Rat tail static compression model mimics extracellular matrix metabolic imbalances of matrix metalloproteinases, aggrecanases, and tissue inhibitors of metalloproteinases in intervertebral disc degeneration

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Abstract

Introduction: The longitudinal degradation mechanism of extracellular matrix (ECM) in the intervertebral disc remains unclear. Our objective was to elucidate catabolic and anabolic gene expression profiles and their balances in intervertebral disc degeneration using a static compression model.

Methods: Forty-eight 12-week-old male Sprague-Dawley rat tails were instrumented with an Ilizarov-type device with springs and loaded statically at 1.3 MPa for up to 56 days. Experimental loaded and distal-unloaded control discs were harvested and analyzed by real-time reverse transcription-polymerase chain reaction (PCR) messenger RNA quantification for catabolic genes [matrix metalloproteinase (MMP)-1a, MMP-2, MMP-3, MMP-7, MMP-9, MMP-13, a disintegrin and metalloproteinase with thrombospondin motifs (ADAMTS)-4, and ADAMTS-5], anti-catabolic genes [tissue inhibitor of metalloproteinases (TIMP)-1, TIMP-2, and TIMP-3], ECM genes [aggrecan-1, collagen type 1-α1, and collagen type 2-α1], and pro-inflammatory cytokine genes [tumor necrosis factor (TNF)-α, interleukin (IL)-1α, IL-1β, and IL-6]. Immunohistochemistry for MMP-3, ADAMTS-4, ADAMTS-5, TIMP-1, TIMP-2, and TIMP-3 was performed to assess their protein expression level and distribution. The presence of MMP- and aggrecanase-cleaved aggrecan neoepitopes was similarly investigated to evaluate aggrecanolytic activity.

Results: Quantitative PCR demonstrated up-regulation of all MMPs and ADAMTS-4 but not ADAMTS-5. TIMP-1 and TIMP-2 were almost unchanged while TIMP-3 was down-regulated. Down-regulation of aggrecan-1 and collagen type 2-α1 and up-regulation of collagen type 1-α1 were observed. Despite TNF-α elevation, ILs developed little to no up-regulation. Immunohistochemistry showed, in the nucleus pulposus, the percentage of immunopositive cells of MMP-cleaved aggrecan neoepitope increased from 7 through 56 days with increased MMP-3 and decreased TIMP-1 and TIMP-2 immunopositivity. The percentage of immunopositive cells of aggrecanase-cleaved aggrecan neoepitope increased at 7 and 28 days only with decreased TIMP-3 immunopositivity. In the annulus fibrosus, MMP-cleaved aggrecan neoepitope presented much the same expression pattern. Aggrecanase-cleaved aggrecan neoepitope increased at 7 and 28 days only with increased ADAMTS-4 and ADAMTS-5 immunopositivity.

Conclusions: This rat tail sustained static compression model mimics ECM metabolic imbalances of MMPs, aggrecanases, and TIMPs in human degenerative discs. A dominant imbalance of MMP-3/TIMP-1 and TIMP-2 relative to ADAMTS-4 and ADAMTS-5/TIMP-3 signifies an advanced stage of intervertebral disc degeneration.
Introduction
Low back pain is a global health problem due to its high prevalence and high socioeconomic burden. It affects 70 to 85% of the population during a lifetime, 15 to 45% in a year, and 12 to 30% at any point, and accounts for approximately 13% of sickness absences [1]. Although the cause of low back pain is multifactorial, intervertebral disc degeneration is implicated in more than half of the cases [2].

The intervertebral disc has a complex structure with the nucleus pulposus (NP) encapsulated by endplates and the annulus fibrosus (AF). Intervertebral disc degeneration is biochemically characterized by extracellular matrix (ECM) degradation [3-5]. ECM consists primarily of proteoglycans – principally aggrecan – and collagens – mainly type 2 in the NP and type 1 in the AF [6]. ECM metabolism is regulated by the balance between degradative enzymes, matrix metalloproteinases (MMPs) and aggreganases, and their natural inhibitors, tissue inhibitors of metalloproteinases (TIMPs) [7,8]. Aggrecanases are identified as members of a disintegrin and metalloproteinase with thrombospondin motifs (ADAMTS) family [7]. Imbalances of MMPs, ADAMTSs, and TIMPs significantly correlate with cartilage ECM metabolism in patients with osteoarthritis and rheumatoid arthritis [9-11]. In degenerated disc tissue, modified expressions of MMPs, ADAMTSs, and TIMPs have also been detected [12-19]. However, balances of these enzymes and their practical significance in intervertebral disc degeneration remain unclear.

Studying disc degeneration is difficult because of the challenge of reproducing the variety of etiological aspects of the degenerative process: ECM degradation, inflammation, nutrient loss, cell senescence, and apoptotic cell death [20]. Systematic analysis of these etiologies using human specimens is impractical; therefore, reliable animal models of disc degeneration are required.

Rodent tails are popular to assess disc degeneration because of easy accessibility with minimal damage to surrounding tissues and minimal interference with normal physiological functions [21]. Rodents keep notochordal cells in the disc NP throughout their lifetime [21] whereas humans lose them at young ages in somatic development, when discs begin to show first signs of degeneration [22]. Recent evidence has suggested that the change of NP cell phenotype from notochordal to chondrocyte-like plays a significant role in the initiation of disc degeneration [23,24]. Thus, understanding rodent disc degeneration provides an interpretation of the pathogenesis of human disc degeneration.

Many methods to induce degeneration are proposed; mechanical loading provokes chronic degenerative responses unlike annular puncture which provides reliable responses to acute injury [21]. Mounting evidence has revealed that dynamic compression stimulates anabolism whereas static compression accelerates catabolism [25-27]. Static compression induces histomorphological degeneration [28-30], cell apoptosis [28-32], and altered content of proteoglycans [25,28,29,33] and collagens [28,29,34,35]. Static compression thus has the potential to reproduce disc degeneration via cell apoptosis and ECM degradation; this conveys its primary advantage for longitudinal investigation of the degenerative mechanism compared with dynamic compression [21,36]. ECM metabolism under static compression has been partially explained by activation of MMP-2 [37] and up-regulation of MMP-13 and TIMP-1 [34,35]. The authors have previously reported that in vivo sustained static compression leads to progressive and prolonged up-regulation of MMP-3 with the progression of radiological and histomorphological degeneration [38]. However, comprehensive degeneration-related gene expression including MMP, ADAMTS, and TIMP balances has not been profiled. Their ECM degradation potential has not been the focus of investigation. Therefore, longitudinal gene-quantification studies using the static compression model still need to be conducted.

The objectives of this study were to clarify catabolic and anabolic gene expression profiles and to elucidate expression including MMP, ADAMTS, and TIMP balances in ECM metabolism of intervertebral disc degeneration.

Materials and methods
All animal procedures were performed under the approval and guidance of the Animal Care and Use Committee at Kobe University Graduate School of Medicine.

Animals and surgical procedure
Forty-eight 12-week-old male Sprague-Dawley rats (CLEA Japan, Tokyo, Japan), ranging in weight from 452 to 509 g, were used. Rats are reported to reach approximately 90% of skeletal maturity 12 weeks after birth [39]. Rat tails were affixed with an ilizarov-type apparatus with springs between the 8th and 10th coccygeal (C) vertebrae as described in our previous paper (Figure 1) [38]. This loading system was similar to that of Iatridis and colleagues [33]. Under intraperitoneal anesthesia, two-cross 0.7 mm diameter Kirschner wires were inserted percutaneously into each vertebral body perpendicular to the tail’s axis and attached to aluminum rings. Rings were connected longitudinally with four threaded rods. Four 0.50 N/mm calibrated springs were installed over each rod. After instrumentation, axial stress was loaded from the distal side to produce a calculated compressive pressure of 1.3 MPa. This pressure,
corresponding closely to the disc loading force produced by lifting a moderate weight in the human lumbar spine, was shown to induce morphological and biochemical disc degeneration with cell apoptosis by Lotz and colleagues [28,31].

Following surgery, rats were randomly loaded for 0 (sham), 7, 28, or 56 days and euthanized; the data did not consist of repeated measurements over time points, but of single measurements in each time point. Rat tails with the compressive apparatus unloaded for up to 56 days were used as the sham group. In 24 rats, C9-10, the distal loaded disc, and C12-13, the unloaded internal control disc [38], were harvested for messenger RNA (mRNA) quantification following radiographic and magnetic resonance imaging (MRI) assessments (n = 6/time point). To exclude potential level effects [40], those discs in the additional 24 rats were harvested for histomorphological and immunohistochemical assessments (n = 6/time point). Radiological, histomorphological, and cell population data were presented previously [38,41]. No obvious change in adjacent disc levels of the Ilizarov-type device over 56 days was confirmed biochemically [33] and radiologically [38].

**RNA extraction and reverse transcription**

Loaded and unloaded discs were immediately dissected using a scalpel after euthanasia. NP tissue was collected using a curette, pulverized under liquid nitrogen, and total RNA was isolated using RNeasy Mini Kit (Qiagen, Hilden, Germany). Then 0.1 μg RNA was reverse-transcribed in the presence of RT² First Strand Kit (SABiosciences, Frederick, MD, USA) including oligo d (T) primer and random hexamers.

**Quantitative real-time reverse transcription-polymerase chain reaction**

**Catabolic genes**

Relative expression levels of mRNA encoding rat *MMP-1a* (GenBank: NM_001134530), *MMP-2* (GenBank: NM_031054), *MMP-3* (GenBank: NM_133523), *MMP-7* (GenBank: NM_012864), *MMP-9* (GenBank: NM_031055), *MMP-13* (GenBank: NM_133530), *ADAMTS-4* (GenBank: NM_023959), and *ADAMTS-5* (GenBank: NM_198761) were calculated by real-time reverse transcription (RT)-polymerase chain reaction (PCR) using ABI Prism 7500 sequence detection system (Applied Biosystems, Foster City, CA, USA). *MMP-1a* was predicted to encode a rat homolog of human MMP-1 [42] because rodent species appear to lack MMP-1 [43]. *Glyceraldehyde 3-phosphate dehydrogenase* (*GAPDH*) (GenBank: NM_017008) mRNA expression was measured as an endogenous control. Good feasibility of *GAPDH* was confirmed in our previous experiment using this rat tail model [41]. We used a custom-made RT² Profiler PCR Array (SABiosciences, Frederick, MD, USA), which consisted of a set of SYBR green fluorescent dye and multiple pre-designed primers with an arrangement to analyze multiple target gene mRNA expressions of experimental and control samples simultaneously. We separately used primers for *MMP-3* (Takarabio, Shiga, Japan) in order to amplify a specific sequence previously

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**Figure 1** Whole and close-up view of rat tail instrumented with an Ilizarov-type loading device.
published: sense 5'-TGG ACC AGG GAC CAA TGG A-3' and anti-sense 5'-GTC CAA GTT CAT GAG CAG CA-3' [38]. The use of the customized PCR Array system made it possible to precisely measure multiple gene alterations in the same rat samples under the same experimental conditions. The measurement was performed in duplicate. The mRNA expression of each enzyme in the C9-10 loaded disc was converted to a relative number representing the amount of mRNA compared with the C12-13 unloaded disc using the 2−ΔΔCt method [44]. Difference in threshold cycles for target gene and reference gene was calculated as ΔCt = Ct(target gene) - Ct(GAPDH). Difference in threshold cycles for target sample and reference sample was calculated as ΔΔCt = ΔCtC9-10 - ΔCtC12-13. Finally, the mRNA expression fold change of target gene in the loaded relative to unloaded disc was calculated as 2ΔΔCt.

**Anti-catabolic genes**

**TIMP-1** (GenBank: NM_053819), **TIMP-2** (GenBank: NM_021989), and **TIMP-3** (GenBank: NM_012886) mRNA expression levels were quantified following the same PCR array procedure.

**Extracellular matrix genes**

**Aggrecan-1** (GenBank: NM_022190), **collagen type 1-α1** (GenBank: NM_053304), and **collagen type 2-α1** (GenBank: NM_012929) mRNA expression levels were assessed using PCR array.

**Pro-inflammatory cytokines genes**

**TNF-α** (GenBank: NM_012675), **IL-1α** (GenBank: NM_017019), **IL-1β** (GenBank: NM_031512), and **IL-6** (GenBank: NM_012589) mRNA expression levels were evaluated with PCR array.

**Paraffin-embedded disc tissue preparation**

Loaded and unloaded discs were excised, fixed in 4% paraformaldehyde, decalcified in 10% ethylenediaminetetraacetic acid, embedded in paraffin, sectioned from the mid-sagittal plane at 5-μm thickness, and prepared for immunohistochemical analysis.

**Immunohistochemistry**

**Catabolic and anti-catabolic genes**

Immunohistochemical staining was performed to determine the level and distribution of protein expression of MMP-3, ADAMTS-4, ADAMTS-5, TIMP-1, TIMP-2, and TIMP-3. Sections were incubated with 1:75 diluted goat-polyclonal anti-MMP-3 (sc-6839; Santa Cruz Biotechnology, Santa Cruz, CA, USA), 1:100 diluted rabbit-polyclonal anti-ADAMTS-4 (SP4703P; Acris Antibodies, Herford, Germany), 1:20 diluted rabbit-polyclonal anti-ADAMTS-5 (sc-83186; Santa Cruz Biotechnology, Santa Cruz, CA, USA), 1:50 diluted rabbit-polyclonal anti-TIMP-1 (ab78703; Abcam, Cambridge, UK), 1:50 diluted rabbit-polyclonal anti-TIMP-2 (sc-5539; Santa Cruz Biotechnology, Santa Cruz, CA, USA), or 1:20 diluted mouse-monovalent anti-TIMP-3 (ab49670; Abcam, Cambridge, UK) at 4°C overnight, and subsequently treated at room temperature for 30 minutes with a peroxidase-labeled anti-goat, anti-mouse, or anti-rabbit antibody (Nichirei Bioscience, Tokyo, Japan). The signal, a brown reaction product, was developed using peroxidase substrate 3,3'-diaminobenzidine and counterstained with hematoxylin. Parallel sections treated with goat, mouse, or rabbit normal serum (Dako, Glostrup, Denmark) at equal protein concentrations were used as negative controls. The number of immunopositive cells of each NP and AF dissected on the center was counted in five random high-power fields (×400) using BZ-9000 microscope and analysis software (Keyence, Osaka, Japan); positive staining was expressed as a percentage of immunopositive cells to total cell population measured by counting the nuclei.

**Aggrecanolytic activity**

Immunohistochemistry using cleavage site-specific antibodies for aggrecan was performed to detect aggrecan core protein fragments generated by MMP and aggrecanase activity. After deglycosylation of the aggrecan core proteins using chondroitinase ABC (Sigma-Aldrich, St. Louis, MO, USA) and keratanases (Seikagaku Biobusiness, Tokyo, Japan), sections were treated at 4°C overnight with 1:20 diluted mouse-monovalent antibody BC-14 (NB110-6852; Novus Biologicals, Littleton, CO, USA), which recognized the N-terminal neoeptite sequence of 342FFGVG generated by MMP cleavage in the intergrobal domain of aggrecan, or 1:100 diluted mouse-monovalent antibody BC-3 (NB600-504; Novus Biologicals, Littleton, CO, USA), which recognized the N-terminal neoeptite sequence of 374ARGSV generated by aggrecanase cleavage. The secondary antibody was a peroxidase-labeled anti-mouse antibody (Nichirei Bioscience, Tokyo, Japan) used at room temperature for 30 minutes. Brown color-development, counterstaining, cell-counting, and negative control preparation were performed as described above.

**Statistical analysis**

Two-way mixed-design analysis of variance (ANOVA) with the Turkey-Kramer post-hoc test was used to assess changes of mRNA level for the effects of disc level (loaded and unloaded: within-subjects) and time (0, 7, 28, and 56 days: between-subjects). Two-way ANOVA with the Turkey-Kramer post-hoc test was applied to evaluate changes of percentage of immunopositive cells similarly (both factors: between-subjects). Data analyses were performed using PASW Statistics 18 (SPSS, Chicago, IL, USA). Statistical significance was accepted at P < 0.05. All values are expressed as mean ± standard deviation.
Results
All animals tolerated surgery well and gained body weight throughout the experiment: 455 to 526 g at 7 days, 497 to 563 g at 28 days, and 543 to 614 g at 56 days. All springs maintained their compressive length and fully recovered immediately after release, indicating sustained axial loading. No signs of infection, skin necrosis, neurological problems, or instrument failure were observed.

Quantitative real-time reverse transcription-polymerase chain reaction outcomes
For application of the $2^{-\Delta\Delta C_t}$ method, the efficiencies of all measured gene amplifications were first examined using diluted samples and confirmed to be appropriately equal. In addition, no amplification of primer dimers and other non-specific products were detected.

Catabolic genes
High-throughput mRNA quantification of the loaded relative to unloaded disc demonstrated significant up-regulation of all examined catabolic genes except ADAMTS-5 (Figure 2a). MMP-3, MMP-7, MMP-9, and MMP-13 showed significant up-regulation from seven days of loading, MMP-1a from 28 days, and MMP-2 from 56 days ($P < 0.05$). Furthermore, MMP up-regulation significantly progressed over time with compression ($P < 0.05$). In aggrecanases, ADAMTS-4 exhibited similar significant up-regulation from seven days ($P < 0.05$) while ADAMTS-5 underwent no significant up-regulation throughout the study duration. The increasing tendency was more remarkable in MMP-1a, MMP-2, and MMP-3 than MMP-7, MMP-9, MMP-13, and ADAMTS-4. In the mRNA expression of these enzymes at seven days, MMP-3 showed the most notable increase.

Anti-catabolic genes
All TIMPs demonstrated no obvious up-regulation during the loading period (Figure 2b). TIMP-1 exhibited no significant change but did show an increasing trend toward up-regulation. TIMP-2 showed significant down-regulation at seven days ($P < 0.05$) and it recovered at 28 days ($P < 0.05$). Meanwhile, TIMP-3 was significantly and continuously down-regulated from seven days of loading ($P < 0.05$).

Extracellular matrix genes
Aggrecan-1 and collagen type 2-α1 were significantly down-regulated from seven days ($P < 0.05$) whereas collagen type 1-α1 was up-regulated with significance at 56 days ($P < 0.05$) (Figure 2c).

Pro-inflammatory cytokine genes
TNF-α showed significant up-regulation at seven days ($P < 0.05$) and subsequently maintained high expression levels through 56 days (Figure 2d). However, IL expression demonstrated little to no elevation; significant down-regulation of IL-1α at 7 and 28 days ($P < 0.05$), no significant change of IL-1β throughout, and significant up-regulation of IL-6 only at 56 days ($P < 0.05$) were observed (Figure 2d).

Immunohistochemical outcomes
The number of disc cells progressively decreased with compression. In the NP, while large vacuolated notochordal cells were frequently observed at 0 day but apparently disappeared from seven days, smaller round chondrocyte-like cells clustered and collapsed elliptically but were found throughout the study. In the AF, evenly distributed fibroblast-like cells were observed at 0 day but subsequently decreased and larger round chondrocyte-like cells appeared.

Catabolic and anti-catabolic genes
Immunoreactivity for all examined genes was predominantly localized in the cytoplasm of disc cells and higher in NP cells than AF cells; particularly, large NP cells with vacuoles, suspecting notochordal origin, showed strong immunoreactivity (Figures 3a and 4a). Immunopositivity for all studied genes was detected in sham and unloaded discs and also generally higher in the NP than the AF (Figures 3b and 4b). No immunopositivity was detected in the matrix. IgG controls were negative, and positive controls showed strong positive stainings.

The percentage of immunopositive cells of MMP-3 in the NP significantly increased from 7 to 56 days ($P < 0.05$) despite a slight decrease at 56 days. The percentage of immunopositive cells of MMP-3 in the AF significantly increased from seven days ($P < 0.05$) and retained the enhanced expression through 56 days. The percentage of immunopositive cells of TIMP-1 in the NP had no significant change through 28 days but significantly decreased at 56 days ($P < 0.05$). The percentage of immunopositive cells of TIMP-1 in the AF did not show any significant change over the loading duration. The percentage of immunopositive cells of TIMP-2 in the NP significantly decreased at 28 and 56 days ($P < 0.05$). The percentage of immunopositive cells of TIMP-2 in the AF did not show any significant change throughout (Figure 3b).

The percentage of immunopositive cells of ADAMTS-4 and ADAMTS-5 in the NP showed no significant change through 28 days but significantly decreased at 56 days ($P < 0.05$). The percentage of immunopositive cells of ADAMTS-4 and ADAMTS-5 in the AF exhibited a transient increase at 28 days ($P < 0.05$). The percentage of immunopositive cells of TIMP-3 in the NP, despite possessing detectable staining in 0-day non-degenerated discs, consistently decreased until levels were nearly undetectable ($P < 0.05$). The percentage of immunopositive cells of...
Figure 2 Real-time reverse transcription-polymerase chain reaction gene expression profile. The messenger RNA (mRNA) expression of target gene normalized to glyceraldehyde 3-phosphate dehydrogenase (GAPDH) is represented by fold change in the loaded relative to unloaded disc (control value = 1). *P < 0.05 when compared between loaded and unloaded conditions. †P < 0.05 when compared between different time points. (a) Relative mRNA expression at 0, 7, 28, and 56 days of: catabolic matrix metalloproteinase (MMP)-1α = 0.9, 1.6, 6.1, respectively; MMP-2 = 1.2, 0.8, 3.4, 10.8, respectively; MMP-3 = 1.0, 5.0, 9.8, respectively; MMP-7 = 1.0, 1.9, 2.0, respectively; MMP-9 = 1.1, 1.3, 1.5, respectively; MMP-13 = 1.0, 1.3, 1.5, respectively; ADAMTS-4 = 1.0, 1.6, 19, respectively; ADAMTS-5 = 1.0, 1.1, 1.1, respectively. (b) Relative mRNA expression at 0, 7, 28, and 56 days of: anti-catabolic tissue inhibitor of metalloproteinases (TIMP)-1 = 1.0, 1.2, 1.4, respectively; TIMP-2 = 0.9, 0.5, 1.2, 1.4, respectively; TIMP-3 = 1.0, 0.6, 0.8, respectively. (c) Relative mRNA expression at 0, 7, 28, and 56 days of: extracellular matrix aggrecan-1 = 1.0, 0.3, 0.5, respectively; collagen type 1α1 = 0.9, 0.6, 0.3, 0.2, respectively; collagen type 2α1 = 0.9, 0.6, 0.3, 0.2, respectively. (d) Relative mRNA expression at 0, 7, 28, and 56 days of: pro-inflammatory cytokine tumor necrosis factor (TNF)-α = 1.0, 1.9, 2.2, 2.0, respectively; interleukin (IL)-1α = 1.0, 0.7, 0.8, 0.9, respectively; IL-1β = 1.1, 1.1, 1.1, 1.2, respectively; IL-6 = 0.9, 1.0, 1.2, 1.2, respectively.
TIMP-3 in the AF remained low throughout the study period with a significant decrease at 56 days ($P < 0.05$) (Figure 4b).

**Aggrecanolytic activity**

Few positive stainings for MMP-generated and aggrecanase-generated aggrecan neoepitopes were detected in the 0-day NP and AF (Figure 5a). The percentage of immunopositive cells surrounded by MMP cleavage aggrecan neoepitope-positive matrix significantly increased from 7 to 56 days ($P < 0.05$) in the NP and AF. The percentage of immunopositive cells surrounded by aggrecanase cleavage aggrecan products significantly increased at 7 and 28 days ($P < 0.05$) but significantly decreased at 56 days ($P < 0.05$) in the NP and AF (Figure 5b).

**Discussion**

Our results with the static compression model may be summarized in five major findings. First, real-time RT-PCR demonstrated up-regulated MMPs, lesser up-regulated ADAMTSs, and unchanged or down-regulated TIMPs during 56 days. Second, immnohistochemistry showed that the percentage of immunopositive cells of MMP-3 increased from 7 to 56 days while those of TIMP-1 and TIMP-2 were unchanged or decreased in the NP and AF. Third, in the NP, the percentage of immunopositive cells of ADAMTS-4 and ADAMTS-5 were unchanged through 28 days but decreased at 56 days while those of TIMP-3 consistently decreased from seven days. Fourth, in the AF, the percentage of immunopositive cells of ADAMTS-4 and ADAMTS-5 transiently increased at 28 days while those of TIMP-3 remained low throughout the study. Fifth, the percentage of immunopositive cells of MMP-cleaved aggrecan neoepitope increased from 7 to 56 days while those of aggrecanase-cleaved aggrecan neoepitope increased at 7 and 28 days but decreased at 56 days in the NP and AF. These findings indicate this rat tail sustained static compression model mimics ECM metabolic imbalances of MMPs, aggrecanases, and TIMPs in human degenerative discs. A dominant imbalance of MMP-3/TIMP-1 and TIMP-2 relative to ADAMTS-4 and ADAMTS-5/TIMP-3 signifies an advanced stage of intervertebral disc degeneration.

We previously reported that sustained static compression induced disc height loss in radiographs and lower NP intensity in T2-weighted MRIs [38]. Histological sections showed cell decrease with altered detectable phenotypes, scar formation, tissue defect, and a decrease of proteoglycans in Safranin-O staining [38,41]. In the current study, sustained static compression further induced aggrecan-1 and collagen type 2-$\alpha 1$ mRNA down-regulation and collagen type 1-$\alpha 1$ mRNA up-regulation (Figure 2c). These macro, micro, and biological findings...
have a good accordance with human degenerated discs [3,5,22,45] and other static compression models [25,28-30,34,35].

Our rat tail MMP mRNA up-regulation paralleled degeneration (Figure 2a), consistent with human disc findings [12-16,19]. A significant correlation between elevated MMP-2 and MMP-9 levels and degenerative disc grades was reported by Crean and colleagues [13]. Weiler and colleagues found that MMP-1, MMP-2, and
MMP-3 expression highly correlated with cleft and scar formation in degenerated discs [15]. The gene expression study by Bachmeier and colleagues revealed that MMP-3 up-regulation highly dependent on histological evidence of disc degeneration [19]. Furthermore, genomic anomalies of MMPs have significant involvement in disc degeneration [46-49]. A polymorphism in the promoter of MMP-1 [48], MMP-2 [47], MMP-3 [46], and MMP-9 [49] genes, which enhance promoter activity, is associated with a higher prevalence of lumbar disc degeneration in Japanese elderly patients [46] and Chinese adult cohorts [47-49]. Thus, the progression of disc degeneration strongly correlates with MMP up-regulation. The static compression model simulates MMP expression in intervertebral disc degeneration.

Our rat tail ADAMTS-4 mRNA up-regulation accompanied degeneration (Figure 2a), consistent with human discs [16-18]. On the other hand, our rat tail model exhibited no significant mRNA change of ADAMTS-5 (Figure 2a). ADAMTS-5 expression is controversial in human discs [17,18]. Pockert and colleagues reported ADAMTS-5 up-regulation correlated with histological grades of degenerative discs [18] whereas Patel and colleagues noted ADAMTS-5 expression level did not differ between histological grades [17]. In osteoarthritis cartilage, ADAMTS-5 is thought to have a more influential role than ADAMTS-4 [50,51]. ADAMTS-5 thus has the potential to play a role in intervertebral disc degeneration; however, further investigations are needed.

Our rat tail model demonstrated no consistent mRNA change of TIMP-1 and TIMP-2 and consistent mRNA down-regulation of TIMP-3 (Figure 2b). TIMP expression is also controversial in human discs [12,16,18,19]. Bachmeier and colleagues described TIMP-1 and TIMP-2 mRNA up-regulation in lumbar discs with degeneration [19]. In the investigation by Le Maitre and colleagues, up-regulation of TIMP-1 and TIMP-2, but not TIMP-3, correlated with the severity of histological degeneration [16]. No correlation of TIMP-3 with degeneration in the NP and inner AF and a negative correlation in the outer AF were observed by Pockert and colleagues [18]. Partially supporting these human data, the current animal model results indicate a differential expression pattern of TIMP-3 in disc degeneration. In ECM metabolism, much evidence for TIMP functions has been accumulated [7,8]; TIMPs inhibit MMPs by 1:1 interaction with zinc-binding site [8]; TIMP-1 is the main inhibitor of MMP-3 [52], TIMP-2 inhibits MMP-2 [53] and MMP-14 [54]; TIMP-3 is a potent inhibitor of ADAMTS-4 and ADAMTS-5 [55], and TIMP-1, TIMP-2, and TIMP-4 do not inhibit ADAMTS-4 [56]. Therefore, ratios of MMPs/TIMP-1 and TIMP-2 and ADAMTS-4 and ADAMTS-5/TIMP-3 must remain balanced to maintain matrix homeostasis.

In human degenerative discs, it is difficult to profile these balances in vivo; however, the static compression model facilitates detailed longitudinal analysis of these balances in disc ECM metabolism during degeneration.

Our rat tail model showed mRNA up-regulation of TNF-α but not ILs (Figure 2d); however, human disc tissues frequently demonstrate up-regulation of TNF-α [57,58], IL-1α [59], IL-1β [58,59], and IL-6 [60]. In human non-herniated degenerative discs, large cleft formation and immunocompetent CD68-positive cells around cleft are observed [61]. In the repeated stab model, TNF-α, IL-1β, and IL-8 production in p38-positive cells is detected around the stab wound [62]. However, the static compression model does not present any large cleft formation or radial wound from the NP through the AF [28-30,38], potentially differing from physiological degeneration in the production mechanism of pro-inflammatory cytokines. TNF-α stimulates collagenase gene transcription by prolonged activation of Jun gene expression in fibroblasts [63]. TNF-α induces the production of nitric oxide [64], which can inhibit proteoglycan synthesis [65] and enhance MMP activity [66] in chondrocytes. TNF-α reduces the gene for Sox9 [67] which is required for the expression of chondrocyte-specific markers aggrecan-1 and collagen type 2-a1 [68]. Thus, TNF-α up-regulation in this model may indicate TNF-α contribution to the pathogenesis of disc degeneration.

To further characterize this cellular pathobiology, immunohistochemical study for MMPs, ADAMTSs, and TIMPs and their aggrecanolytic activity was performed. We investigated MMP-3 immunopositivity as representative of MMPs, judging from published evidence highlighting the significant importance of MMP-3 in disc degeneration [12,19]. MMP-3 is a major aggrecan-degrading MMP and activates many other pro-MMPs such as MMP-1, MMP-7, MMP-8, MMP-9, and MMP-13 [8]. In the NP, the prominent MMP-3/TIMP-1 and TIMP-2 imbalance (Figures 3a and 3b) with a persistent increase of MMP-cleaved aggrecan neoepitope (Figures 5a and 5b) was observed. Meanwhile, the modest ADAMTS-4 and ADAMTS-5/TIMP-3 imbalance (Figures 4a and 4b) with a transient increase of aggrecanase-cleaved aggrecan neoepitope (Figures 5a and 5b) was detected. Human disc studies have shown controversial findings of MMP-generated and aggrecanase-generated aggrecan fragments during degeneration [14,17,18,69]. Aggrecan neoepitopes cleaved by MMPs and aggrecanases are more frequently detected in degenerated than non-degenerated discs [18]. However, with advancing degeneration, the MMP-cleaved neoepitope abundance is constant in the NP and AF [69], whereas the aggrecanase-cleaved neoepitope abundance is unchanged [17] or decreased [69] in the NP but remains...
constant in the AF [17,69]. Aggrecanase-cleaved neoepti-
tope is less abundant than MMP-cleaved neoeptiopoe in
degenerated discs [14]. The static compression model
recapitulates these pieces of evidence regarding disc
matrix aggrecan degradation; advancing degeneration
does not necessarily imply increased aggrecan fragments,
particularly by aggrecanase cleavage.

In this study, gene expression results with real-time
RT-PCR and immunohistochemistry were partially mis-
mached. In the MMP-3/TIMP-1 and TIMP-2 imbal-
ance, MMP-3 mRNA and protein up-regulation was
consistent while TIMP-1 and TIMP-2 protein down-reg-
ulation at 28 to 56 days was inconsistent with their con-
stant mRNA expression. These TIMP protein findings
are not in agreement with those reported by Le Maitre
and colleagues [16]; however, they do corroborate those
by Kanemoto and colleagues where 78.1% of cervical
spondylosis and 93.3% of lumbar spondylosis specimens
were MMP-3 positive and TIMP-1 negative [12]. A pro-
longed catabolic shift might lead to decreased TIMP-1
and TIMP-2 protein. In the ADAMTS-4 and ADAMTS-
5/TIMP-3 imbalance, TIMP-3 mRNA and protein
down-regulation was consistent while no obvious change
of ADAMTS-4 protein expression at 7 and 28 days was
inconsistent with its mRNA up-regulation. Immunohis-
tochemistry shows localization, but not productive
quantity; therefore, it may be difficult to detect small
expression change. This ADAMTS-4 finding possibly
indicates that the ADAMTS-4 and ADAMTS-5/TIMP-3
imbalance is primarily due to TIMP-3 down-regulation
in the NP. ADAMTS-4 and ADAMTS-5 protein down-
regulation at 56 days was further inconsistent with their
elevated or constant mRNA expression. These
ADAMTS findings possibly correlate with decreased
aggrecanase-cleaved aggrecan neoepitope. The observed
discrepancy between mRNA and protein expression in
the NP might be explained by prolonged, prominent
MMP-induced aggrecanolysis and shortened, modest
aggrecanase-induced aggrecanolysis; however, further
investigations are required to understand the regulation
mechanism of MMPs, ADAMTSs, and TIMPs at the
post-transcriptional level.

The pathomechanism of AF degradation has not been
clarified in detail. The annulus matrix comprises col-
lagen, proteoglycans, and elastic fibers – elastin and
microfibril such as glycoprotein fibrillins [70]. Elastin is
readily degraded by MMP-2, MMP-3, MMP-7, MMP-9,
MMP-10, and MMP-12 [8,71]. Fibrillins are degraded by
MMP-2, MMP-3, MMP-9, MMP-12, MMP-13, and
MMP-14 [72]. This study lacked real-time RT-PCR ana-
lysis of AF tissue, which is an inherent limitation. In
immunohistochemistry, the prominent MMP-3/TIMP-1
and TIMP-2 imbalance (Figures 3a and 3b) with a per-
sistent increase of MMP-cleaved aggrecan neoepitope
(Figures 5a and 5b) was observed in the AF as well as in
the NP. Meanwhile, the ADAMTS-4 and ADAMTS-5/
TIMP-3 imbalance in the AF had a pattern different
from that in the NP; more remarkable ADAMTS-4 and
ADAMTS-5 up-regulation than TIMP-3 down-regula-
tion was detected (Figures 4a and 4b). This imbalance
appeared to produce a transient increase of aggrecanase-
cleaved aggrecan neoepitope at much the same time as
the NP (Figures 5a and 5b). The histomorphological
study by Boos and colleagues showed the NP was more
severely degenerated in the same age group than the AF
[22]; however, our biological findings indicate AF degen-
eration occurs simultaneously with NP degeneration.

Our rat tail immunohistochemical results are summa-
rized in Figure 6. Both imbalances of MMP-3/TIMP-1
and TIMP-2 and ADAMTS-4 and ADAMTS-5/TIMP-3
work in the early to middle stages of 7 and 28 days;
however, the MMP-3/TIMP-1 and TIMP-2 imbalance is
more severe than the ADAMTS-4 and ADAMTS-5/
TIMP-3 imbalance at the late stage of 56 days. This pro-
vides a sound argument for low aggrecanase activity in
the discs with advanced degeneration. In ECM metabo-
lism, the relative importance of MMPs and aggrecanases
has long been debated. Little and colleagues reported
catabolic aggrecan degradation in normal and osteoar-
thritis cartilage primarily involved cleavage by aggreca-
nase and not by MMPs [73]. It was found by Karsdal
and colleagues that MMP-mediated degradation of
aggrecan and collagen type 2 caused irreversible damage
for cartilage, while aggrecanase-mediated degradation of
aggrecan was fully reversible [74]. Integrated with these
reports, our findings show that a state of dominant
MMP-3/TIMP-1 and TIMP-2 imbalance relative to
ADAMTS-4 and ADAMTS-5/TIMP-3 imbalance may
indicate an irreversible stage of intervertebral disc
degeneration.

This rodent disc study provides information not only
about staged processes involved in disc degeneration but
also about its possible pathogenesis. In particular, the
loss of NP notochordal cells induced by sustained static
compression should be noted. In our previous study,
immunofluorescence exhibited that the number of NP
cells decreased to 43.0% during 56 days of loading [41]. In
the in vivo study by Guehring and colleagues, sus-
tained static compression induced more remarkable
decrease of notochordal cells than of chondrocyte-like
cells with a total cell decrease of more than 50% over 56
days [75]. Severity of cell decrease is similar in both stu-
dies; although we did not identify notochordal cells
using immunostaining for the markers, it is speculated
that our rat tail disc significantly loses notochordal cells
following compression. In seven-day loaded disc sec-
tions, we observed an apparent decrease of large vacu-
olated cells suspecting notochordal origin, supporting the

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proposal by Guehring and colleagues that notochordal cells are less resistant to mechanical stress than chondrocyte-like cells [75].

We further found more distinct immunoreactivity for MMP, ADAMTS, and TIMP enzymes in notochordal cells than in chondrocyte-like cells (Figures 3a and 4a). In 0-day discs, large vacuolated NP cells showed strong immunoreactivity despite little detection of aggregan fragments. In seven-day loaded discs, many NP cells were chondrocyte-like and demonstrated generally weaker immunoreactivity with increased detection of aggregan neoepitopes, which was irrespective of immunopositivity affected by loading. This finding gives rise to the possibility that notochordal cells play an important role in matrix homeostasis, providing a plausible explanation for the observed higher baseline expression levels of MMPs, ADAMTSs, and TIMPs in the NP than in the AF and differential imbalanced pattern of ADAMTS-4 and ADAMTS-5/TIMP-3 between the NP and AF. Notochordal cells produce a larger amount of proteoglycans than chondrocyte-like cells [76,77] and stimulate chondrocyte-like cells to produce proteoglycans [78,79]. Proteoglycan loss is an early, significant biochemical change to occur in disc degeneration [3,4].

Human disc NP tissues have a higher content of glycosaminoglycans than AF in donors aged 25 years or younger but thereafter lose them; further, NP specimens have a higher level of newly synthesized aggregan than AF in donors five years or younger however lose it markedly by 5 to 15 years [5], corresponding well with the disappearance of notochordal cells [22]. In keeping with these findings, our study results indicate that the loss of notochordal cells is potentially linked with the initiation of disc degeneration.

A major limitation of this study is that the rat tail static compression model does not perfectly reflect all aspects of human disc degeneration. This model’s unphysiological situation – immobility, extended pressure, and absence of trauma – could lead to some contradictions to humans, such as lack of radial tears, mucous degeneration, and granular changes in histology and little involvement of inflammation. It could also explain the observed simultaneous NP and AF degeneration.

Further studies using the static compression model should be conducted to detect the turning point from reversible to irreversible degeneration, providing key information to prevent degenerative disc diseases.
Conclusions
This rat tail sustained static compression model mimics ECM metabolic imbalances of MMPs, aggrecanases, and TIMPs in human intervertebral disc degeneration. In early stages, MMP-3/TIMP-1 and TIMP-2 and ADAMTS-4 and ADAMTS-5/TIMP-3 imbalances exist. In the late stage, the MMP-3/TIMP-1 and TIMP-2 imbalance dominates. These imbalances and their effects on aggrecanolysis are common in the NP and AF. A dominant imbalance of MMP-3/TIMP-1 and TIMP-2 relative to ADAMTS-4 and ADAMTS-5/TIMP-3 is a possible indication of an advanced, irreversible stage of intervertebral disc degeneration.

Abbreviations
ADAMTS: a disintegrin and metalloproteinase with thrombospondin motifs; AF: annulus fibrosus; ANOVA: analysis of variance; C: coccyeal; ECM: extracellular matrix; GAPDH: glyceraldehyde 3-phosphate dehydrogenase; IL: interleukin; MMP: matrix metalloproteinase; MRI: magnetic resonance imaging; mRNA: messenger RNA; NP: nucleus pulposus; PCR: polymerase chain reaction; RT: tissue inhibitor of metalloproteinases; TNF: tumor necrosis factor.

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Authors’ contributions
TY conceived the study, secured funding, participated in its conception and design, acquisition of data, and analysis and interpretation of data, and drafted the manuscript. TT participated in analysis and interpretation of data. KK participated in analysis and helped to draft the manuscript. TS participated in acquisition of data and drafted the manuscript. TT participated in analysis and interpretation of data. TY conceived the study, secured funding, participated in its conception and analysis and interpretation of data. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

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