Title
Treatment of intervertebral disc degeneration

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ABSTRACT

The present application discloses a method for preventing or retarding degeneration of intervertebral disc at an intervertebral disc defect site, which includes injecting a mammalian connective tissue cell into the intervertebral disc defect site.
Invention Title: Treatment of intervertebral disc degeneration

The following statement is a full description of this invention, including the best method of performing it known to us:
TREATMENT OF INTERVERTEBRAL DISC DEGENERATION

CROSS REFERENCE TO RELATED APPLICATIONS

[0001] This application is a divisional of Australian patent application no. 2009225419, the entire disclosure of which is incorporated herein by reference.

BACKGROUND OF THE INVENTION

[0001a] Field of the Invention:

[0002] The present invention relates to prevention or retardation of intervertebral disc degeneration. The present application also relates to treating degenerating disc by preventing or retarding intervertebral disc degeneration. The present invention also relates to methods of using chondrocytes for introduction into injured intervertebral disc region and preventing or retarding degeneration of the intervertebral disc. The present invention also relates to a method of introducing at least one gene encoding a member of the transforming growth factor β superfamily into at least one mammalian connective tissue cell for use in preventing or retarding degeneration of intervertebral disc in the mammalian host. The present invention also relates to a method of using a mixture of chondrocytes and connective tissue cells containing a gene encoding a member of the transforming growth factor β superfamily into injured intervertebral disc region and preventing or retarding degeneration of the intervertebral disc.

SUMMARY OF THE INVENTION

[0003] In one aspect, the present invention is directed to a method for preventing or retarding degeneration of intervertebral disc at an intervertebral disc defect site, which includes injecting a mammalian connective tissue cell into the intervertebral disc defect site. The process preferably does not use a scaffolding or any supporting structure for the cells. Preferably, chondrocyte or fibroblast is used, and the subject is preferably a human being. If a chondrocyte is being used, the chondrocyte is preferably a non-disc chondrocyte or preferably a juvenile chondrocyte, meaning that the cells are isolated from a child who is less than two years old.
In other aspects, the chondrocyte may be primed chondrocytes. In particular, the connective tissue cell may be allogeneic relative to the mammalian subject sought to be treated.

[0004] In one aspect, the present invention relates to methods of using allogeneic juvenile chondrocytes or allogeneic non-disc chondrocytes for introduction into injured intervertebral disc region and preventing or retarding degeneration of the intervertebral disc.

[0005] In one aspect, the present invention is used to prevent or retard further degeneration of an area in the intervertebral disc that has been injured, torn or herniated.

[0006] In another aspect, the invention is directed to a method for preventing or retarding degeneration of intervertebral disc at an intervertebral disc defect site of a mammal, which method includes a) inserting a gene encoding a protein having intervertebral disc regenerating function into a mammalian cell, and b) transplanting the mammalian connective tissue cell into the intervertebral disc defect site. The process preferably does not use a scaffolding or any supporting structure for the cells. In this method, the gene may belong to TGF-β superfamily, such as TGF-β, and preferably TGF-β1. The connective tissue cell may be a chondrocyte or fibroblast. More preferably, chondrocyte is used, and the mammalian subject is preferably a human being. The connective tissue cell may allogeneic relative to the mammalian subject.

[0007] In yet another aspect, the invention is directed to method for preventing or retarding degeneration of intervertebral disc at an intervertebral disc defect site of a mammal, which includes a) inserting a gene encoding a protein having intervertebral disc regenerating function into a first mammalian connective tissue cell, and b) transplanting a mixture of the mammalian connective tissue cell of a) and unmodified second mammalian connective tissue cell into the intervertebral disc defect site. The process preferably does not use a scaffolding or any supporting structure for the cells. In this method, the gene may belong to TGF-β superfamily, such as TGF-β, and preferably TGF-β1. The first and second mammalian connective tissue cell may be chondrocyte or fibroblast. In the case of chondrocyte, the
chondrocyte may be non-disc chondrocyte or juvenile chondrocyte. In particular, the chondrocyte for the second mammalian connective tissue cell may be a primed chondrocyte. In another aspect, either or both of the first or second connective tissue cell may be allogeneic relative to the mammalian subject or to each other.

BRIEF DESCRIPTION OF THE DRAWINGS

FIGS. 1A-1F show a slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a rabbit spine four (4) weeks after surgery in which (i) the disc at L1/2 was injured and TGF-β1-producing chondrocytes were injected, (ii) no puncture and no treatment is seen at spine locus L2/3, and (iii) disc at L3/4 was injured and mixture of TGF-β1-producing chondrocytes and untransduced human chondrocytes in 1:3 ratio were injected; arrows point to L1/2 and L3/4 disc region. (C) shows MRI radiograph of a rabbit spine eight (8) weeks after surgery in which (i) the disc at L1/2 was injured and TGF-β1-producing chondrocytes were injected, (ii) no puncture and no treatment control at spine locus L2/3, and (iii) disc at L3/4 was injured and mixture of TGF-β1-producing chondrocytes and untransduced human chondrocytes in 1:3 ratio were injected; arrows point to L1/2 and L3/4 disc region. (D) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (E) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc. (F) shows X-ray radiograph of the rabbit described in (C) above, which is used to obtain a disc height index of the intervertebral disc. Mixed cell treatment in particular, has an intervertebral anti-degenerating effect.

FIGS. 2A-2F show a slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a rabbit spine four (4) weeks after surgery in which (i) the disc at L1/2 was injured and TGF-β1-producing chondrocytes were injected, (ii) no puncture and no treatment is seen at spine locus L2/3, and (iii) disc at L3/4 was injured and mixture of TGF-β1-producing chondrocytes and untransduced human chondrocytes in 1:3 ratio were injected; arrows point to L1/2 and L3/4 disc region. (D) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (E) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc. (F) shows X-ray radiograph of the rabbit described in (C) above, which is used to obtain a disc height index of the intervertebral disc. Mixed cell treatment in particular, has an intervertebral anti-degenerating effect.
β1-producing chondrocytes were injected, (ii) no puncture and no treatment control at spine locus L2/3, and (iii) disc at L3/4 was injured and mixture of TGF-β1-producing chondrocytes and untransduced human chondrocytes in 1:3 ratio were injected; arrows point to L1/2 and L3/4 disc region. (C) shows MRI radiograph of a rabbit spine eight (8) weeks after surgery in which (i) the disc at L1/2 was injured and TGF-β1-producing chondrocytes were injected, (ii) no puncture and no treatment is seen at spine locus L2/3, and (iii) disc at L3/4 was injured and mixture of TGF-β1-producing chondrocytes and untransduced human chondrocytes in 1:3 ratio were injected; arrows point to L1/2 and L3/4 disc region. (D) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (E) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc. (F) shows X-ray radiograph of the rabbit described in (C) above, which is used to obtain a disc height index of the intervertebral disc. Mixed cell treatment in particular, has an intervertebral anti-degenerating effect.

[0010] FIGS. 3A-3D show a slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a rabbit spine four (4) weeks after surgery in which (i) the disc at L1/2 was injured and TGF-β1-producing chondrocytes were injected, (ii) no puncture and no treatment control at spine locus L2/3, and (iii) disc at L3/4 was injured and mixture of TGF-β1-producing chondrocytes and untransduced human chondrocytes in 1:3 ratio were injected; arrows point to L1/2 and L3/4 disc region. (C) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (D) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc. Mixed cell treatment in particular, has an intervertebral anti-degenerating effect.
FIGS. 4A-4D show a slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a rabbit spine four (4) weeks after surgery in which (i) the disc at L1/2 was injured and mixture of TGF-β1-producing chondrocytes and untransduced human chondrocytes in 1:3 ratio were injected, (ii) no puncture and no treatment control at spine locus L2/3, and (iii) disc at L3/4 was injured and TGF-β1-producing chondrocytes were injected; arrows point to L1/2 and L3/4 disc regions. (C) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (D) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc. TGF-β1-producing chondrocytes treatment in particular, has an intervertebral anti-degenerating effect.

FIGS. 5A-5D show a slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a rabbit spine four (4) weeks after surgery in which (i) the disc at L1/2 was injured and mixture of TGF-β1-producing chondrocytes and untransduced human chondrocytes in 1:3 ratio were injected, (ii) no puncture and no treatment control at spine locus L2/3, and (iii) disc at L3/4 was injured and TGF-β1-producing chondrocytes were injected; arrows point to L1/2 and L3/4 disc regions. (C) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (D) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc. TGF-β1-producing chondrocytes treatment and mixed cell treatments in particular, have an intervertebral anti-degenerating effect.

FIGS. 6A-6D show a slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a rabbit spine four (4) weeks after surgery in which (i) the disc at L1/2 was injured and cell
culture media DMEM was injected, (ii) no puncture and no treatment control at spine locus L2/3, and (iii) disc at L3/4 was injured and untransduced chondrocytes were injected; arrows point to L1/2 and L3/4 disc regions. (C) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (D) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc. Untransduced chondrocytes treatment has an intervertebral anti-degenerating effect.

FIGS. 7A-7F show a slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a rabbit spine four (4) weeks after surgery in which (i) the disc at L1/2 was injured and cell culture media DMEM was injected, (ii) no puncture and no treatment control at spine locus L2/3, and (iii) disc at L3/4 was injured and untransduced chondrocytes were injected; arrows point to L1/2 and L3/4 disc regions. (C) shows MRI radiograph of a rabbit spine eight (8) weeks after surgery in which (i) the disc at L1/2 was injured and cell culture media DMEM was injected, (ii) no puncture and no treatment control at spine locus L2/3, and (iii) disc at L3/4 was injured and untransduced chondrocytes were injected; arrows point to L1/2 and L3/4 disc regions. (D) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (E) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc. (F) shows X-ray radiograph of the rabbit described in (C) above, which is used to obtain a disc height index of the intervertebral disc. Untransduced chondrocytes treatment has an intervertebral anti-degenerating effect.

FIGS. 8A-8F show a slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a rabbit spine four (4) weeks after surgery in which (i) the disc at T12/L1 was injured by needle
puncture and no injection, (ii) no puncture and no treatment control at spine locus L1/2, and (iii) disc at L2/3 was injured and untransduced chondrocytes were injected; arrows point to T12/L1 and L2/3 disc regions. (C) shows MRI radiograph of a rabbit spine eight (8) weeks after surgery in which (i) the disc at T12/L1 was injured by needle puncture and no injection, (ii) no puncture and no treatment control at spine locus L1/2, and (iii) disc at L2/3 was injured and untransduced chondrocytes were injected; arrows point to T12/L1 and L2/3 disc regions. (D) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (E) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc. (F) shows X-ray radiograph of the rabbit described in (C) above, which is used to obtain a disc height index of the intervertebral disc. Untransduced chondrocytes treatment has an intervertebral anti-degenerating effect.

FIGS. 9A-9D show a slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a rabbit spine eight (8) weeks after surgery in which (i) the disc at L2/3 was injured and cell culture media DMEM was injected, (ii) no puncture and no treatment control at spine locus L3/4, and (iii) disc at L4/5 was injured and primed chondrocytes were injected; arrows point to L2/3 and L4/5 disc regions. (C) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (D) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc. Primed chondrocyte treatment has an intervertebral anti-degenerating effect.
DETAILED DESCRIPTION OF THE INVENTION

[0017] As used herein, the term “biologically active” in reference to a nucleic acid, protein, protein fragment or derivative thereof is defined as an ability of the nucleic acid or amino acid sequence to mimic a known biological function elicited by the wild type form of the nucleic acid or protein.

[0018] As used herein, the term “connective tissue” is any tissue that connects and supports other tissues or organs, and includes but is not limited to a ligament, a cartilage, a tendon, a bone, and a synovium of a mammalian host.

[0019] As used herein, the term “connective tissue cell” or “cell of a connective tissue” include cells that are found in the connective tissue, such as fibroblasts, cartilage cells (chondrocytes), and bone cells (osteoblasts/osteocytes), which secrete collagenous extracellular matrix, as well as fat cells (adipocytes) and smooth muscle cells. Preferably, the connective tissue cells are fibroblasts, chondrocytes, or bone cells. More preferably, the connective tissue cells are chondrocytes cells. It will be recognized that the invention can be practiced with a mixed culture of connective tissue cells, as well as cells of a single type. It is also recognized that the tissue cells may be treated with an agent such as by chemical or radiation so that the cells stably express the gene of interest, preferably TGF-β1. Preferably, the connective tissue cell does not cause a negative immune response when injected into the host organism. It is understood that allogeneic cells may be used in this regard, as well as autologous cells for cell-mediated gene therapy or somatic cell therapy.

[0020] As used herein, “connective tissue cell line” includes a plurality of connective tissue cells originating from a common parent cell.

[0021] As used herein, "hyaline cartilage" refers to the connective tissue covering the joint surface. By way of example only, hyaline cartilage includes, but is not limited to, articular cartilage, costal cartilage, and nose cartilage.
In particular, hyaline cartilage is known to be self-renewing, responds to alterations, and provides stable movement with less friction. Hyaline cartilage found even within the same joint or among joints varies in thickness, cell density, matrix composition and mechanical properties, yet retains the same general structure and function. Some of the functions of hyaline cartilage include surprising stiffness to compression, resilience, and exceptional ability to distribute weight loads, ability to minimize peak stress on subchondral bone, and great durability.

Grossly and histologically, hyaline cartilage appears as a slick, firm surface that resists deformation. The extracellular matrix of the cartilage comprises chondrocytes, but lacks blood vessels, lymphatic vessels or nerves. An elaborate, highly ordered structure that maintains interaction between chondrocytes and the matrix serves to maintain the structure and function of the hyaline cartilage, while maintaining a low level of metabolic activity. The reference O’Driscoll, J. Bone Joint Surg., 80A: 1795-1812, 1998 describes the structure and function of hyaline cartilage in detail, which is incorporated herein by reference in its entirety.

As used herein, “injectable” composition refers to a composition that excludes various three-dimensional scaffold, framework, mesh or felt structure, which may be made of any material or shape that allows cells to attach to it and allows cells to grow in more than one layer, and which structure is generally implanted, and not injected. In one embodiment, the injection method of the invention is typically carried out by a syringe. However, any mode of injecting the composition of interest may be used. For instance, catheters, sprayers, or temperature dependent polymer gels also may be used.

As used herein, “juvenile chondrocyte” refers to chondrocyte obtained from a human being who is less than two years old. Typically, the chondrocyte is obtained from preferably the hyaline cartilage region of an extremity of the body, such as a finger, nose, ear
lobe and so forth. Juvenile chondrocytes may be used as donor chondrocytes for allogeneic treatment of defected or injured intervertebral disc.

[0026] As used herein, the term "mammalian host” includes members of the animal kingdom including but not limited to human beings.

[0027] As used herein, “mixed cell” or a “mixture of cells” or “cell mixture” refers to the combination of a plurality of cells that include a first population of cells that are transfected or transduced with a gene of interest and a second population of cells that are untransduced.

[0028] In one embodiment of the invention, mixed cells may refer to the combination of a plurality of connective tissue cells that include cells that have been transfected or transduced with a gene or DNA encoding a member of the transforming growth factor β superfamily and cells that have not been transfected or transduced with a gene encoding a member of the transforming growth factor β superfamily. Typically, the ratio of cells that have not been transfected or transduced with a gene encoding a member of the transforming growth factor β superfamily to cells that have been transfected or transduced with a TGF superfamily gene may be in the range of about 3-20 to 1. The range may include about 3-10 to 1. In particular, the range may be about 10 to 1 in terms of the number of cells. However, it is understood that the ratio of these cells should not be necessarily fixed to any particular range so long as the combination of these cells is effective to treat injured intervertebral disc by slowing or retarding degeneration of defected intervertebral disc.

[0029] As used herein, “non-disc chondrocyte” refers to chondrocytes isolated from any part of the body except for intervertebral disc cartilage tissue. Non-disc chondrocytes of the present invention may be used for allogeneic transplantation or injection into a patient to treat defected or injured intervertebral disc.

[0030] As used herein, the term "patient" includes members of the animal kingdom including but not limited to human beings.
As used herein, the term “primed” cell refers to cells that have been activated or changed to express certain genes.

As used herein, “slowing” or “prevention” of intervertebral disc degeneration refers to the retention of volume of intervertebral disc or height of the disc over time compared with the volume or height level that would normally be found at the site of injury leading to normal degeneration over a given time. This may mean a percentage increase of volume or height, such as about 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, or 90% compared with the normal expected degeneration levels at a given time, or may mean lessening of damage or depletion of volume or height of the intervertebral disc at the locus.

As used herein, the “transforming growth factor-β (TGF-β) superfamily” encompasses a group of structurally related proteins, which affect a wide range of differentiation processes during embryonic development. The family includes, Müllerian inhibiting substance (MIS), which is required for normal male sex development (Behringer, et al., Nature, 345:167, 1990), Drosophila decapentaplegic (DPP) gene product, which is required for dorsal-ventral axis formation and morphogenesis of the imaginal discs (Padgett, et al., Nature, 325:81-84, 1987), the Xenopus Vg-1 gene product, which localizes to the vegetal pole of eggs (Weeks, et al., Cell, 51:861-867, 1987), the activins (Mason, et al., Biochem, Biophys. Res. Commun., 135:957-964, 1986), which can induce the formation of mesoderm and anterior structures in Xenopus embryos (Thomsen, et al., Cell, 63:485, 1990), and the bone morphogenetic proteins (BMP’s, such as BMP-2, 3, 4, 5, 6 and 7, osteogenin, OP-1) which can induce de novo cartilage and bone formation (Sampath, et al., J. Biol. Chem., 265:13198, 1990). The TGF-β gene products can influence a variety of differentiation processes, including adipogenesis, myogenesis, chondrogenesis, hematopoiesis, and epithelial cell differentiation (for a review, see Massague, Cell 49:437, 1987), which is incorporated herein by reference in its entirety.
The proteins of the TGF-β family are initially synthesized as a large precursor protein, which subsequently undergoes proteolytic cleavage at a cluster of basic residues approximately 110-140 amino acids from the C-terminus. The C-terminal regions of the proteins are all structurally related and the different family members can be classified into distinct subgroups based on the extent of their homology. Although the homologies within particular subgroups range from 70% to 90% amino acid sequence identity, the homologies between subgroups are significantly lower, generally ranging from only 20% to 50%. In each case, the active species appears to be a disulfide-linked dimer of C-terminal fragments. For most of the family members that have been studied, the homodimeric species has been found to be biologically active, but for other family members, like the inhibins (Ung, et al., Nature, 321:779, 1986) and the TGF-β's (Cheifetz, et al., Cell, 48:409, 1987), heterodimers have also been detected, and these appear to have different biological properties than the respective homodimers.

Members of the superfamily of TGF-β genes include TGF-β3, TGF-β2, TGF-β4 (chicken), TGF-β1, TGF-β5 (Xenopus), BMP-2, BMP-4, Drosophila DPP, BMP-5, BMP-6, Vgr1, OP-1/BMP-7, Drosophila 60A, GDF-1, Xenopus Vgf, BMP-3, Inhibin-βA, Inhibin-βB, Inhibin-α, and MIS. These genes are discussed in Massague, Ann. Rev. Biochem. 67:753-791, 1998, which is incorporated herein by reference in its entirety.

Preferably, the member of the superfamily of TGF-β genes is TGF-β1, TGF-β2, TGF-β3, BMP-2, BMP-3, BMP-4, BMP-5, BMP-6, or BMP-7.

**Intervertebral Disc**

The intervertebral discs make up one fourth of the spinal column's length. There are no discs between the Atlas (C1), Axis (C2), and Coccyx. Discs are not vascular and therefore depend on the end plates to diffuse needed nutrients. The cartilaginous layers of the end plates anchor the discs in place.
The intervertebral discs are fibrocartilaginous cushions serving as the spine's shock absorbing system, which protect the vertebrae, brain, and other structures (i.e. nerves). The discs allow some vertebral motion: extension and flexion. Individual disc movement is very limited – however considerable motion is possible when several discs combine forces.

Intervertebral discs are composed of an annulus fibrosus and a nucleus pulposus. The annulus fibrosus is a strong radial tire–like structure made up of lamellae; concentric sheets of collagen fibers connected to the vertebral end plates. The sheets are orientated at various angles. The annulus fibrosus encloses the nucleus pulposus.

Although both the annulus fibrosus and nucleus pulposus are composed of water, collagen, and proteoglycans (PGs), the amount of fluid (water and PGs) is greatest in the nucleus pulposus. PG molecules are important because they attract and retain water. The nucleus pulposus contains a hydrated gel–like matter that resists compression. The amount of water in the nucleus varies throughout the day depending on activity. As people age, the nucleus pulposus begins to dehydrate, which limits its ability to absorb shock. The annulus fibrosus gets weaker with age and begins to tear. While this may not cause pain in some people, in others one or both of these may cause chronic pain.

Pain due to the inability of the dehydrating nucleus pulposus to absorb shock is called axial pain or disc space pain. One generally refers to the gradual dehydration of the nucleus pulposus as degenerative disc disease. When the annulus fibrosus tears due to an injury or the aging process, the nucleus pulposus can begin to extrude through the tear. This is called disc herniation. Near the posterior side of each disc, all along the spine, major spinal nerves extend out to different organs, tissues, extremities etc. It is very common for the herniated disc to press against these nerves (pinched nerve) causing radiating pain, numbness, tingling, and diminished strength and/or range of motion. In addition, the contact of the inner nuclear gel, which contains inflammatory proteins, with a nerve can also cause significant pain. Nerve-related pain is called radicular pain.
Herniated discs go by many names and these can mean different things to different medical professionals. A slipped disc, ruptured disc, or a bulging disc can all refer to the same medical condition. Protrusions of the disc into the adjacent vertebra are known as Schmorl's nodes.

**Primed Cell Therapy**

The present invention encompasses administering primed cells to an intervertebral disc region in a mammal to treat injured intervertebral disc by preventing or retarding degeneration of intervertebral disc. Primed cells are typically connective tissue cells, and include chondrocytes or fibroblasts.

By way of example, when a population of primary chondrocytes are passaged about 3 or 4 times, their morphology typically changes to fibroblastic chondrocytes. As primary chondrocytes are passaged, they begin to lose some of their chondrocytic characteristics and begin to take on the characteristics of fibroblastic chondrocytes. When these fibroblastic chondrocytes are incubated or “primed” with a cytokine such as a protein from the TGF-β superfamily, the cells regain their chondrocytic characteristics, which include production of collagen.

Such primed cells include fibroblastic chondrocytes, which have been incubated with TGFβ1, and as a result have reverted to collagen producing chondrocytes. An advantage of using primed cells in retardation of intervertebral disc degeneration is the ease of creating useable chondrocytes for introduction into the intervertebral disc for production of collagen and otherwise maintenance of the cartilaginous matrix.

The cells may include without limitation primary cells or cells which have undergone about one to twenty passages. The cells may be connective tissue cells. The cells may include cells that have undergone a morphogenic change, wherein the priming causes reversion to the characteristics of the original cell. The cells may include without limitation chondrocytes, fibroblasts, or fibroblastic chondrocytes. Priming may occur by incubating the
cells for a period of at least 40 hours, or from 1 to 40 hours, from 2 to 30 hours, from 3 to 25 hours, from 4 to 20 hours, from 5 to 20, from 6 to 18 hours, 7 to 17 hours, 8 to 15 hours, or 9 to 14 hours, with a cytokine, and then optionally separating the cytokine from the cells and injecting the primed cells into a cartilaginous defect site of interest in order to regenerate cartilage, preferably hyaline cartilage. In one aspect, the cytokine may be a member of the superfamily of TGF-β. In particular, the cytokine may be TGF-β, and in particular, TGF-β1.

[0049] The cytokine may be present in the priming incubation mix in an amount to sufficiently “prime” the chondrocyte to be useful in the intervertebral treatment method. In this aspect, the priming incubation mix may contain at least about 1 ng/ml of the cytokine. In particular, the mix may contain from about 1 to 1000 ng/ml, from about 1 to 750 ng/ml, from about 1 to 500 ng/ml, from about 1 to 400 ng/ml, from about 1 to 300 ng/ml, from about 1 to 250 ng/ml, from about 1 to 200 ng/ml, from about 1 to 150 ng/ml, from about 1 to 100 ng/ml, from about 1 to 75 ng/ml, from about 1 to 50 ng/ml, from about 10 to 500 ng/ml, from about 10 to 400 ng/ml, from about 10 to 300 ng/ml, from about 10 to 250 ng/ml, from about 10 to 200 ng/ml, from about 10 to 150 ng/ml, from about 10 to 100 ng/ml, from about 10 to 75 ng/ml, from about 10 to 50 ng/ml, from about 15 to 500 ng/ml, from about 15 to 400 ng/ml, from about 15 to 300 ng/ml, from about 15 to 250 ng/ml, from about 15 to 200 ng/ml, from about 15 to 150 ng/ml, from about 15 to 100 ng/ml, from about 15 to 75 ng/ml, from about 15 to 50 ng/ml, from about 20 to 500 ng/ml, from about 20 to 400 ng/ml, from about 20 to 300 ng/ml, from about 20 to 250 ng/ml, from about 20 to 200 ng/ml, from about 20 to 150 ng/ml, from about 20 to 100 ng/ml, from about 20 to 75 ng/ml, from about 20 to 50 ng/ml, from about 25 to 500 ng/ml, from about 25 to 400 ng/ml, from about 25 to 300 ng/ml, from about 25 to 250 ng/ml, from about 25 to 200 ng/ml, from about 25 to 150 ng/ml, from about 25 to 100 ng/ml, from about 25 to 75 ng/ml, from about 25 to 50 ng/ml, from about 30 to 500 ng/ml, from about 30 to 400 ng/ml, from about 30 to 300 ng/ml, from about 30 to 250 ng/ml, from about 30 to 200 ng/ml, from about 30 to 150 ng/ml, from about 30 to 100 ng/ml, from
about 30 to 75 ng/ml, from about 30 to 50 ng/ml, from about 35 to 500 ng/ml, from about 35 to 400 ng/ml, from about 35 to 300 ng/ml, from about 35 to 250 ng/ml, from about 35 to 200 ng/ml, from about 35 to 150 ng/ml, from about 35 to 100 ng/ml, from about 35 to 75 ng/ml, from about 35 to 50 ng/ml, from about 40 to 500 ng/ml, from about 40 to 400 ng/ml, from about 40 to 300 ng/ml, from about 40 to 250 ng/ml, from about 40 to 200 ng/ml, from about 40 to 150 ng/ml, from about 40 to 100 ng/ml, from about 40 to 75 ng/ml, or from about 40 to 50 ng/ml.

[0050] One method of practicing the invention may include incubating the cells with a cytokine for a certain length of time to create primed cells and optionally separating the cytokine from the cells, and injecting the primed cells into intervertebral disc or the site of interest near it. Alternatively, the cells may be incubated with the cytokine of interest for a time and the combination may be administered to the site of defect without separating out the cytokine.

[0051] It is to be understood that while it is possible that substances such as a scaffolding or a framework as well as various extraneous tissues may be implanted together in the primed cell therapy protocol of the present invention, it is also possible that such scaffolding or tissue not be included in the injection system of the invention. In a preferred embodiment, in the inventive somatic cell therapy, the invention is directed to a simple method of injecting a population of primed connective tissue cells to the intervertebral disc space.

[0052] It will be understood by the artisan of ordinary skill that the source of cells for treating a human patient may be the patient's own connective tissue cells, such as autologous fibroblast or chondrocyte cells, but that allogeneic cells as well as xenogeneic cells may also be used without regard to the histocompatibility of the cells. Alternatively, in one embodiment of the invention, allogeneic cells may be used having matching histocompatibility to the mammalian host. To describe in further detail, the histocompatibility of the donor and the patient are determined so that histocompatible cells are administered to
the mammalian host. Also, juvenile chondrocytes may also be used allogeneically without necessarily determining the histocompatibility of the donor and the patient.

[0053] Gene Delivery

[0054] In one aspect the present invention discloses *ex vivo* and *in vivo* techniques for delivery of a DNA sequence of interest to the connective tissue cells of the mammalian host. The *ex vivo* technique involves culture of target connective tissue cells, *in vitro* transfection of the DNA sequence, DNA vector or other delivery vehicle of interest into the connective tissue cells, followed by transplantation of the modified connective tissue cells to the target area of the mammalian host, so as to effect *in vivo* expression of the gene product of interest.

[0055] It is to be understood that while it is possible that substances such as a scaffolding or a framework as well as various extraneous tissues may be implanted together in the protocol of the present invention, it is preferred that such scaffolding or tissue not be included in the injection system of the invention. In a one embodiment, the invention is directed to a simple method of injecting a TGF superfamily protein or a population of cultured, untransfected/untransduced or transfected/transduced connective tissue cells or a mixture thereof to the intervertebral disc space so that the exogenous TGF superfamily protein is expressed or is active in the space.

[0056] It will be understood by the artisan of ordinary skill that one source of cells for treating a human patient is the patient’s own connective tissue cells, such as autologous chondrocyte cells. Another source of cells includes allogeneic cells without regard to the histocompatibility of the cells to the patient sought to be treated.

[0057] More specifically, this method includes employing a gene product that is a member of the transforming growth factor β superfamily, or a biologically active derivative or fragment thereof, or a biologically active derivative or fragment thereof.
In another embodiment of this invention, a compound for parenteral administration to a patient in a therapeutically effective amount is provided that contains a TGF-β superfamily protein and a suitable pharmaceutical carrier.

Another embodiment of this invention provides for a compound for parenteral administration to a patient in a prophylactically effective amount that includes a TGF-β superfamily protein and a suitable pharmaceutical carrier.

In therapeutic applications, the TGF-β protein may be formulated for localized administration. Techniques and formulations generally may be found in Remington’s Pharmaceutical Sciences, Mack Publishing Co., Easton, Pa., latest edition. The active ingredient that is the TGF protein is generally combined with a carrier such as a diluent of excipient which may include fillers, extenders, binding, wetting agents, disintegrants, surface-active agents, erodable polymers or lubricants, depending on the nature of the mode of administration and dosage forms. Typical dosage forms include, powders, liquid preparations including suspensions, emulsions and solutions, granules, and capsules.

The TGF protein of the present invention may also be combined with a pharmaceutically acceptable carrier for administration to a subject. Examples of suitable pharmaceutical carriers are a variety of cationic lipids, including, but not limited to N-(1-2,3-dioleyloxy)propyl)-n,n,n-trimethylammonium chloride (DOTMA) and dioleoylphosphotidyl ethanolamine (DOPE). Liposomes are also suitable carriers for the TGF protein molecules of the invention. Another suitable carrier is a slow-release gel or polymer comprising the TGF protein molecules.

The TGF beta protein may be mixed with an amount of a physiologically acceptable carrier or diluent, such as a saline solution or other suitable liquid. The TGF protein molecule may also be combined with other carrier means to protect the TGF protein and biologically active forms thereof from degradation until they reach their targets and/or
facilitate movement of the TGF protein or biologically active form thereof across tissue barriers.

[0063] A further embodiment of this invention includes storing the connective tissue cell prior to transferring the cells. It will be appreciated by those skilled in the art that the connective tissue cell may be stored frozen in 10 percent DMSO in liquid nitrogen.

[0064] In the present application, a method is provided for regenerating or preventing degeneration of intervertebral disc by injecting an appropriate mammalian cell that is transfected or transduced with a gene encoding a member of the transforming growth factor-beta (TGF-β) superfamily, including, but not limited to, BMP-2 and TGF-β 1, 2, and 3.

[0065] In another embodiment of the present application, a method is provided for preventing or retarding degeneration of intervertebral disc by injecting an appropriate mammalian cell that is not transfected or transduced with a gene encoding a member of the transforming growth factor-beta (TGF-β) superfamily or that is not transfected or transduced with any other gene. In another aspect, the invention is directed to treating injured or degenerated intervertebral disc by preventing or retarding degeneration of the intervertebral disc by using the above-described method.

[0066] In another embodiment of the present application, a method is provided for preventing or retarding degeneration of intervertebral disc by injecting an appropriate mammalian cell that is transfected or transduced with a gene encoding a member of the transforming growth factor-beta (TGF-β) superfamily. In another aspect, the invention is directed to treating injured or degenerated intervertebral disc by preventing or retarding degeneration of the intervertebral disc by using the above-described method.

[0067] In another embodiment of the invention, a method is provided for preventing or retarding degeneration of intervertebral disc by injecting a combination of or a mixture of an appropriate mammalian cell that is transfected or transduced with a gene encoding a member of the transforming growth factor-beta (TGF-β) superfamily and an appropriate mammalian
cell that is not transfected or transduced with a gene encoding a member of the transforming growth factor-beta (TGF-β) superfamily or that is not transfected or transduced with any other gene. In another aspect, the invention is directed to treating injured or degenerated intervertebral disc by preventing or retarding degeneration of the intervertebral disc by using the above-described method.

[0068] In an embodiment of the invention, it is understood that the cells may be injected into the area in which degeneration of the intervertebral disc is to be sought to be prevented or retarded by using the cell above-described composition with or without scaffolding material or any other auxiliary material, such as extraneous cells or other biocompatible carriers. That is, the modified cells alone, unmodified cells alone, or a mixture or combination thereof may be injected into the area in which the degeneration of the intervertebral disc is sought to be prevented or retarded.

[0069] The following examples are offered by way of illustration of the present invention, and not by way of limitation.

EXAMPLES

[0070] EXAMPLE I - MATERIALS AND METHODS

[0071] Plasmid Construction

[0072] The plasmid pMTMLVβ1 was generated by subcloning a 1.2-kb Bgl II fragment containing the TGF-β1 coding sequence and a growth hormone poly A site at the 3' end into the Bam HI site of pMTMLV. pMTMLV vector was derived from the retroviral vector MFG by deleting entire gag and env sequences as well as some of ψ packaging sequence.

[0073] Cell Culture and Transduction - The TGF-β cDNA cloned in retroviral vectors were individually transduced into chondrocytes (hChon-TGF-β1). They were cultured in Dulbecco's Modified Eagle's Medium (GIBCO-BRL, Rockville, MD) with 10% concentration of fetal bovine serum.
To select the cells with the transduced gene sequence, neomycin (300 μg/ml) was added into the medium. The cells with TGF-β1 expression were sometimes stored in liquid nitrogen and cultured just before the injection.

pMTMLVβ1 and pVSVG plasmid DNAs were cotransfected into GP2-293 cells by Fugene transfection methods. After 48 hr of culture, the supernatant was filtered through a 0.45 um polysulfone filter. The filtered culture supernatant was then diluted 2-fold with medium plus 10% FBS and used to infect human chondrocytes. Human chondrocytes seeded in 60 mm culture dish 18 hr prior to infection were infected with the filtrate plus polybrene (8 μg/ml, Sigma, St. Louis, MO). After 4 hr of incubation, the medium was replaced with fresh medium. Infection was repeated 24 hr later with the saved viral supernatant. Transduced cells were cultured in Dulbecco’s modified Eagle’s medium containing 10% fetal bovine serum from 48 hr after the second infection. Selected colonies were transferred to 24 or 6 well plates and TGF-β1 production was measured using a Quantikine ELISA assay kit (R&D system, Minneapolis, MN).

Radiographic Analysis of Disc Height

Radiographs were taken after administration of ketamine hydrochloride (25 mg/kg) and Rompun (1 mg/kg) at various week intervals after the puncture. Extreme care was taken to maintain a consistent level of anesthesia during radiography of each animal and at each time to obtain a similar degree of muscle relaxation, which may affect the disc height. Therefore, the preoperative radiograph was always used as a baseline measurement. Efforts were also made to keep the spine in a slightly flexed position. To decrease the error from axial rotation of the spine and beam divergence, radiographs were repeated at least twice on each animal in the lateral decubitus position, with the beam centered at 4cm from the rabbit iliac crest. Radiographs were digitally scanned and digitally stored using an Image Capture software.
Using digitized radiographs, measurements, including the vertebral body height and IVD height, were analyzed using the public domain image analysis. The data were transported to Excel software, and the IVD height was expressed as the DHI using the method of Lu et al. “Effects of chondroitinase ABC and chymopapain on spinal motion segment biomechanics. An in vivo biomechanical, radiologic, and histologic canine study”, Spine 1997;22:1828–34. Average IVD height (DHI) was calculated by averaging the measurements obtained from the anterior, middle, and posterior portions of the IVD and dividing that by the average of adjacent vertebral body heights. Changes in the DHI of injected discs were expressed as percent DHI and normalized to the measured preoperative IVD height (percent DHI = postoperative DHI/preoperative DHI X 100). The within-subject standard deviation (Sw) was calculated using the equation:

\[ \sqrt{\frac{\sum(x_1 - x_2)^2}{2n}} \]

Where \( X_1 \) is the first measurement value, \( X_2 \) is the second measurement value, and \( n = 450 \). The percent coefficient of variance (percent CV) was calculated as (Sw/means of all measurements X 100). The intraobserver error of DHI measurements was estimated to be minimal (Sw: 0.001800316; percent CV: 3.13). The interobserver error was also reported to be small (Sw: 0.003227; percent CV: 9.6)

**MRI Assessments**

MRI examinations were performed on all rabbits in the study using a 0.3-T imager (Airis II, version 4.0 A; Hitachi Medical System America, Inc.) with a quadrature extremity coil receiver. After sacrifice, the spinal columns with surrounding soft tissue were isolated and subjected to MRI analysis. T2-weighted sections in the sagittal plane were obtained in the following settings: fast spin echo sequence with TR (time to repetition) of 4000 milliseconds and TE (time to echo) of 120 milliseconds; 256(h) X 128 (v) matrix; field of view of 260; and 4 excitations. The section thickness was 2mm with a 0-mm gap. A blinded
observer using the modified Thompson classification based on changes in the degree and area of signal intensity from grade 1 to 4 (1 = normal, 2 = minimal decrease of signal intensity but obvious narrowing of high signal area, 3 = moderate decrease of signal intensity, and 4 = severe decrease of signal intensity) evaluated MRIs. The intraobserver and interobserver reliability correlation coefficients of MRI grading based on 2 evaluations were excellent (K = 0.98, 0.90, respectively), as determined by the Cohen kappa correlation coefficient.

EXAMPLE II – EXPERIMENTAL METHODS AND RESULTS

Preventing Degeneration of Injured Intervertebral Disc

New Zealand white male rabbits were used. An open surgical technique was used. Three intervertebral levels in the lumbar spine: L2-3, L3-4, L4-5 were experimentally treated or observed as a control in each animal. Treatments were assigned to levels in a balanced manner with multiple sites/discs per rabbit observed. Within subject design, pre-post surgery comparisons, change across disc levels were used as controls.

EXAMPLE III

Preventing Degeneration Of Injured Intervertebral Disc Using Untransduced Chondrocyte Alone, TGF-β1-Producing Chondrocyte Alone, Or With Mixed-Cells (Human Chondrocytes And TGF-β1-Producing Chondrocyte) Injection In Rabbits

All of the chondrocytes used in Examples I-V are non-disc chondrocytes and are juvenile chondrocytes, obtained from the hyaline cartilage portion of a finger of a less than two year old child.

Needle puncture was produced in the intervertebral discs of the lumbar spine. After this needle puncture, TGF-β1-producing chondrocytes, primary untransduced human chondrocytes, mixture of TGF-β1-producing chondrocytes and primary untransduced human chondrocytes, primed untransduced human chondrocytes or carrier/media are injected. Several controls are used. Experimental conditions are listed below Table I.

Table I
Briefly, a needle puncture injury is produced in the intervertebral discs of the lumbar spine of rabbit or a pig. After this needle puncture, rabbits are left to heal for 4 weeks. Then in a second surgical procedure, experimental treatment composition, which includes TGF-β1-producing chondrocytes and/or primary untransduced human chondrocytes (~5 x 10⁶) is injected or control conditions observed (Table I).

After endotrachial intubation and general anesthesia is achieved such as by administration of ketamine hydrochloride and Rompun®, the animal is placed in supine position. Lactated ringers are used at about (5 ml/kg/hr). The area of incision is shaved and prepped and draped in the usual sterile fashion with alternating betadine scrubs and alcohol wipes (> three times). Bland ophthalmic ointment is placed on the eyes. A left retroperitoneal approach is used to expose the right anterior aspect of the disc from L2-L5 (the rabbit has 6 to 7 lumbar vertebra). Various preparation schemes are used and treatment schema is applied to each disc level. For 'Needle Puncture' preparation of the disc, a 18-gauge needle is used to place a puncture in the disc at the depth of 5 mm (Aoki et al., “Nerve fiber ingrowth into scar tissue formed following nucleus pulposus extrusion in the rabbit anular-puncture disc degeneration model: effects of depth of puncture.” Spine.
2006;31(21):E774-80). After puncture, the test materials listed in Table I are injected. Treatment composition is applied to any one of L1-2, L2-3, L3-4, L4-5 region of each rabbit.

Monthly radiographs are used to monitor any disc changes. Animals are sacrificed at 2, 8, and 24 weeks after surgery.

Radiographs/MRI. Healing is indicated by a detectable radiographic change of increased disc height from same disc at baseline (pre op) compared to disc at other disc levels. Other discs are compared before and after needle puncture only, and disc before and after no needle puncture yielding an index of normal degeneration over time.

Retro-Transcription PCR. Retro-transcription PCR is performed to assay relative quantity of surviving transfected chondrocytes.

Histology. Also histology is used to confirm characterization of the collagen type I and type II and the gross appearance and evaluation of de novo chondrocytes.

Western Blot analysis and or ELISA. Quantitative expression of collagen type I and type II, and proteoglycan concentration, Smads 2/3, Sox-9. Additionally ELISA is used to evaluate TGFβ-1, BMP2, BMP7, GDF5 and other related growth factors where there are available antibodies.

Apoptosis is examined in the other tissue structures of the intervertebral disc via observing the expression of Capase-3.

EXAMPLE IV

Results

The results are as shown in the Figures and the description of the Figures of the present application. Punctured intervertebral disc treated with untransduced chondrocytes alone, transduced chondrocytes alone, primed chondrocyte alone or a mixture of transduced and untransduced chondrocytes, show beneficial effects in preventing or retarding disc degeneration compared with vehicle control.
Example IV-1 - Mixed-Cell (Transduced and Untransduced Chondrocytes)

Treatment of Punctured Intervertebral Disc in Rabbit

Mixed cell treatment has an intervertebral anti-degenerating effects when tested on rabbits. The effect is seen in a variety of experiments in FIGS. 1-4. FIGS. 1A-1F show a slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a rabbit spine four (4) weeks after surgery in which (i) the disc at L1/2 was injured and TGF-β1-producing chondrocytes were injected, (ii) no puncture and no treatment is seen at spine locus L2/3, and (iii) disc at L3/4 was injured and mixture of TGF-β1-producing chondrocytes and untransduced human chondrocytes in 1:3 ratio were injected; arrows point to L1/2 and L3/4 disc region. (C) shows MRI radiograph of a rabbit spine eight (8) weeks after surgery in which (i) the disc at L1/2 was injured and TGF-β1-producing chondrocytes were injected, (ii) no puncture and no treatment control at spine locus L2/3, and (iii) disc at L3/4 was injured and mixture of TGF-β1-producing chondrocytes and untransduced human chondrocytes in 1:3 ratio were injected; arrows point to L1/2 and L3/4 disc region. (D) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (E) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc. (F) shows X-ray radiograph of the rabbit described in (C) above, which is used to obtain a disc height index of the intervertebral disc.

FIGS. 2A-2F show a slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a rabbit spine four (4) weeks after surgery in which (i) the disc at L1/2 was injured and TGF-β1-producing chondrocytes were injected, (ii) no puncture and no treatment control at spine locus L2/3, and (iii) disc at L3/4 was injured and mixture of TGF-β1-producing chondrocytes and untransduced human chondrocytes in 1:3 ratio were injected; arrows point to L1/2 and L3/4 disc region.
L3/4 disc region. (C) shows MRI radiograph of a rabbit spine eight (8) weeks after surgery in which (i) the disc at L1/2 was injured and TGF-β1-producing chondrocytes were injected, (ii) no puncture and no treatment is seen at spine locus L2/3, and (iii) disc at L3/4 was injured and mixture of TGF-β1-producing chondrocytes and untransduced human chondrocytes in 1:3 ratio were injected; arrows point to L1/2 and L3/4 disc region. (D) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (E) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc. (F) shows X-ray radiograph of the rabbit described in (C) above, which is used to obtain a disc height index of the intervertebral disc.

[00104] FIGS. 3A-3D show a slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a rabbit spine four (4) weeks after surgery in which (i) the disc at L1/2 was injured and TGF-β1-producing chondrocytes were injected, (ii) no puncture and no treatment control at spine locus L2/3, and (iii) disc at L3/4 was injured and mixture of TGF-β1-producing chondrocytes and untransduced human chondrocytes in 1:3 ratio were injected; arrows point to L1/2 and L3/4 disc region. (C) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (D) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc.

[00105] Example IV-2 - Transduced Chondrocyte Treatment of Punctured Intervertebral Disc in Rabbit

[00106] TGF-β1-producing chondrocytes treatment has an intervertebral anti-degenerating effect. The effect is seen in FIGS. 4A-4D, which show a slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a rabbit spine four (4) weeks after surgery in which (i) the disc at
L1/2 was injured and mixture of TGF-β1-producing chondrocytes and untransduced human chondrocytes in 1:3 ratio were injected, (ii) no puncture and no treatment control at spine locus L2/3, and (iii) disc at L3/4 was injured and TGF-β1-producing chondrocytes were injected; arrows point to L1/2 and L3/4 disc regions. (C) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (D) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc.

Example IV-3 - Transduced Chondrocyte Treatment and Mixed-Cell Treatment of Punctured Intervertebral Disc in Rabbit

TGF-β1-producing chondrocytes treatment and mixed cell treatments have an intervertebral anti-degenerating effect. The effect is seen in FIGS. 5A-5D, which show a slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a rabbit spine four (4) weeks after surgery in which (i) the disc at L1/2 was injured and mixture of TGF-β1-producing chondrocytes and untransduced human chondrocytes in 1:3 ratio were injected, (ii) no puncture and no treatment control at spine locus L2/3, and (iii) disc at L3/4 was injured and TGF-β1-producing chondrocytes were injected; arrows point to L1/2 and L3/4 disc regions. (C) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (D) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc.

Example IV-4 - Untransduced Chondrocyte Treatment of Punctured Intervertebral Disc in Rabbit

Untransduced chondrocyte treatment has an intervertebral anti-degenerating effect. The effect is seen in a variety of experiments in FIGS. 6-8. FIGS. 6A-6D show a
slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a rabbit spine four (4) weeks after surgery in which (i) the disc at L1/2 was injured and cell culture media DMEM was injected, (ii) no puncture and no treatment control at spine locus L2/3, and (iii) disc at L3/4 was injured and untransduced chondrocytes were injected; arrows point to L1/2 and L3/4 disc regions. (C) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (D) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc.

FIGS. 7A-7F show a slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a rabbit spine four (4) weeks after surgery in which (i) the disc at L1/2 was injured and cell culture media DMEM was injected, (ii) no puncture and no treatment control at spine locus L2/3, and (iii) disc at L3/4 was injured and untransduced chondrocytes were injected; arrows point to L1/2 and L3/4 disc regions. (C) shows MRI radiograph of a rabbit spine eight (8) weeks after surgery in which (i) the disc at L1/2 was injured and cell culture media DMEM was injected, (ii) no puncture and no treatment control at spine locus L2/3, and (iii) disc at L3/4 was injured and untransduced chondrocytes were injected; arrows point to L1/2 and L3/4 disc regions. (D) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (E) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc. (F) shows X-ray radiograph of the rabbit described in (C) above, which is used to obtain a disc height index of the intervertebral disc.

FIGS. 8A-8F show a slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a
rabbit spine four (4) weeks after surgery in which (i) the disc at T12/L1 was injured by needle puncture and no injection, (ii) no puncture and no treatment control at spine locus L1/2, and (iii) disc at L2/3 was injured and untransduced chondrocytes were injected; arrows point to T12/L1 and L2/3 disc regions. (C) shows MRI radiograph of a rabbit spine eight (8) weeks after surgery in which (i) the disc at T12/L1 was injured by needle puncture and no injection, (ii) no puncture and no treatment control at spine locus L1/2, and (iii) disc at L2/3 was injured and untransduced chondrocytes were injected; arrows point to T12/L1 and L2/3 disc regions. (D) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (E) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc. (F) shows X-ray radiograph of the rabbit described in (C) above, which is used to obtain a disc height index of the intervertebral disc.

[00113] Example IV-5 - Untransduced Primed Chondrocyte Treatment of Punctured Intervertebral Disc in Rabbit

[00114] Primed chondrocyte treatment has an intervertebral anti-degenerating effect. The effect is seen in FIGS. 9A-9D, which show a slowing, retardation or prevention of degeneration of injured disc. (A) shows MRI radiograph of rabbit spine pre-surgery; (B) shows MRI radiograph of a rabbit spine eight (8) weeks after surgery in which (i) the disc at L2/3 was injured and cell culture media DMEM was injected, (ii) no puncture and no treatment control at spine locus L3/4, and (iii) disc at L4/5 was injured and primed chondrocytes were injected; arrows point to L2/3 and L4/5 disc regions. (C) shows X-ray radiograph of the rabbit described in (A) above, which is used to obtain a disc height index of the intervertebral disc to measure its morphology, its level of degeneration or regeneration. (D) shows X-ray radiograph of the rabbit described in (B) above, which is used to obtain a disc height index of the intervertebral disc.
EXAMPLE V

Source of Human Chondrocytes

Primary human chondrocytes were grown from cartilage tissue obtained from the surgical excision of a polydactyl finger from a one-year-old female human donor. The polydactyl tissue was harvested in a surgical room. The following procedure for chondrocyte isolation was performed in a biosafety cabinet. The plastic bottle containing the cartilage tissue was swiped with alcohol and the cartilage tissue was washed with sterile PBS (1X) using a pipette. A collagenase solution was prepared by dissolving 7 mg of collagenase (Gibco BRL) in 10 mL of DMEM (containing 10% FBS) and filtering through a 0.2 µm syringe filter (Corning). The washed cartilage tissue was treated with the collagenase solution for 17 to 18 hrs in a 37°C shaker incubator. On the following day, the bottle was sanitized with alcohol. The collagenase treated material was pipetted up and down several times to separate loose cells from the tissue mass. After pipetting, the supernatant was filtered through 70 µm nylon cell strainer (Falcon). Collagenase treated tissue which had lost its integrity (e.g., loose cells) was able to pass through the filter. The cell filtrate was collected in a 50 mL tube (Falcon) and then centrifuged at 1,500 rpm for 5 minutes. Two thirds of the supernatant was discarded and the pellet washed with 10 ml of sterile PBS (1X). The resuspended cells were again centrifuged at 1,500 rpm for 5 minutes and, after removal of two-thirds of the supernatant, washed with 10 ml of sterile PBS (1X). The cells were again centrifuged at 1,500 rpm for 5 minutes and then resuspended in DMEM (containing 10% FBS). The resuspended cells were then transferred to four uncoated 25 cm² flasks and cultured for four days at 37°C with 5% CO₂. The cells were then transferred into two uncoated 185 cm² flasks. The cells were cultured for two weeks and then collected, washed and resuspended in a cryopreservative media of DMEM, FBS and DMSO in a 5:4:1 ratio. The cells were aliquotted in to cryovials containing 1 mL of cell suspension at 4 x 10⁵ cells/mL. The cells were held in vapor phase liquid nitrogen storage.
What is claimed is:

1. A method for preventing or retarding degeneration of intervertebral disc at an intervertebral disc defect site comprising: injecting a mammalian connective tissue cell into the intervertebral disc defect site.

2. The method according to claim 1, wherein the connective tissue cell is allogeneic relative to the mammal.

3. The method according to claim 1, wherein the cell is a chondrocyte.

4. The method according to claim 3, wherein the chondrocyte is non-disc chondrocyte or juvenile chondrocyte.

5. The method according to claim 3, wherein the chondrocyte is primed chondrocyte.

6. The method according to claim 1, wherein the mammal is human.

7. A method for preventing or retarding degeneration of intervertebral disc at an intervertebral disc defect site of a mammal comprising:
   a) inserting a gene encoding a protein having intervertebral disc regenerating function into a mammalian cell, and
   b) transplanting the mammalian connective tissue cell into the intervertebral disc defect site.

8. The method according to claim 7, wherein said gene belongs to TGF-β superfamily.

9. The method according to claim 8, wherein said gene encodes TGF-β1.

10. The method according to claim 7, wherein the connective tissue cell is allogeneic relative to the mammal.
11. The method according to claim 7, wherein the connective tissue cell is a chondrocyte.

12. The method according to claim 7, wherein the mammal is human.

13. A method for preventing or retarding degeneration of intervertebral disc at an intervertebral disc defect site of a mammal comprising:
   a) inserting a gene encoding a protein having intervertebral disc regenerating function into a first mammalian connective tissue cell, and
   b) transplanting a mixture of the mammalian connective tissue cell of a) and unmodified second mammalian connective tissue cell into the intervertebral disc defect site.

14. The method according to claim 13, wherein said gene belongs to TGF-β superfamily.

15. The method according to claim 13, wherein said first and second mammalian connective tissue cell is chondrocyte.

16. The method according to claim 15, wherein the chondrocyte is non-disc chondrocyte or juvenile chondrocyte.

17. The method according to claim 13, wherein the chondrocyte for the second mammalian connective tissue is primed chondrocyte.

18. The method according to claim 13, wherein the first or second connective tissue cell is allogeneic relative to the mammal.

19. A method of treating degenerated or injured intervertebral disc in a patient comprising employing the method according to claim 1 to a subject in need thereof.

20. A method of treating degenerated or injured intervertebral disc in a patient comprising employing the method according to claim 7 to a subject in need thereof.
21. A method of treating degenerated or injured intervertebral disc in a patient comprising employing the method according to claim 13 to a subject in need thereof.
FIG. 3

C

D

L1
L2
L3

M (L2)

L1
L2
L3
L4
L5
L6
FIG. 4

C

D

L1

L2

L3

M (L2)

L1

L2

L3

L4

L5
FIG. 6

A

B

L1

L2

L3

L4

L5

L6
FIG. 7

A

B

C