensional tissue.

22. The method of claim 21 in which the ischemic heart tissue is contacted concurrently with said first and at least second cultured three-dimensional tissues.

23. The method of claim 1, 2, 16, 17, or 18 in which the cultured three-dimensional tissue comprises fibroblasts.

24. The method of claim 1, 2, 16, 17, or 18 in which the cultured three-dimensional tissue comprises smooth muscle cells.

25. The method of claim 24 in which the smooth muscle cells are vascular smooth muscle cells.

26. The method of claim 25 in which the vascular smooth muscle cells are aortic smooth muscle cells.
27. The method of claim 1, 2, 16, 17, or 18 in which the cultured three-dimensional tissue comprises cardiac muscle cells.

28. The method of claim 1, 2, 16, 17, or 18 in which the cultured three-dimensional tissue comprises stem cells.

29. The method of claim 1, 2, 16, 17, or 18 in which the cultured three-dimensional tissue comprises a plurality of cell types, each of the plurality of cell types independently selected from the group consisting of fibroblasts, smooth muscle cells, cardiac muscle cells, endothelial cells, mesenchymal stem cells, pericytes, macrophages, monocytes, leukocytes, plasma cells, mast cells and/or adipocytes.

30. The method of claim 16, 17, or 18 in which the ischemic region of the patient's heart is the epicardium.

31. The method of claim 16, 17, or 18 in which the ischemic region of the patient's heart is the myocardium.

32. The method of claim 16, 17, or 18 in which the ischemic region of the patient's heart is the endocardium.

33. The method of claim 1, 2, 16, 17, or 18 in which the cultured three-dimensional tissue comprises one or more WNT proteins.

34. The method of claim 33 in which the one or more Wnt proteins are selected from the group consisting of Wnt5a, Wnt7a, and/or Wnt1.

35. The method of claim 1 or 2 in which the cells of the cultured three-dimensional tissue are attached to a scaffold comprising a degradable material.

36. The method of claim 35 in which the degradable material comprises polyglycolic acid, polylactide, polylactide-co-glycolic acid, catgut sutures, cellulose, gelatin, collagen, or dextran.

37. The method of claim 1 or 2 in which the cells of the cultured three-dimensional tissue are attached to a scaffold comprising a non-degradable material.

38. The method of claim 37 in which the non-degradable material comprises a polyamide, a polyester, a polystyrene, a polypropylene, a polyacrylate, a polyvinyl, a polycarbonate, a polytetrafluorethylene, or a nitrocellulose compound or cotton.
39. The method of claim 1 or 2 in which the cultured three-dimensional tissue is attached to a mesh scaffold.

40. The method of claim 1 or 2 in which the cultured three-dimensional tissue is attached to a scaffold comprised of microparticles.

41. The method of claim 1 or 2 in which the cultured three-dimensional tissue is obtained directly from a fresh culture.

42. The method of claim 1 or 2 in which the cultured three-dimensional tissue has been cryopreserved.

43. The method of claim 1 in which the ischemic tissue is selected from the group consisting of liver, ulcerated intestinal tissue, pancreas, kidney, and/or bone marrow.
Figure 1
Figure 2A

Figure 2B
**Figure 3**

Cardiac Output in Dogs at Rest/Stress at 30 days

* P <0.05 vs. Resting Groups 2, 3, 4

**Figure 4**

Cardiac Output in Dogs at Rest/Stress at 90 days

* P <0.05 vs. Resting Groups.
Figure 5

Figure 6
**Figure 7**

**Figure 8**
Figure 9

Figure 10
Figure 11

Figure 12
A. CLASSIFICATION OF SUBJECT MATTER
INV. C12N5/06 C12N5/08 A61L27/38

According to International Patent Classification (IPC) onto both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
A61L C12N

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the International search (name of data base and, where practical, search terms used)
EPO-Internal, WPI Data, PAJ, BIOSIS

C. DOCUMENTS CONSIDERED TO BE RELEVANT

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Date of the actual completion of the international search: 31 May 2006

Date of mailing of the international search report: 21/06/2006

Name and mailing address of the ISA/ European Patent Office, P B 5818 Patentlaan 2 NL- 2280 HV Rijswijk Tel (+31-70) 340-2040, Tx 31 651 epo nl, Fax (+31-70) 340-3016

Authorized officer: Peri s Antol i , B
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