A Comparison of Two Ovine Lumbar Intervertebral Disc Injury Models for the Evaluation and Development of Novel Regenerative Therapies

Chris D. Daly, MBBS, MPhil, PhD\textsuperscript{1,2}, Peter Ghosh, DSc, PhD, FRSC\textsuperscript{1,3}, Tanya Badal, MSc\textsuperscript{4}, Ronald Shimmon, PhD\textsuperscript{4}, Graham Jenkin, PhD\textsuperscript{1}, David Oehme, MBBS, PhD, FRACS\textsuperscript{5}, Justin Cooper-White, PhD, BE (Chem)(Hons)\textsuperscript{6}, Idrees Sher, BAappSc-MRS(DR), BMedSc, MBBS(Hons)\textsuperscript{1,2}, Ronil V. Chandra, MBBS, FRANZCR\textsuperscript{1,2}, and Tony Goldschlager, MBBS, PhD, FRACS\textsuperscript{1,2}

Abstract

Study Design: Large animal research.

Objective: Lumbar discectomy is the most commonly performed spinal surgical procedure. We investigated 2 large animal models of lumbar discectomy in order to study the regenerative capacity of mesenchymal stem cells following disc injury.

Methods: Twelve adult ewes underwent baseline 3-T magnetic resonance imaging (MRI) followed by lumbar intervertebral disc injury by either drill bit (n = 6) or annulotomy and partial nucleotomy (APN) (n = 6). Necropsies were performed 6 months later. Lumbar spines underwent 3-T and 9.4-T MRI prior to histological, morphological and biochemical analysis.

Results: Drill bit-injured (DBI) and APN-injured discs demonstrated increased Pfirrmann grades relative to uninjured controls (\(P < .005\)), with no difference between the 2 models. Disc height index loss was greater in the APN group compared with the DBI group (\(P < .005\)). Gross morphology injury scores were higher in APN than DBI discs (\(P < .05\)) and both were higher than controls (\(P < .005\)). Proteoglycan was reduced in the discs of both injury models relative to controls (\(P < .005\)), but lower in the APN group (\(P < .05\)). Total collagen of the APN group disc regions was higher than DBI and control discs (\(P < .05\)). Histology revealed more matrix degeneration, vascular infiltration, and granulation in the APN model.

Conclusion: Although both models produced disc degeneration, the APN model better replicated the pathobiology of human discs postdiscectomy. We therefore concluded that the APN model was a more appropriate model for the investigation of the regenerative capacity of mesenchymal stem cells administered postdiscectomy.

Keywords

animal model, intervertebral disc, discectomy, regeneration

Introduction

Lower back pain causes more global disability than any other condition worldwide.\textsuperscript{1} Lower back pain commonly results from degenerative lumbar disc disease causing discogenic pain.\textsuperscript{2} Lumbar disc degeneration is a complex process manifested by changes in cellular, matrix, endplate, and the neurovascular components of the intervertebral disc. Intervertebral disc herniation is a common outcome of lumbar disc degeneration, while lumbar discectomy is the most commonly performed

\textsuperscript{1} Monash University, Clayton, Victoria, Australia
\textsuperscript{2} Monash Medical Centre, Clayton, Victoria, Australia
\textsuperscript{3} Proteobioactives, Pty Ltd, Sydney, New South Wales, Australia
\textsuperscript{4} University of Technology Sydney, Broadway, New South Wales, Australia
\textsuperscript{5} St Vincent’s Hospital, Fitzroy, Victoria, Australia
\textsuperscript{6} University of Queensland, St Lucia, Queensland, Australia

Corresponding Author:
Chris D. Daly, The Ritchie Centre, Hudson Institute of Medical Research, Monash University, 246 Clayton Road, Clayton, Victoria, 3168, Australia.
Email: chrisddaly@gmail.com

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spinal surgical procedure.3 Lumbar discectomy successfully treats radicular symptoms associated with neural compression in more than 80% of patients.4 However, the procedure fails to address the underlying pathophysiology of intervertebral disc degeneration responsible for the syndrome. Moreover, following lumbar discectomy up to one-third of patients report low back pain.5 In addition, up to 18% of patients experience recurrent disc herniation6 with 12% undergoing reoperation within 4 years.7 Ultimately, 40% of these patients will undergo spinal fusion.7

Given the significant disease burden resulting from disc degeneration and lower back pain numerous animal models have been developed to further understand the pathobiology of disc degeneration and examine potential modalities for its treatment.8 There are, however, relatively few reports of large animal models of lumbar discectomy.9-12 Given the clinical ubiquity of discectomy, the inherent anatomical challenges to disc repair and the opportunity presented to initiate regenerative therapy at the time of surgical intervention, we sought to develop a suitable large animal model of discectomy that could be used to evaluate potential tissue regenerative therapies, such as transplantation of stem cells.

Limited annular injury to ovine discs have been widely used to generate a model of disc degeneration.13-15 The ovine species has also been used to test implant devices and in the preclinical investigation of cellular therapies to support spinal fusion and disc reconstitution.9,16-18 Ovine discs, like human discs, undergo chondroid metaplasia with skeletal maturation,19 because of the loss of their notochordal cell remnants.19,20 Additionally, the ovine disc is closer in size and cellular phenotype20 to the human intervertebral disc than small animal models, important characteristics given the nutritional limitations associated with the central regions of the disc. Furthermore, despite its quadrupedal conformation, the sheep spine has been shown to exhibit significant biomechanical similarities to the human spine.21

We have previously described the use of an ovine annulotomy and partial nucleotomy (APN) model for investigation of the potential of mesenchymal progenitor cells (MPCs) formulated with the pharmaceutical agent, pentosan polysulfate (PPS), embedded in a biodegradable gelatin scaffold to promote intervertebral disc regeneration following lumbar discectomy in a pilot study.3 In this APN model, a full-thickness 3 × 5 mm annulotomy was performed with a scalpel blade and 200 mg of annular and nuclear tissue removed with a pituitary rongeur. PPS was used as it was known to enhance MPC viability and promote their differentiation to a chondrogenic phenotype whilst also inhibiting osteogenesis.22 Our pilot study demonstrated the feasibility of the modified APN model and provided positive outcomes on the efficacy of the MPC + PPS formulation. However, prior to further investigations of other potential therapeutic modalities of lumbar disc repair that required the use of a liquid hydrogel, we sought to determine the most appropriate large animal model for such applications.

An earlier publication by Zhang et al23 reported that disc degeneration could be induced in goat lumbar discs by using a drill bit to penetrate the annulus fibrosus (AF) through to the nucleus pulposus (NP). Using a subjective histological grading system, this model was reported to provide more reliable degenerative changes than insertion of a horizontal surgical blade along the same path. In principle, the Zhang et al23 model offers advantages in facilitating the injection of regenerative liquid hydrogels/cell combinations into the disc without the use of a solid scaffold, which was a requirement of using the Oehme discectomy model.9 However, the study of Zhang et al23 was performed in goats and did not include biochemical analysis of the injured intervertebral discs, thereby limiting the ability to directly compare these 2 models directly.

The drill bit injury (DBI) model produces injury of both the annulus and nucleus. We hypothesized that modification of this model by increasing the depth of penetration to the midpoint of the nucleus pulposus and using the maximum drill bit diameter without inducing endplate injury, that is, slightly less than disc height, may provide a readily replicable model of the postdiscectomy intervertebral disc. This model has the advantage of being a single step procedure with a highly standardized annular and nuclear defect. Furthermore, the ovine DBI model could also serve as a model of intervertebral disc herniation, in which nuclear injury is associated with a full thickness annular injury. In the present study, we evaluated the APN and drill bit methods of surgically inducing disc failure using a homogeneous group of adult sheep and monitoring the relative outcomes 6 months later using both subjective and objective methods of assessment.

Material and Methods

Surgical Procedure

With ethics approval from the Monash Medical Centre Animal Ethics Committee and conforming to the Australian code of practice for the care and use of animals for scientific purposes (eighth edition, 2013), 12 adult (2–4 years of age) Border Leicester Merino cross-bred ewes underwent preoperative 3-T magnetic resonance imaging (MRI; Siemens Skyra Widebore 3T MRI, Siemens, Erlangen, Germany) under general anesthetic. Ewes were used in this study because of their better temperament than males—castrate or intact. Sheep were raised in open pastures and ambulated freely prior to the trial. All sheep were fasted for 24 hours prior to surgery and anesthetized using intravenous thiopentone (10–15 mg/kg) (Bayer Australia Ltd, Pymble, New South Wales, Australia) followed by intubation and isoflurane inhalation (Pharmachem, Eagle Farm, Queensland, Australia) (2%-3% in oxygen). Sheep were placed in the right lateral position. Local anesthetic (bupivacaine 0.5%) (AstraZeneca Australia, Macquarie Park, New South Wales, Australia) was administered subcutaneously and the L2-3 and L3-4 lumbar intervertebral discs exposed via left lateral retroperitoneal approach, as previously described.24,25 Intraoperative lateral radiographs (Radlink, Atomscope HF200A, Redondo Beach, CA, USA) were performed to confirm the correct levels. Six sheep underwent microdiscectomy.
APN injury, performed by the creation of a 3 mm × 5 mm annular window followed by disc resection using pituitary rongeurs. The disc tissues collected (200.0 ± 3.0 mg) consisted mainly of AF with some NP. The adjacent L1-2 and L4-5 discs served as untreated controls. DBI was performed on the L2/3 and L3/4 intervertebral discs of the remaining 6 sheep using a 3.5-mm Brad point drill bit (Carbatec, Melbourne, Victoria, Australia) with a drill bit stop applied at 12 mm drill bit length (Drill Warehouse, Amazon, Seattle, WA, USA) as described previously.[25]

Following intervertebral disc injury, the wound was closed using a routine layered procedure performed using absorbable sutures (Vicryl, Ethicon, Somerville, NJ, USA). Animals received a fentanyl patch (Duragesic 75 μg/h, Jannsen LLC., North Ryde, New South Wales, Australia) and intravenous paracetamol (Pfizer Ltd, West Ryde, New South Wales, Australia) for postoperative analgesia. Following recovery, animals were returned to the pen with other sheep and allowed free ambulation. Sheep were returned to open pasture 1 week postsurgery.

**Necropsy**

Six months postsurgery, animals were euthanized by intravenous injection of 150 mg/kg of pentobarbital (Sigma-Aldrich, Castle Hill, New South Wales, Australia). The lumbar spines were then removed en bloc, a segment was isolated from the mid-sacrum to the thoracolumbar junction and transferred to Monash Biomedical Imaging for MRI analysis. Spinal columns were then transected in the horizontal plane through their vertebral bodies, using a band saw, to provide spinal segments consisting of a complete lumbar disc with half of the adjacent vertebral bodies attached. Subsequent gross morphological, biochemical, and histological analysis of discs were undertaken using these spinal segments as described below. Spinal segments containing discs destined for histological analysis were transferred to phosphate buffered formalin.

**Radiological Analysis**

Using standardized methods, disc height index (DHI) measurements were calculated and recorded by an observer blinded to the treatment regimen, using standard digital processing software (Osiris MD v8.0.2, Pixmeo, Geneva, Switzerland). Sagittal 3-T (Siemens Skyra Widebore 3T MRI, Siemens, Erlangen, Germany) T2-weighted MRI sequences of the entire lumbar spine explant were obtained for each animal. Axial 9.4-T (Agilent 9.4T MRI Small Animal Scanner Agilent/Varian, Santa Clara, CA, USA) T1 and T2 MRI sequences of the control and intervention lumbar intervertebral discs were taken for each animal. Using sagittal 3-T T2-weighted sequences and 9.4-T T2 sagittal reconstructions (Osiris MD v 8.0.2), 4 blinded observers (a neuroradiologist, neurosurgeon and 2 neurosurgery residents blinded to the treatment regimen) determined the Pfirrmann MRI disc degeneration scores for all lumbar discs.

**Biochemical Analysis**

After collection of the digital images of discs for morphological analysis all tissue regions shown in Figure 1 were subjected to biochemical analysis. The individual AF and NP from each region were separated from each other and their vertebral attachments by careful dissection using the boundaries shown in Figure 1. Tissues from each region were finely diced, frozen in liquid nitrogen, and powdered in a liquid nitrogen-cooled ball-mill. The powdered tissues were transferred to preweighed Eppendorf vials and weighed, lyophilized, and reweighed to constant weight to determine their anhydrous weights. Triplicate aliquots of the dehydrated tissues were solubilized using a papain digestion buffer (50 mM sodium acetate [pH 6.0]) containing 2 mg/mL papain (Sigma-Aldrich Chemicals, Sydney, New South Wales, Australia) by incubation at 60°C for 16 hours.[28] The digested tissues were centrifuged at 3000 × g for 15 minutes and supernatants diluted to standard volumes (the stock digest solution). Aliquots of the stock solution were analyzed for sulfated glycosaminoglycan (S-GAG) (an index of
proteoglycan content) levels using the dimethylmethylen blue (DMMB) assay and hydroxyproline assay (to derive collagen content). The results of biochemical analyses were normalized and were expressed as percentage of tissue dry weight for S-GAG and collagen.

**Histological Analysis**

The individual disc segments, consisting of the intervertebral disc with attached hemisected vertebral bodies were in 10% neutral buffered formalin for 8 days then stored in 70% ethanol. The volume of vertebral bone was reduced to the growth plate using a fine diamond saw. Prior to paraffin based tissue embedding, decalcification of the remaining vertebral bone was undertaken with multiple changes of 10% formic acid. Coronal paraffin sections of the entire disc segments for the APN sheep and axial sections for the drill injured sheep were cut using a standard rotary microtome and stained using hematoxylin and eosin. Axial sections were taken from the drill injury disc to allow for visualization of the entire DBI tract. The APN-injured discs were only subjected to standard coronal plane sectioning.

**Statistical Analysis**

All data analyses and storage were performed using Microsoft Excel for Mac (Version 15.33, Microsoft, Redmond, WA, USA) and Prism 7.0c for Mac (GraphPad Software Inc, La Jolla, CA, USA). Parametric data was analyzed using 1-way analysis of variance, and Tukey’s multiple comparison test was performed when significant differences in means were observed. Nonparametric data was analyzed using Kruskal-Wallis test of median values followed by Dunn’s multiple comparison test. Groups were compared using the 2-tailed Student’s t test followed by Mann-Whitney U tests. A P value <.05 was considered statistically significant.

**Results**

**Disc Height Index**

The APN-injured discs demonstrated significantly greater loss of height than the drill injured and control discs (P < .005). However, both the APN and drill injured discs demonstrated greater loss of height than control discs (P < .005) (Figure 2A).

Baseline preoperative 3-T MRIs of all animals revealed no evidence of underlying disc degeneration at control or intervention levels (L1/2 to L4/5). Scoring of drill bit– and APN-injured discs demonstrated significantly increased Pfirrmann grades relative to control discs (both P < .005); however, there was no significant difference in Pfirrmann grades between the 2 injury groups (Figure 2, A, B, and D).

**9.4-T MRI**

The horizontal images obtained by 9.4-T MRI allowed ready appreciation of the extent of annular disruption observed in the APN versus drill injured intervertebral discs (Figure 3, B and C) and correlated well with gross morphological observations. However, the 9.4-T MRI Pfirrmann grades were consistent with 3-T grades once sagittal reconstruction and grading was performed (Figure 2, B and D). Significantly increased Pfirrmann grades were observed in the drill- and APN-injured discs compared with control discs (P < .005) with no significant difference between the 2 injury models.

**Gross Morphology**

Gross morphological analysis was completed on a regional basis, with the data presented in Figure 4D showing the aggregate morphological scores. Representative gross morphological digital images are displayed in Figure 4A-C and their respective scores, determined using the criteria described in Table 1 and shown in Figure 4D. Drill injured discs generally demonstrated a more focal annular and nuclear tract injury (Figure 4B), with minimal NP disorganization, in comparison with the APN-injured discs, which showed more widespread
changes and blood degradation product staining (Figure 4C). Nonetheless, total disc gross morphological scores were significantly increased in both the drill bit– and APN-injured intervertebral discs relative to control ($P < .005$). APN-injured discs had significantly increased Pfirrmann grades relative to control discs; however, there was no significant difference in Pfirrmann grades between injury groups. APN, annulotomy and partial nucleotomy. *$P < .05$, **$P < .005$.

**Proteoglycan Content as Determined by Sulfated-Glycosaminoglycan (S-GAG) Analysis**

S-GAG content was significantly reduced in the injured (NP1) (Figure 5A) and contralateral NP region (NP2) (Figure 5B) in both the drill- and APN-injured discs relative to control ($P < .005$). APN-injured intervertebral disc gross morphological scores were, however, significantly elevated relative to drill-injured discs ($P < .05$) (Figure 4D).

**Collagen Content**

The collagen content of the APN injury site AF (AF1) was significantly higher than both the control and drill-injured discs ($P < .05$) (Figure 6A). Furthermore, the drill injury AF1 collagen content was significantly lower than control discs ($P < .05$). The AF adjacent to the injury site (AF4) also demonstrated a significant increase in collagen content in the APN group relative to controls ($P < .01$) (Figure 6B). The nucleus pulposus, both ipsilateral and contralateral to the injury site, demonstrated significantly higher collagen content in the APN-injured disc relative to both the control and drill-injured discs ($P < .005$ and $P < .05$, respectively) (Figure 6, C and D). These differences were also reflected in the total NP and total disc S-GAG content.
(NP + AF) collagen content of the APN group compared with both the drill-injured and control discs (Figure 6, E and F) ($P < .001$ and $P < .005$, respectively).

**Histology**

Qualitative histological analysis revealed differences between the injury models with regard to the disc ultrastructure and extent of vascular infiltration and granulation tissue deposition. Control discs demonstrate intact annulus fibrosi displaying multiple lamellae abutting the proteoglycan-rich nucleus pulposus (Figures 7A and 8A [axial] and Figures 7B and 8B [coronal]). Axial hematoxylin and eosin–stained slides of drill-injured disc demonstrate the focal nature of the disruption of AF and NP tissues induced by this injury model (Figure 7, C and D). Relatively little vascular infiltration is evident. Reduced proteoglycan content is evident on the drill-injured disc (Figure 8C) relative to the control disc (Figure 8A) on the Safranin O and Fast Green–stained slides. The APN-injured disc (Figure 7E) demonstrates extensive disruption of lamellae with deposition of granulation tissue and infiltration within the injury site. High-power microscope images highlighted the extent of vascular proliferation occurring at the injury site of the AF (Figure 7F). Marked reduction in proteoglycan content is also evident at the APN injury site (Figure 8D).

**Discussion**

The results of the present study demonstrated that both the modified APN injury model originally described by Oehme et al$^9$ and a modification of the DBI model described by Zhang et al$^{23}$ induced degeneration in sheep lumbar discs 6 months following the surgical intervention. Furthermore, as assessed from the disc height indices, gross morphological, biochemical, and histological analyses, the APN injury provoked more extensive degenerative changes in injured discs, than observed with the DBI procedure.

We suggest that the extent and nature of the degenerative changes induced in the discs of each model was determined by the relative magnitude of several time-dependent mechanical and biological events that are known to dictate the pathology of disc degeneration.$^{31-36}$ For example, surgical compromise of AF hoop stress tensile function and reduction in NP hydroelasticity would impose an immediate disturbance in the nature of

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**Figure 3.** Axial magnetic resonance images (9.4-T) (A) Control disc demonstrating intact AF and hyperintense NP. (B) Drill injury disc demonstrating injury tract (arrow) extending to NP. (C) APN disc demonstrating broader injury tract (arrow) and decreased NP hyperintensity. (D) 9.4-T MRI Pfirrmann grades: drill-injured and APN-injured discs demonstrated significantly higher 9.4-T Pfirrmann grades than control discs with no significant difference between injured groups. AF, annulus fibrosus; NP, nucleus pulposus; APN, annulotomy and partial nucleotomy. $^*P < .05$, $^{**}P < .005$. 

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the mechanical stresses acting across the spinal unit and its adjacent structures. A secondary event elicited, would be an inflammation and an early repair response at the site of injury, with proliferation of the capillaries serving the outer AF, accompanied by deposition of granulation and fibrotic tissue within the defect site. Such events have been described in other animal models and are the sequelae of human disc herniation.

Loss of proteoglycan and its associated water is a relatively early event in the pathophysiology of intervertebral disc degeneration. The loss of these matrix components leads to marked changes in the mechanical properties of the intervertebral disc and adjacent structures, as intradiscal pressure is reduced and the ability of the disc to tolerate compressive loading diminishes. Biomechanical studies have demonstrated such changes in the mechanical behavior of discs that were proportional to the size of the annular defects.

The compromise of AF integrity and loss of NP material initiated in the APN model closely mirrors the clinical condition of symptomatic lumbar intervertebral disc herniation. In the clinical setting, radicular symptoms are caused by herniated intervertebral disc material producing neural compression. This herniated AF and NP, and any additional loose NP, is excised during conventional lumbar microdiscectomy. The creation of a full thickness APN, with removal of 200 mg of annular and nuclear tissue, differentiates this ovine APN model from those recently reported by Shu et al and most other ovine annular injury models in the literature in which partial thickness annular incisions are performed. Such models may be more representative of spontaneous intervertebral disc degeneration, in which annular tears are often observed, than the post–lumbar discectomy intervertebral disc.

The model of Zhang et al demonstrated the ability of the DBI to induce disc degeneration 2 months following surgery, confirmed histologically but not biochemically. Furthermore, the degeneration observed histologically did not produce correlative MRI changes such as increased 1.5-T MRI Pfirrmann grades. Many prior studies of the intervertebral disc, including that of Zhang et al, were performed with 1.5-T MRI scanners. With the advance of technology, 3-T MRI scanners are commonly available in the clinical setting and the 9.4-T MRI scanner is now available for preclinical studies. Despite our use of both 3-T and 9.4-T MRI the MRI Pfirrmann grade findings did not directly parallel our histological, biochemical,
The increase in collagen observed in the injury site AF and NP of the APN-injured disc reflects the normal tissue response to traumatic injury, with inflammation, capillary invasion, and fibrotic granulation tissue deposition representing key matrix events. The repair tissues that normally accumulate at these early injury sites, consist mainly of type I collagen, and are accompanied by the loss of proteoglycans as was observed for the APN model but not drill bit model. In healthy discs and the early stages of disc degeneration, the chondrocyte-like cells of the NP synthesize predominantly type II collagen. However, as the disc degeneration progresses, these cells undergo a transition and synthesize type I collagen and less proteoglycans. Therefore, it is possible that the endogenous NP cells of the injured discs in the APN model also contributed to the deposition of fibrotic tissue. Additional studies of the collagen subtypes resident in the various regions of the injured discs from the APN model are required to resolve this important question. The observed lack of a significant difference in collagen content between the drill-injured disc and control discs, with the exception of AF1, is indicative of the relatively low-

![Figure 5](image-url)

**Figure 5.** Sulfated glycosaminoglycan (S-GAG) content (% dry weight). (A) NP1 GAG is significantly lower in APN-injured than drill-injured discs, which is in turn less than control discs. (B) NP2 demonstrates the same relationship with APN-injured discs lower than drill-injured discs \((P < .05)\), which in turn are less than control discs. (C) NP Total GAG and (D) total disc GAG demonstrates the same series of relationships. APN, annulotomy and partial nucleotomy. *\(P < .05\), **\(P < .005\).

The failure of the MRI Pfirrmann grading system to adequately reflect differences in the extent of intervertebral disc degeneration has been observed for lower resolution 1.5-T and 3-T MRI scanners in both preclinical and clinical studies. We suggest that the discrepancy between the MRI and our other findings may be attributed to the relatively low sensitivity of conventional MRI Pfirrmann grades for detecting subtle degenerative differences relative to other quantitative and semiquantitative modalities. The 9.4-T MRI axial images (Figure 3) provided superior resolution of anatomical detail compared to conventional 1.5-T and 3-T MRI scanners; however, this additional data was not captured in the Pfirrmann grading system when the axial images were converted to the sagittal plane. Advanced quantitative radiological methods, such as T1-weighted MRI, may provide a more sensitive method of measuring early intervertebral disc degeneration radiologically using the clinical magnets currently available. Furthermore, our group is investigating novel methods to fully utilize the increased resolution this imaging modality affords.

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grade degeneration changes induced in these tissues by either surgical approach.

As a result of the extended time required to completely decalcify the vertebral bone, the quality of the disc tissue sections was unfortunately unacceptable for quantitative histological scoring. Nevertheless, all histological sections were reviewed qualitatively. On histological examination, vascular invasion was noted in the APN-injured discs that was relatively

Figure 6. Collagen content (% dry weight). (A) AF1 APN group discs had significantly more collagen than both control and drill-injured discs. (B) AF4 APN group collagen was significantly higher than control discs. (C) NP1 APN group collagen was significantly higher than both control and drill injury collagen. (D) NP2 collagen demonstrated the same pattern. (E) NP total also demonstrated an increase in collagen in the APN group relative to the control group. (F) Total disc collagen was significantly increased in the APN group relative to both control and drill-injured discs. AF, annulus fibrosus; NP, nucleus pulposus; APN, annulotomy and partial nucleotomy. *P < .05. **P < .005.
absent from the DBI discs. Vascular invasion is consistent with histological observations from herniated and degenerate intervertebral discs in the clinical context. In addition, the APN-injured discs demonstrated disorganization of the lamella pattern of the AF and adjacent NP. In contrast, the DBI disc demonstrated a focal lesion with minimal disruption of the AF and NP. The absence of significant tissue ingrowth observed in DBI specimens is in agreement with the lack of a significant increase in collagen content in the DBI AF.

There are, however, important differences between these preclinical models and the human intervertebral disc that contribute to the limitations of this study. A limitation common to large animal models of intervertebral disc degeneration is the inability to assess low back pain. The only animal models allowing for low back pain assessment are rodent models. Such models present challenges with regard to their differences to the human disc in size, anatomy, the presence of notochordal cells, and ultimately, translatability. Furthermore, in the clinical situation, the underlying disc demonstrates preexisting degeneration culminating in disc prolapse and neural compression. The models we present are of normal intervertebral discs subjected to injury to induce intervertebral disc degeneration. Discs with preexisting degeneration may have reduced capacity to repair or regenerate or may demonstrate a higher extent of
degeneration following injury induction than the previously healthy discs included in this trial. This is an important consideration in the context of investigating regenerative therapies for disc degeneration in the future.

**Conclusion**

The ovine intervertebral disc drill bit and APN injury models both produce intervertebral disc degeneration at 6 months following injury. However, the severity of degeneration associated with the APN model was greater, as assessed by MRI, gross morphology, biochemical and histological analysis, than the drill bit model. Moreover, the ovine APN injury model better replicated the postdiscectomy lumbar intervertebral disc mechanistically and importantly, more closely reproduced the known pathology of disc tissues examined post–human disc herniation.52,53

On the basis of the aforementioned findings, we concluded that the ovine APN model provides a more suitable animal model for the evaluation of novel cellular modalities that targeted disc repair than the drill bit model. In this regard, we have subsequently used the ovine APN model of microdiscectomy to compare the relative efficacy of 2 mesenchymal stem cell preparations in their ability to restore disc integrity 6 months postsurgery.26

**Declaration of Conflicting Interests**

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**Figure 8.** Sections stained with Safranin O and Fast Green. (A) Control disc (axial section) demonstrating intact annulus fibrosus and adjacent nucleus pulposus. (B) Control disc (coronal section) demonstrating the coronal plane. (C) Drill-injured disc demonstrating reduced Safranin O stain (indicative of reduced proteoglycan content) in the drill-injured region of the nucleus pulposus (arrow). (D) APN-injured disc demonstrating significant reduction in Safranin O stain in the injured region of the intervertebral disc (arrow). Scale bar = 200 μm. APN, annulotomy and partial nucleotomy. 


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