



Deposited via The University of Sheffield.

White Rose Research Online URL for this paper:

<https://eprints.whiterose.ac.uk/id/eprint/205605/>

Version: Published Version

Article:

Lama, P., Tiwari, J., Mutreja, P. et al. (2023) Cell clusters in intervertebral disc degeneration: an attempted repair mechanism aborted via apoptosis. *Anatomy & Cell Biology*, 56 (3). pp. 382-393. ISSN: 2093-3665

<https://doi.org/10.5115/acb.23.067>

Reuse

This article is distributed under the terms of the Creative Commons Attribution-NonCommercial (CC BY-NC) licence. This licence allows you to remix, tweak, and build upon this work non-commercially, and any new works must also acknowledge the authors and be non-commercial. You don't have to license any derivative works on the same terms. More information and the full terms of the licence here:

<https://creativecommons.org/licenses/>

Takedown

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.



Cell clusters in intervertebral disc degeneration: an attempted repair mechanism aborted via apoptosis

Polly Lama¹, Jerina Tiwari¹, Pulkit Mutreja¹, Sukirti Chauhan¹, Ian J Harding², Trish Dolan², Michael A Adams², Christine Le Maitre³

¹Department of Anatomy, Sikkim Manipal Institute of Medical Sciences, Sikkim Manipal University, Sikkim, India, ²Centre for Clinical Anatomy, University of Bristol, Bristol, ³Biomolecular Research Centre, Sheffield Hallam University, Sheffield, UK

Abstract: Cell clusters are a histological hallmark feature of intervertebral disc degeneration. Clusters arise from cell proliferation, are associated with replicative senescence, and remain metabolically, but their precise role in various stages of disc degeneration remain obscure. The aim of this study was therefore to investigate small, medium, and large size cell-clusters. For this purpose, human disc samples were collected from 55 subjects, aged 37–72 years, 21 patients had disc herniation, 10 had degenerated non-herniated discs, and 9 had degenerative scoliosis with spinal curvature <45°. 15 non-degenerated control discs were from cadavers. Clusters and matrix changes were investigated with histology, immunohistochemistry, and Sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE). Data obtained were analyzed with spearman rank correlation and ANOVA. Results revealed, small and medium-sized clusters were positive for cell proliferation markers Ki-67 and proliferating cell nuclear antigen (PCNA) in control and slightly degenerated human discs, while large cell clusters were typically more abundant in severely degenerated and herniated discs. Large clusters associated with matrix fissures, proteoglycan loss, matrix metalloproteinase-1 (MMP-1), and Caspase-3. Spatial association findings were reconfirmed with SDS-PAGE that showed presence to these target markers based on its molecular weight. Controls, slightly degenerated discs showed smaller clusters, less proteoglycan loss, MMP-1, and Caspase-3. In conclusion, cell clusters in the early stages of degeneration could be indicative of repair, however sustained loading increases large cell clusters especially around microscopic fissures that accelerates inflammatory catabolism and alters cellular metabolism, thus attempted repair process initiated by cell clusters fails and is aborted at least in part via apoptosis.

Key words: Intervertebral disc, Intervertebral disc degeneration, Disc herniation, Cell clusters, Apoptosis

Received March 8, 2023; 1st Revised April 5, 2023; 2nd Revised April 24, 2023; 3rd Revised April 28, 2023; 4th Revised May 18, 2023; Accepted May 22, 2023

Introduction

Back pain is the most common, chronic, and discomforting condition associated with aging and consequent progressive joint dysfunction [1, 2]. Back pain arises from functionally impaired intervertebral discs [3], which are load-bearing viscocartilaginous tissue lying between the consecutive vertebral bodies allowing moderate spinal movements [4]. Anatomically, each intervertebral disc has three integrated

Corresponding author:

Polly Lama 
Department of Anatomy, Sikkim Manipal Institute of Medical Sciences,
Sikkim Manipal University, Sikkim 737102, India
E-mail: pollylama@smims.smu.edu.in

Copyright © 2023. Anatomy & Cell Biology

This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (<http://creativecommons.org/licenses/by-nc/4.0/>) which permits unrestricted non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited.

parts: the centrally placed gelatinous nucleus pulposus (NP), the laminated annulus fibrosus (AF), and the hyaline cartilage endplate regions [5-7]. The NP, IAF and outer annulus fibrosus (OAF) regions contain two distinct cell types which are described as rounded cells of the nucleus pulposus and the inner annulus regions, and the highly flattened cells of the outer annulus [8, 9]. Recent and past evidence suggests that the nucleus pulposus cells proliferate and may have progenitor capabilities with tissue repair functions [10]. However, cell clusters are also a conventional hallmark of disc degeneration and catabolism, as the disc cells, particularly those in clusters produce cytokines that can trigger catabolic activity without the involvement of immune cells [11, 12]. Cell clusters undergo stress induced premature replicative senescence involving a brief phase of proliferation followed by growth arrest while remaining metabolically active during disc degeneration [13, 14]. Therefore, the activity of nucleus pulposus and the inner annulus cell phenotypes is uncertain, and a clearer distinction between proliferating reparative disc cells, non-proliferating, or proliferating degenerative cells still lacks, and a thorough investigations could assist in development of strategies that can promote endogenous cell-based repair techniques.

In young and mature non-degenerated intervertebral discs, nucleus pulposus cells appear singly, in pairs, or in small cell clusters, whereas with age and degeneration single cells or small cell cluster often progress to form medium and then large sized clusters [15] which is considered as the most common histological sign of disc degeneration. These large cell clusters are confined to the nucleus pulposus and inner annulus regions [9], and may contain as many as 60–100 individual cells in a single large sized cluster [8] presumed to arise from proliferation [9, 16], although a few studies suggest that cell migration may be another process involved with formation of clusters [17, 18], furthermore, proliferating or migrating cells, are influenced by cytokines and chemokines overexpressed by the cell themselves during degeneration to initiate catabolism [19].

Animal models of osteoarthritis shows, chondrocyte cell proliferation is also associated with increased proteoglycan synthesis representing a repair response to tissue damage [20, 21]. Similarly, cyclic mechanical compression increases proteoglycan and collagen production by the nucleus pulposus cells [20] allowing cell clustering [22, 23]. There is also evidence that disc cell proliferation is initiated to facilitate repair due to stimulation by diffusible growth factors [24], which

is in contrast to stimulation through neuropeptide such as Substance-P that accelerates the release of cytokines [25, 26].

Spatial associations studies between disc cell clusters and histological degenerative features indicates [27] that highly crosslinked disc matrix is beyond the repair capacities by the cells, as the main anabolic mediator produced by these cells are proteoglycan molecules and fibrillar collagen which are structurally weak. This problem could be exacerbated by the relative avascularity, accelerated nutritional depletion, and accumulation of toxic metabolites such as lactate [28, 29] promoting premature replicative senescence and apoptosis, of the otherwise viable cells. Large clusters of senescent cells can also possibly synthesize sufficient proteolytic enzymes to cause focal matrix breakdown and allow fissure formation [11].

Cells of the nucleus pulposus, in contrast, has also been shown to form colonies ex-vivo and are immunopositive to stem and progenitor cell markers such as GD 2, a disialoganglioside, identified as a marker for bone marrow and umbilical cord mesenchymal stem cells. Tie-2, a tyrosine kinase receptor is localised in the disc cell membrane, that forms ligand with angiopoietin-1 to maintain the progenitor potential of the cell and protects it from apoptosis [30, 31]. Such progenitor nucleus pulposus cells differentiate into adipocytes, osteocytes, and chondrocytes in cell culture [32], and can be extracted from the intact discs [33]. This afore-said evidence indicate that phenotypic heterogeneity exists within the nucleus pulposus cells, which may differ during early, intermediate, and advanced stages of disc degeneration. These undefined parameters may influence the ontogeny and function of the nucleus pulposus cells. Therefore, cell proliferation markers such as proliferating cell nuclear antigen (PCNA) and Ki-67 can identify proliferating cells clusters and define the differences in cluster sizes seen during the various grades of disc degeneration. Thus, we objectively performed an immunohistochemical and protein quantification sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE) investigation to analyse such differences.

Materials and Methods

Human tissue

Ethical clearances were obtained from National Research Ethics Service (NRES17/SW/0112), Frenchey Hospital, North Bristol NHS Trust, UK, and Sikkim Manipal Institutional

Ethics committee (IEC/SMIMS/2018/174-24), Ministry of Health and Research (F.N.P.L.19015/53/97-IH) Govt., of India, prior to the human tissue sample collection process. Experimental protocol/methods were approved by Institutional Research Committee of the University of Bristol, UK and Sikkim Manipal University, India. The method described is in accordance with the guidelines mentioned by the World Medical Association, Declaration of Helsinki for medical research. Written informed consents were obtained from the patients for collecting the waste disc tissue. Fifty-five human disc tissues were collected from patients aged 37–72 years undergoing spinal surgeries. 19 patients had slightly degenerated non-herniated discs, or degenerative scoliosis disc with $<45^\circ$ deformity; 21 patients had severe degenerated discs with herniation; and 15 were non-degenerated cadaveric control discs. An anonymous clinical data sheet was obtained for each patient, and magnetic resonance imaging (MRI) scans were graded 1–5 as according to the clinically relevant Pfirrmann (pf) grade of disc degeneration [34]. Data are summarized in Table 1.

Semi-quantitative histological analysis of human discs

Each disc was divided into two halves in the sagittal plane. One-half was snap-frozen in isobutane, embedded in optimal cutting tissue medium (OCT), five-micron thin sequential sections were obtained on a Leica S-320 cryostat, and fixed in 10% neutral buffered formalin. Ehrlich's Haematoxylin & Eosin stain was used for the assessment of cells and the matrix. Consecutive sections were stained with toluidine blue to assess proteoglycans loss [35]. Outer annulus, inner annulus, and nucleus pulposus tissue was distinguished from each

other following a published protocol [8]. Using the modified criteria of Boos et al. [36] histological features were graded on an ordinal scales: cell clustering (0–5), inflammatory cells (0–3), proteoglycan staining score (0–3), and presence of tears/fissures in the matrix (0–3). Inflammatory cells were identified by observing its characteristic shape, organization, and localization around blood vessels. In each case, '0' and '3' (or 5) were referred to as 'absence' or 'abundance' of the scored variables as analyzed with Table 2. Scores were averaged across three fields of view, and then across specimen groups.

Quantitative immunohistochemical analysis

Five-micron thin sections were fixed in acetone for 10 minutes at -20°C , washed in phosphate buffer saline (PBS) and all non-specific binding sites were blocked by application of 5% of rabbit or donkey serum at 4°C for one hour (1:5 dilutions in PBS). Sections were washed in PBS and applied with monoclonal primary antibodies to matrix metallopro-

Table 1. Summary data for human discs used in this study

Disc details	Non-degenerated cadaveric 'control' discs (n=15)	Slightly degenerated disc (n=19)	Severely degenerated and herniated disc (n=21)
Age (yr)	Mean 53 (range 42–72)	Mean 53 (range 35–74)	Mean 55 (range 37–75)
Spinal level (n)	L2–3 (4), L3–4 (3), L4–5 (4), L5–S1 (4)	L2–3 (1), L3–4 (2), L4–5 (4), L5–S1 (12)	T12–L1 (2), L1–2 (3), L2–3 (2), L3–4 (5), L4–5 (2), L5–S1 (7)
Sex	M (6), F (9)	M (6), F (13)	M (9), F (12)
pf grade	Mean 3.0 (range 2–4)	Mean 3.6 (range 3–4)	Mean 3.4 (range 3–4)
Pain (mo)	No pain	Mean 14 (range 2–60)	Mean 17 (range 2–60)

M, male; F, female; pf, Pfirrmann.

Table 2. Variables for histological assessment

Variables for histological assessment	Grades/scores
Neovascularity/blood vessels	0=absent
Newly formed blood vessels with distinct endothelial cell lining located with inflammatory cells	1=rarely present 2=present in intermediate amount 3=abundant
Tears/fissures	0=absent
Splitting of collagen fibre bundles in the annulus fibrosus and nucleus regions creating a gap or break within the tissue	1=rarely present 2=present in intermediate amount 3=abundant
Proteoglycan staining	0=absent
Oval or irregular areas with variations in staining intensity for sulphated glycosaminoglycans/proteoglycans (glycosaminoglycans rich matrix usually appears dark blue to purple with toluidine blue staining)	1=rarely present 2=present in intermediate amount 3=abundant
Inflammatory cells	0=absent
Immunocytes with small irregular nuclei, often in swarms along the edges of/surfaces of disc tissue	1=rarely present 2=present in intermediate amount 3=abundant
Cell clustering	0=no cells/absent
Multiple rounded disc cells in small groups or clusters of small, medium, and large sizes sharply demarcated by a rim of territorial matrix	1=single cells only 2=paired cells only 3=small clusters with 3–7 cells 4=medium clusters with 8–15 cells 5=large clusters with >15 cells

Based on modified Boos et al. [36].

teinas-1 (MMP-1) (Ab52631; Abcam), Caspase-3 (31A1067; Novus Biologicals), Ki-67 (Novus biologicals, MK167-8D5), and PCNA (PC10; Novus biologicals) at 1:40 dilutions in PBS, and PBS-immunoglobulin G (IgG) alone as negative controls. Sections were incubated overnight to allow maximum interaction between the antigen of interest and primary antibodies. The next day, sections were washed in PBS and incubated with appropriate Alexa fluorophore secondary antibody at 1:100 dilutions in PBS for one hour, after PBS washes auto-fluorescence was quenched using 0.1% Sudan black B for 12 minutes. Sections were counterstained with Vecta Shield Dapi (H-1200; Vector), and secured with a cover slip [37].

Quantitative analysis was done with Leica-Olympus DP72 (Olympus) camera (12.8 megapixels) with a fixed frame view of 650 μm $\times$ 500 μm . Positively stained cells in clusters for MMP-1, Caspase-3, Ki-67, and PCNA were counted across the entire section at a magnification of 20 $\times$, with an average of 8 fixed frames per section. Ten 5- μm sections were analyzed for each disc sample. Immuno-positive images were counted with the cell sense software package.

Protein extraction and SDS-PAGE analysis

The remaining half of each human disc was rinsed in phosphate buffered saline, finely diced and proteins were extracted at 4°C under continuous agitation for 72 hours using 15 volumes of 4 M guanidine hydrochloride (Sigma Aldrich), 50 mM sodium acetate and 10 mM EDTA (Sigma Aldrich). The solution was adjusted to 5.8 pH, and protease inhibitor was added before introducing the diced disc samples into the extraction buffer. The extract was separated from the tissue by centrifugation in 13,000 g, and aliquots of 8 μl were prepared for SDS-PAGE. This solution was digested with Keratanase (Ambsio LLC) at 1 mU/25 μl for 6 hours, further treated with 100 mM Tris-HCL, pH 7.4 and digested overnight with chondroitinase ABC (Ambsio LLC) at 50 mU/25 μl of the extract.

SDS-PAGE

Lamelli buffer containing 2 $\times$ sodium dodecyl sulfate, 2 mercaptoethanol, and milliQ double filtered water was added to the sample and heated in a water bath at 95°C for 10 minutes before being fractionated on 10% polyacrylamide gels at 94 V (constant voltage) for 1 hour 40 minutes. Proteins of interest were transferred to nitrocellulose membranes (0.45 μm) using transfer buffer with 20% methanol at 100 V (con-

stant voltage) for 1 hour. Membrane was blocked with 5% skim milk powder in 0.01M Tris-HCL, 0.15 M NaCl, 0.1% Tween20 (TBST), pH 7.6. Monoclonal antibodies to MMP-1, PCNA, Ki-67, and Caspase-3 diluted in 1: 1,000 TBST with 1% BSA, were applied to the blots for 2 hours, followed by TBST washes. Appropriate secondary conjugated antibody to horseradish peroxidase (Thermo Fisher Scientific) was applied at 1:1,000 dilutions, followed by TBST washes. Protein bands were developed by chemiluminescence with an ECL mixture (GE Healthcare). Blots were also analyzed omitting primary antibody to check for IgG species in the tissue extracts.

Quantification of western blot bands with Image J

Protein blot images were saved at 600 dpi in gray scale, each protein lane (row-wise) was then measured by selecting a single region of interest with a rectangle tool from the Image J measurement tool bar (<https://imagej.nih.gov/ij/download.html>) to define a frame around the largest band of that row. The ratio of each protein band to the loading control was analysed by inverting the pixel density of the band as 255-X, where X was the value recorded for each protein band and the loading control.

Statistical analysis

Spearman's rank correlation was used to examine associations between histological variables. One and two-way ANOVA was used to compare mean values of cell clusters within the NP, IAF, and OAF regions, and to compare cell counts of positively stained Ki-67, PCNA, MMP-1, and Caspase-3 in the patient and control groups. Statistical tests were performed using SPSS software version 19 package (IBM Co.). $P < 0.05$ was considered as statistically significant.

Results

Frequency of cell clustering in NP, IAF, and OAF regions

The OAF region of the control discs comprised of parallel and crimped collagen fibers along with elongated and extremely flattened fibroblast-like cells, with almost indistinct cytoplasm (Fig. 1A). Cell clusters were always absent in the outer annulus of the control tissue but were a prominent feature of severely degenerated and herniated discs with structural defects, inflammatory cells, and disrupted collagen lamellae (Figs. 1B, C, 2B, D). Infiltrating inflammatory

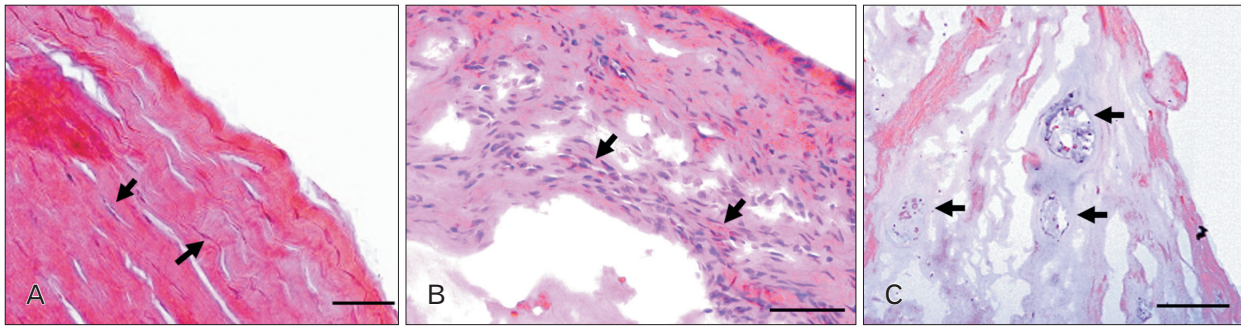


Fig. 1. Comparisons between (A) control non-degenerated cadaveric human disc with extremely flattened annulus fibrosus (AF) cells (arrows) and (B) herniated disc, arrows shows inflammatory cells, and (C) degenerated disc with displaced lamellae and large cell clusters (arrows). Scale bar: 100 µm.

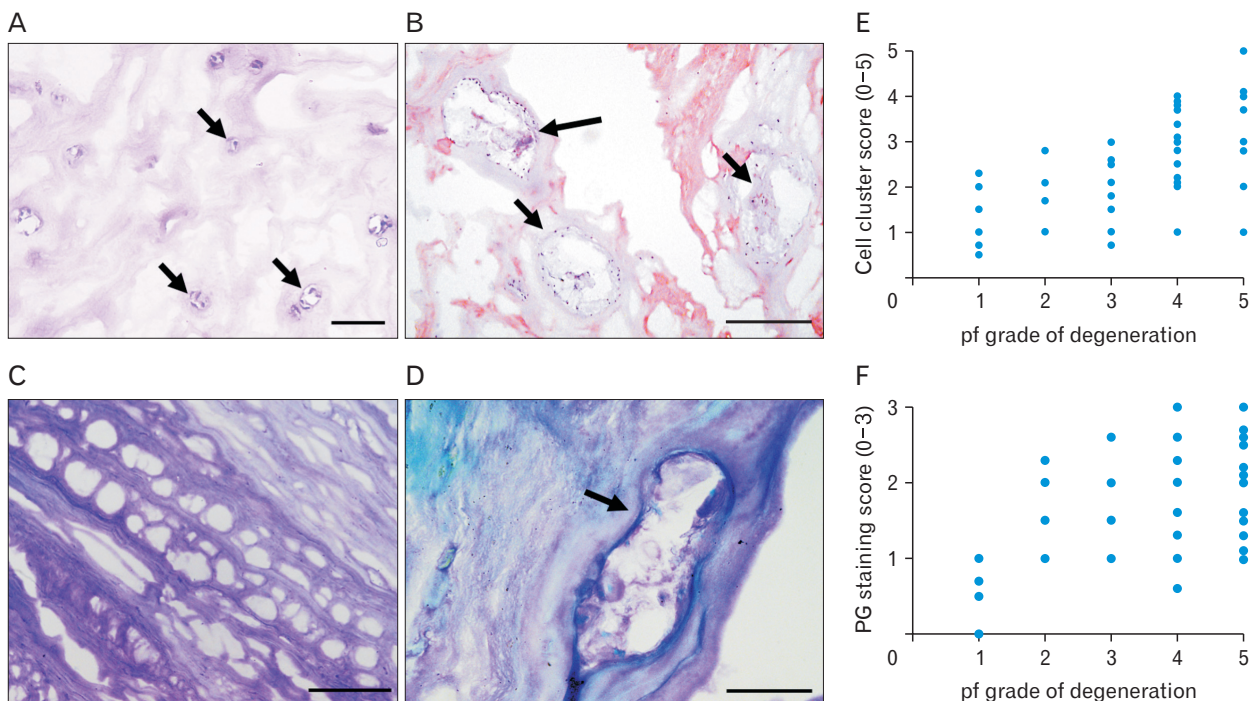


Fig. 2. Cell clusters comparisons between (A) control non-degenerated human cadaveric disc with small size cell clusters (2, 3, 6 cells) and (B) herniated disc with large sized cell clusters (50–60 cells, arrow). Scale bar: 100 µm (A) and 50 µm (B). (C) Toluidine blue stained control NP/IAF region without clusters and (D) herniated disc tissue with large cluster (arrow) stained with toluidine blue. Scale bar: 50 µm. (E) Cell cluster scores (0–5) and increasing pf grade of disc degeneration. (F) Proteoglycan staining score (0–3) and increasing pf grade of disc degeneration. NP, nucleus pulposus; IAF, inner annulus fibrosus; pf, Pfirrmann grade of degeneration; PG, proteoglycans.

cells and blood vessels were localized around the vicinity of fissured areas in herniated mixture of nucleus and annulus tissue (Fig. 1B). Nucleus pulposus and inner annulus regions showed single cells, cells in pairs, in small size clusters (3–7 cells), medium size clusters (8–15 cells) in non-degenerated control cadaveric disc (Fig. 2A, C). Large-sized clusters appeared in a meshwork of loosely organized collagen fibres without any crimp, and these large clusters (>15 cells) were a

feature of herniated, severely degenerated discs, localised in the NP and IAF junction (Fig. 2B, D). Clustering cells identified through histological scoring increased with pf grade of disc degeneration (Fig. 2F) in the examined tissue sections ($r_s=0.316$, $P<0.04$).

Toluidine blue staining to assess proteoglycans loss

Proteoglycan staining score was greatest in control discs,

the NP and IAF regions, and decreased with severity of degeneration, Table 3. Staining intensity reduced around invading inflammatory cells, large clusters, especially in pf grade 4, 5 degenerated discs (Fig. 2D, F). Proteoglycans loss scores correlated with the presence of fissures in the OAF ($r_s=0.85$, $P<0.001$), IAF ($r_s=0.53$, $P<0.001$) and NP regions ($r_s=0.74$, $P<0.001$). Large clusters showed little or no purple-stained proteoglycan matrix, and often exhibited nuclear shrinkage or 'pyknosis' (Fig. 2B), a characteristic feature of cells that had lost its ability to divide. Herniated discs showed greatest proteoglycans loss in comparison to moderately degenerated and control discs (Fig. 2C, D, Table 3).

Immunohistochemical localization and quantification of MMP-1, Ki-67, PCNA, and Caspase-3

Patient discs showed selective cells in clusters that were positive to MMP-1, Caspase-3, Ki-67, and PCNA, Table 3.

MMP-1 positive cells were selectively stained in clusters (Figs. 3A, 4B) and localized around the vicinity of disrupted tissue region in the NP-IAF junction as determined by phase-contrast (Fig. 4A, B) in severely degenerated and herniated discs. MMP-1 was also found to be localized around infiltrating inflammatory cells (Fig. 4B). Control discs showed only a few MMP-1 positive cells (Fig. 3A).

Caspase-3 positive clusters, of all sizes, increased with pf grade of degeneration, were most common in herniated and degenerated discs (Figs. 3B, 4A), frequently located in disrupted inner annulus region (Fig. 4C). Very few clusters (4%)

were positive in control discs.

Ki-67 positive clusters were a feature of moderately degenerated discs and controls. Herniated and severely degenerated discs showed fewer Ki-67 and PCNA positive cells (Fig. 3C). Ki-67 and PCNA positive clusters did not increase with pf grade of disc degeneration.

Ki-67 staining was visualized as non-nuclear, which is in contrast to the general acceptance that Ki-67 and PCNA proliferative markers must always be a nuclear stain, however, recent evidence indicates non-nuclear cytoplasmic staining of Ki-67 and PCNA is common in mature neutrophils [38], breast carcinoma cells [39], and hyalinizing trabecular adenomas of the thyroid and sclerosing haemangiomas of the lung [40]. Such non-nuclear localization can be explained as cells in discs tissue are predominantly non-mitotic and don't proliferate, but during degeneration to compensate for proteoglycans loss, cells begins to form clusters of various sizes and relatively non-dividing cells proliferate to as many as 20–100 cells within a large clusters which can translocate Ki-67 and PCNA to subcellular, cytoplasmic, and membrane bound locations either due to point mutation or deletions of the nuclear export signals resulting in molecular remodeling of its cellular and subcellular components [41].

All patient discs additionally also contained fewer numbers of single, paired cells, and flattened annulus cells, that were positive to MMP-1, Caspase-3, Ki-67, and PCNA.

Table 3. Summary of results for human discs, comparing three tissue regions NP, IAF, and OAF in three groups

Variable for assessment	Non-degenerated cadaveric 'control' discs (n=15)			Slightly degenerated discs (n=11)			Severely degenerated and herniated discs (n=21)		
	NP (n=15)	IAF (n=15)	OAF (n=15)	NP (n=8)	IAF (n=11)	OAF (n=8)	NP (n=17)	IAF (n=21)	OAF (n=12)
Tears/fissures (scale 0–3)	0.7±0.5	1.1±0.3	0.9±0.4	1.5±1.0*	1.5±0.7*	1.3±0.9*	1.8±1.0*	2.2±1.5*	2.2±0.7*
PG score (scale 0–3)	1.0±0.6	1.3±0.5	0.7±0.5	0.9±0.7**	1.8±0.9**	1.0±0.4**	1.4±0.8	2.1±0.4	1.8±1.0
Cell clusters (scale 0–5)	1.1±0.3**	1.2±0.9**	0**	1.2±1.0**	2.4±1.2**	0.2±0.3**	1.4±0.8**	2.5±1.0**	0**
Inflammation and blood vessels (scale 0–3)	0	0	0	0**	0.0±0.1**	0.4±0.9**	0*	1.0±0.9**	1.4±1.2**
MMP-1 cell count/mm ²	6.2±5.9	13.5±9.7	8.2±9.4	35.7±49.0	55.4±47.3	20.9±37.5	41.7±52.4**	99.8±71.5**	12.1±11.1
Ki-67 cell count/mm ²	56.8±21.1	63.14±24.1	38.21±17.7	13.0±15.3	35.5±21.9*	27.4±10.8*	21.1±23.7	33.3±26.7*	15.8±19.5*
PCNA cell count/mm ²	8.0±8.43	11.1±11.0	3.7±3.8	26.4±22.0	58.6±29.0*	17.0±22.6*	31.1±28.5**	84.4±79.1**	17.3±11.1**
Caspase-3 cell count/mm ²	8.4±5.9	10.0±6.3	7.1±5.0	61.9±48.2	68.5±32.7	12.2±11.9	29.5±30.3**	67.2±45.2**	45.1±15.1**

Values refer to the mean±SD.

NP, nucleus pulposus; IAF, inner annulus fibrosus; OAF, outer annulus fibrosus; PG, proteoglycan; MMP-1, matrix metalloproteinase-1; PCNA, proliferating cell nuclear antigen.

'n' values indicate total number and reflects the absence of some tissue types for the surgically removed patient disc samples.

Significant differences between three tissue regions NP, IAF, and OAF in non-degenerated cadaveric control, slightly degenerated and severely degenerated and herniated discs are shown with significance levels denoted. * $P<0.05$, ** $P<0.01$.

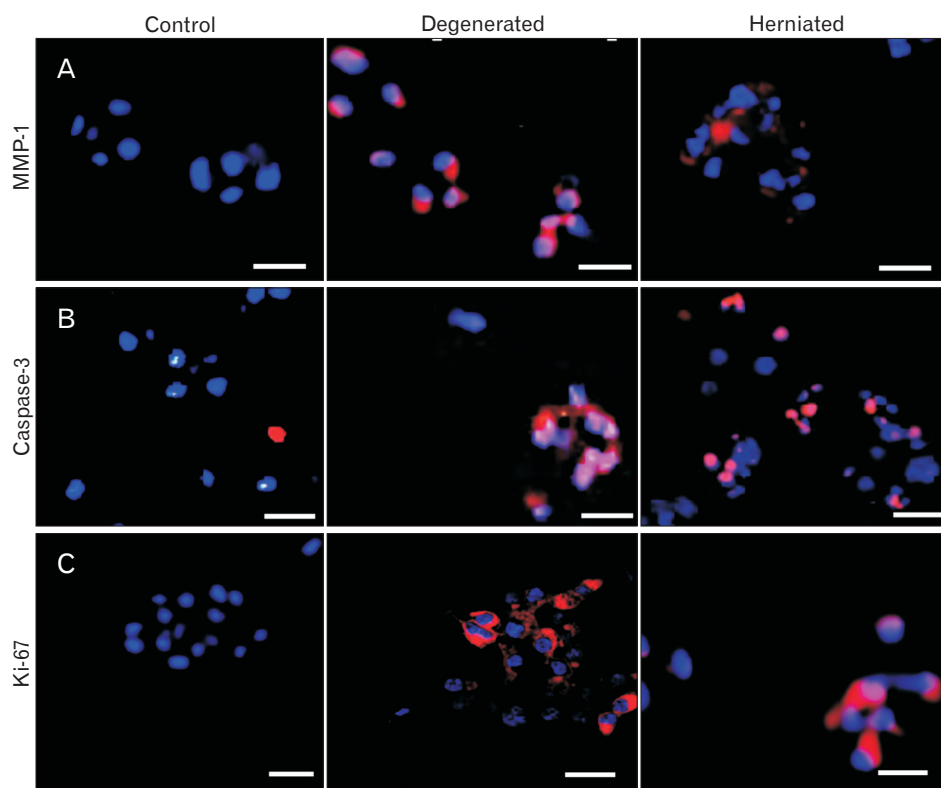


Fig. 3. Small, medium, and large sized cell clusters of the 'control' and its comparisons to 'degenerated' and 'herniated' discs for (A) MMP-1, (B) Caspase-3, and (C) Ki-67. Scale bar: 50 μ m. MMP-1, matrix metalloproteinase-1.

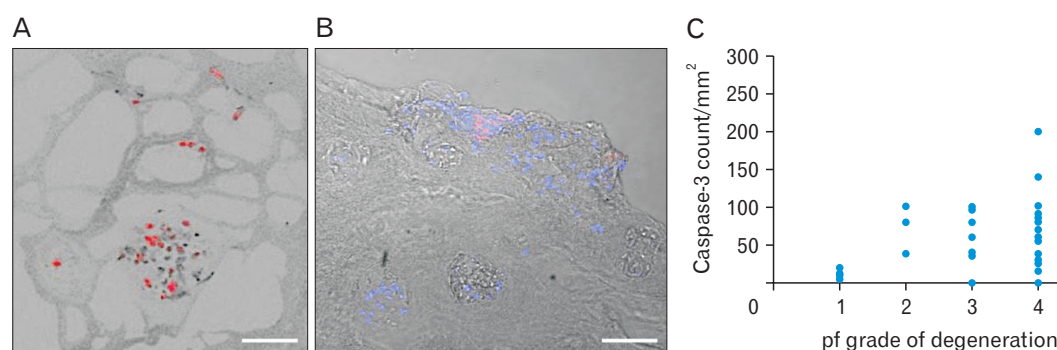


Fig. 4. Medium and large sized cell clusters stained with (A) Caspase-3 (red stain, blue indicates DAPI stained cell nuclei), (B) MMP-1 (red stain, blue indicates DAPI stain cell nuclei) with phase contrast background. Scale bar: 100 μ m. (C) Caspase-3 positive cell clusters count and increasing pf grade of disc degeneration. MMP-1, matrix metalloproteinase-1; DAPI, 4',6-diamidino-2-phenylindole; pf, Pfirrmann.

SDS-PAGE Quantitative analysis for MMP-1, Ki-67, PCNA, and Caspase-3

Protein extracts from human disc tissues along with non-degenerated control and glyceraldehyde 3-phosphate dehydrogenase (GAPDH) as loading control were analyzed on a weight per volume basis by gel electrophoresis. Image J analysis software were used to quantify the bands.

MMP-1 was expressed as a weak band in control discs, while degenerated and herniated discs showed prominent bands at 54 kDa (Fig. 5A). MMP-1 expressions gradually

increased with pf grades of degeneration and were most expressed in grade 3 discs (Fig. 5B), followed by decline in grade 4 and grade 5 discs. Control, grade 1 and grade 2 disc also had similar low MMP-1 expression profile perhaps to prevent the tissue from self-proteolysis.

Ki-67, PCNA, and Caspase-3 were expressed moderately in control discs (Fig. 5 C-H). Ki-67 appeared as a 350 kDa band and were prominently expressed in grade 1 and 2 discs followed by grade 3, 4, 5 discs (Fig. 5C, D). PCNA, appeared at 36 kDa and followed the same expression pattern as Ki-67

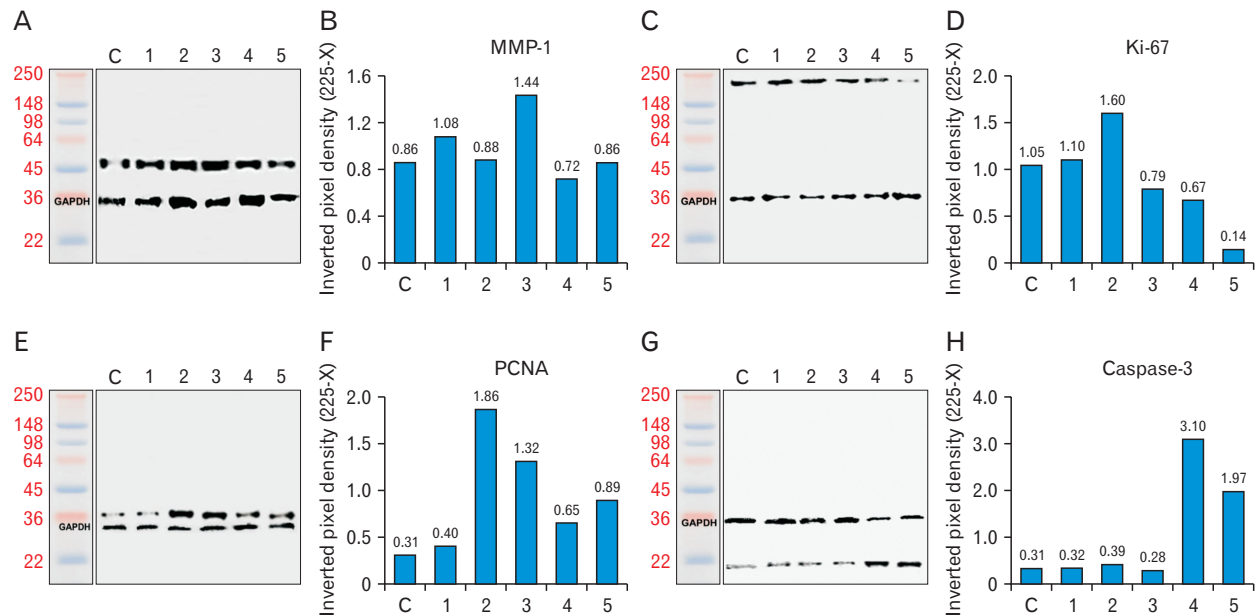


Fig. 5. Western blot analysis on a 10% gel. Lane were loaded as C, 'non-degenerated cadaveric disc' followed by disc specimens categorised according to pf grades degeneration in the subsequent lanes as pf grade 1, 2, 3, 4, and 5 discs. (A) MMP-1 core protein at 54 kDa. (B) Expression level of control 'c', and grade '1–5 discs' shown with grade 3 discs at the highest expression level. (C) Ki-67 core protein with a molecular weight of 250 kDa. (D) Weak expression level for degenerated grade 3–5 discs. (E) PCNA core protein with a molecular weight of 36 kDa. (F) Bands expressed higher values for grade 2 and 3 discs in comparisons to other grades. (G) Caspase-3 core protein at 22 kDa. (H) Strongly expressed for the severely degenerated grade 4 and 5 discs. GAPDH as loading control is at 36 kDa. C, control; pf, Pfirrmann; MMP-1, matrix metalloproteinase-1; PCNA, proliferating cell nuclear antigen; GAPDH, glyceraldehyde 3-phosphate dehydrogenase.

(Fig. 5E, F) suggesting that grade 2, 3 degenerated disc tissue had greater quantity of viable cells. Caspase-3, appeared as a 22 kDa band, and were minimally expressed in controls, grade 1–3 discs, followed by increased expression in severely degenerated grade 4, 5 discs (Fig. 5G, H).

Discussion

Summary of results and gap analysis

The present study shows small and medium-sized clusters in control and slightly degenerated discs express Ki-67 and PCNA markers whereas large clusters localised around disrupted proteoglycan-depleted tissue in the NP/IAF junction of severely degenerated, herniated discs are positive to MMP-1 and Caspase-3. These findings were confirmed with SDS-PAGE that shows these target markers can be extracted from the relatively dense collagenous matrix. SDS-PAGE investigation allowed segregation and identification of these protein of interest through its molecular weight, thereby eliminating any interferences which may be caused due to non-specific binding charges introduced by immuno-histochemical methodologies. Small and medium-sized cell clusters had higher

proportions of cells positive to proliferating markers, while caspase-3 activity in large clusters were common in end-stages of disc degeneration. Control disc showed least number of cell clusters, proteoglycan loss score, and immunopositivity towards MMP-1 and Caspase-3. Large clusters, in contrast, were a characteristic feature of severely degenerated and/or herniated disc with increased Caspase-3 activity [42].

This study compared pathological human discs with age-matched cadaveric controls. Ki-67 and PCNA antigen were the operational markers used for determining cellular proliferation. Ki-67 is not expressed by quiescent or resting cells in the G0-phase [43] and PCNA markers are expressed only during the S phase of the cell cycle [44], hence a combination of these two markers facilitated the assessment of cell proliferation phase in the clusters of various sizes. MMP-1, is the main enzymatic marker involved in initiation of extracellular matrix degradation and is suggested to play a vital role in increased proteolysis of the disc matrix [42] and leukocyte migration. MMP-1 cleaves collagen types I, II, III and proteoglycans which forms the main microstructural framework of the intervertebral discs [45]. Similarly, other ubiquitous matrix metalloproteinases such as MMP-2, 3, 5, 7, 9, 13 [46] are also

shown to be involved in diverse functions associated with tissue injury, repair, inflammation, blood vessels and nerve ingrowth [8, 47].

Caspase-3, is involved in the activation cascade of apoptotic cell death pathway in human intervertebral discs [43]. At the onset of apoptosis, it proteolytically cleaves poly (ADP-ribose) polymerase at '216-Asp-Gly-217' bond which shows caspase 3 is a key cysteine protease that executes programme cell death. Caspase 3 can also activate caspases 6, 7, and 9 that overlap with the necrotic mode of cell death [47, 48], a prominent feature of inflammation and degeneration. Caspase-3 antibody used in this study identified both the pro-caspase form (32 kDa) and the active caspase form (14–22 kDa).

This study made comparisons only between human tissue, and even though the results are applicable to the human population, a controlled animal tissue study along with human discs samples would perhaps be a more insightful prospect, however animal are quadrupled species and unlike humans their discs are more complex to load and would involve ethical concerns for *in vivo* loading.

Relationship to other studies

With age chondrocyte-like cells of the nucleus and inner annulus regions become more and more rounded or even irregular in shape, develop a dense pericellular matrix, and frequently proliferate to form cell clusters [15, 49]. Each small clusters consists of 3–4 cells within a lacunae that isolate them from the rest of the matrix [24] large cell clusters with <8–15 cells are considered to be a histological hallmark feature of cartilage tissue degeneration [9], but cell cluster are also hypothesized to represent an attempted repair response to tissue damage, where remodeling of matrix occur via chondrocyte division or proliferation of cells, swelling of the chondrons, and distension of the surrounding microenvironment [50]. Since cell clusters are abundantly located in regions that show loss of fibrillar orientation [51], or are located mostly near lesions in the experimental animal model [52] they could be indicative of a repair response. Cell proliferation are considered to be an essential process for maintaining hydration of tissue, and organ culture studies of human discs show that native disc cells produce large proteoglycan aggregates [53] that play a crucial role in maintaining swelling pressure of the disc tissue [54], but with aging, accelerated degeneration, and pain, changes in the morphological and biochemical structure of the collagen lamellae, and vital

cellular or non-cellular protein components increases cell clustering. Anabolic factor such as the proteoglycans molecules produced by the cells and clusters, as a repair response, can easily leach out of the disc due to excessive tissue swelling and loss in restrain especially in grade 4 and 5 discs [11]. Other significant problem associated with tissue repair is decline in the number of viable disc cells due to poor metabolite transport [55], subsequently increasing the proportions of cells that undergoes senescence [13], followed by apoptosis [56]. Thus, large, and medium sized cell clusters formed during degeneration, and in aged individuals, may contribute towards increased release of matrix-degrading enzymes [37], capable of altering the mechano-transduction pathways [29]. Moreover, unavoidable daily loading stress, age-related changes of the intrinsic and extrinsic cell-matrix environment, and genetic factors may progressively weaken the disc matrix, and structural changes, such as fissure formed, may not be easily repaired by the temporarily proliferating cell. Thus, with age and injury, cell clustering is perhaps unavoidable, especially near the nucleus pulposus and inner annulus junction where clusters were abundant [8, 11], and immuno-reactivity towards MMP-1 and Caspase-3 are located within the same region. This finding are additionally in agreement with mechanical studies that show the inner annulus is weak and prone to structural disruptions due to repetitive loading and ageing [3]. Current and past studies on humans, porcine and ovine discs confirm structural failure always leads to progressive degenerative biochemical changes [52], but are not seen in non-degenerated disc [57]. Thus, findings from this study suggest cell clustering is initiated as tissue repair or remodeling process that fails due to unsupportive metabolic, biomechanical, and biochemical responses, and these cell cluster eventually undergoes apoptosis.

Explanation of results and conclusion

Proteoglycan-depleted tissue regions with large cell clusters indicates phenotypic alteration perhaps due to abnormal mechano-transduction pathways [58], and degenerated, painful discs exhibit negative regulation to nuclear factor- κ B pathway [59] influencing the proliferation process. Increased MMP expression contributes towards focal tissue disruptions by allowing enzymatic fragmentation of matrix that may activate TLR receptors [60, 61], and these fragmented molecules can acts as damaged associated molecular pattern (DAMPS) which acts in a similar ways as the pathogen associated molecular pattern (PAMPS) to initiate a sterile

inflammatory catabolic cascade. Therefore, to conclude, distinctive microscopic changes and mechanical loading can cause the discs to enter 'vicious cycle' of interconnected mechanisms such as focal swelling that accelerates proteoglycan loss and increase cell clustering in an attempt to compensate proteoglycan loss, but as this attempt fails, clusters increase the expression of MMP-1 and caspase 3. Therefore, clusters appearing during late stages degeneration can't remodel the matrix and 'senolytic drugs' proposed to clear out non-functioning cells could possibly benefit the end stages of disc degeneration.

ORCID

Polly Lama: <https://orcid.org/0000-0002-0756-674X>
 Jerina Tiwari: <https://orcid.org/0000-0002-9112-9955>
 Pulkit Mutreja: <https://orcid.org/0000-0002-6994-710X>
 Sukirti Chauhan: <https://orcid.org/0000-0003-4103-7814>
 Ian J Harding: <https://orcid.org/0000-0003-1322-417X>
 Trish Dolan: <https://orcid.org/0000-0001-8561-675X>
 Michael A Adams: <https://orcid.org/0000-0003-0435-8651>
 Christine Le Maitre: <https://orcid.org/0000-0003-4489-7107>

Author Contributions

Conceptualization: PL, MAA. Data acquisition: JT, PM, SC, IJH. Data analysis or interpretation: PL, MAA, TD, CLM. Critical revision of the manuscript: PL, MAA, CLM. Approval of the final version of the manuscript: all authors.

Conflicts of Interest

No potential conflict of interest relevant to this article was reported.

Funding

Research Grants from Indian Council of Medical Research (No. ICMR/5/7/1290-2015 RCH), Dr TMA Pai Grant (No. SMU/131/REG/TMAPURK/164/2020) for Polly Lama and Action Medical Research Grant (No. MS0549089/2016) for Trish Dolan and Michael A Adams.

References

1. Karppinen J, Shen FH, Luk KD, Andersson GB, Cheung KM,

Samartzis D. Management of degenerative disk disease and chronic low back pain. *Orthop Clin North Am* 2011;42:513-28, viii.

2. Newell N, Little JP, Christou A, Adams MA, Adam CJ, Masouros SD. Biomechanics of the human intervertebral disc: a review of testing techniques and results. *J Mech Behav Biomed Mater* 2017;69:420-34.
3. Adams MA, Freeman BJ, Morrison HP, Nelson IW, Dolan P. Mechanical initiation of intervertebral disc degeneration. *Spine (Phila Pa 1976)* 2000;25:1625-36.
4. Adams MA, Roughley PJ. What is intervertebral disc degeneration, and what causes it? *Spine (Phila Pa 1976)* 2006;31:2151-61.
5. Adams A, Roche O, Mazumder A, Davagnanam I, Mankad K. Imaging of degenerative lumbar intervertebral discs; linking anatomy, pathology and imaging. *Postgrad Med J* 2014;90:511-9.
6. Adams MA, Dolan P. Intervertebral disc degeneration: evidence for two distinct phenotypes. *J Anat* 2012;221:497-506.
7. Roberts S, Menage J, Urban JP. Biochemical and structural properties of the cartilage end-plate and its relation to the intervertebral disc. *Spine (Phila Pa 1976)* 1989;14:166-74.
8. Lama P, Le Maitre CL, Dolan P, Tarlton JF, Harding IJ, Adams MA. Do intervertebral discs degenerate before they herniate, or after? *Bone Joint J* 2013;95-B:1127-33.
9. Johnson WE, Eisenstein SM, Roberts S. Cell cluster formation in degenerate lumbar intervertebral discs is associated with increased disc cell proliferation. *Connect Tissue Res* 2001;42:197-207.
10. Lyu FJ, Cheung KM, Zheng Z, Wang H, Sakai D, Leung VY. IVD progenitor cells: a new horizon for understanding disc homeostasis and repair. *Nat Rev Rheumatol* 2019;15:102-12.
11. Lama P, Claireaux H, Flower L, Harding IJ, Dolan T, Le Maitre CL, Adams MA. Physical disruption of intervertebral disc promotes cell clustering and a degenerative phenotype. *Cell Death Discov* 2019;5:154.
12. Le Maitre CL, Freemont AJ, Hoyland JA. The role of interleukin-1 in the pathogenesis of human intervertebral disc degeneration. *Arthritis Res Ther* 2005;7:R732-45.
13. Le Maitre CL, Freemont AJ, Hoyland JA. Accelerated cellular senescence in degenerate intervertebral discs: a possible role in the pathogenesis of intervertebral disc degeneration. *Arthritis Res Ther* 2007;9:R45.
14. Horner HA, Roberts S, Bielby RC, Menage J, Evans H, Urban JP. Cells from different regions of the intervertebral disc: effect of culture system on matrix expression and cell phenotype. *Spine (Phila Pa 1976)* 2002;27:1018-28.
15. Trout JJ, Buckwalter JA, Moore KC. Ultrastructure of the human intervertebral disc: II. Cells of the nucleus pulposus. *Anat Rec* 1982;204:307-14.
16. Sharp CA, Roberts S, Evans H, Brown SJ. Disc cell clusters in pathological human intervertebral discs are associated with increased stress protein immunostaining. *Eur Spine J* 2009;18:1587-94.
17. Kim KW, Lim TH, Kim JG, Jeong ST, Masuda K, An HS. The

- origin of chondrocytes in the nucleus pulposus and histologic findings associated with the transition of a notochordal nucleus pulposus to a fibrocartilaginous nucleus pulposus in intact rabbit intervertebral discs. *Spine (Phila Pa 1976)* 2003;28:982-90.
18. Kim JH, Deasy BM, Seo HY, Studer RK, Vo NV, Georgescu HI, Sowa GA, Kang JD. Differentiation of intervertebral notochordal cells through live automated cell imaging system *in vitro*. *Spine (Phila Pa 1976)* 2009;34:2486-93.
 19. Phillips KL, Chiverton N, Michael AL, Cole AA, Breakwell LM, Haddock G, Bunning RA, Cross AK, Le Maitre CL. The cytokine and chemokine expression profile of nucleus pulposus cells: implications for degeneration and regeneration of the intervertebral disc. *Arthritis Res Ther* 2013;15:R213.
 20. Ashton IK, Eisenstein SM. The effect of substance P on proliferation and proteoglycan deposition of cells derived from rabbit intervertebral disc. *Spine (Phila Pa 1976)* 1996;21:421-6.
 21. Henriksson HB, Svala E, Skioldebrand E, Lindahl A, Brisby H. Support of concept that migrating progenitor cells from stem cell niches contribute to normal regeneration of the adult mammal intervertebral disc: a descriptive study in the New Zealand white rabbit. *Spine (Phila Pa 1976)* 2012;37:722-32.
 22. Gardner DL. Problems and paradigms in joint pathology. *J Anat* 1994;184(Pt 3):465-76.
 23. Lotz MK, Otsuki S, Grogan SP, Sah R, Terkeltaub R, D'Lima D. Cartilage cell clusters. *Arthritis Rheum* 2010;62:2206-18.
 24. Bryant PJ. Growth factors controlling imaginal disc growth in *Drosophila*. *Novartis Found Symp* 2001;237:182-94; discussion 194-202.
 25. Li FX, Xu F, Lin X, Wu F, Zhong JY, Wang Y, Guo B, Zheng MH, Shan SK, Yuan LQ. The role of substance P in the regulation of bone and cartilage metabolic activity. *Front Endocrinol (Lausanne)* 2020;11:77.
 26. Opolka A, Straub RH, Pasoldt A, Grifka J, Grässel S. Substance P and norepinephrine modulate murine chondrocyte proliferation and apoptosis. *Arthritis Rheum* 2012;64:729-39.
 27. Ogunlade B, Fidelis OP, Adelakun SA, Adedotun OA. Grape seed extract inhibits nucleus pulposus cell apoptosis and attenuates annular puncture induced intervertebral disc degeneration in rabbit model. *Anat Cell Biol* 2020;53:313-24.
 28. Horner HA, Urban JP. 2001 Volvo Award Winner in Basic Science Studies: effect of nutrient supply on the viability of cells from the nucleus pulposus of the intervertebral disc. *Spine (Phila Pa 1976)* 2001;26:2543-9.
 29. Mavrogenatou E, Papadimitriou K, Urban JP, Papadopoulos V, Kletsas D. Deficiency in the $\alpha 1$ subunit of Na⁺/K⁺-ATPase enhances the anti-proliferative effect of high osmolality in nucleus pulposus intervertebral disc cells. *J Cell Physiol* 2015;230:3037-48.
 30. Sakai D, Nakamura Y, Nakai T, Mishima T, Kato S, Grad S, Alini M, Risbud MV, Chan D, Cheah KS, Yamamura K, Masuda K, Okano H, Ando K, Mochida J. Exhaustion of nucleus pulposus progenitor cells with ageing and degeneration of the intervertebral disc. *Nat Commun* 2012;3:1264.
 31. Tessier S, Risbud MV. Understanding embryonic development for cell-based therapies of intervertebral disc degeneration: toward an effort to treat disc degeneration subphenotypes. *Dev Dyn* 2021;250:302-17.
 32. Mizrahi O, Sheyn D, Tawackoli W, Ben-David S, Su S, Li N, Oh A, Bae H, Gazit D, Gazit Z. Nucleus pulposus degeneration alters properties of resident progenitor cells. *Spine J* 2013;13:803-14.
 33. Sakai D, Schol J, Bach FC, Tekari A, Sagawa N, Nakamura Y, Chan SCW, Nakai T, Creemers LB, Frauchiger DA, May RD, Grad S, Watanabe M, Tryfonidou MA, Gantenbein B. Successful fishing for nucleus pulposus progenitor cells of the intervertebral disc across species. *JOR Spine* 2018;1:e1018.
 34. Pfirrmann CW, Metzendorf A, Zanetti M, Hodler J, Boos N. Magnetic resonance classification of lumbar intervertebral disc degeneration. *Spine (Phila Pa 1976)* 2001;26:1873-8.
 35. Schmitz N, Laverty S, Kraus VB, Aigner T. Basic methods in histopathology of joint tissues. *Osteoarthritis Cartilage* 2010;18 Suppl 3:S113-6.
 36. Boos N, Weissbach S, Rohrbach H, Weiler C, Spratt KF, Nerlich AG. Classification of age-related changes in lumbar intervertebral discs: 2002 Volvo Award in basic science. *Spine (Phila Pa 1976)* 2002;27:2631-44.
 37. Baschong W, Suetterlin R, Laeng RH. Control of autofluorescence of archival formaldehyde-fixed, paraffin-embedded tissue in confocal laser scanning microscopy (CLSM). *J Histochem Cytochem* 2001;49:1565-72.
 38. Bouayad D, Pederzoli-Ribeil M, Mocek J, Candalh C, Arlet JB, Hermine O, Reuter N, Davezac N, Witko-Sarsat V. Nuclear-to-cytoplasmic relocalization of the proliferating cell nuclear antigen (PCNA) during differentiation involves a chromosome region maintenance 1 (CRM1)-dependent export and is a prerequisite for PCNA antiapoptotic activity in mature neutrophils. *J Biol Chem* 2012;287:33812-25.
 39. Faratian D, Munro A, Twelves C, Bartlett JM. Membranous and cytoplasmic staining of Ki67 is associated with HER2 and ER status in invasive breast carcinoma. *Histopathology* 2009;54:254-7.
 40. Cserni G. Analysis of membranous Ki-67 staining in breast cancer and surrounding breast epithelium. *Virchows Arch* 2018;473:145-53.
 41. Remnant L, Kochanova NY, Reid C, Cisneros-Soberanis F, Earnshaw WC. The intrinsically disorderly story of Ki-67. *Open Biol* 2021;11:210120.
 42. Le Maitre CL, Freemont AJ, Hoyland JA. Localization of degradative enzymes and their inhibitors in the degenerate human intervertebral disc. *J Pathol* 2004;204:47-54.
 43. Miller I, Min M, Yang C, Tian C, Gookin S, Carter D, Spencer SL. Ki67 is a graded rather than a binary marker of proliferation versus quiescence. *Cell Rep* 2018;24:1105-12.e5.
 44. Sasaki K, Kurose A, Ishida Y. Flow cytometric analysis of the expression of PCNA during the cell cycle in HeLa cells and effects of the inhibition of DNA synthesis on it. *Cytometry* 1993;14:876-82.
 45. Van Doren SR. Matrix metalloproteinase interactions with col-

- lagen and elastin. *Matrix Biol* 2015;44-46:224-31.
46. Vo NV, Hartman RA, Yurube T, Jacobs LJ, Sowa GA, Kang JD. Expression and regulation of metalloproteinases and their inhibitors in intervertebral disc aging and degeneration. *Spine J* 2013;13:331-41.
47. Tschöke SK, Hellmuth M, Hostmann A, Robinson Y, Ertel W, Oberholzer A, Heyde CE. Apoptosis of human intervertebral discs after trauma compares to degenerated discs involving both receptor-mediated and mitochondrial-dependent pathways. *J Orthop Res* 2008;26:999-1006.
48. Jiang LB, Liu HX, Zhou YL, Sheng SR, Xu HZ, Xue EX. An ultrastructural study of chondroptosis: programmed cell death in degenerative intervertebral discs *in vivo*. *J Anat* 2017;231:129-39.
49. Lama P, Kulkarni JP, Tamang BK. The role of cell clusters in intervertebral disc degeneration and its relevance behind repair. *Spine Res* 2017;3:15.
50. Sandell LJ, Aigner T. Articular cartilage and changes in arthritis. An introduction: cell biology of osteoarthritis. *Arthritis Res* 2001;3:107-13.
51. Pearle AD, Warren RF, Rodeo SA. Basic science of articular cartilage and osteoarthritis. *Clin Sports Med* 2005;24:1-12.
52. Melrose J, Roberts S, Smith S, Menage J, Ghosh P. Increased nerve and blood vessel ingrowth associated with proteoglycan depletion in an ovine annular lesion model of experimental disc degeneration. *Spine (Phila Pa 1976)* 2002;27:1278-85.
53. Buckwalter JA. Aging and degeneration of the human intervertebral disc. *Spine (Phila Pa 1976)* 1995;20:1307-14.
54. Urban JP, Maroudas A. Swelling of the intervertebral disc *in vitro*. *Connect Tissue Res* 1981;9:1-10.
55. Urban JP, Smith S, Fairbank JC. Nutrition of the intervertebral disc. *Spine (Phila Pa 1976)* 2004;29:2700-9.
56. Zhao CQ, Wang LM, Jiang LS, Dai LY. The cell biology of intervertebral disc aging and degeneration. *Ageing Res Rev* 2007;6:247-61.
57. Lotz JC, Chin JR. Intervertebral disc cell death is dependent on the magnitude and duration of spinal loading. *Spine (Phila Pa 1976)* 2000;25:1477-83.
58. Le Maitre CL, Frain J, Millward-Sadler J, Fotheringham AP, Freemont AJ, Hoyland JA. Altered integrin mechanotransduction in human nucleus pulposus cells derived from degenerated discs. *Arthritis Rheum* 2009;60:460-9.
59. Wuertz K, Vo N, Kletsas D, Boos N. Inflammatory and catabolic signalling in intervertebral discs: the roles of NF- κ B and MAP kinases. *Eur Cell Mater* 2012;23:103-19; discussion 119-20.
60. Vogl T, Eisenblätter M, Völler T, Zenker S, Hermann S, van Lent P, Faust A, Geyer C, Petersen B, Roebrock K, Schäfers M, Bremer C, Roth J. Alarmin S100A8/S100A9 as a biomarker for molecular imaging of local inflammatory activity. *Nat Commun* 2014;5:4593.
61. Haglund L, Bernier SM, Onnerfjord P, Recklies AD. Proteomic analysis of the LPS-induced stress response in rat chondrocytes reveals induction of innate immune response components in articular cartilage. *Matrix Biol* 2008;27:107-18.