A repeated strike loading organ culture model for studying compression-associated chronic disc degeneration

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ABSTRACT

Mechanical stress has been viewed as one of the key risk factors in accelerating the intervertebral disc degeneration process. The goal of the present study was to employ a repeated strike loading bovine caudal disc system to elucidate the pathophysiological impacts of cumulative mechanical stress on the disc. The discs in the model groups were subjected to two different mechanical stresses: one strike loading or repeated strike loading. The following indices were analyzed: histological morphology, glycosaminoglycan release, disc height, cell viability, apoptosis-related protein expression, and catabolism-related gene expression. Both mechanical stress modes induced degenerative changes in the discs by day 11, such as clefts and delamination of the annulus fibrosus; they increased glycosaminoglycan release. Cell viability was significantly decreased and catabolic gene expression was significantly up-regulated in the degenerative loading group and repeated strike loading group by day 9. These alterations remained evident in the annulus fibrosus tissue of the repeated strike loading group on day 11. Our data suggests that the repeated strike loading model adopted in this study could lead to degenerative changes in the disc organ model. Annulus fibrosus cells displayed a more noticeable response to mechanical stress damage and a slower recovery process, suggesting that the annulus fibrosus serves as a pivotal factor in disc degeneration due to mechanical stress injuries. The study also indicates that due to the gradual self-repair of intervertebral disc cells after injury, it is necessary to apply repeated strike loading on the disc at specific intervals when researching the repair of chronic disc injuries.

Keywords: Intervertebral disc degeneration; mechanical stress; organ culture model.
INTRODUCTION

Intervertebral disc degeneration (IDD) is one of the primary contributing factors of low back pain, which causes a substantial socioeconomic burden worldwide[1]. In Western countries, approximately 80% of the population is affected by low back pain, with its incidence rising with advancing age[2]. Acute pain from disc degeneration costs up to $100 billion annually in the U.S.[3]. Though significant progress has been made in the exploration of the pathogenesis of IDD, the molecular mechanisms still hold some ambiguity.

The intervertebral disc (IVD) is an avascular organ positioned between neighboring vertebral bodies and is responsible for load transmission during daily activity. It is composed of three fundamental tissues: the gelatinous inner nucleus pulposus (NP), which is encircled by the annulus fibrosus (AF) and interposed between two cartilaginous endplates (CEPs)[4]. The CEP is a layer of hyaline cartilage that is crucial for the nutrient and metabolite exchange between the IVD and vertebral body[5]. The NP contains large amounts of proteoglycans and glycosaminoglycans (GAGs), with high hydration characteristics, allowing for the conversion of compressive pressure into an evenly distributed expansion pressure. Studies have shown that the degeneration of NP tissue reduces the disc's ability to absorb and distribute loads evenly across the vertebral column[6]. This alteration in mechanical properties leads to increased mechanical stress on other components of the disc, accelerating the degeneration process[7]. The AF is rich in highly organized type I collagen bundles that are ideal for generating tensile strength to balance the expansion pressure from the NP[8]. Degeneration of the AF leads to a decrease in its structural integrity. This deterioration can cause the AF to become more susceptible to tears and fissures, allowing NP material to herniate or extrude, contributing to spinal instability and pain[9].

A vast body of research has been devoted to the study of IVD degeneration. The selection of an appropriate experimental model is crucial, both cell and animal models are instrumental yet present notable limitations. Primarily, these models fail to fully encapsulate the complex and multifaceted nature of human IDD pathophysiology due to significant differences in cell population, tissue composition, and biomechanical properties across species [10]. Furthermore, induced degenerative changes in animal models may not accurately reflect the human condition, limiting their applicability and the translation of findings to clinical scenarios [11]. Consequently, while these models contribute valuable insights into disc biology, caution must be exercised when extrapolating results to human IDD treatment and
prevention strategies. Given these considerations, the use of IVD organ culture has been identified as a compelling method to mitigate the limitations inherent in both in vivo and in vitro models. Bovine caudal discs have been recognized for their pronounced similarities to mature human discs in critical areas including cellular composition, microstructural mechanics, and extracellular matrix composition[12, 13]. These similarities render bovine caudal discs an invaluable model for preclinical research, facilitating studies that are both relevant and translatable to human conditions.

The IVD is subjected to diverse physiological and pathological mechanical stress, including compressive dynamic/static loads, tensile strains and osmotic stress. Previous epidemiological and clinical research has identified strike loading as a significant contributor to the initiation and progression of IDD[14]. Repeated mild lumbar injury is not an uncommon occurrence in our daily lives, especially for those that perform manual work and athletes. Epidemiological evidence has pointed out that repeated lumbar injury exposure is an important risk factor for IDD[15, 16]. Additionally, a recent study found that progressive degenerative changes in rabbit lumbar discs were induced by repeated axial loading[17]. An animal model of accelerated disc degeneration demonstrated that repeated stab injury initiated a vicious cycle of inflammation and matrix damage[18]. We have proved in a strike-loading disc degeneration model that a short-term degeneration reaction was initiated after one strike loading. This reaction was self-limited and typically ceased within a few days[19]. As most studies have focused on exploring the timely pathophysiological consequence of the mechanical stress on IVD, few studies have investigated whether cumulative mechanical stress could induce persistent molecule response in the IVD.

To better clarify the relationship between repeated acute strike loading and IDD, we employed a fully developed IVD organ culture system in this study. Specifically, the deleterious effects of one strike loading and repeated strike loading on the IVD were assessed. We hypothesized that repeated strike loading would result in persistent IVD degeneration.

**MATERIALS AND METHODS**

**Separation and nurture of IVDs**

Bovine tails were attained from 18 month old cows within 6 hours of slaughter. Since we used residues from an abattoir, there was no need for ethical approval. IVDs were harvested using the method described earlier on Day 0[20]. In short, the peripheral tissues were
dissected with a scalpel blade, and then IVDs accompanied by the endplates were separated using a sawing machine. The IVDs underwent cleaning by an irrigator with PBS and then washed with PBS and 10% penicillin/streptomycin (Gibco) in an orbital shaker at 37 °C for 15 minutes. IVDs were cultured at 37 °C in a 5% CO2 incubator, and the medium was changed daily. The culture medium was composed of Dulbecco’s modified Eagle’s medium (Sigma–Aldrich) supplemented with 10% fetal bovine serum, 1% penicillin/streptomycin, 1% ITS+1 (Sigma–Aldrich), 50 µg/mL L-ascorbic acid (Sigma–Aldrich) and 0.1% Primocin.

**Strike loading application**

IVDs from the same animal were randomized into 4 groups: control (CON) group, degenerative (Deg) group, one strike (OS) group, repeated strike (RS) group. To maintain cell viability, IVDs in the CON and strike groups underwent physiological sinusoidal loading (20–200 KPa; 0.2 Hz; 1 hr/day) in our customized chamber (Fig. 1A) with fresh high glucose (4.5 g/L) medium using our custom designed bioreactor (Fig. 1C) and then cultured fresh in high glucose medium overnight. IVDs in the Deg group underwent degenerative sinusoidal loading (320–500 KPa; 5 Hz; 2 hr/day) in our customized chamber with fresh low glucose (2 g/L) medium using our custom designed bioreactor and then cultured in fresh low glucose medium overnight[20]. IVDs in OS group were only subjected to strike loading (40% strain in one second) in DAY 1. IVDs in RS loading group were subjected to strike loading (35% strain in one second) in Days 1, 4 and 7. The IVDs underwent an initial 3-minute preloading of 10 N to preclude over-hydration and to affirm certain connect among the IVD, the customized chamber, and the specially designed mechanical testing machine (Fig. 1D). Then the mechanical testing machine was employed to subject the IVD to strike loading for a duration of 1 second, inducing strains equivalent to 40% or 35% of the IVD.[7]. IVDs in the CON group, Deg group and OS group were collected on Day 11 for further analysis. IVDs in the RS group were collected at Day 9 and Day 11.

**Cell viability analysis**

After dissection of one endplate, whole IVDs were snap frozen in optimal cutting temperature compound (Sakura). Ten-micrometer transverse cryosections were cut with a microtome (Leica). Cell viability was measured by double labeling the cells with lactate dehydrogenase (LDH, cell stained blue) and ethidium homodimer-1 (EthD-1, nucleus emitted red fluorescence)[21]. Cell viability was assessed in the NP and AF. Images were visualized at 20× magnification using a DM6B fluorescence microscope (Leica). Pure red fluorescence represents dead cells. Blue or blue/red fluorescence represents live cells. Image-J software
was employed to count the living and dead cells as following: briefly, click on each cell to mark it with a point, each click will place a mark and keep a count.

**Immunofluorescence**

IVD tissue was fixed in fresh 4% paraformaldehyde overnight before dehydration in graded sucrose solutions and embedding in OCT mounting medium. 10 um thick sections were cut with a microtome (Leica). The following antibodies were used: CASPASE3 (1:400, Proteintech) and BCL2 (1:200, Proteintech). Briefly, IVD tissue sections were rinsed in PBS and permeabilized in 0.3% Triton X-100 at room temperature for 30 minutes. The sections were blocked for 1 hour in 5% bovine serum albumin (BSA) and 0.1% Triton X-100. The sections were then incubated with antibodies overnight at 4°C in a dark humidified chamber. After washing, the sections were incubated with an Alexa Flu-or-594-conjugated anti-rabbit secondary antibody (Jackson ImmunoResearch) for 60 minutes. Finally, sections were rinsed with TBST and DAPI staining for 10 minutes. Images were visualized at 20× magnification using a DM6B fluorescence microscope (Leica). Immunofluorescence was evaluated using ImageJ software.

**qPCR**

The endplates of the IVD were dissected, and roughly 150 mg of NP or AF tissue was gathered. The tissue specimens underwent a one-hour digestion with 2 mg/mL pronase at 37 °C, followed by rapid freezing, pulverization in liquid nitrogen, and homogenization utilizing a TissueLyser. RNA was isolated with TRIzol (Invitrogen) and reverse-transcribed with SuperScript VILO (Invitrogen). qPCR was conducted on a Real Time System (Bio–Rad).

Cycle conditions: 50°C for 2 min, 95°C for 2 min, followed by 40 cycles of 15 s at 95°C, and 1 min at 60°C. RPLP0 served as the internal reference gene, and the relative gene expression was determined utilizing the 2^−ΔΔCt approach. The Primer 6.0 software was used to design primer sequences, which are listed in Table 1. The main selection parameters for screening the primers were as follows: primers should be 18-25 nucleotides long; Tm is between 58-62°C; GC content of the primers should be between 40%-60%; Avoid having consecutive G or C bases at the 3’ end; qPCR product should be between 100-300 bp.

**1,9-Dimethylmethylene blue (DMMB) analysis**

A modified DMMB method was used to measure the cumulative release of GAGs in IVD culture medium. DMMB solution was prepared by dissolving 16 mg of DMMB, 3.04 g of glycine and 2.37 g of NaCl in 1 L of distilled water, stirring overnight at room temperature,
and covering with tin foil. The pH value of the DMMB solution was adjusted to 3.0 with 1 M HCl (approximately 9.5 mL) so that the absorbance at 525 nm (A525) was 0.31. The solution was stored at room temperature away from light. Then, 20 µL of diluted IVD culture medium was blended with 0.2 ml of DMMB solution, and the A525 was measured instantly with a microplate reader. A series of chondroitin 4-sodium sulfate dilutions was prepared, exhibiting a halving concentration gradient from 125 µg/mL to 3.90625 µg/mL. The standard GAG reference curve was obtained by mixing 20 µL of diluent with 200 µL of DMMB solution.

**Histomorphological analysis**

IVDs were fixed in 4% paraformaldehyde for 48 hours at 4 °C and embedded in O.C.T. (Sakura). Transverse cryosections (10 µm) were cut with a microtome (Leica). The sections were stained with Safranin O-Fast Green (Sigma–Aldrich, S-O) and pictured by a digital pathology system.

**Measurement of IVD height and dynamic compressive stiffness (DCS)**

Mean IVD height was measured using a Vernier caliper. Each disc was measured at 2 separate locations to determine IVD height, and the mean value was normalized to the Day 1 dimension. The height of the IVD measured on Day 1 after overnight culture, before loading, served as the standardized baseline. The percent IVD height change was calculated.

DCS was measured after overnight culture. The discs were initially subjected to a preloading force of 10 N for 3 minutes, followed by 10 cycles of sinusoidal compression, with strain levels ranging from 5% to 15%. The calculation formula is as follows: \((F_{\text{max}} - F_{\text{min}})/S\). In this context, \(F_{\text{max}}\) and \(F_{\text{min}}\) represent the peak and nadir forces experienced in each cycle, while \(S\) denotes the cross section area of the IVD. Following this, the calculation of the DCS is based on the mean value of 10 rounds. Stiffness measured on Day 1 served as the standardized baseline.

**Statistical analysis**

GraphPad Prism 8.0 (San Diego, CA, USA) was used to perform statistical analysis. Data are shown as mean ± SEM. To determine whether the data were normally distributed, data was first checked by the Shapiro–Wilk normality test. The Student’s t-test was applied to data that was normally distributed, whereas the Mann–Whitney U test was employed for datasets that deviated from a normal distribution. Data yielding a p-value below 0.05 were regarded as statistical significance.
RESULTS

Cell viability
The IVD samples were collected, and frozen sections were prepared for LDH/EthD-1 staining. The images of LDH/EthD-1 staining in each group are shown in (Fig. 2A), where blue or blue-red double staining represents live cells, and red staining alone represents dead cells. Compared with the CON group at day 11, the viability of the AF cells in the model groups was significantly decreased (p < 0.01), but there was no significant difference among the Deg, OS and RS groups at day 11. Compared with the RS group at day 11, AF cell viability was further reduced at day 9 (Fig. 2B). Deg culture condition and RS group at day 9 significantly reduced NP cell viability, whereas the RS group at day 11 did not alter NP cell viability (Fig. 2C).

Immunofluorescence
Immunofluorescence was used to explore the relationship between cell viability and apoptosis. In NP tissue, Deg culture condition and RS at day 9 up-regulated CASPASE3, and the expression of BCL2 showed a trend of down-regulation in the two model groups. In AF tissue, Deg culture condition and RS down-regulated BCL2 and up-regulated CASPASE3 (Fig. 3 A-E).

Gene expression
In the NP tissue, compared with the CON group, the expression levels of the catabolic gene ADAMTS5 and MMP3 was significantly up-regulated in the Deg group and RS group at day 9. Deg culture condition and RS at day 9 also down-regulated the COL2 gene expression. Furthermore, IL-6 a proinflammatory gene expression marker was significantly overexpressed in the Deg group and RS group at day 9. However, there was no significant difference observed in the OS group and RS group at day 11 (Fig. 4).

In the AF, the expression level of the catabolic genes MMP1 and MMP3 was significantly up-regulated in the RS group at day 9. Gene expression level of MMP3 was up-regulated in the Deg group. Proinflammatory gene IL-6 was significantly overexpressed in the RS group at day 9. Genes expression remained unaltered in the OS group and RS group at day 11 (Fig. 5).
GAG release
The DMMB method was used to analyze the cumulative GAG release from the IVDs during the study period. Conditioned medium was collected daily following free-swelling culture and physiological loading culture. The content of released GAG measured on Day 1 following free-swelling culture was used as a baseline for standardization. The DMMB assay demonstrated that GAG release increased in the Deg group and strike loading groups compared with the CON group (p < 0.05), but there was no significant difference among the Deg, and OS and RS groups (Fig. 6).

Histomorphological changes
S-O staining was used to show the morphological changes in the IVD tissue (Fig. 7). Compared with the IVD in the Day 0 group, the microstructures of the IVDs in the CON group did not change significantly. In the Day 0 and CON groups, slight to mild clefts were observed in the AF. The majority of the clefts were parallel to the AF lamellae, without anomalous distortions, and little delamination was found. In the Deg, OS and RS groups, modest and severe clefts were observed in the AF. Additionally, more clefts perpendicular to the AF lamellae were observed. Evident delamination was found in the model groups.

IVD height and DCS
The results showed that the height of the IVD significantly decreased in all groups after physiological loading culture on Day 1 (p < 0.01). On Day 11, all IVD heights showed no significant difference compared to that on Day 1 after overnight free-swelling culture (Fig. 8A).

After 11 days of culture, the IVDs in CON group showed an increase trend in DCS compared to Day 1, but there was no significant difference (p > 0.05). The IVDs in Deg, OS and RS groups showed a significant decrease in DCS compared to the CON group (p < 0.05), but there was no significant difference among the Deg, and OS and RS groups (p > 0.05) (Fig. 8B).

DISCUSSION
This study is based on previous work in which we cultured bovine IVDs in a whole organ culture system with physiological loading or one strike loading. In that study, cell viability was maintained for 10 days after subjection to physiological loading. Although strike loading alone caused a substantial reduction in cell viability and gene expression changes in the short...
term, several of these degradation changes were abolished in the long-term experiment[22]. The goal of the present study was to explore whether repeated strike loading could induce persistent IVD degeneration. Our results indicate that repeated strike loading, one strike loading and degenerative loading culture could lead to IVD degeneration, such as decreased cell viability, increased release of GAGs, decreased DCS, and matrix structure destruction. Repeated strike loading has a long-term effect on the catabolic and apoptotic gene expression of AF cells. In response to mechanical stress damage, NP cells demonstrate a rapid recovery to their normal state, highlighting their robust adaptability and resilience. In contrast, AF cells exhibit a more prolonged response to mechanical stress, characterized primarily by cell viability and alterations in gene expression. Overall, these findings reveal different responses of NP and AF cells to mechanical stress injury, highlighting AF cells as a potentially more vulnerable and critical element in IVD degeneration and damage.

Our study corroborates previous research on the adverse effects of mechanical stress on intervertebral disc degeneration. Croft et al. demonstrated that more intense dynamic loading regimes lead to more pronounced IDD levels[23]. Dudli et al. found that endplate fractures significantly promote disc degeneration, while impact loading without fractures causes minimal changes in cell viability, apoptosis, and inflammation markers[24]. Illien-Jünger et al. examined high-frequency loading under various nutritional conditions over seven days, emphasizing the harmful effects of combined mechanical and nutritional stress[25]. While their study highlighted the detrimental effects of combined mechanical and nutritional stress, our study emphasizes the role of repeated mechanical stress over a longer duration, highlighting the cumulative effects on disc degeneration. Zhou et al., using a one-strike loading model, observed early degenerative changes post-trauma[22]. Our repeated strike loading model, however, provides additional insights into how chronic repetitive loading further exacerbates these degenerative processes over time. Alkhatib et al. indicated that a single hyperphysiological compression event causes endplate fractures and significant cell death[26]. In contrast, our repeated strike loading model shows the cumulative impact of mechanical stress on disc degeneration, with significant increases in apoptosis, matrix remodeling, and pro-inflammatory responses. These findings highlight the critical role of repetitive mechanical stress in accelerating degenerative processes and emphasize the importance of considering the removal of mechanical loads in preventing and treating intervertebral disc degeneration.
At present, experimental studies on the relationship between abnormal mechanical stress and IVD degeneration mostly use chronic sustained mechanical injury or acute violent injury models[27-29]. Chronic sustained mechanical injury can lead to persistent IVD degeneration, while acute abnormal mechanical stress injury can induce transient and short-term IVD degeneration [30]. Previous investigations have identified that impact loading can precipitate degeneration within the IVDs[22, 28, 31]. However, within such models, significant disruptions occur, including the destruction of disc endplates, tears in the AF, and alterations to the low oxygen and nutrient environments in the NP, alongside changes in mechanical characteristics. It remains uncertain whether impact loading results in irreversible damage to the IVD, rendering it incapable of recovery. Consequently, these models may be unsuitable for the study of early-stage IVD degeneration. Another study using the dynamic compression model of disc degeneration incorporates factors of nutritional limitation, unsuited to accurately assess the impact of mechanical stress on disc degeneration[25]. Additionally, the model necessitates daily compression-induced damage, diverging from the natural human response to disc injury, thereby limiting the observation of therapeutic effects on discs following the cessation of injury.

There are few studies on the effects of repeated injuries on IVD degeneration. Jill et al. used a no. 11 surgical blade to puncture and damage rat caudal IVDs on Days 1, 3 and 6 of the experiment and found that repeated injuries resulted in a continuous inflammatory response and persistent degeneration of IVDs[18]. As the integrity of the IVD becomes damaged, the NP loses its protective isolation environment and triggers a molecular cascade immune response. Therefore, this model is not suitable to explore the early impact of repeated injury on IVD degeneration. In this study, we used the strike loading IVD injury model from previous reports and performed repeated strike loading injuries to the IVD[19, 22]. Since the heights of the IVDs in each group did not change significantly after swelling, the strike loading IVD degeneration model used in this experiment has maintained relatively normal mechanical properties. This disc degeneration model induced by strike loading, is suitable for studying the early stages of disc degeneration, especially in the absence of significant mechanical stable structure destruction. In the study of IVDs with severe degeneration, the endplate or AF of the IVD organ model should be destroyed to simulate the structural defect and mechanical instability of the disc during severe IDD.

In the context of IVD degeneration, it is observed that a reduction in hydration is a concomitant phenomenon of the process, contrasting with the high hydration state inherent to
a healthy disc[32]. The water content in healthy NP tissue can be as high as 90%, and in AF tissue, it is approximately 65-70%[33, 34]. The difference in GAG content was the main reason for the difference in water content between the two tissues. GAGs are negatively charged and have a fixed charge density, attracting positive charges into the NP matrix[35]. Therefore, the NP can absorb much more water than its weight and lock the water molecules through polar interactions in the tissue, which makes the IVD in vitro culture have a strong tendency to swell and cause swelling stress. During the disc culture cycle in this study, neither mechanical stress injury mode resulted in disc height reduction after swelling culture. Since GAGs are primarily located in the center of the IVD, the narrowing of the space caused by the catabolism of the small amount of GAGs is filled by additional water absorbed by the remaining GAGs, resulting in no significant change in IVD height. Andreas et al. also employed a bovine organ culture model with a bioreactor and reported a significant loss of relative IVD height in the model groups after one week of loading[23]. They viewed that IVD height losses as a sign of IDD.

These inferences are supported by the DCS of the discs. Measurements of the DCS of the discs can indirectly reflect the changes in GAG contents in tissues[36]. The IVD GAG content was directly correlated with tissue swelling stress. As the amount of IVD proteoglycan decreases with age and degeneration and the total fixed charge density of the tissue decreases, leading to a decrease in water retention. The swelling stress of degenerative discs was significantly lower than that of normal disc tissue[36]. In this experiment, we measured the DCS of the discs organ. DCS is a measure of the average amount of stress generated by the disc in the 10% range of deformation. The results showed that both mechanical stress injury modes reduced the DCS of the discs. The dynamic stiffness of the physiological CON group on the 11th day increased compared with that on the 1st day. This may be related to the disc organ absorbing more water in the in vitro medium, leading to an increase in dissolution stress and subsequently an increase in dynamic stiffness.

There are several limitations that should be highlighted in the present research. First, the disc organ culture model did not incorporate blood vessels or the immune system, both critical components in disc degeneration processes. The absence of a vascular and immune response in organ culture models underscores the necessity for a multifaceted research approach to fully elucidate the mechanisms of disc degeneration. Second, the structural integrity of the IVD is preserved, with the NP remaining encapsulated within the AF. This model, both from a mechanical and biological perspective, is specifically suited for investigating responses
associated with the early stages of IVD degeneration. In addition, despite their applicability in simulating intervertebral disc degeneration, bovine tail and human intervertebral discs differ significantly in anatomy, biomechanics, cellular composition, and degeneration manifestations. These differences must be considered when using bovine discs as models. Finally, the IVDs were not subjected to extended culturing periods, precluding the exploration of prolonged degenerative mechanisms. Future research could benefit from focusing on long-term culture studies to elucidate the progressive degenerative processes and explore potential regenerative therapies.

CONCLUSION

In conclusion, our results provide evidence that repeated strike loading could induce degenerative changes in the IVD. As continuous mechanical stress is required to induce long lasting molecular changes, repeated strike loading is apt for studying chronic IDD. The results also indicated that getting out from abnormal mechanical condition is a very effective way to halt or even reverse IDD progression. In addition, we found that AF is more susceptible to mechanical stress, this may provide a new perspective on IDD research.

ACKNOWLEDGMENTS

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homeostasis throughout the annulus fibrosus: A finite element analysis of healthy and
TABLE 1. Primers used for qPCR.

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<th>Gene ID</th>
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<td>286806</td>
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(F)

- **CON**
- **OS**
- **RS**
- **Deg**

IVD height measured on day 1 and day 11

- 40% strain in one second, Day 1
- 35% strain in one second, Days 1, 4 and 7
- Physiological loading 20–200 KPa; 0.2 Hz; 1 hr/day
- Degenerative loading 320–500 KPa; 5 Hz; 2 hr/day

IVD height measured on day 1 and day 11

- Cultured in high glucose (4.5 g/L) medium
- Cultured in low glucose (2 g/L) medium
<table>
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(G)

**Figure 1.** (A) Customized chamber (B) the parts of the customized chamber (C) custom designed bioreactor (D) specially designed mechanical testing machine (E) Representative sinusoidal stress curves of the IVDs physical loading pattern using the custom designed bioreactor. (F) Flowchart of grouping IVDs into different models. (E) The culture time and the outcome parameters. CON: control group, OS: one strike group, RS: repeated strike group, Deg: degenerative group.
**Figure 2.** IVD cell death was induced by strike loading. LDH/EthD-1 staining was employed to assess cell viability. In the representative images, viable cells appear in blue or blue/red, whereas dead cells are shown in red. CON: control group on day 11, OS: one strike group on day 11, Deg: degenerative group on day 11, RS day 9: repeated strike group measure on day 9, RS day 11: repeated strike group measure on day 11 (A). Cell viability in the IVD cells is detailed as follows: (B) represents the NP region, and (C) represents the AF region. n = 4, means ± SEM, Scale bars: 50 µm. *P < 0.05, **P < 0.01.
Figure 3. The expression levels of apoptosis-related proteins were examined using immunofluorescence staining. (A-B) Represent NP region, (C-D) Represent AF region. (E) Relative expression levels of apoptosis-related proteins. CON: control group on day 11, OS: one strike group on day 11, Deg: degenerative group on day 11, RS day 9: repeated strike group measure on day 9, RS day 11: repeated strike group measure on day 11. Scale bars, 100 μm. n = 3, means ± SEM, *P < 0.05, **P < 0.01.
**Figure 4.** The genes expression levels in NP tissue were examined using qPCR. CON: control group on day 11, OS: one strike group on day 11, Deg: degenerative group on day 11, RS day 9: repeated strike group measure on day 9, RS day 11: repeated strike group measure on day 11. n = 3, means ± SEM, *P < 0.05, **P < 0.01.
Figure 5. The genes expression levels in AF tissue were examined using qPCR. CON: control group on day 11, OS: one strike group on day 11, Deg: degenerative group on day 11, RS day 9: repeated strike group measure on day 9, RS day 11: repeated strike group measure on day 11. n = 3, means ± SEM, *P < 0.05, **P < 0.01.
Figure 6. Relative cumulative release of GAGs into IVD culture medium. CON: control group on day 11, OS: one strike group on day 11, Deg: degenerative group on day 11, RS: repeated strike group measure on day 11. n = 3, means ± SEM, *P <0.05.
Figure 7. The microstructure of the IVD was examined using Safranin O-Fast Green. Black arrow indicates cleft. Day 0: on the day that IVDs were harvested, CON: control group on day 11, OS: one strike group on day 11, Deg: degenerative group on day 11, RS: repeated strike group measure on day 11.
Figure 8. (A) IVDs height changes after different culture conditions. In loading culture conditions, IVDs in the CON, OS and RS group were measured after physiological loading (20–200 KPa; 0.2 Hz; 1 hr), IVDs in the Deg group were measured after degenerative loading (320–500 KPa; 5 Hz; 2 hr/day). (B) Changes in the DCS of the IVDs after 11 days of culture. DCS measured on Day 1 served as the standardized baseline. CON: control group, OS: one strike group, Deg: degenerative group, RS: repeated strike group measure. n = 3, means ± SEM, *P < 0.05, **P < 0.01.