Research Article (Member)

Nondestructive, Indirect Assessment of the Biomechanical Properties of the Rat Intervertebral Disc using Contrast-Enhanced µCT

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Running Title: IVD Mechanics Correlate to CE-µCT

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Author contributions

All authors have made substantial contributions to the conception and design of the study, acquisition of data, and/or analysis and interpretation of data, as well as manuscript preparation and final approval of the submitted manuscript. TM takes responsibility for the integrity of the work as a whole.
Abstract

Mechanical characterization of the intervertebral disc involves labor-intensive and destructive experimental methodology. Contrast-enhanced micro-computed tomography is a nondestructive imaging modality for high-resolution visualization and glycosaminoglycan quantification of cartilaginous tissues. The purpose of this study was to determine whether anionic and cationic contrast-enhanced micro-computed tomography of the intervertebral disc can be used to indirectly assess disc mechanical properties in an *ex vivo* model of disc degeneration. L3/L4 motion segments were dissected from female Lewis rats. To deplete glycosaminoglycan, samples were treated with 0 U/mL (Control) or 5 U/mL papain. Contrast-enhanced micro-computed tomography was performed following incubation in 40% Hexabrix (anionic) or 30 mg I/mL CA<sup>4+</sup> (cationic) for 24 hours (*n*=10/contrast agent/digestion group). Motion segments underwent cyclic mechanical testing to determine compressive and tensile modulus, stiffness, and hysteresis. Glycosaminoglycan content was determined using the dimethylmethylene blue assay. Correlations between glycosaminoglycan content, contrast-enhanced micro-computed tomography attenuation and mechanical properties were assessed via the Pearson correlation. The predictive accuracy of attenuation on compressive properties was assessed via repeated random sub-sampling cross validation. Papain digestion produced significant decreases in glycosaminoglycan content and corresponding differences in attenuation and mechanical properties. Attenuation correlated significantly to glycosaminoglycan content and to all compressive mechanical properties using both Hexabrix and CA<sup>4+</sup>. Predictive linear regression models demonstrated a predictive accuracy of attenuation on compressive modulus and stiffness of 79.8-86.0%. Contrast-enhanced micro-computed tomography was highly
predictive of compressive mechanical properties in an *ex vivo* simulation of disc degeneration and may represent an effective modality for indirectly assessing disc compressive properties.

**Keywords**: intervertebral disc; contrast-enhanced micro-computed tomography; disc biomechanics; degenerative disc disease

**Introduction**

Intervertebral disc (IVD) degeneration, and its resulting disability, represent an enormous global health burden. Low back pain, which can result from degenerative disc disease (DDD), is responsible for over $200 billion in health expenditures and lost wages in the United States alone\(^1\). IVD degeneration and its root cause have been difficult to define. Genetic predisposition, aging, impedance of nutrient transport, changes to extracellular matrix (ECM) structure and composition, acute injury, and heavy loading have all been identified as potential factors in the onset and progression of DDD\(^2\). Here, we focus on one of the early biochemical indicators of DDD, - the loss of sulfated glycosaminoglycan (GAG) from the intervertebral disc (IVD). It is unknown whether GAG loss is a causative factor in DDD, or a symptom of degeneration initiated by some other physiological process. Regardless, depletion of negatively-charged GAG contributes to an irreversible degenerative cascade by reducing hydrostatic pressure in the nucleus pulposus (NP) and compromising axial load distribution throughout the IVD, leading to focal overloading, damage, and, ultimately, the progressive degeneration of IVD tissues\(^3; 4\).

The biomechanical properties of the IVD are largely dependent on the structure and function of the GAG-rich extracellular matrix (ECM). During normal activity, the spine is
subjected to highly variable axial loads – human L3/L4 discs have been reported to experience loads equivalent to 250% of total body weight – and the IVD is uniquely adapted to withstand this loading. A healthy IVD is adequately hydrated, containing up to 80% water in the NP, where the majority of GAG is concentrated. When axially compressed, the hydrostatic internal pressure in the NP redistributes the load radially to the collagen-rich lamellar fibers of the annulus fibrosus (AF), which resist radial tension, resulting in effective load transmission across the IVD. The inner AF contains a considerable concentration of GAG and is also capable of some direct resistance to axial compression via its own internal pressure. Studies of cadaveric IVDs have demonstrated significant correlations of GAG content to NP swelling stress and aggregate modulus, and decreases in GAG content are associated with increasing age and increasing IVD degeneration score. In preclinical models of IVD degeneration, IVD injury has been shown to reduce compressive properties, and GAG concentration in the NP has been shown to significantly correlate to compressive stiffness and range of motion in cyclic tension/compression tests.

Small animal models of DDD have played an increasingly important role in understanding pathology, as well as testing new regenerative medicine-based strategies. The experimental determination of IVD mechanical properties in these small animal models is both labor-intensive, and destructive. The process usually involves careful dissection to isolate the motion segment, irreversible embedding of the vertebral bodies, meticulous mounting on a materials testing system, and a potentially-lengthy testing protocol. Given the destructive and time-intensive nature of IVD mechanical testing, a minimally-invasive, rapid assessment of IVD mechanical properties is of significant interest for both preclinical and clinical applications.
Contrast-enhanced micro-computed tomography (CE-µCT) is a nondestructive tool for high-resolution visualization and GAG quantification of cartilaginous tissues\textsuperscript{13-17}. Our group has recently demonstrated the use of both anionic and cationic contrast agent-enhanced µCT for the 3D visualization and molecular characterization of the IVD\textsuperscript{18}, and we have previously shown that anionic CE-µCT is a sensitive tool for the \textit{ex vivo} assessment of DDD in a rabbit model\textsuperscript{14}. Lakin \textit{et al} have demonstrated a local correlation of CE-µCT attenuation to compressive modulus and coefficient of friction in articular cartilage, exhibiting the utility of CE-µCT for the mechanical characterization of cartilaginous tissues\textsuperscript{19-21}. GAG content plays a similarly important role in the compressive mechanics of articular cartilage and IVD tissue. However, compressive load distribution across the entire vertebra-disc-vertebra motion segment is much more mechanically complex compared to the local compression of articular cartilage tissue, and thus it is unknown whether a similar correlation between CE-µCT attenuation and compressive properties exists in the IVD. As such, the purpose of this study was to assess the use of CE-µCT to nondestructively characterize the mechanical properties of the IVD by establishing correlations of anionic and cationic CE-µCT attenuation to experimentally-determined biomechanical properties in healthy and degenerate rat lumbar IVDs.

\textbf{Methods}

\textit{Specimen Preparation and Enzyme-Mediated Digestion}

IACUC approval was not required for this study, as all described methodology was performed \textit{post mortem}. All specimens were obtained from animals used in an unrelated, IACUC-approved protocol. Skeletally mature, female Lewis rats aged 14 – 20 weeks (~200 – 220 g) were euthanized via CO\textsubscript{2} asphyxiation. Lumbar spines were dissected \textit{en bloc} and fresh frozen until dissection according to a previously described protocol\textsuperscript{18}. To ensure that the spines
would fit uniformly into a custom cutting jig, the transverse processes and pedicle fragments were removed using a die grinder equipped with a sanding drum attachment, which was able to remove bone without inducing concomitant fractures. Each specimen was then secured in the cutting jig and transverse cuts were made through the L3 and L4 vertebral bodies to isolate the L3/L4 motion segment. The cutting jig facilitated two parallel cuts through the vertebral bodies to ensure that applied loads were uniformly axial and consistent between specimens. Specimens were then randomized to one of two contrast agent groups (Hexabrix and CA\textsuperscript{4+}) and further randomized to one of two enzymatic digestion groups (0 or 5 units/mL) (n=10 per group per enzyme concentration). Hexabrix (ioxaglate) was acquired commercially (Hexabrix 320, Guerbet, Inc., Bloomingdale, IN, USA), while CA\textsuperscript{4+} was synthetically prepared using a protocol adapted from Joshi et al\textsuperscript{22} (Supplemental Information). To deplete GAG content, enzymatic digestion was induced by incubation in phosphate-buffered saline (PBS, pH = 7.4) containing 0 or 5 units/mL papain (Sigma-Aldrich, St. Louis, MO, USA) for 24 h at 37°C, as previously described\textsuperscript{18, 23, 24}. In an earlier study, 5 units/mL papain was determined to induce significant reduction in GAG concentration while leaving the NP and AF structures intact and distinguishable via \( \mu \)CT and gross observation under a microscope\textsuperscript{18}.

\textbf{Contrast Agent Incubation and \( \mu \)CT Imaging}

Contrast agent solutions containing 40% Hexabrix (Guerbet, LLC, Bloomington, IN, USA) and 30 mg I/mL CA\textsuperscript{4+} were prepared according to a previously-described protocol\textsuperscript{18}. These contrast agent concentrations were previously determined to provide optimal contrast enhancement for morphological and compositional analysis of rat IVDs. Immediately following enzyme-mediated digestion, specimens were incubated in contrast agent for 24 h at room temperature. The motion segments were secured in a custom sample holder containing
humidifying silica beads to maintain a 70% humid environment, and the IVDs were imaged on a 
µCT imaging system (µCT-40, Scanco Medical AG, Brüttisellen, Switzerland) at 55 kVp, 145 
µA, 250 ms integration time, with an isotropic voxel size of 20 µm. The cross-sectional area 
(CSA) of each IVD was calculated from a three-dimensional (3D) volume of interest (VOI) 
manually contoured by an experienced investigator (SEH, KG) using a custom MATLAB 3-
plane viewing interface. Attenuation data was calculated from the IVD VOI along with an 
additional NP VOI that was manually contoured. Contours were checked for accuracy by two 
additional investigators (MDN, TM). Individual disc heights were calculated using a custom 
MATLAB program to manually delineate the endplate borders and calculate the average distance 
between consistent anatomical locations.

Mechanical Testing and Data Analysis

Non-destructive, uniaxial, cyclic compression-tension testing was performed to determine 
the mechanical properties of the IVDs. Using cyanoacrylate, motion segments were secured 
between rigid, parallel aluminum platens on a materials testing system equipped with a 250 N 
load cell (Insight 5, MTS, Eden Prairie, MN). Each end of the motion segment was secured 
within a shallow trough within each platen, enabling the cyanoacrylate to pool around the ends of 
the specimen and create a pot to ensure secure fixation. Preliminary testing confirmed that this 
fixation scheme was able to support both compressive and tensile loads at up to double those 
experienced during testing. During mechanical testing, the IVDs were immersed in room 
temperature saline supplemented with 1X protease inhibitors and 1% penicillin/streptomycin 
with fungizone (Thermo Fisher Scientific, Waltham, MA, USA). Mechanical loading was 
performed as previously described\(^4;\,\,\,12;\,\,\,25\). A 1 N preload was applied for 30 minutes, followed by 
20 sinusoidal loading cycles from 4.5 N compression to 3 N tension at 0.1 Hz. Data from the 20\(^{th}\)
cycle was analyzed. Load and extension values were converted to stress and strain using individually-determined IVD CSA and height. Compressive and tensile moduli were defined as the slope of the linear compressive and tensile regions of the stress/strain curve, respectively, which were isolated using MATLAB. An algorithm was employed which facilitated manual selection of the compressive and tensile regions of the $20^{th}$ testing cycle, as well as the linear region of each stress/strain curve. The neutral point delineating the compressive and tensile regions of the cycle was defined as the midpoint between the two points on the cycle at which zero stress was recorded.\textsuperscript{26} Compressive, tensile, and total hysteresis was calculated as the area under the stress/strain curve for the respective portion of the cycle.

**Biochemical Quantification of GAG Content**

Prior to GAG quantification, each IVD was rinsed for 20 h in room-temperature PBS to remove residual contrast agent. IVDs were then meticulously dissected from the vertebral endplates using microscopy-guided dissection and lyophilized for 2 hours. The desiccated IVDs were kept in storage at -80°C prior to batch processing using a 1,9-dimethylmethylene blue (DMMB) assay. IVDs were digested in a 50 µg/ml proteinase K solution for 20 hours and the DMMB assay was performed as previously described.\textsuperscript{18} A standard curve was generated using solutions containing 0-70 µg/mL chondroitin sulfate (CS).

**Statistical Analysis**

All statistical analysis was performed using SPSS (v22.0, IBM, Armonk, NY, USA). Correlations between variables were assessed using the Pearson product moment correlation. To assess differences between testing groups, a two-way analysis of variance (ANOVA) was applied as follows: normality and equality of variances were assessed using Shapiro-Wilk and
Levene’s tests, respectively. There were no significant violations of normality. However, several variables exhibited highly unequal variances. Variables which met the assumption of equal variances were assessed using a standard two-way ANOVA. Variables which significantly violated the assumption of equal variances underwent log transformation, were reassessed to confirm that the transformed data exhibited equality of variances, and then the transformed data was analyzed via two-way ANOVA. Log transformation was performed on the following variables: IVD, NP, and AF attenuation, tensile stiffness, tensile modulus, total strain, and compressive, tensile, and total hysteresis. Regardless of whether log transformation was employed for statistical analysis, all aggregate data was reported in terms of the untransformed mean and 95% confidence interval.

The predictive accuracy of contrast-enhanced attenuation on compressive mechanical properties was assessed across 100 iterations of repeated random sub-sampling cross validation. At each iteration, datasets from each contrast agent were randomly split into evenly-sized training and validation sets (n=10 training samples; n=10 validation samples). Within each set, there was an even distribution of undigested and digested samples to avoid model bias. The mean absolute percentage error (MAPE) between predicted and actual values was calculated as follows:

$$MAPE = \frac{100}{n} \sum \left| \frac{y_{pred} - y_{exp}}{y_{exp}} \right|$$

MAPE was averaged across the 100 iterations of cross validation and reported for each model combination.

Results
There were no significant differences in CSA (Hexabrix: 7.51 [7.36, 7.65] mm\(^2\); CA\(^{4+}\): 7.60 [7.35, 7.84] mm\(^2\), \(P = 0.526\)) or post-preload IVD height (Hexabrix: 1.04 [0.98, 1.09] mm; CA\(^{4+}\): 1.04 [0.99, 1.08] mm, \(P = 0.945\)) between samples in the Hexabrix and CA\(^{4+}\) groups. Furthermore, there were no significant differences in CSA between undigested (0 U/mL papain) and digested samples (5 U/mL papain) in the Hexabrix group (0 U/mL: 7.60 [7.38, 7.81] mm\(^2\); 5 U/mL: 7.41 [7.19, 7.65], \(P = 0.362\)) or CA\(^{4+}\) group (0 U/mL: 7.70 [7.35, 8.05] mm\(^2\); 5 U/mL: 7.49 [7.09, 7.88] mm\(^2\), \(P = 0.263\)). Within the Hexabrix group, post-preload IVD height was significantly lower in digested IVDs in the Hexabrix group (0 U/mL: 1.10 [1.06, 1.13] mm; 5 U/mL: 0.98 [0.95, 1.01] mm, \(P < 0.001\)), but no significant differences were observed within the CA\(^{4+}\) group, although digested IVDs exhibited slightly lower mean height (0 U/mL: 1.05 [1.01, 1.09] mm; 5 U/mL: 1.02 [0.96, 1.08] mm, \(P = 0.224\)). Enzymatic digestion induced significant decreases in total GAG content in both contrast agent groups (Table 1). As expected, corresponding changes in the attenuation profile of the whole IVD and the NP were measured due to digestion in both contrast agent groups, with IVDs in the Hexabrix group exhibiting significant increases in attenuation and IVDs in the CA\(^{4+}\) group exhibiting significant decreases in attenuation (Table 1). In the Hexabrix group, digested IVDs exhibited 36.6% higher whole IVD attenuation and 33.6% higher NP attenuation compared to undigested IVDs (Figure 1A, B). In the CA\(^{4+}\) group, digested IVDs exhibited 28.5% lower whole IVD attenuation and 31.5% lower NP attenuation compared to undigested IVDs (Figure 1C, D). Enzymatic digestion induced significant changes in the mechanical properties of IVDs in both contrast agent groups; digested IVDs had significantly lower compressive modulus and stiffness, tensile modulus and stiffness, and significantly higher compressive and total hysteresis compared to undigested IVDs (Table
2). Digested IVDs in the Hexabrix group also exhibited significantly higher tensile hysteresis, though this difference was not observed in the CA$^4+$ group (Table 2).

Strong linear correlations were observed between GAG content, µCT attenuation, and mechanical properties in both contrast agent groups. Total GAG content correlated significantly to whole IVD attenuation and NP attenuation in both Hexabrix and CA$^4+$ groups (Figure 2), and significantly to compressive modulus in both groups separately (Hexabrix: $r = 0.914, P < 0.001$; CA$^4+$: $r = 0.811, P < 0.001$) and when aggregating all IVDs together (Figure 3A). Weaker, albeit significant, correlations were observed between GAG content and tensile modulus (Hexabrix: $r = 0.853, P < 0.001$; CA$^4+$: $r = 0.565, P = 0.009$).

Both whole IVD and NP attenuation correlated strongly to several mechanical properties (Table 3). As the same trends were observed in both whole IVD and NP attenuation, and NP attenuation was more strongly correlated to most mechanical parameters, only correlations to NP attenuation will be discussed here. In the Hexabrix group, NP attenuation correlated strongly to compressive (Figure 4A) and tensile modulus, compressive and tensile stiffness (Table 3), and compressive, tensile, and total hysteresis (Figure 4B, Table 3). In the CA$^4+$ group, NP attenuation correlated strongly to compressive modulus (Figure 4A), compressive stiffness (Table 3), compressive and total hysteresis (Figure 4B, Table 3), but interestingly there was only a weak (though significant) correlation to tensile modulus, a weak, non-significant correlation to tensile stiffness, and no appreciable correlation to tensile hysteresis (Table 3).

MAPE values of predictive linear regression models using Hexabrix ranged from 14.05% to 17.63% (Table 4), and MAPE values of linear regression models using CA$^4+$ ranged from 16.79% to 20.23% (Table 4), demonstrating an overall predictive accuracy of the use of CE-µCT to determine the compressive stiffness or modulus of the IVD to be ~80 – 86%. Using NP
attenuation to predict compressive properties generally yielded lower MAPE values compared to using whole-IVD attenuation.

**Discussion**

The biomechanical assessment of the IVD is important for disease characterization and the development of potential treatment strategies, but it is ultimately limited by its complex, time-consuming, and often destructive nature. In this study, the ability to employ CE-µCT for the nondestructive assessment of rodent IVD mechanical properties was demonstrated. Strong, highly-significant linear correlations between CE-µCT attenuation and IVD mechanical properties were observed in an *ex vivo* model of IVD degeneration. CE-µCT attenuation correlated strongly to compressive mechanical properties and moderately, though significantly, to tensile properties. Linear regressions were established, which predicted disc mechanical properties based on IVD and NP attenuation, and the accuracy of these regressions as predictors of compressive stiffness and modulus was calculated using cross validation and found to be 79.8-86%.

The transmission and absorption of intervertebral/transvertebral compressive loading is a primary function of the IVD, and correspondingly CE-µCT attenuation correlated most strongly to compressive properties of the disc. Compressive modulus and stiffness of the IVD are primarily derived from hydrostatic pressure produced by GAG-mediated water retention, and its interaction with the highly organized lamellar structure of the AF\[^5;7\]. Hysteretic behavior in cartilaginous tissues such as the IVD arises from their inherent viscoelasticity, which has been suggested to arise in part from water extravasation and imbibition during loading\[^27;28\]. Thus, compressive modulus, stiffness, and hysteresis are all directly related to GAG and water content.
within the IVD. Both modalities of CE-µCT utilized in this study are highly sensitive to IVD GAG content\textsuperscript{14;18}, and we demonstrated that CE-µCT attenuation correlated strongly to both compressive modulus and compressive stiffness of the IVD. Using cross validation modeling, we found that CE-µCT attenuation of the IVD was 80-86\% accurate in predicting compressive stiffness and modulus, indicating a potential future role for CE-µCT as a surrogate for direct mechanical measurement in pathophysiological and regenerative studies. As our investigation employed an \textit{ex vivo} model of enzymatic digestion to mimic the degenerative state of the IVD, future studies need to confirm this level of predictive accuracy in \textit{in vivo} models of IVD degeneration.

CE-µCT attenuation also correlated significantly to IVD tensile properties – primarily tensile modulus – though predictably, these correlations were not as strong as those observed with compressive mechanical properties. The relationship between IVD tensile behavior and GAG content is not as well-established as in the case of compressive behavior. Furthermore, while the tensile properties of individual AF fibers and the tensile properties responsible for resisting radial expansion are well understood, uniaxial tensile properties of the whole IVD are not well characterized. Tensile properties of the IVD derive from the highly organized fiber structure of the AF, along with the Sharpey’s fibers which anchor the IVD to the endplates, and also likely arise in part from IVD interstitial pressure and are therefore affected by GAG depletion, dehydration, and depressurization\textsuperscript{29-32}. In addition to their role in water retention, GAG molecules can serve as matrix linkers\textsuperscript{7}, and thus GAG digestion may directly destabilize the AF matrix. Finally, the broad-spectrum activity of papain utilized in our study likely also results in partial digestion of other AF matrix molecules, diminishing both compressive and tensile properties, though previous work suggests that the concentration of papain used in this
study largely preserve AF structure\textsuperscript{23}. Interestingly, CA\textsuperscript{4+} attenuation correlated markedly poorer to tensile properties compared to Hexabrix attenuation, though no difference was noted in compressive correlations. It is possible that these effects arise from the incomplete washout of CA\textsuperscript{4+} from cartilaginous tissues as previously described by Joshi \textit{et al}\textsuperscript{33}.

Aside from the present study, there is minimal data correlating quantitative imaging techniques to mechanical properties of the IVD. Ellingson \textit{et al} correlated T2* MRI to pure moment bending mechanics in cadaveric IVDs and found significant correlations of T2* imaging parameters to stiffness, range-of-motion, and neutral zone length\textsuperscript{34}. Their study further determined that multiple T2* parameters, which indirectly measure GAG content via free water content, were more predictive of IVD mechanics than Pfirrmann damage grade, corroborating the direct correlation of GAG content to IVD mechanical properties observed in the present study. In articular cartilage, Lakin \textit{et al} have established correlations between CE-\(\mu\)CT attenuation and mechanical properties of cartilage\textsuperscript{19-21}. In their study, bovine osteochondral plugs underwent CE-\(\mu\)CT imaging followed by unconfined compressive stress-relaxation and torsional tests, and CE-\(\mu\)CT attenuation was found to correlate highly to both equilibrium compressive modulus and coefficient of friction\textsuperscript{20}. Follow-up studies in articular cartilage of the human metacarpal joint and murine tibial plateau produced similar findings\textsuperscript{19; 21}. Though the loading pattern differs considerably between articular cartilage and the IVD, the compressive mechanical properties of both tissues are highly dependent on GAG-mediated water retention, and in that context, this data agrees with the findings of the present study.

CE-\(\mu\)CT of the IVD has been previously demonstrated using both Hexabrix and CA\textsuperscript{4+}\textsuperscript{18}. The contrast agent concentrations used in this study were independently optimized to maximize tissue contrast and thus are not equal between contrast agents. At the concentrations used in this
study, $\text{CA}^{4+}$ attenuation was more sensitive to changes in the GAG content of the NP, while Hexabrix was more sensitive to changes in the GAG content of the whole IVD, based on the slopes of the linear regressions. This is consistent with the mechanism of contrast enhancement of each agent – the greater affinity of $\text{CA}^{4+}$ to the GAG-rich NP affords greater uptake and a much higher dynamic range of measurement in the NP compared to the AF, whereas the partial exclusion of Hexabrix from both the NP and AF produces a more comparable dynamic range in both tissues. Due to its higher net charge, and thereby higher interaction with negatively charged GAG, $\text{CA}^{4+}$ is known to provide greater contrast at a given concentration compared with Hexabrix\textsuperscript{33; 35}.

This study was conducted in a rat model of the IVD, and our results should be interpreted in the context of this model. As the mechanical role of GAG in the IVD is relatively well-conserved across species, the findings of this study should generally be applicable in larger species as well. However, translation of CE-$\mu$CT imaging into large animal and cadaveric human models would enable the indirect assessment of IVD mechanics described in this study to be applied in more clinically relevant scenarios. To date, CE-$\mu$CT has generally been applied as a research tool for rodent disease models, and while applications to articular cartilage in cadaveric and bovine models have been described\textsuperscript{19; 20}, applications to the IVD have thus far been limited to rabbits\textsuperscript{14} and rats\textsuperscript{18; 36}. Translation into larger models is primarily limited by the increase in diffusion distance and subsequent increase in incubation time required to achieve equilibrium. This could potentially be circumvented via the use of mechanical convection\textsuperscript{37}, incubation at higher temperature, or by other means.

This study should be interpreted in light of several limitations. An \textit{ex vivo} enzymatic digestion model of DDD was used to induce GAG depletion. Though GAG depletion is the
hallmark biochemical change associated with DDD, this model does not account for active remodeling of IVD tissue, cellular activity, nor for the concomitant changes to the vertebral endplates and vertebral bodies which are known to occur during DDD progression\textsuperscript{2, 38}. Future studies should confirm the observed correlations in \textit{in vivo} models of DDD to account for these variables. As we did not perform \textit{in vivo} \textit{µCT}, and motion segments were extensively dissected prior to testing and analysis, determination of \textit{in situ} IVD height was not possible given the absence of native muscle, tendon, and body weight forces acting on the dissected motion segment. To convert load/displacement to stress/strain data, we used \textit{µCT} data and mechanical preconditioning data to calculate IVD height. This height, therefore, represents the “gauge length” of the final mechanical cycle and not the \textit{in situ} height of the IVD. We performed whole motion segment mechanical testing as opposed to isolated characterization of the intrinsic properties of IVD substructures/tissues. While this represents a more practical and translational approach for studies assessing pathology and regeneration, future studies may wish to specifically assess the inherent compressive properties of the NP to establish correlations to CE-\textit{µCT}. Uniaxial cyclic testing was employed in this study. Though uniaxial loading is an important component of overall IVD loading, we did not assess other important motions, including bending and torsion. Asymmetric loading of the disc has been shown to induce decreased annular cellularity and subsequent repair in the disc region subjected to bending, and multiple studies have linked IVD bending to structural instability and perpetuation of IVD degeneration\textsuperscript{39, 40}. Future studies correlating CE-\textit{µCT} to mechanical properties derived from these clinically relevant motion profiles are warranted.

\textbf{Conclusion}
The purpose of this study was to determine whether CE-µCT attenuation of the IVD can be used as an indirect measure of IVD mechanical properties in an ex vivo model of DDD. The results of this study demonstrate that CE-µCT may represent an effective indirect and nondestructive modality for assessing the compressive properties of the IVD with high predictive accuracy.

Acknowledgements

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References


Pain Practice 8:18-44.


Figure Legends

Table 1. CE-µCT Attenuation and GAG content of IVDs. Data expressed as Mean [95% Confidence Interval].

Figure 1. Representative colormaps of contrast-enhanced IVD attenuation imaged with either Hexabrix (A, B) or CA⁴⁺ (C, D) following a 24-hour incubation in either saline (A, C) or 5 U/mL papain (B, D).

Table 2. Mechanical properties of IVDs. Data expressed as Mean [95% Confidence Interval].

Figure 2. Correlation of GAG content, as determined by a DMMB assay, to IVD attenuation (A) and NP attenuation (B), shown with 95% confidence intervals. Correlation coefficients and P-values were determined using the Pearson product-moment correlation.

Figure 3. Representative stress-strain of IVDs with (0 U/mL) or without (5 U/mL) enzymatic digestion in papain (A). Correlation of GAG content, as determined by DMMB assay, to
compressive modulus (B), shown with 95% confidence intervals. Correlation coefficients and P-values were determined using the Pearson product-moment correlation.

Table 3. Correlation statistics between CE-µCT attenuation and IVD mechanical properties.
Correlation coefficients and P-values were determined via the Pearson product-moment correlation.

Figure 4. Correlation of NP attenuation to compressive modulus (A) and compressive hysteresis (B), shown with 95% confidence intervals. Correlation coefficients and P-values were determined via the Pearson product-moment correlation.

Table 4 – Mean absolute percentage error (MAPE) values from 100 iterations of repeated random sub-sampling cross validation of linear regression models between CE-µCT attenuation and IVD mechanical properties

Table 1. CE-µCT Attenuation and GAG content of IVDs. Data expressed as Mean [95% Confidence Interval].

<table>
<thead>
<tr>
<th>Contrast Agent</th>
<th>Papain (U/mL)</th>
<th>Mean CE-µCT Attenuation (HU)</th>
<th>GAG Content (µg CS)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Whole IVD</td>
<td>NP</td>
<td>AF</td>
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<tr>
<td>Hexabrix 0</td>
<td>3559 [3430, 3688]</td>
<td>2807 [2691, 2923]</td>
<td>3784 [3643, 3926]</td>
</tr>
</tbody>
</table>

Table 2. Mechanical properties of IVDs. Data expressed as Mean [95% Confidence Interval].

<table>
<thead>
<tr>
<th>Contrast Agent</th>
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<td>AF</td>
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<td>Hexabrix 0 v. 5</td>
<td>&lt;0.001</td>
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<td>&lt;0.001</td>
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<tr>
<td>Contrast Agent</td>
<td>Papain (U/mL)</td>
<td>Modulus (MPa)</td>
<td>Stiffness (N/mm)</td>
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<td>----------------</td>
<td>--------------</td>
<td>---------------</td>
<td>-----------------</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Compressive</td>
<td>Tensile</td>
</tr>
<tr>
<td>Hexabrix</td>
<td>0 v. 5</td>
<td>&lt; 0.001</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>CA4+</td>
<td>0 v. 5</td>
<td>&lt; 0.001</td>
<td>0.029</td>
</tr>
</tbody>
</table>

Table 3. Correlation statistics between CE-µCT attenuation and IVD mechanical properties. Correlation coefficients and P-values were determined via the Pearson product-moment correlation.

<table>
<thead>
<tr>
<th>Mechanical Parameter</th>
<th>Hexabrix Attenuation</th>
<th>CA4+ Attenuation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Whole IVD</td>
<td>NP</td>
</tr>
<tr>
<td>Modulus</td>
<td>Compressive</td>
<td>-0.884</td>
</tr>
<tr>
<td></td>
<td>Tensile</td>
<td>-0.777</td>
</tr>
<tr>
<td>Stiffness</td>
<td>Compressive</td>
<td>-0.868</td>
</tr>
<tr>
<td></td>
<td>Tensile</td>
<td>-0.667</td>
</tr>
<tr>
<td>Hysteresis</td>
<td>Compressive</td>
<td>0.876</td>
</tr>
<tr>
<td></td>
<td>Tensile</td>
<td>0.824</td>
</tr>
<tr>
<td>Total</td>
<td>0.868</td>
<td>&lt; 0.001</td>
</tr>
</tbody>
</table>

Table 4 – Mean absolute percentage error (MAPE) values from 100 iterations of repeated random sub-sampling cross validation of linear regression models between CE-µCT attenuation and IVD mechanical properties.

<table>
<thead>
<tr>
<th></th>
<th>Hexabrix Attenuation</th>
<th>CA4+ Attenuation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Whole IVD</td>
<td>NP</td>
</tr>
<tr>
<td>Compressive Modulus</td>
<td>17.63 % ± 3.23</td>
<td>14.85 % ± 2.92</td>
</tr>
<tr>
<td>Compressive Stiffness</td>
<td>14.42 % ± 2.70</td>
<td>14.05 % ± 2.43</td>
</tr>
</tbody>
</table>
FIG1 - colormaps.
FIG2 - GAG scatter.

<table>
<thead>
<tr>
<th></th>
<th>Hexabrix</th>
<th>CA**+</th>
</tr>
</thead>
<tbody>
<tr>
<td>Slope (HU/µg CS)</td>
<td>-36.6</td>
<td>30.7</td>
</tr>
<tr>
<td>r</td>
<td>-0.903</td>
<td>0.951</td>
</tr>
<tr>
<td>P</td>
<td>&lt; 0.001</td>
<td>&lt; 0.001</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>Hexabrix</th>
<th>CA**+</th>
</tr>
</thead>
<tbody>
<tr>
<td>Slope (HU/µg CS)</td>
<td>-76.2</td>
<td>46.3</td>
</tr>
<tr>
<td>r</td>
<td>-0.873</td>
<td>0.960</td>
</tr>
<tr>
<td>P</td>
<td>&lt; 0.001</td>
<td>&lt; 0.001</td>
</tr>
</tbody>
</table>
FIG3 - mechanics digestion.
FIG4 - NP to mechanics scatter.