Orientation anisotropy of quantitative MRI relaxation parameters in ordered tissue

Nina Hänninen, Jari Rautiainen, Lassi Rieppo, Simo Saarakkala & Mikko Johannes Nissi

In highly organized tissues, such as cartilage, tendons and white matter, several quantitative MRI parameters exhibit dependence on the orientation of the tissue constituents with respect to the main imaging magnetic field (B0). In this study, we investigated the dependence of multiple relaxation parameters on the orientation of articular cartilage specimens in the B0. Bovine patellar cartilage-bone samples (n = 4) were investigated ex vivo at 9.4 Tesla at seven different orientations, and the MRI results were compared with polarized light microscopy findings on specimen structure. Dependences of T2 and continuous wave (CW)-T1ρ relaxation times on cartilage orientation were confirmed. T2 (and T2*) had the highest sensitivity to orientation, followed by T1RAFF2 and adiabatic T2ρ. The highest dependence was seen in the highly organized deep cartilage and the smallest in the least organized transitional layer. Increasing spin-lock amplitude decreased the orientation dependence of CW-T1ρ. T1 was found practically orientation-independent and was closely followed by adiabatic T1ρ. The results suggest that T1 and adiabatic T1ρ should be preferred for orientation-independent quantitative assessment of organized tissues such as articular cartilage. On the other hand, based on the literature, parameters with higher orientation anisotropy appear to be more sensitive to degenerative changes in cartilage.

Orientation anisotropy describes the dependence of a measured parameter on the orientation of the measured object or material. In the case of magnetic resonance imaging (MRI), nuclear magnetic properties of tissue depend not only on its chemical characteristics and macroscopic structure, but also on the physical organization of macromolecules. Highly organized tissues, such as skeletal muscle, tendon, white matter and cartilage, exhibit different properties when MRI measurements are conducted at different tissue depths or at different physical orientations with respect to the primary magnetic field. Multiple underlying physical or chemical properties of tissue may act as a source for the anisotropy in different MRI parameters. While the anisotropy of relaxation affects various tissues, as listed above, in this study we focus on the orientation dependence of relaxation parameters in articular cartilage. Specifically the deep cartilage, which has a uniform and highly organized structure, may serve as a generic model for ordered tissue.

At microscopic scale, articular cartilage consists of a network of collagen fibres and proteoglycan molecules, which together comprise the extracellular matrix. Orientation anisotropy of MRI parameters in cartilage arises from the structure of the collagenous meshwork in which the fibres have a specific depth-wise orientation and organization. Non-calcified cartilage can be divided into three structural zones on the basis of the depth-wise change of collagen fibre orientation. Starting from the articular surface, the first zone is the superficial or tangential zone (SZ) with fibres running mostly parallel to the surface. Below this, the fibres have a more random orientation in the transitional (intermediate) zone (TZ), and finally in the radial (deep) zone (RZ), the orientation has turned perpendicular to the surface. In the deep zone, the structure of cartilage is the most organized, resulting also in the highest anisotropy of its properties.

Articular cartilage functions as a shock-absorbing tissue and enables low-friction during joint movement. Osteoarthritis (OA) is a disease which affects a growing number of people, and results in progressive degeneration of articular cartilage and the surrounding tissues in joints. There is no cure for advanced OA, but the
progression of the disease may be slowed down by dietary choices and weight management. Thus, diagnosing OA at the earliest possible stage is essential in preventing the progression and minimizing the symptoms. While numerous quantitative MRI methods have been proposed and established for early diagnosis of OA, a significant issue persists that the results of many of those (particularly T2 relaxation time) are susceptible to orientation anisotropy.

In almost every joint of the human body, articular cartilage is naturally at variable orientations, causing different regions of cartilage to be scanned at different angles with respect to the MRI scanner's magnetic field; thus yielding potentially variable results for the different regions of otherwise similar cartilage. In practice, the measurement geometry due to joint shapes cannot be controlled and, therefore, orientation-independent MRI methods or detailed understanding of the dependence would be necessary to avoid possible false diagnoses and to allow realistic analysis of the structure and state of the health of cartilage.

The sensitivity of different quantitative MRI (qMRI) parameters to relaxation anisotropy varies. T1 has been shown not to be sensitive to orientation, whereas T2 and continuous wave (CW)-T1rho have been shown to have excessive sensitivity to orientation. For CW-T1rho, reduction of orientation sensitivity in cartilage has been reported for increasing spin-lock powers.

Orientation dependence of T2 relaxation time in articular cartilage has been ascribed to residual dipolar interactions (RDI) of water protons due to their restricted spatial arrangement within the collagen fibres. In addition to water proton intramolecular dipolar interactions, intermolecular dipolar couplings between water and biopolymer protons have also been suggested as a source of T2 anisotropy. Dipolar interaction between two nuclei is directly proportional to the factor (3 cos^2 θ – 1), where θ is the angle between the direction of the magnetic field and a vector joining the nuclei. Dipolar interaction is one of the predominant relaxation mechanisms and is generally noticed as a signal reduction except at θ angles near or at so called “magic angle” of 54.74° where (3 cos^2 θ – 1) = 0 and the dipolar interaction vanishes. Thus, qMRI relaxation time parameters affected by dipolar interaction reach maximum values at this specific angle and show a (3 cos^2 θ – 1)-dependence on the sample orientation.

Practical value of qMRI parameters depends on their potential and accuracy for diagnosis of diseases, such as OA. Recently, T2 and T2*, and rotating frame relaxation (RFR) methods including adiabatic T1rho, adiabatic T2, and TRAFF2, have been reported to be sensitive for cartilage degeneration. Orientation sensitivities of these parameters have been investigated in a preliminary study; adiabatic T2 and TRAFF2 were observed to have a similar orientation sensitivity as T2*, whereas adiabatic T1rho was shown to be less sensitive to orientation than T2, both in ex vivo and in vivo measurements.

The aim of the experimental part of this study was to investigate the orientation dependence of several quantitative MRI parameters (T1, T2, T2*, CW-T1rho, adiabatic T1rho, adiabatic T2, and TRAFF2) and evaluate the findings against the collagen fibre orientation and anisotropy, measured by the gold standard reference technique, i.e., quantitative polarized light microscopy (qPLM). As a secondary aim, the usefulness of the aforementioned parameters for OA diagnostics, as described and reported in the literature, was evaluated against the orientation sensitivity determined in the experimental part.

**Results**

qPLM revealed the typical tri-laminar collagen orientation in all the samples, starting with fibres oriented along the cartilage surface and arching towards radial orientation at the cartilage-bone interface (Fig. 1a). Similar to qPLM anisotropy, optical retardation was lowest in the transitional zone and increased towards the deep tissue, indicating increasing anisotropy (Fig. 1b,c).

Different sensitivities of the parameters to sample orientation with respect to B0 were observed (Figs 2 and 3). Orientation dependence was absent or minimal for T1, adiabatic T1rho, with HS1 pulse and CW-T2, at 2 kHz spin-lock amplitude. On the other hand, using HS4 or HS8 pulses for adiabatic T1rho or decreasing the spin-lock power of CW-T1rho increased the orientation dependence. T2, T2*, adiabatic T2, and TRAFF2 had the highest sensitivity to orientation, visualized by the largest changes over the orientation, especially in the deep tissue. Identical behaviour was observed with all the samples. The regional minimum, maximum and mean values of the relaxation times over all the measurements reflected the same observations on orientation sensitivities (Table 1).

Analysis of the depth-wise MR anisotropy, as determined using the Michelson contrast parameter, revealed distinctive differences between the relaxation times (Fig. 4). For those parameters that demonstrated sensitivity, the anisotropy changed as a function of depth: the minimum was observed at the transitional zone, higher relaxation anisotropy at the surface and the maximum in the radial zone, closely resembling the qPLM anisotropy of the collagen network (Fig. 1c). Relaxation anisotropy of the MRI parameters was assessed as a bulk value in the radial zone with the expected most uniform collagen architecture and the highest qPLM anisotropy. T1 relaxation anisotropy in this region had the lowest correlation with qPLM anisotropy (r = 0.03) and retardation (r = −0.10), followed by adiabatic T1rho with HS1 pulse (r = 0.09/−0.10), whereas the anisotropies of T2 and T2* relaxation times had the highest correlations (r = 0.87/−0.88 and r = 0.87/−0.86, respectively) (Table 2).

Assessing the anisotropies of the MRI parameters in the radial zone enabled sorting the parameters by their respective sensitivities to the specimen orientation in the main field (Fig. 5, Table 2). Based on values reported in the literature, relative differences between “normal” or “intact” and variable degenerated or degraded (“OA”) animal and human tissue were evaluated and plotted together with the estimated orientation anisotropies (Fig. 5). Broadly, parameters with higher sensitivity to orientation anisotropy also demonstrated largest relative differences between intact and degenerated articular cartilage (Fig. 5).
Discussion

Orientation dependence of several quantitative traditional and rotating frame MRI parameters in ordered tissue was investigated in this study. The findings on the MR anisotropy of several quantitative relaxation time parameters were compared to measurements of structural anisotropy of cartilage tissue by qPLM. Furthermore, literature-reported relative differences for the investigated MR parameters between intact and degenerated tissue were evaluated against the assessed orientation sensitivities.

In this study, we selected articular cartilage as a model tissue that has both highly anisotropic and also relatively isotropic regions. The current findings confirmed earlier reports on the orientation sensitivity of $T_1$ and $T_2$ relaxation times, and the reduction of the orientation sensitivity with increasing spin-lock power for $\text{CW-}T_1$ and $\text{CW-}T_2$. Orientation dependence was observed also for other relaxation parameters, and was the most significant for $T_2^*$, adiabatic $T_1$ and $T_{\text{RAFF2}}$.

As previously demonstrated in multiple studies, $T_2$ relaxation time along with $T_2^*$ was the most sensitive to orientation of the sample, a phenomenon ascribed to the residual dipolar interaction in cartilage and other similar highly organized tissues. $T_2$ relaxation time is sensitive to dipolar interactions and, thus, this was an expected finding. $T_2^*$, on the other hand, might be sensitive also to field inhomogeneity contributions from anisotropic susceptibility of the collagen fibres in the cartilage, since it carries not only $T_2$ but also the inhomogeneity related component. However, very similar relaxation anisotropy to $T_2^*$ was noted for $T_2^*$.

The orientation dependencies of adiabatic $T_1$, adiabatic $T_2$ and $T_{\text{RAFF2}}$ have been briefly investigated previously for articular cartilage. The present results demonstrated the orientation dependence of $T_{\text{RAFF2}}$ and adiabatic $T_2$ to be very similar to that of $T_2$ or $T_2^*$, as was noted in the preliminary findings. Furthermore, an increase in the dependence for adiabatic $T_1$ was observed with increasing pulse stretching factor. Three different pulse modulations using hyperbolic secant function with stretching factors of 1, 4 and 8 were tested. To allow comparison, the RMS powers and bandwidths of the pulse modulations were kept the same (approximately equal to 906 Hz CW-pulse power). As the pulse is stretched further, the shape approaches that of a CW-pulse and the time the magnetization spends in the vicinity of the xy-plane is increased, i.e. the contributions from off-resonance $T_1$ relaxation are reduced and more dipolar relaxation is allowed, explaining the observed change in anisotropy. However, even with the HS8 pulse modulation, the relaxation was nearly orientation-independent.

Native (non-contrast enhanced) $T_1$ relaxation time and adiabatic $T_1$ with HS1 pulses were found to be essentially independent of the tissue orientation. Adiabatic $T_1$ and $T_2$ have been shown to be sensitive to changes in cartilage even in vivo, and adiabatic $T_1$ seems promising also with clinical MRI equipment. The independence of $T_1$ and adiabatic $T_1$ parameters on the orientation of the tissue promotes them as suitable parameters to be applied in vivo for assessing tissue status. For quantitative MR imaging of articular cartilage, this also implies that the dGEMRIC method, while requiring exogenous contrast agent, should provide orientation-independent assessment. Similar orientation-independence is expected for other $T_1$-based contrast-enhanced imaging methods.

Figure 1. PLM results presented as a map for one representative sample and profiles of all the specimens: (a) orientation; (b) retardation; (c) anisotropy. Articular surface and cartilage-bone interface are marked on the maps with arrowheads. Profiles were calculated separately for each sample along cartilage depth and averaged point-by-point for mean profile.
Assuming dipolar interaction is the main contributor to spin relaxation in cartilage, both longitudinal $T_1$ and transverse $T_2$ relaxation depend on molecular fluctuations introducing field perturbations driving the relaxations. Compared with $T_1$, $T_2$ relaxation has an additional contribution from the secular component of the dipole-dipole interaction (relaxation depending on the spectral density at zero frequency, which does not affect the $T_1$ relaxation). This additional component brings about the orientation sensitivity to $T_2$ relaxation. In sufficiently organized matter, such as cartilage, this orientation dependent component may not completely vanish by random molecular motion and represents itself as orientation-dependent $T_2$ relaxation. Since $T_1$ relaxation does not have this component, it is not orientation dependent. This also suggests an explanation for the orientation insensitivity of the adiabatic $T_1^\rho$ relaxation. With an adiabatic pulse, the magnetization follows the trajectory of the RF field; this results in the magnetization relaxing along the RF field. The RF modulation of the adiabatic...
HS-pulse is such that the effective field starts along the Z-axis, halfway through the pulse traverses through the transverse plane and ends along the opposite direction of the Z-axis. Thus the relaxation during an adiabatic HS-pulse can be viewed as a combination of longitudinal and transverse relaxations, with the amounts of the respective components dependent on the pulse shape. This is also evidenced by the increased orientation sensitivity of adiabatic $T_1^\rho$ when the pulse is stretched, making the effective field (and the magnetization) spend more time in the transverse plane and thus experiencing more $T_2$-like relaxation.

For all the parameters with clear anisotropy along the cartilage depth, the minimal orientation dependence was detected in the transitional zone, at approximately 12–14% of depth from the articular surface. This is the zone in which the anisotropy of the collagen fibres is also at its minimum. Above this zone, in the superficial cartilage, the anisotropy is higher due to the preferential arrangement of the fibres along the surface. However, collagen fibres in the superficial cartilage are also distributed at multiple orientations along the plane parallel to the surface. Thus, orientation dependent relaxation times depend also on the rotation angle about the axis of the surface normal. In the radial zone, however, the collagen fibres are more uniformly oriented along the axis of the surface normal and relaxation is less affected by the rotation about this axis. This also explains the observed maximum of the qMRI anisotropy in the radial zone. The overall average relaxation times and their ranges in the different zones generally reflect the same observations, although the averages can only be considered indicative and descriptive of the measurements due to the different orientation sensitivities of the parameters.

Since the deep cartilage has the most uniform collagen fibre structure, it was chosen as a region to represent the overall orientation sensitivity of the relaxation parameters. Orientation anisotropy in the radial zone was calculated for all the parameters as the average value from the depth of 40% to 80%. According to

Figure 3. Interpolated profile maps for relaxation parameters (one representative sample). Articular surface is on the left, bone interface towards right, with the profiles at different orientations stacked on top of each other and then interpolated. SZ, TZ and RZ denote the approximate locations of the histological zones.
qPLM data, this region is safely within the limits of approximately constant fibre angle, and represents only the deep cartilage without the influence of transitional zone or cartilage-bone interface.

The results of the qPLM orientation and retardation are in accordance with the results of Xia et al.\(^5\) and Rieppo et al.\(^1\) on mature articular cartilage. On average, the difference between orientations of fibres in the superficial and radial zone was approximately 75 degrees, which is less than the theoretical value of 90 degrees. This likely reflects the actual variations in the fibre orientation in the deep zone, as well as the possibility that the primary orientation can differ from the angle exactly perpendicular to the cartilage-bone interface or cartilage surface\(^5\).

The change in orientation anisotropy for CW-$T_1^\rho$ followed the previously reported change: increasing spin-lock amplitude reduced the sensitivity\(^2\), with the conclusion that spin-lock amplitude greater than 500 Hz starts to overpower the RDI at 1.5 T. Here it was found that 500 Hz already reduces the sensitivity to orientation (i.e. to RDI), but does not remove it; the sensitivity is further reduced with 1 kHz and 2 kHz amplitudes (see also Fig. 4). Mlynarik et al. tested CW-$T_1^\rho$ at two different $B_0$ field strengths and concluded that dipolar interaction is a major factor in $T_1^\rho$ relaxation especially at lower field strengths\(^2\). However, in practice the increasing SAR values of increasing spin-lock amplitude (and often also hardware limitations) prevent the clinical use at high frequencies (typically up to ~500 Hz is clinically applicable)\(^3\).

The maximum values for the relaxation times (Fig. 3) were found at sample orientations of 59°, 56°, 69° and 72°. Theoretically, the dipolar relaxation vanishes at the magic angle, which is 54.7°\(^8\). The observed small deviations are probably reflective of the small variations in the actual fibre angles in deep cartilage with respect to the sample itself, as the orientations of the specimens were determined from the scout images based on the

|        | SZ    | TZ    | RZ    | SNR | Scan time |
|--------|-------|-------|-------|-----|-----------|
| $T_1$  | 1425  | 1534  | 1367  | 34.8| 5:50      |
| Adiabatic $T_1^\rho$ (HS1) | 193 (150/232) | 236 (200/278) | 189 (167/224) | 70.8 | 4:11      |
| Adiabatic $T_1^\rho$ (HS4) | 126 (94/174) | 155 (129/191) | 113 (80/145) | 62.4 | 4:11      |
| Adiabatic $T_1^\rho$ (HS8) | 129 (102/162) | 160 (136/192) | 117 (84/151) | 63.1 | 4:11      |
| $T_2$  | 42 (23/71) | 60 (38/98) | 26 (6/68) | 32.5 | 4:11      |
| $T_2^\ast$ | 38 (20/73) | 52 (30/79) | 22 (5/55) | 37.0 | 2:14      |
| Adiabatic $T_2^\rho$ | 58 (33/91) | 75 (48/115) | 35 (10/80) | 48.0 | 3:30      |
| RAPF2  | 69 (40/110) | 85 (57/135) | 40 (14/96) | 54.8 | 6:51      |
| CW-$T_1^\rho$ (250 Hz) | 60 (29/97) | 77 (37/129) | 39 (14/85) | 41.1 | 3:23      |
| CW-$T_1^\rho$ (500 Hz) | 73 (51/106) | 91 (63/125) | 53 (27/90) | 44.9 | 3:23      |
| CW-$T_1^\rho$ (1000 Hz) | 83 (62/112) | 103 (83/125) | 72 (47/97) | 48.6 | 3:23      |
| CW-$T_1^\rho$ (2000 Hz) | 95 (69/118) | 115 (92/138) | 88 (70/108) | 51.3 | 3:23      |

Table 1. Mean and min/max values of the relaxation parameters (in ms) in the different cartilage zones and SNR values through all samples and all orientations. Scan times (in minutes) are for one sample per single orientation.

Figure 4. Depth-wise MR anisotropy profiles for relaxation parameters (average of four samples). Smallest anisotropy is noted at the location of the transitional zone, while the maximum anisotropy is observed in the deep zone. Boxed area shows the depth used for calculating deep cartilage average values.
Recognizing this possibility, there was no specific attempt to scan the specimens exactly at the magic angle, potentially also allowing missing of the true maximum, which might lie between the measured angles.

As a step forward from standard anatomical imaging, qMRI has a huge potential to provide improved diagnostics and objective, quantitative information of tissue properties at the molecular level. However, in clinical MRI, the measurement geometry often cannot be altered. Thus, orientation independent MRI methods, or understanding of the dependence, is necessary to avoid possible false diagnoses and to allow realistic analysis of the structure and of the state of the tissue being imaged. The correlation (or lack of it) of the anisotropy of the qMRI parameters with the PLM anisotropy effectively indicates the sensitivity of the different parameters to orientation and further to the geometry of the scan setup. To precisely characterize relaxation changes in ordered tissue, more orientations are required for those parameters that correlate with structural anisotropy. From the literature comparison (Fig. 5) it can be seen that the parameters with the best sensitivity in detecting tissue changes related to OA also tend to have the highest sensitivity to orientation anisotropy. This suggests that the methods sensitive to the orientation are thus also sensitive to changes in the orientation, i.e. sensitive to the properties of the collagen fibre network, which is one of the primary components of articular cartilage. Thus, sensitivity to orientation anisotropy may have a role in the sensitivity of the parameters for detecting differences between “intact” or “normal” and “degenerated” tissue. However, the optimal qMRI parameter would exhibit zero sensitivity to orientation while having the maximum sensitivity to tissue degeneration or changes in tissue, i.e., parameters approaching the lower right corner in Fig. 5 would be generally preferred. While a number of different MRI parameters were

| Parameter | Average MRI Anisotropy | MRI Anisotropy vs. PLM Anisotropy | MRI Anisotropy vs. PLM Retardation |
|-----------|------------------------|-----------------------------------|-----------------------------------|
| $T_1$     | 80.0 (77.8/85.3)       | 0.87 (0.82/0.90)                 | −0.88 (−0.95/−0.77)              |
| $T_1^*$   | 79.4 (75.3/86.9)       | 0.87 (0.71/0.93)                 | −0.86 (−0.94/−0.68)              |
| Adiabatic $T_1$ | 71.3 (66.0/81.4) | 0.89 (0.86/0.91)                 | −0.87 (−0.93/−0.82)              |
| RAFF2     | 69.6 (64.2/76.5)       | 0.90 (0.81/0.96)                 | −0.86 (−0.92/−0.77)              |
| CW-$T_1$ (250 Hz) | 64.6 (55.9/77.0) | 0.84 (0.71/0.92)                 | −0.77 (−0.89/−0.54)              |
| CW-$T_1$ (500 Hz) | 43.6 (30.2/54.5) | 0.80 (0.63/0.88)                 | −0.72 (−0.81/−0.60)              |
| CW-$T_1$ (1000 Hz) | 22.6 (16.9/32.5) | 0.69 (0.52/0.83)                 | −0.58 (−0.65/−0.48)              |
| Adiabatic $T_1$ (HS4) | 17.6 (14.0/21.8) | 0.55 (0.24/0.88)                 | −0.46 (−0.66/−0.18)              |
| Adiabatic $T_1$ (HS8) | 17.3 (11.7/24.9) | 0.52 (0.13/0.80)                 | −0.46 (−0.58/−0.28)              |
| CW-$T_1$ (2000 Hz) | 12.0 (9.4/15.7) | 0.34 (−0.12/0.66)                | −0.35 (−0.59/0.13)               |
| Adiabatic $T_1$ (HS1) | 6.8 (6.0/8.4) | 0.09 (−0.19/0.44)                | −0.10 (−0.25/0.08)               |
| $T_1$     | 1.9 (1.1/2.5)         | 0.03 (−0.06/0.14)                | −0.10 (−0.17/−0.06)              |

Table 2. Average anisotropy of the relaxation parameters in deep cartilage (average of four samples and range (min/max)) and the coefficients of correlation for the depth-wise MRI anisotropy with depth-wise PLM anisotropy and PLM retardation. Averages for correlation coefficients have been calculated using Fisher’s z transform. The parameters are ordered in the table based on their anisotropy.

![Figure 5](https://example.com/image.png)

Figure 5. Average anisotropy of relaxation parameters in deep cartilage versus relative difference of relaxation parameters between “normal” and “OA” in different models as reported in literature (○ = human in vivo, □ = human ex vivo, △ = in vivo animal OA model with imaging done for samples, ◊ = ex vivo animal model). Shaded area represents deviation in anisotropy and range of relative difference values. For $T_1^*$, Adiabatic $T_1$ with HS8 pulses, and CW-$T_1$ 2000 Hz no reference values were found. The optimal parameter would lie in the lower right corner.
studied, yet more possibilities exist: for example, magnetization transfer experiments have been demonstrated to have minimal or no orientation dependence\(^1\). On the other hand, sensitivity to orientation changes could provide an alternative means to analyse tissue structure and properties at the molecular level.

**Conclusion**

In conclusion, adiabatic T\(_1\), also reported sensitive to cartilage degeneration\(^24,33,35\), appeared as a promising, minimally orientation-dependent quantitative MRI parameter for the articular cartilage. Additionally, adiabatic pulses provide flexibility with respect to overcoming SAR issues. While limited and somewhat variable results have been reported for native T\(_1\), it appears as the most promising parameter out of the investigated in terms of SAR and orientation (in)dependence. This finding also suggests that other measurements based on estimating T\(_1\) relaxation time (such as contrast enhanced methods) would be promising. The practical usefulness of qMRI parameters depends on how well they can be used in diagnosis of diseases, such as OA. Based on the literature findings and the measured relaxation anisotropies, parameters with higher sensitivity to orientation anisotropy generally also demonstrated larger relative differences between intact and degenerated articular cartilage. The best-suited MRI parameter for diagnostics in ordered tissues remains uncertain.

**Methods**

**Sample Preparation and MRI Measurements.** Cylindrical osteochondral plugs (\(n = 4\), diameter = 6 mm) from bovine patella were prepared (bovine knee joints were obtained from a local abattoir; the exact age of the animals is not known, but the typical age at the time of slaughter is approximately 18 months). The samples were immersed in perfluoropolyether (Galden HS 240, Solvay Solexis, Italy) in a custom-built holder, which allowed (manual) rotation of the specimens i.e. rotation of cartilage surface normal axis with respect to the main imaging magnetic field (B\(_0\)) from outside the scanner. MRI was performed at 9.4 T using a 19 mm quadrature RF volume transceiver and Varian Agilent Direct Drive console. The samples were imaged at room temperature at seven different orientations, spanning 0–90 degrees at approximately 15 degree intervals with respect to B\(_0\). The orientation was confirmed from scout images and later precisely measured (cartilage surface normal vs. B\(_0\)) from 3-D gradient echo (GRE) data acquired for every orientation. Relaxation time measurements were realized using a global preparation block coupled to a single slice fast spin echo readout (TR = 5 s, ESP = 5.5 ms, ETL = 8 with centric echo ordering, matrix = 256 × 64, FOV = 16 × 16 mm and 1 mm slice, yielding a resolution of 62.5 μm along cartilage depth). A single imaging slice was positioned at the centre of the specimen, perpendicular to the axis of specimen rotation, and was rotated with the specimen. The measurements included T\(_1\) relaxation time with inversion recovery (TI = 0.2, 0.5, 0.8, 1.1, 1.4 and 3 s); T\(_2\) with spin echo preparation (TE = 10, 20, 40, 80, 100 and 128 ms); CW-T\(_1\) with four spin-lock amplitudes (\(\gamma B_1/2\pi = 250, 500, 1000\) and 2000 Hz)\(^26\); adiabatic T\(_1\), with pulses from the hyperbolic secant HSe family (HS1, HS4 and HS8 pulses, \(\gamma B_1\) = 4.5 ms, and \(\gamma B_1\) = 2.5, 1.2 and 1.04 kHz, respectively, to match the R/M ratio between the pulse shapes (equal to approximately 906 Hz CW-pulse))\(^31\); adiabatic T\(_2\) with HS1 pulse train embedded between adiabatic half passages and T\(_{RAFF2}\) (\(\gamma B_1\) = 250 Hz, \(\tau_\rho = 9\) ms). CW-T\(_1\) was measured with spin-lock durations of 0, 24, 48, 96 and 192 ms; both adiabatic T\(_1\) and T\(_2\) were measured with pulse trains of 0, 4, 8, 12 and 24 pulses using MLEV4 phase cycling, and T\(_{RAFF2}\) with trains of 0, 2, 4, 6 and 8 pulses acquired with and without an inversion preparation\(^35,59\). Finally, T\(_2\) relaxation time was measured in the same slice using a multi-echo GRE sequence (TE = 2.5, 7.5, 12.5, 17.5, 22.5, 27.5, 32.5 and 37.5 ms). Scan times together with the overall SNR values for the sequences are given in Table 1. All measurements were repeated for every orientation. After the MRI studies, the samples were fixed in 10% neutral buffered formalin for 48 hours and then decalcified in EDTA for histology processing.

**Histology and quantitative polarized light microscopy (qPLM).** After decalcification, the samples were cut in the same plane that was used in MR imaging utilizing 3-D reconstructions of the MRI data as visual guides. Subsequently, the samples were dehydrated and embedded in paraffin. Unstained histological sections were prepared from the locations of the MRI slices and placed on the standard microscope slides. The sections were digested with hyaluronidase enzyme for 18 hours to remove proteoglycans before quantitative polarized light microscopy imaging (qPLM) measurements (pixel size 2.53 × 2.53 μm). The optical retardation and collagen fibre orientation in the sections were measured according to Rieppo et al.\(^{13}\) using quantitative Abrio PLM imaging system. Abrio PLM system (CRI, Inc., Woburn, MA, USA) is mounted on a conventional light microscope (Nikon Diaphot TMD, Nikon, Inc., Shinagawa, Tokyo, Japan) and consists of a green bandpass filter, a circular polarizer, a computer-controlled analyser composed of two liquid crystal polarizers and a CCD camera.

**Data Analysis.** In MRI analyses, to obtain relaxation time maps, relaxation time constants were fitted in a pixel-wise manner using the respective signal equations: T\(_{EXOT}\) was fitted using exponential decay and approach to steady state\(^{20}\), CW-T\(_1\), using 2-parameter exponential with baseline and the rest of the parameters were fitted using 2-parameter monoeponential with noise floor subtraction before fitting, using in-house developed Matlab (Matlab R2015b, Mathworks, Natick, MA, USA) plugin functions for Aedes (http://aedes.uef.fi) (examples of fit curves and values can be found in the supplementary Figure S1 and Table S1). Depth-wise profiles of 1.75 mm width from the cartilage surface to bone interface were calculated at the centres of the specimens. The profiles were then depth-normalized for comparison and averaging between specimens.

In qPLM analyses, anisotropy of the collagen fibres was calculated by applying an entropy filter of size 5 × 5 pixels (entropyfilt, Matlab) to the orientation angle images obtained from the qPLM. The anisotropy is represented by the inverse of the entropy + 1 to constrain the values to the range [0, 1]. The orientation, retardation and anisotropy profiles were calculated separately for each sample along cartilage depth and averaged point-by-point for mean profile and depth-normalized for comparison. Average values and ranges for the relaxation times over the samples and orientations were calculated in the three structural zones, SZ, TZ and RZ (regions shown in Fig. 3).
Parallelism, or anisotropy of the collagen fibres, can be defined as Michelson contrast\textsuperscript{13, 38}:

\[
A_i = \frac{R_{i}^{\text{max}} - R_{i}^{\text{min}}}{R_{i}^{\text{max}} + R_{i}^{\text{min}}},
\]

where min and max are the minimum and maximum measured intensity \( R \) over the different physical orientations of the specimen. Here, this formalism was used to calculate the depth-wise MR anisotropy profiles for the relaxation parameters using the relaxation rates at different sample orientations as the measured intensities.

To quantitatively analyse relaxation anisotropy of the MRI parameters as a bulk property, an average value for the depth of 40\% to 80\% was calculated, since the deep cartilage, i.e., radial zone has relatively constant fibre orientation with the highest anisotropy (Fig. 4, ROI region shown with dashed line).

Correlations between the MR anisotropy profiles and PLM measurements were calculated per sample and then averaged using Fisher’s z transform\textsuperscript{39}. Matlab was used for all calculations and analyses.

To assess the orientation dependence versus relative sensitivity of the parameters, the average anisotropies of the MR parameters were plotted against relative difference of the same parameters between early OA and advanced OA, or intact and degenerated tissue, taken from multiple previous studies reporting different OA cases or OA models\textsuperscript{24, 33, 65, 66, 67}.

Data Availability. All of the raw data, documentation and analysis codes of the study are available for download at Zenodo (doi:10.5281/zenodo.519752).

References

1. Henkelman, R. M., Stanisz, G. J., Kim, J. K. & Bronskill, M. J. Anisotropy of NMR Properties of Tissues. Magn. Reson. Med. 32, 592–601 (1994).
2. Cleveland, G. G., Chang, D. C., Hazlwood, C. F. & Rorschach, H. E. Nuclear magnetic resonance measurement of skeletal muscle: anisotropy of the diffusion coefficient of the intracellular water. Biophys. J. 16, 1043–53 (1976).
3. FULLERTON, G., CAMERON, I. & ORD, V. Orientation of tendons in the magnetic field and its effect on T2 relaxation times. Radiology 155, 433–435 (1985).
4. NAVON, G., ELIA, U., DEMCIO, D. E. & BLMICH, B. Study of order and dynamic processes in tendon by NMR and MRI. J. Magn. Reson. Imaging 25, 362–380 (2007).
5. BENDER, B. & KLOSE, U. The in vivo influence of white matter fiber orientation towards B0 on T2* in the human brain. J. Magn. Reson. Biomed. 23, 1071–1076 (2010).
6. RUDKO, D. A. et al. Origins of R* orientation dependence in gray and white matter. Proc. Natl. Acad. Sci. USA 111, E159–67 (2014).
7. XIA, Y. Relaxation anisotropy in cartilage by MRI microscopy (μMRI) at 14-μm resolution. Magn. Reson. Med. 39, 941–949 (1998).
8. ERICKSON, S. J., PROST, R. W. & TIMINS, M. E. The “Magic Angle” Effect: Background Physics and Clinical Relevance. Radiology 23–25 (1993).
9. LEE, J. et al. T2*-based fiber orientation mapping. NeuroImage 57, 225–234 (2011).
10. NIEMINEN, M. T., NISSI, M. J., HANNI, M. & XIA, Y. in Biophysics and Biochemistry of Cartilage by NMR and MRI (eds Xia, Y. & MOMOT, K. L.) 147–175 (Royal Society of Chemistry). doi:10.1039/9781782623363-00145 (2017).
11. BUCKWALTER, J. A. & MANKIN, H. J. Articular cartilage I. Tissue design and chondrocyte-matrix interactions. J. Bone Jt. Surg 79, 612–632 (1997).
12. RENNINGHOFF, A. Form und Bau der Gelenkknorpel in ihren Beziehungen zur Funktion. Zeitschrifft für Zelforsch. und Mikroskopische Anat 2, 783–862 (1925).
13. RIEPPO, J. et al. Practical considerations in the use of polarized light microscopy in the analysis of the collagen network in articular cartilage. Microsc. Res. Tech. 71, 279–287 (2008).
14. WEISS, C., ROSENBERG, L. & HELLET, A. J. Ultrastuctural Study Articular of Normal Cartilage. J. Bone Jt. Surg 50A, 663–674 (1968).
15. JEFFERY, A., BLUMEN, G., ARCHER, C. & BENLEY, G. Three-dimensionnal collagen architecture in bovine articular cartilage. J. Bone Jt. Surg 73, 795–801 (1991).
16. HUBER, M., TRATTNIG, S. & LINTNER, F. Anatomy, Biochemistry, and Physiology of Articular Cartilage. Invest. Radiol. 35, 573–580 (2000).
17. XIA, Y., MOODY, J. B. & ALHADLAQ, H. Oriational dependence of T2 relaxation in articular cartilage: A microscopic MRI (μMRI) study. Magn. Reson. Med. 48, 460–469 (2002).
18. MEACHIN, G. & Stockwell, R. in Adult articular cartilage (ed. Freeman, M.) 69–144 (Pitman Medical). doi:10.1002/art.1780230134 (1979).
19. GOLDRING, M. B. & GOLDRING, S. R. Articular cartilage and subchondral bone in the pathogenesis of osteoarthritis. Ann. N. Y. Acad Sci 1192, 230–237 (2010).
20. CHU, C. R., WILLIAMS, A. A., COYLE, C. H. & BOWERS, M. E. Early diagnosis to enable early treatment of pre-osteoarthritis. Arthritis Res. Ther. 14, 212 (2012).
21. CHANG, E. Y., MA, Y. & DU, J. MR Parametric Mapping as a Biomarker of Early Joint Degeneration. Sport. Heal. A Multidiscip. Approach 8, 405–411 (2016).
22. GOLD, G. E. et al. MRI of articular cartilage in OA: novel pulse sequences and compositional/functional markers. Osteoarthr. Cartil 14, 76–86 (2006).
23. NIEMINEN, M. T. et al. Prediction of biomechanical properties of articular cartilage with quantitative magnetic resonance imaging. J. Biomech. 37, 321–328 (2004).
24. RAUTIAINEN, J. et al. Multiparametric MRI assessment of human articular cartilage degeneration: Correlation with quantitative histology and mechanical properties. Magn. Reson. Med. 74, 249–259 (2014).
25. PETO, S. & GILLIS, P. Fiber-to-field angle dependence of proton nuclear magnetic relaxation in collagen. Magn. Reson. Imaging 8, 705–712 (1990).
26. MYLNARIK, V., SZOMOLANYI, P., TOFFANIN, R., VITTOR, F. & TRATTNIG, S. Transverse relaxation mechanisms in articular cartilage. J. Magn. Reson. 169, 300–307 (2004).
27. AKELLA, S. S. V., REGATTE, R. R., WHEATON, A. J., BORTHAKUR, A. & REDDY, R. Reduction of residual dipolar interaction in cartilage by spin-lock technique. Magn. Reson. Med. 52, 1103–1109 (2004).
28. WANG, N. & XIA, Y. Dependencies of multi-component T2 and T1p relaxation on the anisotropy of collagen fibrils in bovine nasal cartilage. J. Magn. Reson. 212, 124–132 (2011).
29. SHAO, H. et al. Magic angle effect plays a major role in both T1rho and T2 relaxation in articular cartilage. Osteoarthr. Cartil. 1–9. doi:10.1016/j.joca.2017.01.013 (2017).
Financial support from the Academy of Finland (grants #285909, #293970, #268378 and #303786) and the European Research Council under the European Union’s Seventh Framework Programme (FP/2007-2013)/ ERC Grant Agreement no. 336267 are gratefully acknowledged. The authors wish to thank Mr. Hassaan Elsayed for the sample-rotation holder and Dr. Olli Gröhn for helpful discussions.

Author Contributions M.J.N., J.R. and N.H. conceived and designed the experiments. J.R. and M.J.N. performed the MRI experiments and data collection. L.R. and S.S. performed histology experiments. N.H. and M.J.N. analysed the data, prepared figures and wrote the main manuscript text. All authors reviewed and approved the manuscript.

Additional Information Supplementary information accompanies this paper at doi:10.1038/s41598-017-10053-2
