Intervertebral Disc Degeneration and Low Back Pain: Molecular Mechanisms and Stem Cell Therapy

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Abstract

BACKGROUND: Low back pain (LBP) mostly caused by disc degeneration, reflects to a tremendous of health care system and economy. More knowledge about these underlying pathologies will improve the opportunities that may represent critical therapeutic targets.

CONTENT: Basic research is advancing the understanding of the pathogenesis and management of LBP at the molecular and genetic levels. Cytokines such as matrix metalloproteinases, phospholipase A2, nitric oxide, and tumor necrosis factor-α are thought to contribute to the development of LBP. Mesenchymal stem cells (MSCs) transplant to cartilage-like cells and secrete extracellular matrix and encourage nucleus pulposus (NP) cell activity inhibiting NP cell apoptosis, together with some chemical mediators such as cytokines and growth factors become a safe and effective new strategy for intervertebral disc degeneration (IDD) treatment and regeneration.

SUMMARY: IDD occurs where there is a loss of homeostatic balance with a predominantly catabolic metabolic profile. A basic understanding of the molecular changes occurring in the degenerating disc is important for practicing clinicians to help them to inform patients to alter lifestyle choices, identify beneficial or harmful supplements, or offer new biologic, genetic, or stem cell therapies.

KEYWORDS: low back pain, intervertebral disc, degeneration, nucleus pulposus, annulus fibrosus, extracellular matrix, genetic, stem cells


Introduction

The lumbar intervertebral discs (IVDs) play important roles for the support and mobility of spine.(1-3) These remarkable tissues is able to maintain stability under a large variety of loading conditions, while still permitting intersegmental motion.(1,4) Disc herniation and IVD degeneration (IDD) are two of the most common causes of low back pain (LBP) which is targeted for intervention.(5) Disc degeneration (DD) is a multifactorial process characterized by cellular and biochemical alternations in disc tissue which result in structural failure.(6) While DD is a part of normal aging, a significant number of people with indications of DD on magnetic resonance imaging (MRI) are actually asymptomatic, with no history of pain or disability.(7,8) The risk of back pain is increasing with the severity of DD.(9,10) Biological changes like proteoglycan and water loss is not really related with back pain, but back pain is more related to structural alternations, such as endplate defects and annulus height loss.(11-15) Most closely linked to pain are radial fissures in the annulus, whether or not they cause disc herniation.(14-17)

Another feature of discogenic back pain is the ingrowth of nerves and blood vessels.(18-20) Within some degenerated discs, nerves become sensitized by
inflammatory-like reactions, so that they can signal pain after minimal mechanical stimulation in animal experiments and in pain-provocation studies on humans.\(^{(21,22)}\) The different knowledge of pain-sensitization processes may explain why some degenerated discs are painful, whereas the others are not.\(^{(23)}\) More insights into the pathogenesis of DD may establish new paradigms for early or differential diagnostics of degeneration using new techniques such as systemic biomarkers. Research on the mechanobiology of disease also enriches the development of therapeutics for disc repair, with potential to reduce pain and disability associated with DD.\(^{(24)}\) More recently, some studies about the application of mesenchymal stem cells (MSC) from many source for example bone marrow, synovial membrane and adipose tissues showed a promising results for IDD.\(^{(25-27)}\)

### The Lumbar IVD, Structure and Functions

The healthy IVD is composed of some concentrically arranged layers of fibrocartilage which surround and restrain an amorphous, well-hydrated, inner core of proteoglycan gel.\(^{(1-3)}\) Strongly bound to the vertebral bodies and cartilaginous vertebral endplates, the composite make-up of the IVD creates a hydraulic system that can absorb and transmit various combinations of compression, shear, and tensile forces.\(^{(6,28,29)}\) The healthy disc creates a “spacer effect” which maintains sufficient vertical distance between the vertebrae (disc height) to provide ligamentous tension, alignment of the facet joints, and adequate space for the passage of neurovascular structures within the vertebral foramina.\(^{(1)}\) The IVDs lie between the vertebral bodies, linking them together (Figure 1). They are the main joints of the spinal column and occupy one-third of its height.

The foremost function of the IVD is mechanical: it transfers loads, dissipates energy and facilitates joint mobility. The IVDs are complex structures that consist of a thick outer ring of fibrous cartilage termed the annulus fibrosus (AF), which surrounds a more gelatinous core known as the nucleus pulposus (NP). The NP is sandwiched inferiorly and superiorly by cartilage endplates.\(^{(30)}\) The NP and the AF structures act synergistically to distribute and transmit loads between the vertebral bodies (Figure 2).\(^{(31,32)}\) The central NP containing collagen fibers and elastin fibers (Figure 3).\(^{(30)}\) The vertebral endplate containing hyaline cartilage bonded to the perforated cortical bone of the vertebral body and collagen fibers of the annulus and the nucleus (Figure 4).\(^{(30)}\) When the disc is compressed, hydrostatic pressure is generated within the NP, which is constrained peripherally by the AF, generating tensile circumferential stresses within the lamellar structure.\(^{(31,32)}\) Compressive loads are also supported directly by the inner AF, which is rich in proteoglycans.\(^{(33,34)}\) The angle- ply structure and nonlinear properties of the AF facilitate both joint mobility and stability in multiple modalities, including bending and rotation, and combinations thereof.\(^{(35-38)}\) They provide flexibility to this, allowing bending, flexion, and torsion.\(^{(39,40)}\)

Cells in the anulus are elongated parallel to the collagen fibers, rather like fibroblasts. Cells in the nucleus are initially notochordal but are gradually replaced during
childhood by rounded cells resembling the chondrocytes of articular cartilage. Anulus cells synthesize mostly collagen type I in response to deformation, whereas nucleus cells respond to hydrostatic pressure by synthesizing mostly proteoglycans and fine collagen type II fibrils. Cell density declines during growth (41), and in the adult is extremely low, especially in the nucleus (42,43). In adult discs, blood vessels are normally restricted to the outmost layers of the anulus. Metabolite transport is done by both diffusion, which is important for small molecules, and by bulk fluid flow, which is important for large molecules. (42,44) Low oxygen tension in the center of a disc could causes anaerobic metabolism, which results in a high concentration of lactic acid and low pH. (42) In vitro experiments indicate that a chronic lack of oxygen causes nucleus cells to become quiescent, meanwhile a chronic lack of glucose can kill them. (45) Deficiencies in metabolite transport are known to limit both the density and metabolic activity of disc cells. (42) As a result, discs have limited capability to recover from any metabolic or mechanical injury. During growth and the process of aging, it is normal for endplate permeability and also disc metabolite transport to decrease, but it increase in the presence of DD and following endplate damage. (46)

Degradative enzymes, such as matrix metalloproteinases (MMPs) and a disintegrin and metalloproteinase (ADAM), are produced by disc cells to synthesize their matrix and break down the existing matrix. (47-53) Molecular markers of matrix turnover are naturally found most abundant during growth, but usually decline thereafter. (54) The major structural alternations to the disc occur during fetal and juvenile growth, when the nucleus changes in consistency from a translucent fluid to a soft amorphous tissue, caused mainly by an increase in collagen content. (6,55) Collagen turnover time in articular cartilage is approximately 100 years (56) and could be even longer in the disc. Meanwhile the proteoglycan turnover is faster, possibly around 20 years (57), and some regeneration of NP is possible in young animals (58). Injuries that affect the inner anulus or endplate decompress the nucleus and healing processes are then overtaken by severe degenerative changes. (59,60)

### Molecular Mechanisms of IDD

Many evidence show that the aging of the disc is related to damage from oxidative stress. Oxidative stress is known to be a driver of cellular senescence and apoptosis. Higher levels of oxidized proteins and transcription factors activated by oxidative stresses have been found in older discs compared with young discs. (61) Presence of glycalation end products, which are molecules made by non-enzymatic glycosylation and oxidation of proteins and lipids, are another prove of of age-related oxidative damage in the disc. (62,63) The most common advanced glycalation end products...
in the disc are pentosidine and carboxymethyl-lysine. Pentosidine crosslinks collagen molecules and increases collagen stiffness as well as decreasing the synthesis of matrix proteins and proteoglycans. (62, 64) Additionally, notochordal cells, the cells that persist in the NP, which are of notochordal origin, are greatly affected by oxidative stressors and activate both intrinsic and extrinsic pathways of apoptosis. (65) Without aging, there is reduced catabolic activity of NP cells and decreased NP cell number. (66)

Structural disruption of the IVD is manifested by a loss of the hydrostatic capacity of the nucleus that occurs when its surrounding connective tissues cannot provide adequate restraint. (6) This may occur following an injury to, or disruption of, the vertebral endplate and/or annulus. (67, 68) Structural changes often initially occur in small and localized regions of the IVD. (69) Figure 5 shows the classification of internal disc disruption from grade 0 to grade 5, based on the Modified Dallas Classification.

Over time, tissue disruption can spread diffusely throughout the disc and cause a reduction in stiffness and loss of fluid pressure, also some various combinations of bulging, herniation, and decreased disc height. (6, 70, 71) The most significant biochemical change to occur in DD is loss of proteoglycan. (72) The aggrecan molecules become degraded, with smaller fragments being able to leach from the tissue more readily than larger portions. (30)

Decreasing aggrecan content in the NP leads to reduced hydration (73), leading in turn to impaired mechanical function (74, 75). A less hydrated, more fibrous NP is unable to evenly distribute compressive forces between the vertebral bodies. The forces are instead transferred non-uniformly to the surrounding AF (28), which can result in altered AF mechanical properties (76, 77) and progressive structural deterioration, including the formation of circumferential and radial tears (78). On occasion, radial tears can progress to a posterior radial bulge or herniation of NP material (78), resulting in painful symptoms. Decreased disc height is also commonly associated with advanced DD (12) and results in painful compression of surrounding structures.

The vertebral endplate is believed to play a critical role in the transport of nutrients into the IVD and in the removal of waste products. (42, 79, 80) One of the main cause of DD is thought to be failure of the nutrient supply to the disc cells. (81) Just like all cell types, the cells of the disc need nutrients, such as glucose and oxygen, to remain alive and active. The activity of disc cells is very sensitive to extracellular oxygen and pH in in vitro experiment, with matrix synthesis rates falling steeply at acidic pH and at low oxygen concentrations and the cells do not survive prolonged exposure to low pH or glucose concentrations. (45, 82, 83)

Decrease in nutrient supply that leads to a lowering of oxygen tension or of pH, which arising from raised lactic acid concentrations, thus could cause the ability of disc cells to synthesize and maintain the disc’s extracellular matrix and could ultimately lead to DD. Thus, endplate disruption can lead to impaired diffusion (46), disruption in nutrient supply (45), and/or cell death within the IVD, resulting from excessive tissue loading. (29, 84, 85)

Abnormal mechanical loads are also thought to provide a pathway to DD. For many decades, it was suggested that a major cause of back problems is injury, often work-related, that causes structural damage. It is believed that such an injury initiates a pathway that leads to DD and finally to clinical symptoms and back pain. (86) Although intense exercise does not appear to affect discs adversely (87) and discs are reported to respond to some long-term loading regimens by increasing proteoglycan content (88), experimental overloading (89) or injury to the disc (90, 91) can induce degenerative changes.

Figure 5. The classification of internal disc disruption from grade 0 to grade 5, based on the Modified Dallas Classification. (30) (Adapted with permission from John Wiley & Sons, Inc.)
Significant numbers of recent works assume that the factors that lead to DD may have important genetic components. Several studies have reported a strong familial predisposition for DD and herniation.(92-94) Findings from two different twin studies conducted during the past decade showed heritability exceeding 60%. (95,96) MRI in identical twins were very similar with respect to the spinal columns and the patterns of DD. (97) Genes associated with DD have been identified. Individuals with a polymorphism in the aggregan gene were found to be at risk for early DD. Studies of transgenic mice have demonstrated that mutations in structural matrix molecules such as aggregan (98), collagen II (99) and collagen IX (100) can lead to DD. Mutations in genes other than those of structural matrix macromolecules have also been associated with DD. (101-103)

There is increasing evidence supporting the role of the inflammatory cytokine interleukin (IL)-1 in the processes which leads to degeneration. (104-110) During this process, there is an increase in the production of the IL-1 agonists (IL-1α and IL-1β) and their active receptor IL-1 receptor I (IL-1RI), without a concordant increase in the natural inhibitor, IL-1 receptor antagonist (IL-1Ra) or the decoy receptor IL-1RII within the cells of the NP and inner AF. IL-1 induces the expression of a number of MMPs and ADAM with thrombospondin motifs (ADAMTS) family members (109,111), and reduces the expression of normal matrix genes. (109) Neurotrophic factor expression is modulated by IL-1 (110), and has been linked to induction of senescence in articular chondrocytes (112-113) and fibroblasts (114), all of which are features associated with IDD. (115-118)

The aforementioned evidence shows that many different influences are at work in old and DD, including genetic inheritance, impaired metabolite transport, altered levels of enzyme activity, cell senescence and death, changes in matrix macromolecules and water content, structural failure, and neurovascular ingrowth. (6)

LBP Pathophysiology

LBP is related to ageing, mechanical stresses (119) and genetic predisposition (120), and it is attributed to DD in around 40% of patients (121,122). It appears that alteration in biomechanical properties of the disc structure, sensitization of nerve endings by release of chemical mediators, and neurovascular ingrowth into the degenerated discs may all contribute to the development of pain. The loss of disc structure also alters the loading response and alignment of the rest of the spinal column, including that of the facet joints, ligaments, and paraspinal muscles, which eventually may become additional pain generators. (123)

The development of pain might be the result of the presence of macrophage and mast cells that propagates the inflammatory cascade. Macrophages increase the levels of multiple inflammatory mediators, especially IL-6 and IL-8, nitric oxide, tumor necrosis factor (TNF)-α and IL-1β. (124) The concentration levels of the said cytokines have correlated with pain intensity, and persistent activation of sensory fibers upregulates nitric oxide synthase, therewith increasing the level of nitric oxide, suggesting a possible positive feedback loop of pain generation. (125)

The onset of discogenic pain is characterized by nerve fiber ingrowth into an otherwise aneural tissue (Figure 6). (20,126-128) The interplay between inflammatory cytokines and neurotrophins, produced by disc cells and infiltrating immunocytes as well as neurotrophin receptors and their modulators may guide this process. Then neuronal tissue will develop after the development of vascularized granulation tissue. Degenerated disc cells secrete brain-derived growth factor (BDNF), which promotes neuronal development. (129) The release of proinflammatory cytokines IL-1β and TNF-α from the surrounding tissues also upregulates nerve growth factor (NGF) and expression of its receptors on the disc tissue. (130) Progressively small nerve fibers form along with the granulation tissue. (131) NGF promotes the collateral sprouting of additional peripheral sensory nerves into

Figure 6. A series of events occur during DD that are proposed to cause discogenic pain. (126) (Springer International Publishing AG). NP: nucleus pulposus; AF: annulus fibrosus; BDNF: brain-derived growth factor; NGF: nerve growth factor; FGF: fibroblast growth factor; TGF-β1: transforming growth factor beta 1; TNF-α: tumor necrosis factor alpha; IL: interleukin; NO: nitric oxide.
the inner AF and the NP, increases nerve survival, and increases the action and sensitivity of nociceptive sensory neurons.(18-20,130,132-135)

Of note, nerve fibers that innervate disc tissue are categorized as nociceptive and thought to be derived from the dorsal root ganglia. They express acetylcholinesterase, protein gene product (PGP) 9.5, substance P (SP), BDNF, transient receptor potential cation channel subfamily V member 1 (TrpV1), calcitonin gene related peptide (CGRP), and neurofilament protein (NFP).(136-140) These relationships suggest a direct linkage between inflammatory cytokines, neurotrophins and nociception.(141) Mechanical stimuli which are normally innocuous to disc nociceptors can, in certain circumstances, generate an amplified response which has been termed ‘peripheral sensitization’. This may explain why some degenerative discs are painful and others not. There is growing evidence that these pain receptors in painful disc are peripherally sensitized by the activity of sympathetic efferent which may initiate a pain impulse in response to ischemia, pressure changes or in amatory irritation.(142,243)

Knowing why nerves and blood vessels grow into AF may lead to effective strategies to hinder the process, or render it less painful. AF may reflects a microenvironment of low mechanical stress within a tissue which normally exhibits a fluid pressure many times greater than systolic blood pressure.(144,145) Normally, fluid pressure will extend from the nucleus into the inner and middle annulus (28), where it would be expected to collapse any blood vessel. Reduced pressure within a fissure is able to provide a route for ingrowth of blood vessels and accompanying nerves. In addition, proteoglycans can inhibit the growth of nerves (146) and blood vessels in vitro (147), and any loss of proteoglycans from within an annulus fissure may increase their attractiveness to ingrowing vessels.(23)

### Gene Therapy for IDD

The recent standard of care for LBP due to degenerative disc changes includes non-operative approaches, such as pain management, and operative approaches. The non-operative management aims primarily in symptomatic pain relief, while permitting possible endogenous recovery such as resolution of herniation (148,149) or the repairment of structural damage (150,151). The main target of non-operative LBP management is analgesia. It is accomplished by a combination of nonsteroidal anti-inflammatory drugs (NSAIDs) and physical therapy to strengthen core muscles among other programs. Surgical management may start with epidural injections of local anesthetic, steroids, or a combination of both prior to more invasive surgical approach.(24)

The current treatment options for IDD and the pathology associated with it are not the underlying pathophysiology. (152-155) With advances in molecular and cellular biology, researchers have start to characterize the pathophysiological pathways associated with DD and thus provided targets for potential biological treatments to augment or reverse the course of IDD.(156) Although the certain pathophysiology of DD still has not completely understood, however it is known to be affected by the interaction between various genetic, biologic and biomechanical factors.(95,157-159)

The hallmark of DD is the progress loss of proteoglycans which coincides with decreases in oxygen tension, free radial accumulation, decreased pH, and the increased activity of aberrant proteolytic enzymes.(73,160,161) With the loss of proteoglycans the NP cannot maintain normal physiologic hydrostatic pressure, resulting in in the dehydration of the NP.(162) There is also a progressive fibrosis of the NP as the ratio between type I to type II collagen increases.(163) The NP and AF lose their morphological diversity as the degeneration happens, which ultimately distracts the finely balance biomechanics of the disc and spine as whole.(162,164)

Numerous risk factors, such as age, abnormal physical loading, and genetics, may lead to the development of IDD (Figure 7).(165) The homeostasis of IVD tissues is biologically regulated by the active maintenance of a balance between the anabolism and catabolism of disc cells. This is achieved through a complex and precise coordination of a variety of substances, including cytokines, growth factors, enzymes and enzyme inhibitors, in a paracrine or/and autocrine fashion.(166,167) The latest therapeutic strategies for DD have included some efforts in upregulating the production of key matrix proteins (e.g., aggrecan), or downregulating the catabolic events induced by the pro-inflammatory cytokines, IL-1 and TNF-α.(109,168-174)

To deliver these therapeutic agents, some approaches such as protein injection and viral or non-viral gene transfer have been suggested and preclinically tested. (167,175-177) The most direct approach to regenerate or repair a degenerated IVD is by injecting anabolic factors. However, there are some issues that need to be consider, such as the half-life and solubility of the factors, the proper carrier, the presence of inhibitors and some other factors.
The stimulation of matrix synthesis by cytokines or growth factors changes IVD homeostasis by shifting cellular metabolism to its anabolic state. It was demonstrated that the rate of synthesis of proteoglycans by IVD cells increases several-fold following the addition of transforming growth factor (TGF)-α and epidermal growth factor (EGF). Insulin-like growth factor (IGF)-1 also stimulates IVD cell proliferation and matrix synthesis in vitro. Members of the bone morphogenic protein (BMP) family, osteogenic protein (OP)-1 and BMP-2, have both been found to enhance the propylene glycol metabolism of IVD cells. OP-1 strongly stimulates the production and formation of the extracellular matrix by rabbit IVD cells, as well as by human IVD cells in vitro. OP-1 was also found to be effective in the replenishment of a matrix rich in proteoglycans and collagens after depletion of the extracellular matrix following exposure of IVD cells to IL-1 or chondroitinase ABC. BMP-2 is known to facilitates the expression of the chondrogenic phenotype by human IVD cells, increases proteoglycan synthesis and up-regulates the expression of aggrecan, collagen type I, and collagen type II mRNA, compared to untreated control levels. Both recombinant human BMP (rhBMP)-2 and -12 increased human NP cell proteoglycan and collagen synthesis while having minimal effects on AF cells.

Another member of the BMP family, namely growth and differentiation factor-5 (GDF-5), was also found to stimulate propylene glycol and type II collagen expression in mouse IVD cells. Moreover, the recombinant human GDF-5 (rhGDF-5) enhances cell proliferation and matrix synthesis and accumulation by both bovine NP and AF cells. Some epidemiologic studies highlighted that DD may be caused to a large degree by hereditary factors with apparently a relatively minor effects of environmental and behavioral risk factors, which indicated that genetic factors might play an important role in the pathogenesis of IDD.

Recently, Mayer, et al., reviewed the literature and found that the genetic polymorphisms of 21 genes have been associated with IDD, including vitamin D receptor (VDR), GDF5, aggrecan, collagen types I, IX, and XI, fibronectin, hyaluronan and proteoglycan link protein 1 (HAPLN1), thrombospondin, cartilage intermediate layer protein (CILP), asporin, MMPs1, 2, and 3, parkinson protein 2, E3 ubiquitin protein ligase (PARK2), proteosome subunit β type 9 (PSMB9), tissue inhibitor of metalloproteinase (TIMP), cyclooxygenase-2 (COX2), and IL-1α, IL-1β, and IL-6. The idea of gene therapy originated as a means to repair the heritable genetic disorders by replacing defective genes with functional genes, which then able to cure the underlying disorder. The recent concept of gene therapy has broadened to include the transfer of exogenous genes encoding therapeutic proteins into cells to treat disease. Gene therapy changes host cell DNA, which then
and adipogenic lineages but not the hematopoietic lineage. MSCs are a heterogeneous population of multipotent cells capable to differentiate along the chondrogenic, osteogenic, adipogenic lineages but not the hematopoietic lineage. (197-199) Beside the dependence of sustained expression of the therapeutic gene, the success of gene therapy also lie on the efficiency of the genetic material transfer to the host cell. With very little exceptions naked plasmid DNA alone is not an effective means of gene transfer. Therefore the use of vectors is necessary to facilitate the transfer of genetic information to host cell. There are some types of vectors, which later classified into either viral or nonviral vectors (include liposomes, gene guns, DNA-ligand complexes, and microbubble enhanced ultrasound). Liposomes are phospholipid vesicles which deliver the genetic material into the cell by fusing with the host’s cellular membrane. Viral vectors use the natural ability of viruses to infect host cells and thus transfer the viral genetic information into the host. Viral vectors are very efficient at transducing the desired genetic material to the host cell, even into slowly dividing senescent cellular populations like those of the IVD. Viral vectors which is used for the gene therapy applications include adenovirus, adeno-associated virus, herpes simplex virus, lentivirus, retrovirus and also pox virus. Each viral vector is associated with specific advantages and disadvantages. Therefore proper selection of vector is critical to successful gene therapy. (156)

In addition to the selection of the appropriate gene and vector, another notable consideration with gene therapy applications is the delivery strategy utilized. There are currently two basic strategies for the delivery of exogenous therapeutic genes into target cells. The in vivo strategy involves the direct transfer of the gene-vector complex to the targeted cellular population within the living host. The ex vivo strategy differs significantly as the targeted cells are isolated and removed from the living host. These cells are then cultured with transduction of the therapeutic gene occurring in vitro. The final step includes the re-implantation of the genically altered cells back into the host. (156)

Many different sources of MSCs have been identified and studied, for example bone marrow, synovial membrane and adipose tissues. (25-27) Studies with MSC have been particularly promising. Co-culture of MSC with NP cells stimulates both NP cells proliferation and MSC differentiation toward the chondrogenic lineage. (200-203) Increased production of cytokines, particularly TGF-β favors these transformations. (203-205) The NP contains MSC that are similar to the MSC recovered from bone marrow (206), and studies in animal models of DD have shown that MSC injected in the NP area not only survive for months but also proliferate in canine (207,208), porcine (209), and rabbit models (210). Moreover, the transplanted MSC induced production of extracellular matrix proteins, including aggrecan and other proteoglycans, and types I and II collagens. (207,209,210)

A major limitation of using stem cells as a therapy for IDD is an appropriate delivery method that will not cause further injury to the IVD. The most direct route is an injection into the affected IVD, ensuring a localized therapeutic effect. However, in vivo studies suggest that needle injection into the IVD may cause further degeneration. (211-213) In fact, many studies use needle puncture as a model system to study DD in animals. (214-216) Recently, a population of stem cells isolated from human umbilical cord blood, multipotential stem cells (MPSCs), was reported to exhibit expanded multipotency with the ability to differentiate into cells of mesoderm, endoderm and ectoderm lineage. (217) Importantly, these cells were reported to home to sites of injury (218), and engraftment at the injured site following an intravenous injection of these cells. Contrasting to direct injection, intravenous injection neither improved the degeneration status, nor preserve disc height, however, both delivery methods increased glycosaminoglycan (GAG) protein and Acan gene expression relative to controls, suggesting possible paracrine effects. (219)

The mechanism for the inhibition of DD by MSCs most likely follows two aspects. First, MSCs can transplant to cartilage-like cells and secrete extracellular matrix. Second, MSCs can encourage NP cell activity and inhibit NP cell apoptosis. (205) Based on reported animal studies, a systematic review has showed that the use of MSCs for the treatment of DD is largely safe and effective. With the exception of 2 reports out of 24 controlled trials, no further complications were noted. According to previous studies in a rabbit model noted osteopyte formation anterolaterally to the disc space, which was attributed to leakage of the MSCs. (220,221) MSC treatment seems to have a more persistent

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**Stem Cell Therapy for IDD**

One of the available sources for cell-based repair of the disc that recently as been developed is MSC. (197-199) MSCs are a heterogeneous population of multipotent cells capable to differentiate along the chondrogenic, osteogenic, and adipogenic lineages but not the hematopoietic lineage.
and consistent quality of regenerative effect. In a clinical setting, injection of MSCs has the benefit of minimizing invasiveness of secondary surgery in comparison with installation of mechanical devices which also requires tertiary surgery to remove device after treatment. (216)

All the combined evidences support the application of bone marrow MSCs for regeneration of IVD and that long-term survival of injected cells in the hypoxic disc environment is feasible. In addition to MSC long-term survival in vivo, immediate and trophic effects are of great importance in supporting MSC differentiation into disc cells, contributing to immediate disc repairing. Therefore, future studies can also focus on the methods that support MSC differentiation as adjuvants. It should also be remembered that the trophic effects from MSCs injected into the IVD could potentially contribute to activate endogenous disc or stem cells to enhance the regenerative efficiency.

Extending the concept of stem cell therapy further, investigators have exploited the use of allogenic stem cells. This has the added advantage of off-the-shelf availability. Moreover, as the cause of DD is thought to be multifactorial, the use of allogenic stem cells could eliminate potential autogenic precipitating factors such as genetic predisposition (222-225), or the diminished potency of stem cells due to natural aging (197). In fact, IVD is suggested to be immune-privileged due to its avascular nature. A study, showing allogenic NP cell transplantation did not elicit lymphocyte infiltration, is consistent with this notion. (197) The problem of immune rejection is likely to be even less for allogenic MSCs, since MSCs are capable of escaping alloantigen recognition. (194, 197)

Adipose-tissue-derived stromal cell (ADSC) show potential for restoring degenerative discs and may prove effective in the treatment of IVD. The results of ADSC implantation studies in a DD model were promising, indicating that ADSCs could maintain their viability and proliferate within the rat IVD. (194)

Notochordal cells are the developmental origin of the NP. Yet they are not expressed in adult human IVD. Induced pluripotent stem cells (iPSCs) have demonstrated their ability to differentiate into various cell types. In IVD applications, mouse and human iPSCs have been shown to differentiate into NP-like cells expressing notochordal markers and assumed the possibility that they may be used as a novel cell source for cellular therapy. (200) Notochordal cells have been observed to substantially stimulate biosynthetic activity of NP cells through factors secreted into conditioned medium. (200) These findings support the notion that molecular agents secreted by notochordal cells constitute a promising alternative for disc repair. (24) Results of stem cell studies in IVD are developing and, if delivery obstacles can be overcome, may offer alternative future treatment strategies.

**Conclusion**

DD progresses with age and involves a shift in the metabolic productivity of the IVD. The degenerative and inflammatory changes occurring as the disc degenerates promote increased neural and vascular ingrowth into the disc, potentially accounting for the painful discomfort patients experience with DD. Treatments which utilize inherent growth potential, through growth factors or stem cells, can stimulate tissue repair but may also provide advantages by mitigating inflammation. By knowing the mechanism of IVD contributes an essential piece of the repair puzzle, lead to an optimum integrated management of LBP for new and refined concepts in pathophysiology, earlier detection of disease, and improved developments in tissue engineering for treatment.

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