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Dual contrast in computed tomography allows earlier characterization of articular cartilage over single contrast

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Abstract
Cationic computed tomography contrast agents are more sensitive for detecting cartilage degeneration than anionic or non-ionic agents. However, osteoarthritis-related loss of proteoglycans and increase in water content contrarily affect the diffusion of cationic contrast agents, limiting their sensitivity. The quantitative dual-energy computed tomography technique allows the simultaneous determination of the partitions of iodine-based cationic (CA4+) and gadolinium-based non-ionic (gadoteridol) agents in cartilage at diffusion equilibrium. Normalizing the cationic agent partition at diffusion equilibrium with that of the non-ionic agent improves diagnostic sensitivity. We hypothesize that this sensitivity improvement is also prominent during early diffusion time points and that the technique is applicable during contrast agent diffusion. To investigate the validity of this hypothesis, osteochondral plugs (d = 8 mm, N = 33), extracted from human cadaver (n = 4) knee joints, were immersed in a contrast agent bath (a mixture of CA4+ and gadoteridol) and imaged using the technique at multiple time points until diffusion equilibrium. Biomechanical testing and histological analysis were conducted for reference. Quantitative dual-energy computed tomography technique enabled earlier determination of cartilage proteoglycan content over single contrast. The correlation coefficient between human articular cartilage proteoglycan content and CA4+ partition increased with the contrast agent diffusion time. Gadoteridol normalized CA4+ partition correlated significantly (P < .05) with Mankin score at all time points and with proteoglycan content after 4 hours. The technique is applicable during diffusion, and normalization with gadoteridol partition improves the sensitivity of the CA4+ contrast agent.

Keywords
biomechanics, cartilage, cationic contrast agent, contrast-enhanced computed tomography, dual-energy CT
INTRODUCTION

As a consequence of instantaneous impact (eg, related to sports accident), articular cartilage can become injured, leading to the development of post-traumatic osteoarthritis (PTOA). Erosion of articular cartilage, bone remodeling, and joint inflammation are the major characteristic features of osteoarthritis (OA). Often, only in advanced stages of the disease patients experience symptoms, such as pain and limited mobility. Therefore, PTOA is often diagnosed after irreversible damage to the cartilage has already occurred, limiting any possibility of early intervention. Early detection of cartilage damage could enable pharmaceutical or surgical interventions for preventing the progression of OA. Early OA is characterized by loss of proteoglycans (PGs), leading to decreased cartilage fixed charge density, lower swelling pressure, and subsequently reduced matrix stiffness. Fibrillation, due to collagen network disruption, also leads to decreased stiffness and increased tissue deformation under physiological loading predisposing the tissue to further degeneration.

Today's medical imaging modalities provide multiple methods on how to quantify OA. However, they all suffer from limitations. Ultrasoundography provides real-time image acquisition cost-effectively. However, the challenge in achieving perpendicularity between the ultrasound beam angle and the naturally curving cartilage surface limits accurate diagnosis. Magnetic resonance imaging (MRI) is great for soft tissues (eg, cartilage), but it suffers from relatively long scan times and high costs. Computed tomography (CT) imaging is substantially more accessible and affordable, and the image acquisition is swift, and the resolution superior to MRI. Further, significant advancement has been achieved with dose optimization techniques and imaging strategies in CT to reduce the radiation doses involved. However, poor soft-tissue contrast prevents separating native cartilage tissue from the surrounding synovial fluid, and, thus, requires the use of contrast agents.

Delayed contrast-enhanced CT (CECT) has been applied for imaging human articular cartilage in vivo to assess tissue morphology and composition. The diagnosis is based on the evaluation of an anionic contrast agent distribution within cartilage after intra-articular administration. Recently, a cationic CT agent was introduced. Cartilage fixed negative charge, created by PGs, provides strong electrostatic attraction to cationic agents. These distribute inside the cartilage in direct relation to the cartilage PG content. For this reason, cationic agents offer a more sensitive technique for diagnosing the distribution of PGs in cartilage compared with conventional anionic agents. Higher uptake of the cationic agents in cartilage provides higher X-ray attenuation and improves contrast allowing better visualization of PG distribution and its variation within cartilage. Thus, the detection of subtle changes in PG content is possible at different stages of cartilage degeneration. Contrast agents diffusion in early time points is fast, especially in degenerated cartilage, due to increased permeability. The uptake of a cationic agent depends both on the electrostatic attraction between the positively charged molecule and the negative fixed charge in cartilage, and the passive diffusion controlled by cartilage water content and permeability. Thus, in degenerated cartilage, the uptake of the cationic agent is simultaneously reduced due to the decrease of negatively charged PGs, and enhanced due to the increase in permeability and water content. These opposite effects limit the diagnostic effectiveness of the cationic agents, especially in the first hours of diffusion, which is vital for the clinical feasibility of the agent. After intra-articular administration, contrast agents diffuse into cartilage, while simultaneously, the body clears out the agent from the joint cavity, lowering the concentration as time progresses. The concentration of anionic ioxaglate in joint cavity has been reported to be adequate for delayed-CECT until 2 hours after the administration, while the agent concentration in patellar and femoral cartilage reached the maximum 30 and 60 minutes after the administration, respectively. The molar concentration of cationic and non-ionic agents in cartilage increases faster compared with an anionic agent. Thus, considering the diffusion in cartilage and the clearing out of the contrast agents from the joint cavity, the 30 to 60 minutes imaging time window could be clinically feasible for the application of both cationic and non-ionic agents.

Contrast agent partition in cartilage is quantified as a ratio of contrast agent-induced X-ray attenuation in the cartilage relative to the attenuation in the bath. Normalization (division) of an iodine-based cationic agent (CA4+) partition with an electrically neutral gadolinium-based agent (gadoteridol) partition improved the sensitivity of CA4+ to probe cartilage PG content after 72 hours of diffusion. Because water content and permeability of cartilage control the diffusion of the non-ionic gadoteridol the normalization minimizes the effect of these factors on the diffusion of the cationic agent. In early diffusion time points, contrast agent diffusion flux is high. Further, the agent fluxes are even higher in a degenerated cartilage due to loss of collagen network integrity and reduced PG, resulting in increased permeability. Considering this, we hypothesize that the improvement in the sensitivity of the cationic agent after normalization is even more substantial in a degenerated cartilage at early time points. Here we study the diffusion of the agents at clinically relevant time points (<1 hour after contrast agent administration) and at later diffusion time points close to diffusion equilibrium. Further, we examine the validity of the hypothesis by evaluating the sensitivity of normalized CA4+ partition to reflect variation in histopathological and biomechanical properties of human articular cartilage samples. Improvement in the sensitivity of the cationic agent would enable early detection of minor injuries and lesions, allowing timely selection of treatment, thus reducing the risk for PTOA. This quantitative technique is based on the simultaneous diffusion of two contrast agents (iodine-based CA4+ and gadolinium-based gadoteridol) into cartilage. Accurate simultaneous quantification of concentrations of two contrast agents using single X-ray tube voltage is not possible, as X-ray attenuation of both agents contributes to the attenuation. Hence, as iodine and gadolinium have different x-ray attenuation properties as a function of energy imaging with two separate X-ray tube voltages allows quantitative determination of the concentration of the elements in the mix. Determining the concentration of CA4+ and gadoteridol in cartilage is possible by using the Beer-Lambert law and Bragg’s additive rule for mixtures as described in literature and also in the materials and methods chapter of this paper. As the contrast agents are constantly diffusing...
in cartilage during a scan, CT acquisition at two different X-ray tube voltages must be nearly instantaneous for accurate determination of contrast agent tissue partition. In this study, we also quantify the error in the partition of the contrast agents arising from the ongoing diffusion in the cartilage when the imaging is performed separately with two X-ray tube voltages.

2 | MATERIALS AND METHODS

2.1 | Sample extraction and preparation

Human osteochondral plugs (N = 33, d = 8 mm) were extracted from the lateral and medial tibial plateaus and femoral condyles in left and right knee joints of human cadavers (n = 4, mean age = 71.25 ± 5.18 years). The Research Committee of the Northern Savo Hospital District granted a favorable opinion on collecting the human tissue (Kuopio University Hospital, Kuopio, Finland, Decision numbers: 134/2015 and 58/2013). The samples were stored frozen in phosphate-buffered saline (PBS; −22°C).

2.2 | Biomechanical measurements

Samples were thawed at room temperature. A custom-made, high precision material testing device (resolution: 0.1 µm, 0.005 N, PM500-1 A; Newport, Irvine, CA) was employed for biomechanical testing of the osteochondral plugs. Measurement setup schematics are included in the supplementary material (Figure S6). During the test, the plugs were immersed in PBS containing inhibitors of proteolytic enzymes (5 mM EDTA, VWR International, and 5 mM benzamidine hydrochloride hydrate (Sigma-Aldrich Inc, St. Louis, MO). A flat-ended metallic indenter (d = 728 µm [n = 20] or d = 667 µm [n = 13]) was driven in perpendicular contact with the articular surface. During the experiments, the tip of the indenter was accidentally damaged, and we had to continue the experiment with a spare indenter. The new indenter tip diameter was slightly different from the damaged indenter. However, the difference in the diameter is accounted for in the determination of the moduli values along Hayes et al. A pre-stress of 12.5 kPa defined the contact. Based on literature, the Poisson ratios were set to ν = (0.3(Tibia), 0.2(Femur)) for E_{equilibrium} and ν = 0.5 for E_{instantaneous}. The plugs were then again frozen, cut to two halves, and stored in a freezer (−22°C). One half was thawed for contrast-enhanced microCT imaging experiment, and the other half was prepared for reference histological analyses.

2.3 | MicroCT imaging

The dual-energy microCT set-up was tested and validated by quantifying iodine (I) and gadolinium (Gd) contents in phantoms with known contrast agent mixtures consisting of gadoteridol (20, 12, and 8 mgGd/mL) and CA4+ (0, 10, 20, 30, 40, 50, 60, and 70 mg/mL). Calibration curves and the measurement setup are presented in the supplementary material (Figures S7, S8, and S9). The edges of the osteochondral samples were sealed using cyanoacrylate (Super glue Precision, Loctite, Düsseldorf, Germany) to allow the contrast agent diffusion only through the articulating surface. Before the immersion in contrast agent bath, the plugs were imaged with a high-resolution microCT scanner (Quantum FX, Perkin Elmer) with an isotropic voxel size of 40 × 40 × 40 µm and using a 20 × 20 mm field of view. Three samples were arranged in a sample holder and immersed in a contrast agent bath (15 ml) comprised of iodine-based CA4+ (5.5′,-(malonylbis(azanediyl))bis(N,N′-bis(2-aminoethyl))-2,4,6-triiodosilphenyldiamine, q = +4, M = 1499.88 g/mol) and gadolinium-based gadoteridol (Prohance; Bracco International B. V., Amsterdam, Netherlands, q = 0, M = 559 g/mol) diluted in PBS. The expected partitions of the contrast agents in cartilage were accounted for when designing the concentration to use for the bath, to achieve a similar relative contribution to X-ray attenuation at tube voltages of 50 and 90 kVp. By doing so, the optimum signal to noise ratio was achieved while limiting excessive beam hardening and photon starvation artifacts. Based on these considerations 10 mg/mL (CA4+) and 20 mgGd/mL (gadoteridol) were selected for the immersion bath. To prevent degradation of the samples, proteolytic inhibitors, 5 mM of ethylenediaminetetraacetic acid (EDTA, VWR International, France), 5 mM of benzamidine hydrochloride hydrate (Sigma-Aldrich Inc), and penicillin-streptomycin-amphotericin (Antibiotic Anti-mycotic solution, stabilized; Sigma-Aldrich Inc) were added to the bath. The samples were imaged using the Quantum FX microCT scanner at the following diffusion time points: 10 minutes, 30 minutes, 1, 2, 3, 4, 6, 10, 21, 32, 50, 72 hours. The osmolality of the contrast agent bath was 297 mOsm/kg measured using a commercial osmometer (Advanced Model 3320 micro-osmometer; Advanced Instruments, MA). The contrast agent bath was gently stirred throughout the immersion of the samples. The stirring assembly was placed inside a refrigerator (4°C) to preserve the cartilage and prevent bacterial and fungal growth. The data were then analyzed using the software package (Adobe Photoshop CS5). The image acquisition time with each tube voltage was approximately 2 minute. Due to a human error, three samples were imaged using the same tube voltage at 1 hour diffusion time point. Thus, the partition results for those samples could not be calculated, and the 1 hour time point results of those samples have been excluded.
2.4 | Image analysis

The concentrations of iodine (C_I) and gadolinium-based (C_Gd) contrast agents in cartilage were resolved based on Beer-Lambert law and Bragg’s additive rule of mixtures:

\[ \alpha_E = \mu_{I(E)} C_I + \mu_{Gd(E)} C_{Gd} \]  \hspace{1cm} (1)

where \( \alpha \) is X-ray attenuation in a medium at energy \( E \) (tube voltages of 90 and 50 kVp) as,

\[ C_I = \frac{\alpha_{90kV,\mu_{Gd}(50kV)} - \alpha_{50kV,\mu_{Gd}(90kV)}}{\mu_{Gd}(90kV)\mu_{Gd}(50kV) - \mu_{Gd}(50kV)\mu_{Gd}(90kV)} \]  \hspace{1cm} (2)

\[ C_{Gd} = \frac{\alpha_{90kV,\mu_{Gd}(50kV)} - \alpha_{50kV,\mu_{Gd}(90kV)}}{\mu_{Gd}(90kV)\mu_{Gd}(50kV) - \mu_{Gd}(50kV)\mu_{Gd}(90kV)} \]  \hspace{1cm} (3)

The mass attenuation coefficients for CA4+ (\( \mu_{I} \)) and gadoteridol (\( \mu_{Gd} \)) were determined at both energies by imaging series of contrast agent solutions with known I and Gd concentrations in distilled water, respectively. Segmentation of the articulating surface and bone-cartilage interface was done using Seg3D software (vs. 2.4.0; The University of Utah, Salt Lake City, UT). The volume of interest was defined to be 2800 \( \times \) 2000 \( \mu \)m \( \times \) cartilage thickness. The X-ray attenuation profiles from the surface to deep cartilage were extracted using Matlab (R2016b; The Mathworks Inc, Natick, MA). To avoid partial volume effect arising from background and irregular and undulating surface, and cartilage-bone interface, 3% and 5% of cartilage thickness from the articular surface and cartilage-bone interface, respectively, was excluded from the attenuation profiles. X-ray attenuation profiles of the cartilage before immersion in contrast agent bath were subtracted from the contrast-enhanced cartilage profiles to obtain depth-wise attenuation profiles induced only by the contrast agents. Concentration profiles of I and Gd from the surface to the deep cartilage were calculated using Equations 2 and 3, respectively.

2.5 | Histological analysis and Mankin scoring

The osteochondral samples were decalcified in EDTA. Following dehydration, the EDTA decalcified samples were embedded in paraffin to be cut into 3 \( \mu \)m thick sections from the center of the plug along the coronal plane (from articulating surface to the cartilage-bone interface). After removing the paraffin, the cut sections were stained with Safranin-O. Optical density (OD) of the staining in each section was determined by applying quantitative digital densitometry technique using a light microscope (Nikon Microphot-FXA, Nikon Co., Japan) equipped with a 12-bit CCD camera (ORCA-ER; Hamamatsu Photonics K.K., Japan). For each cartilage sample, three histological sections were measured. The depth-wise OD profiles of the sections were then normalized to the length of 100 points and averaged. Before the measurements, the system was calibrated using neutral density filters (Advanced Optics SCHOTT AG, Mainz, Germany) with OD range between 0 (low) and 3 (high).

Four independent observers (M. Honkanen, R. Shaikh, N. Hänninen, M. Prakash) assessed and assigned histopathological Mankin scoring based on the severity of OA using the Safranin-O stained sections. Mankin score characterizes cartilage based on staining (0-4), tidemark integrity (0-1), abnormality in structure (0-6), and cellularity (0-3). Intact cartilage is assigned score 0, and a severely degenerated sample is scored 14. Mankin scores were calculated by averaging the scores of three sections per sample.

2.6 | Error simulation

Error in contrast agent concentrations arising from the progressing diffusion during the time between image acquisitions with two X-ray tube voltages (90 and 50 kVp) was studied using a numerical simulation. To describe the contrast agents diffusion in cartilage, equation \( C = C_{\text{max}} \times (1 - \exp(-t/\tau)) \) was fitted to the experimental data (all the samples in the present study), where \( C \) represents I and Gd concentrations in mgl/ml and mgD/ml, respectively, \( t \) is the diffusion time (minutes) and \( \tau \) is the time required to reach 63.2% of the maximum concentration (\( C_{\text{max}} \)). The error simulation was implemented in steps, as follows:

Step 1. Fitting was done for each sample individually, after which a mean of the parameters (\( C_{\text{max}} \) vs \( \tau \)) for both contrast agents was calculated (Figure 1).

Step 2. Using equation 1, X-ray attenuation was simulated with both tube voltages, based on the contrast agent concentrations obtained from the fit (step 1). This was done with varying time (2, 5, 10, 15, 30, 45, and 60 minutes) between acquisitions with the 90 and 50 kVp tube voltages. This was done for all the diffusion time points until diffusion equilibrium (72 hours).

Step 3. Using the simulated data gathered in step 2, concentrations of the contrast agents were calculated (Equations 2 and 3).

Step 4. The simulated concentration values were then compared with the true concentration values (from the fit) to get the relative error, as illustrated in Figure 5.

2.7 | Statistical analysis

The statistical analyses were conducted using SPSS (v. 23.0 SPSS Inc; IBM Company, Armonk, NY) statistical software. The reliability in the Mankin scoring between the raters was evaluated by determining the Interclass Correlation coefficient. Shapiro-Wilk test showed the sample data to follow the normal distribution. Therefore, the correlations of contrast agent partitions with histopathological and biomechanical reference parameters were evaluated using a parametric test (Pearson’s correlation analysis) within a selected cartilage region. For all statistical tests, \( P < .05 \) was set as the limit of statistical significance.
RESULTS

The CA4+ and gadoteridol partitions in cartilage increased from 14.4% ± 8.8% and 10.2% ± 5.6% at 10 minutes to 344.0% ± 77.9% and 91.4% ± 9.7% at 72 hours, respectively (Figures 1 and S11). The average $C_{\text{max}}$ was 35 and 17 mg/mL, and $\tau$ was 1032 and 244 minutes, for the iodine (CA4+) and gadolinium (gadoteridol) in the cartilage, respectively. At 72 hours, the uptake of CA4+ was 2.5 times greater in deep cartilage compared with the superficial cartilage compared with the superficial cartilage (Figure 2). The correlation coefficients between the Mankin score and CA+ partition of the full-thickness cartilage increased at all the diffusion time points after normalizing with the partition of the non-ionic gadoteridol (Figure 3). The Mankin score correlated significantly with gadoteridol partition from 10 minutes to 10 hours after the start of the immersion in contrast agent mix. The mean OD and thickness values of the cartilage samples were 1.07 ± 0.26 AU (min, 0.43; max, 1.47 AU), and 2.42 ± 0.68 mm (min, 1.01; max, 4.35 mm), respectively (Figure S12). The correlation coefficient between OD and CA4+ partition increased with the contrast agent diffusion time (Figure 4A). The correlation was significant in the earliest time points for the superficial 10% of cartilage ($P < .029$). The equilibrium modulus correlated significantly with the normalized CA4+ partition after 21 hours of diffusion ($P < .014$) and CA4+ partition after 32 hours of diffusion ($P < .004$) (Table 1). Based on the error simulation, a 2-minute delay between the acquisitions at the 10-minute diffusion time point would result in 74.4% and 23.5% relative error in determined CA4+ and gadoteridol partitions (Figure 5). At the 100-minute time point, the errors were 6.2% and 2.2% for CA4+ and gadoteridol, respectively. The mean Mankin score of all the samples was 6.27 ± 1.27 (min, 2; max, 9). The mean equilibrium modulus value for the samples was 0.26 ± 0.32 MPa (min, 0.01;
The inter-rater reliability in the Mankin score was high (Inter-observer correlation = 0.93, \(P < .01\)). The Kruskal-Wallis test revealed no difference (\(P > .79\)) in the equilibrium modulus values 0.27 ± 0.36 and 0.25 ± 0.25 between the samples measured with indenters having tip diameters of 667 and 728 \(\mu\)m, respectively.

### 4 | DISCUSSION

In this ex vivo study, simultaneous diffusion of two contrast agents (CA4+ and gadoteridol) into cartilage was evaluated using a microCT scanner at multiple time points to probe cartilage composition and structural integrity. Normalization of the CA4+ partition with that of the gadoteridol improved the correlation with the Mankin score at all time points (Figure 3). Assessing cartilage structural integrity using only cationic agents at early time points is challenging as the uptake is comparably high in both intact and degenerated cartilage, due to high PG content, and increased permeability, respectively. This limits the sensitivity of cationic agents to quantify reduced PG content especially during the early points of contrast agent diffusion. Normalizing the CA4+ partition with that of gadoteridol improves its sensitivity to detect PG content. In this study, a similar effect is seen between CA4+ and PG content where the normalization with gadoteridol partition reveals a significant correlation six hours earlier, beginning at the 4-hour diffusion time point (Figure 4B).

During very early diffusion (<1 hour time points), the CA4+ significantly correlates with PG content (ie, OD, \(P < .05\)) in the superficial zone (10% of the cartilage thickness) (Figure 4A). Upon inspecting the first 20% of cartilage thickness, the correlation starts to be significant only after 6 hours of diffusion. In this zone, the normalization does not improve correlation with PG content (Table S1). This is likely due to the partial volume effect. In the full thickness cartilage, the correlation with PG content is relatively weak at early time points. However, with the diffusion of CA4+ into deep cartilage, the correlation becomes stronger and is significant (\(P < .05\)) after 10 hours of diffusion. The partition of CA4+ increases towards the deep cartilage at later diffusion time points (Figure 2). This is due to the increased electrostatic attraction, resulting from high PG content in the deep cartilage (Figure S13). Concurrently, the water content in cartilage decreases towards the cartilage-bone interface. Thus, expecting the gadoteridol partition to follow the trend of water content in cartilage, it is surprising to see the higher partitions in the deeper zones after the 21-hour diffusion time point. We suspect that this is a result of X-ray beam hardening, as very high uptake of the cationic agent is observed post 21-hour imaging time point in the PG rich deep cartilage (Figure 2). Based on the present experiments and the results, we cannot determine whether the high CA4+ flux could have caused drag influencing the gadoteridol diffusion. Additionally, the overall gadoteridol partition is not observed to rise after a 10-hour imaging time-point (Figure 1A).
The correlation between the equilibrium modulus and CA4+ partition was significant and strengthened by the normalization after 21 hours until diffusion equilibrium. This was expected as the cationic agent’s uptake is mostly due to the attraction to PG’s, which controls cartilage biomechanical equilibrium response. Therefore, it is natural that the CA4+ which is attracted by the PGs correlates strongly with equilibrium modulus.

Previously, we applied the QDECT technique to evaluate the cartilage PG and water contents at diffusion equilibrium (ie, after 72 hours of diffusion). In the current study, we demonstrate the simultaneous determination of the solute concentration of two contrast agents in cartilage during diffusion at clinically relevant time points. The precise determination of the contrast agent partitions at early time points is possible with the use of short scan times (Figure 5). In our previous study, reliable measurements during diffusion were impossible due to long imaging acquisition time (total of 28 minutes with two tube voltages) required by the applied microCT scanner (Skyscan 1172; Skyscan, Kontich, Belgium). With dual-contrast method, if the imaging is performed separately, an unavoidable error arises in the determination of partition of the contrast agent in cartilage. This is due to the ongoing diffusion during the imaging (in this study the time difference between acquisitions being ~2 minutes). In the present study, we evaluate this error using numerical simulations. An increase in time difference between the CT scans results in a higher error in the determined contrast agent partition values (Figure 5). The relative error is higher for CA4+ due to the higher diffusion flux of the cationic agent compared with that of the non-ionic gadoteridol. The short scan time enabled attenuation measurements of multiple contrast agents during diffusion, and more importantly, in the clinically relevant time points. With a modern dual-energy full-body CT scanner, the image acquisition at separate energies is simultaneous and practically instantaneous. Hence, for the clinical application of the QDECT, this is not a source of significant error. However, the simulations will aid in the planning of the QDECT studies when imaging at two energies is performed separately.

The authors acknowledge limitations related to this study. The osteochondral plugs were extracted from a limited number of cadavers and from various locations: femur (lateral condyle = 4, medial condyle = 10), tibia (lateral plateau = 8, medial plateau = 7), and trochlea = 4. The availability of the cadaveric samples determined the sample size. As this was an exploratory study, and the effectiveness of the dual-contrast

| TABLE 1 | Pearson’s correlation coefficients between contrast agent partitions (N = 33, at 1 h time point N = 30) and biomechanical moduli |
|---|---|---|---|---|---|---|---|---|---|---|---|---|---|
| | Time, min | | Time, h | | | | | | | | | | |
| 10 | 30 | 1 | 2 | 4 | 6 | 10 | 21 | 32 | 50 | 72 |
| Equilibrium modulus, MPa | | | | | | | | | | | | | |
| CA4+ | 0.008 | -0.129 | 0.156 | 0.071 | 0.162 | 0.186 | 0.209 | 0.311 | 0.493*** | 0.521*** | 0.497*** |
| Normalized CA4+ | 0.212 | -0.139 | 0.122 | 0.032 | 0.260 | 0.330* | 0.293 | 0.422* | 0.654** | 0.730** | 0.648** |
| Gadoteridol | -0.218 | -0.105 | -0.192 | -0.126 | -0.246 | -0.285 | -0.246 | -0.339 | -0.471* | -0.453* | -0.282 |
| Instantaneous modulus, MPa | | | | | | | | | | | | | |
| CA4+ | -0.120 | -0.048 | 0.223 | 0.131 | 0.172 | 0.182 | 0.131 | 0.169 | 0.216 | 0.209 | 0.176 |
| Normalized CA4+ | 0.534** | -0.064 | 0.177 | 0.043 | 0.170 | 0.248 | 0.178 | 0.213 | 0.359* | 0.339* | 0.290* |
| Gadoteridol | -0.281 | 0.022 | -0.124 | 0.035 | -0.085 | -0.140 | -0.147 | -0.176 | -0.441** | -0.344* | -0.272 |

*Indicates that correlation is significant at the level P < .05 (two-tailed).
**Bold value Indicates that correlation is significant at the level P < .01 (two-tailed).

FIGURE 5 Simulation of error in (A) gadoteridol and (B) CA4+ partitions resulting from time (2 to 60 minutes) between acquisitions with the two X-ray tube voltages (90 and 50 kVp) [Color figure can be viewed at wileyonlinelibrary.com]
technique over the use of a single contrast agent in cartilage diagnostics was unknown, a power analysis was not conducted. Author’s acknowledge that a higher number of samples would have enabled a more reliable evaluation of the diagnostic potential of the technique. The contrast agents’ concentrations were selected to achieve the highest signal to noise ratio with the microCT scanner and toxicity issues were not taken into consideration. The development and introduction of a commercial clinical dual contrast application are not within the scope of this study, and the techniques may in the future rely on formulations differing from the ones applied here. The authors acknowledge the possibility of changes in cartilage properties arising from freeze-thaw cycles. However, as all the samples were frozen and thawed following a uniform protocol, any changes in the mechanical/biological state between the samples should be similar.

The samples were immersed in the contrast agent bath maintained at 4°C. This temperature is lower than that during the intended clinical application of the agents in the human body (37°C). The time constant $\tau$ of CA4+ was 1032 minutes, being much higher than the value reported for diffusion in bovine cartilage at room temperature ($\tau=104.4$ minutes, cartilage edges not sealed) (Figure 2).\(^{15,43}\) With an increase in temperature, the diffusion rate of the contrast agent will also increase. Hence, at a warmer temperature, contrast agent diffusion will be faster, and reliable assessment of cartilage integrity may be conducted at earlier diffusion time point.

The challenges associated with the clinical application of the QDECT are yet to be explored. In the future, the QDECT should be tested on full knee joints using a clinical CT device, to obtain quantitative information on the capability of the technique to reveal cartilage matrix water and PG contents. In this study, we have demonstrated that QDECT allows the simultaneous determination of two different contrast agents in cartilage from early diffusion time-point (10 minutes) until diffusion equilibrium (72 hours). Normalization of the cationic contrast agent (CA4+) partition with that of the electrically neutral contrast agent (gadoteridol) enhances correlations with the histopathological and biomechanical characteristics allowing swifter determination of cartilage integrity. Thus, QDECT has the potential for diagnosis of cartilage degeneration at clinically relevant imaging time points.

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CONFLICT OF INTERESTS

Dr. Grinstaff reports a patent pending on CA4+ composition. The other co-authors declare no conflict of interests.

AUTHOR CONTRIBUTIONS

AB, BP, and JT designed the study. HK applied for ethical approval to obtain the cartilage samples. AB, MP, and, MH conducted the biomechanical measurements and analyses. RS, MP and, MH performed digital densitometry measurements and analyses. CA4+ was prepared in the laboratory of MG. Contrast agent diffusion measurements were conducted by AB and BP in the laboratory of HHW. AB, JM, and JT were involved in data analysis and interpretation of the results. AB drafted the manuscript and all co-authors contributed to the critical revision of the manuscript. All authors have read and approved the final version of the submitted manuscript.

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**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section.

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