Sodium and $T_{1\rho}$ MRI for molecular and diagnostic imaging of articular cartilage

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ABSTRACT: In this article, both sodium magnetic resonance (MR) and $T_{1\rho}$ relaxation mapping aimed at measuring molecular changes in cartilage for the diagnostic imaging of osteoarthritis are reviewed. First, an introduction to structure of cartilage, its degeneration in osteoarthritis (OA) and an outline of diagnostic imaging methods in quantifying molecular changes and early diagnostic aspects of cartilage degeneration are described. The sodium MRI section begins with a brief overview of the theory of sodium NMR of biological tissues and is followed by a section on multiple quantum filters that can be used to quantify both bi-exponential relaxation and residual quadrupolar interaction. Specifically, (i) the rationale behind the use of sodium MRI in quantifying proteoglycan (PG) changes, (ii) validation studies using biochemical assays, (iii) studies on human OA specimens, (iv) results on animal models and (v) clinical imaging protocols are reviewed. Results demonstrating the feasibility of quantifying PG in OA patients and comparison with that in healthy subjects are also presented. The section concludes with the discussion of advantages and potential issues with sodium MRI and the impact of new technological advancements (e.g. ultra-high field scanners and parallel imaging methods). In the theory section on $T_{1\rho}$, a brief description of (i) principles of measuring $T_{1\rho}$ relaxation, (ii) pulse sequences for computing $T_{1\rho}$, relaxation maps, (iii) issues regarding radio frequency power deposition, (iv) mechanisms that contribute to $T_{1\rho}$ in biological tissues and (v) effects of exchange and dipolar interaction on $T_{1\rho}$ dispersion are discussed. Correlation of $T_{1\rho}$ relaxation rate with macromolecular content and biomechanical properties in cartilage specimens subjected to trypsin and cytokine-induced glycosaminoglycan depletion and validation against biochemical assay and histopathology are presented. Experimental $T_{1\rho}$ data from osteoarthritic specimens, animal models, healthy human subjects and as well from osteoarthritic patients are provided. The current status of $T_{1\rho}$ relaxation mapping of cartilage and future directions is also discussed.

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OSTEOARTHRITIS

Osteoarthritis (OA) affects more than half of the population above the age of 65 (1, 2) and has a significant negative impact on the quality of life of elderly individuals (3). The economic costs in the USA from OA have been estimated to be more than 1% of the gross domestic product (4). OA is now increasingly viewed as a metabolically active joint disorder of diverse etiologies. The biochemistry of the disease is characterized by the following changes in cartilage: reduced proteoglycan (PG) concentration, possible changes in the size of collagen fibril and aggregation of PG, increased water content and increased rate of synthesis and degradation of matrix macromolecules. The earliest changes in the cartilage due to OA result in a partial breakdown in the proteoglycan matrix with a decrease in total content. The total concentration of collagen remains unaltered in the earliest stages, although there are generally changes in the size and arrangement of the fibers. There is also a small increase in the total water content. All of these changes in the macromolecular matrix lead to an alteration in the mechanical properties of cartilage with the result that it can no longer serve as an effective load-bearing material. Current therapies of the disease have
been ineffective at halting or reversing its course and have largely been directed toward symptomatic relief. As the disease progresses to its late stages, joint surface replacement (arthroplasty) is the only effective treatment.

More recently, there have been efforts in developing novel techniques for the treatment of OA such as chondroprotective drugs and re-population of the cartilage defects by chondrocyte precursor cells with subsequent regeneration of the cartilage. Further, recent research has led to the development of drugs in animals that have shown the potential of protecting the macromolecules in cartilage from breakdown, effectively halting the progression of OA. Because of the long natural history of OA (10–20 years in humans), validating the efficacy of these drugs requires a noninvasive technique that can directly assess their effect on molecular changes associated with early stages of cartilage degeneration that precede morphological changes.

The current lack of adequate methods for quantifying these changes has hampered research directed towards the development of potential disease-modifying agents. An integrated, and noninvasive measurement of molecular (PG, collagen and water) and morphological (tissue volume) changes in cartilage will enable the detection of OA in its early stages. These measurements will help monitor disease progression, evaluate potential strategies for disease management and verify the efficacy of disease modifying drugs. Current diagnostic methods include radiography, arthrography, computed tomography (CT) and magnetic resonance imaging (MRI).

Joint space narrowing determined from conventional radiographs is widely accepted as an indication for early diagnosis of OA. However, it does not yield accurate and quantifiable results on molecular changes that precede morphological changes. Arthrography, which has these attributes, is an invasive technique that causes pain and discomfort to the subjects and is, therefore, not ideal for routine clinical use. CT, while quantitative, has the drawback of not providing biochemical information. MRI, on the other hand, provides excellent soft tissue contrast and superior delineation of intra-articular structures. From a morphological point of view, there has been substantial progress in improving our ability to study cartilage, using MRI. MRI can assess cartilage lesions and provide morphologic information about the cartilage damage. More specifically, changes such as fissuring, partial or full thickness cartilage loss and signal change within the residual cartilage can be detected. Several approaches have been developed and validated to use three-dimensional volumetric data to quantify articular cartilage in joints. Although MRI is one of the best noninvasive tools, conventional MRI ($T_1$, $T_2$, density-weighted and magnetization transfer imaging) has proven to be inadequate in quantifying early-stage molecular changes.

Very recently, studies on transgenic animal models have shown that ADAMTSS, an aggreccanase that cleaves the aggrecan at specific site, is responsible for the breakdown of PG and disease initiation (5,6). Although human studies are needed to confirm these studies from transgenic (tg) mice, they indicate that early molecular changes are reflected in a decrease of PG.

Several MR imaging methods have been advanced to detect and quantify such early molecular changes. Prominent in these are: proton–based methods such as $T_2$ relaxation mapping (7–20), delayed gadolinium-enhanced magnetic resonance imaging contrast (dGEMRIC) (21,22), $T_{1p}$ relaxation mapping (23–26) and the direct MRI of sodium (27–30). $T_2$ is predominantly affected by changes in collagen content and to a smaller extent in PG in the tissue. Since the dominant contribution to $T_2$ relaxation is the dipolar interaction of protons of water associated with collagen, it is primarily useful in quantifying changes associated with collagen component of the ECM. dGEMRIC has been shown to be useful in quantifying changes in PG and this method and $T_2$ mapping have been extensively reviewed in recent review articles (31–34). A complete review of MRI of cartilage is beyond the scope of this article. Recent advances in MRI of cartilage have been reviewed extensively (32–43).

In the following sections, we provide: (i) a brief discussion on the structure of normal cartilage, (ii) theoretical aspects of sodium NMR and sodium multiple quantum NMR and MRI studies of cartilage and (iii) theoretical aspects of $T_{1p}$ weighted MRI, pulse sequences for relaxation mapping and review of $T_{1p}$ relaxation studies of cartilage.

The structure of normal articular cartilage (44)

Articular cartilage is a remarkable type of connective tissue that provides the synovial joints with lubrication and makes normal motion possible. It also serves to absorb mechanical shock and to distribute load over the underlying bone. Although articular cartilage will function over the lifetime of the joint under ideal circumstances, it can be damaged by trauma, osteoarthritis and inflammatory arthritis.

Articular cartilage consists of a small population of specialized cells called chondrocytes within a large extracellular matrix (ECM). The primary components of the ECM are water, collagen (15–20%) and PG (3–10%), which the chondrocytes serve to remodel continuously (Figure 1). The structure of the cartilage varies throughout its depth and consists of four histologic zones or layers. The structure and concentration of collagen and PG vary in these zones. Water is the most abundant component of articular cartilage with concentrations ranging from 80% of the volume on the surface to 65% in the deep zone. However, only a small fraction of the water is bound to collagen molecules. The affinity of cartilage for water arises primarily from the presence of the proteoglycans.
whose negative charges serve to attract free-floating positive ions in solution such as Na\(^{+}\), which in turn attract water molecules through osmotic pressure (an example of the Donnan equilibrium).

Proteoglycans are complex macromolecules that consist of protein and polysaccharides. The most common of proteoglycans, making up 80–90% of the total, is called aggrecan [Figure 2(A)]. It consists of a protein core with a long extended domain to which many glycosaminoglycan (GAG) side chains are attached. Although both chondroitin sulfate (CS) and keratan sulfate (KS) are present in ECM, CS is the predominant GAG molecule found in cartilage [Figure 2(B)]. Several aggrecan molecules are in turn attached to hyaluronate, a long, linear polysaccharide. The resulting structure is described as having a ‘lamp-brush’ type of appearance. A large number of carboxyl and sulfate residues, present on the glycosaminoglycan side chains, are ionized under physiological conditions to give COO\(^{-}\) and SO\(_{4}\)^{2-}. The negative charge density imparted by these groups is referred to as fixed charge density (FCD). These negative ions attract positive counter-ions and water molecules and provide a strong electrostatic repulsive force between the proteoglycans. These osmotic and electrostatic forces are responsible for the swelling pressure of cartilage. The configuration of the PG macromolecules also contributes to the resistance of the matrix to the passage of water molecules and hence affects the mechanics of the cartilage in this fashion.

Collagens are proteins with a characteristic triple-helical structure. Although many different types of collagen are found in cartilage, the most common is collagen type II (90–95% of the overall mass of collagen in cartilage). The collagen molecules aggregate into fibers that are 10–100 nm in diameter. The major role of the collagen II fibers is to provide a tensile force opposing the tendency of the proteoglycans to expand the cartilage and

**Figure 1.** The extracellular matrix (ECM) of cartilage.

**Figure 2.** The aggrecan proteoglycan macromolecule (A) comprising hyaluronic acid (HA), keratan sulfate (KS) and chondroitin sulfate (CS). The molecular structures of chondroitin 6-sulfate and keratan sulfate moieties are shown in (B).
also serve to immobilize the proteoglycans. Since there are no net charges on collagen it does not impart any FCD to the ECM. Collagen fibers have different arrangement across the tissue (Figure 3). In the radial zone collagen arrangement is perpendicular to the surface of the tissue while in the superficial zone it is parallel to the surface. However, in the middle zone the arrangement is almost random. This characteristic arrangement leads to the so-called 'magic angle effect' and laminar appearance in the proton MR images. Many other molecules are present in the ECM in low concentrations, most of whose functions are not clearly understood.

Features of articular cartilage during osteoarthritis

Osteoarthritis can affect virtually any joint that contains cartilage; however it is most commonly seen in the hands, knee, hip and spine, either isolated as localized OA or as generalized OA when it affects three or more joints (45). While specific conditions, such as trauma to the joint and congenital disease, may lead to secondary OA, most cases of OA are classified as idiopathic.

The affected joint is victim to a striking breakdown of the cartilage matrix and eventually a total loss of joint cartilage. The OARSI cartilage OA pathology assessment system provides six grades of OA involvement (46). In grade 1, the cartilage matrix undergoes swelling, abrasion and changes in the structure of the cartilage such as patchy condensation of collagen and gain or loss of chondrocytes. Stains such as Safranin O or Toluidene Blue, which are used semi-quantitatively in many studies as markers for proteoglycan, can be indicative of proteoglycan loss at this early point. While structural MRI is sometimes able to see synovial thickening at this early stage (47), only imaging techniques sensitive to molecular changes such as sodium imaging, $T_1$-weighted imaging and dGEMRIC MRI can hope to regularly detect and track OA at this stage.

Grade 2 OA begins to see exfoliation of ECM from the surface of the joint and in grade 3 fissuring from the surface into the mid zone of the cartilage is observed. It is not until grade 4 that large-scale structural changes begin to occur. Increased fissuring and delamination of the surface leads to erosion and loss of the superficial zone. Chondrocyte death and metaplasia begin at stage 3, but are prominent at this point. It is at this time that larger structural changes can be regularly seen on $T_2$-weighted imaging, CT, and after joint space narrowing with loss of cartilage, X-Ray. Grades 5 and 6 OA exhibit almost a total loss of articular cartilage, and changes to the underlying bone structure such as osteophytes and microfracturing are evident.

The etiology of OA is thought to be a complex interplay of mechanical joint trauma and molecular factors where chondrocytes are unable to repair joint damage, leading to progressive cartilage loss (48). The classification of joint disease as OA is often meant to separate the disease process from other types of arthritis, such as rheumatoid arthritis (RA), that have better elucidated inflammatory components. Nevertheless, it is increasingly recognized that inflammatory or other molecular mediators may have a central role in the progression of OA. While it is beyond the scope of this article to discuss the myriad proteins implicated as markers or factors in OA pathogenesis, certain models of OA deserve attention for later discussion.

Models of osteoarthritis in articular cartilage

Early studies determined that a certain factor isolated from synovial fluid and produced by lymphocytes, later discovered to be interleukin-1 (IL-1), stimulates release of GAG from the ECM and suppresses new GAG synthesis in explant cultures (49). It was not long before IL-1B was being used frequently with in vivo and ex vivo cartilage as a model of arthritis for the testing of
pharmaceuticals and investigations of arthritis signaling (50). Genetic evidence for a role of IL-1β in OA came much more recently with the discovery of polymorphisms in the IL-1 gene cluster that act as risk factors for OA (51–53). In arthritis imaging research, IL-1β is now used to simulate osteoarthritis in animal models in order to track progression of mild disease by emerging techniques (54).

The interleukin-1 super-family includes the agonists IL-1α/IL-1F and IL-1β/IL-1F2 (55). IL-1α and IL-1β are produced as approximately 31 kDa precursor forms and are cleaved by IL-1β converting enzyme (ICE), also known as caspase-1, to forms weighing about 17.5 kDa. However, only IL-1β requires cleavage to have activity, while both the pro-form and the mature form of IL-1α are active (56). Activity is often conserved across higher mammals, as evidenced by homologous biological activity of recombinant human IL-1α and IL-1β on many other species, including cow (57), pig (58) and mouse (59).

Interleukin-1 is thought to stimulate breakdown of the ECM by causing the upregulation of two families of metalloproteinases. Messenger RNA of numerous members of the matrix metalloproteinase (MMP) family, including the collagenases MMP-1, MMP-8 and MMP-13, as well as MMP-14, MMP-3 and MMP-9, have been shown to be upregulated by IL-1 in human chondrocytes in vitro (60). The same study also showed upregulation of the aggrecanases ADAMTS-4 (aggrecanase-1) and ADAMTS-5 (aggrecanase-2) by IL-1. These proteins belong to the disintegrin and metalloprotease with thrombospondin motifs family (61), and are the only enzymes that have been shown to cleave aggrecan at a site that generates the Glu373-Ala374 fragments (62,63) seen in interleukin-1 stimulated bovine explant cultures (64) and human synovial fluid of OA patients (65).

Very recent studies of transgenic animal models suggest that ADAMTS5 may in fact be necessary for the induction or progression of symptomatic OA (5,6). Two groups working independently found that ADAMTS5 knockout mice were protected from experimentally induced arthritis by IL-1β, foreign antigen stimulation and joint instability. This provides an exciting target for development of medications targeting early mediators of arthritis that will slow or arrest the progression of OA. Testing of these new therapies may benefit from confirmation by noninvasive molecular imaging techniques that show a corresponding halt in ECM loss in tissue.

An alternative model of osteoarthritis is the use of trypsin to induce the degeneration of proteoglycan (66–68). Trypsin is a 24 kDa endopeptidase commonly produced in the pancreas for the digestion of dietary amino acids (69). It cleaves a wide range of proteins at the C-terminus of lysine or arginine except when followed by a proline residue. While collagen, due to its large number of prolines, is not susceptible to trypsin digestion under all but very harsh conditions (70), application of trypsin to explant cartilage cultures readily causes the degradation of PG and other extracellular molecules (71). It is unlikely that trypsin itself plays a major role in the pathogenesis of OA; however imaging techniques are frequently tested by their ability to detect trypsin-induced degeneration of cartilage (72,73).

Sodium NMR

Sodium is one of the most ‘NMR-visible’ nuclei in living systems. The NMR relaxation properties of a nucleus depend on its immediate environment and its interactions that perturb the dominant Zeeman Hamiltonian in a significant manner. The most important interaction experienced by sodium nuclei is that between the nonspherically symmetric nucleus and surrounding electric field gradients. This is called the quadrupolar interaction. In general, sodium in solids experience most of these interactions while in liquids, the static quadrupolar interaction is averaged to zero. In the intermediate regimes, e.g. in biological tissues, the quadrupolar interaction results in biexponential relaxation rates. Multiple quantum filtered $^{23}$Na NMR can be used to analyze spectra from such systems. The following sections describe quadrupolar interaction and its effect on the $^{23}$Na NMR spectrum and relaxation rates in greater detail.

The quadrupolar interaction. Sodium is a spin 3/2 nucleus and possesses quadrupole moment (Q), which interacts with electric field gradient (EFG) generated by the electronic distribution around the nucleus. In the absence of external magnetic field, this interaction establishes degenerate energy states in solid state. Transitions induced by appropriate RF pulses (with a frequency that is resonant with the frequency separation of the states) are responsible for the pure nuclear quadrupole resonance (NQR). This has been reviewed extensively in several original articles and books (74–76) and will not be discussed here.

When Zeeman interaction is greater than the quadrupolar interaction, the degeneracy of the energy levels is lifted and the spin dynamics are dictated by the physical status of the material. The theory of NMR of quadrupolar nuclei (74,75,77–80), relaxation-induced sodium single and multiple quantum coherences (MQC) and their applications in the studies of biological systems has been extensively discussed in several original and review articles (81–91). Here a brief overview of steps involved in the calculation of spin dynamics using the density matrix approach is presented.

The nuclear quadrupole interaction is determined by the orientation, magnitude and temporal duration of EFG generated by the surrounding lattice and the electronic configuration around the nucleus and can be described by a tensor. In a system of rapid motion, e.g. fluid, all orientations of the EFG are equally probable. In this ‘isotropic’ system, the quadrupolar interaction is aver-
aged to zero on the time scale of $1/\omega_0$ (Figure 4). However, in an anisotropically oriented system, such as liquid crystals or oriented macromolecules, the sodium nuclei experience a nonzero average EFG. This static quadrupole coupling induces a shift in the energy levels of the spin system and consequently the spectrum shows multiple lines. The frequency separation between these lines provides indirect information about the magnitude of macroscopic ordering in the system.

In the Redfield regime, where the relaxation of spin 3/2 is described by second-order perturbation theory, the spin dynamics can be solved analytically. It is well known that longitudinal relaxation gives information about relatively fast motions (in MHz) and its dispersion can be studied by measuring relaxation rates at different field strengths. Slow dynamics (frequencies in the range of 0 to a few kHz) can be probed by transverse relaxation ($T_2$) or relaxation under the influence of a spin-locking pulse ($T_{1\rho}$) known as the ‘spin-lattice relaxation in the rotating frame’. Here, we closely follow the treatment by van der Marrel (83,84,92). It is convenient to describe the dynamics of spin $I > 1$, the effects of RF pulses and the evolution under the action of static and fluctuating quadrupolar interactions using the irreducible tensor operator ($T_p^I$) representation. Where $I$ represents the rank and $p$ represents the order of the tensor. We use the

\[
T_p^I(s) = (1/\sqrt{5})(T_{p-}^I + T_{p+}^I)
\]

\[
T_p^I(a) = (1/\sqrt{2})(T_{p-}^I - T_{p+}^I)
\]

for spin 3/2, the density operator is expanded in terms of 16 orthonormal basis operators, which are shown in Table 1.

In the principal axis system of the spin 3/2 nucleus and for an axially symmetric electric field gradient, the static quadrupolar Hamiltonian is given by:

\[
H_{QS} = \omega_Q T_0^2
\]

where $\omega_Q$ and represents the static residual quadrupolar interaction (RQI), only the part of the interaction that persists after motional averaging and not the r.m.s. average of the fluctuating part.

Several models have been explored to describe the role of static quadrupolar interaction in biological systems. If this nonaveraged interaction is uniform, as in the case of liquid crystals, then clear quadrupolar splitting is observed in the spectra provided the splitting exceeds the linewidths of the individual peaks. The interaction can be heterogeneous across the sample, as in the case of nonuniformly aligned systems. In this situation, the sample may be modeled as made from individual domains, each characterized by a local director but all possessing the same dynamic properties. The exchange among the domains usually is slow on a time scale of the inverse of the linewidths and the splitting, so that the observed spectrum represents a static average over all domains. Each domain is characterized by a residual quadrupolar coupling (RQC) given by:

\[
\omega_Q = \omega_{Q0} \left( \frac{3 \cos^2 \theta - 1}{2} \right)
\]

![Figure 4. Energy level diagram of spin 3/2 nucleus. The frequency difference between adjacent energy levels is the same, $\omega_0 (A)$. It represents the isotropic solution state, where quadrupolar interaction is averaged to zero. (B) Nonzero static quadrupolar interaction shifts the energy levels, producing three spectral peaks. This represents the situation in solids, liquid crystals and oriented macromolecules. The frequency separation between adjacent energy level are given by: $(\omega_0 - \omega_Q), \omega_0$ and $(\omega_0 + \omega_Q)$.](image)

**Table 1. The 16 orthonormal basis operators of a spin-3/2 system**

<table>
<thead>
<tr>
<th>Operator</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T_0^0$</td>
<td>Identity</td>
</tr>
<tr>
<td>$T_0^0 = (1/\sqrt{5})I_z$</td>
<td>Longitudinal magnetization</td>
</tr>
<tr>
<td>$T_1^1(a)$</td>
<td>Proportional to x- and y-magnetization, respectively</td>
</tr>
<tr>
<td>$T_1^1(s)$</td>
<td>$(1/\sqrt{5})I_y$</td>
</tr>
<tr>
<td>$T_0^0$</td>
<td>Quadrupolar spin polarization</td>
</tr>
<tr>
<td>$T_1^1(s)$</td>
<td>Rank 2 single quantum coherences</td>
</tr>
<tr>
<td>$T_1^1(a)$</td>
<td>Rank 2 double quantum coherences</td>
</tr>
<tr>
<td>$T_2^2(s)$</td>
<td>Octupolar spin polarization</td>
</tr>
<tr>
<td>$T_2^2(a)$</td>
<td>Rank 3 single quantum coherences</td>
</tr>
<tr>
<td>$T_3^3(s)$</td>
<td>Rank 3 double quantum coherences</td>
</tr>
<tr>
<td>$T_3^3(a)$</td>
<td>Rank 3 triple quantum coherences</td>
</tr>
<tr>
<td>$T_4^4(s)$</td>
<td>Rank 4 single quantum coherences</td>
</tr>
<tr>
<td>$T_4^4(a)$</td>
<td>Rank 4 double quantum coherences</td>
</tr>
<tr>
<td>$T_5^5(s)$</td>
<td>Rank 5 single quantum coherences</td>
</tr>
<tr>
<td>$T_5^5(a)$</td>
<td>Rank 5 double quantum coherences</td>
</tr>
<tr>
<td>$T_6^6(s)$</td>
<td>Rank 6 single quantum coherences</td>
</tr>
</tbody>
</table>

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where $\theta$ is the angle between the local director and the
direction of the main magnetic field $B_0$ and $(\omega_Q) = \omega_Q^0$
denotes the maximum splitting measured (in the local
director frame) for $\theta = 0^\circ$.

The frequency of the central transition is unaffected in
the first order approximation of this interaction, but the
satellite transitions are spread out over a large frequency
range depending on the distribution in values of $\theta$. The
broadening of satellites due to an inhomogeneous static
quadrupolar coupling makes it difficult to measure the
position of the satellite transitions using conventional
methods. Several pulse sequences have been developed to
extract this information based on the detection of the
formation of the even rank double quantum coherence
(86). The spin system evolves under the influence of
Zeeman, radiofrequency, static quadrupolar and fluctu-
ating quadrupolar Hamiltonians.

In NMR, the detected signals are equivalent to those
obtained by performing the calculations in the rotating
frame or interaction representation given by:

$$U = e^{-iH_z t}$$

(4)

$$H_z = -\omega_0 T_0$$

(5)

$$\omega_0 = -\gamma B_0$$

(6)
in which the Zeeman Hamiltonian vanishes.

The total interaction Hamiltonian for spin 3/2 in the
interaction representation (represented with $'*$) is:

$$H^*_T = H^*_R + H^*_Q + H^*_Q$$

(7)

where $H^*_R$ represents the RF field applied in the
transverse plane with a phase $\phi$.

$$H'^*_R = -\omega_{R} \left\{ \sqrt{S} \left\{ -T^1_1(a) \sin \phi + iT^1_1(s) \cos \phi \right\} \right\}$$

(8)

$$H'^*_Q = \omega_Q T^2_0$$

(9)

Since $H^*_Q$ commutes with $H^*_R$, it remains unaffected in
the interaction representation:

$$H'^*_Q(t) = C_Q \sum_{m=-2}^{2} (-1)^m T^2_0 e^{im\omega t} [F_{m}^2(t) - < F_{m}^2 >]$$

(10)

where $C_Q = \varepsilon Q / (h \sqrt{6})$, and $Q$ is the quadrupole moment
of the nucleus and EFG components are given by:

$$F_0^2 = (1/2) V_{zz}$$

$$F_1^2 = \pm (1/\sqrt{6}) (V_{zz} \pm i V_{zx})$$

(11)

$$F_2^2 = (1/2\sqrt{6}) (V_{xx} - V_{yy} \pm 2i V_{xy})$$

and $< F_m^2 >$ represents their average values.

Macromolecular motions impart fluctuations and hence
a time dependence in the EFG components. Having
established the Hamiltonians in the interaction represen-
tation, we are ready to discuss the spin dynamics. Spin
dynamics of quadrupolar nuclei can be described by
solving the Liouville equation over a range of motional
regimes and following the time evolution of density
operator $\rho$ in response to RF pulses and in the presence
of quadrupolar relaxation and static quadrupolar inter-
action (75,81–84,93–97). The time evolution of density
operator under a static Hamiltonian, such as the static
quadrupolar Hamiltonian is given by (74):

$$\frac{d \rho^*}{dt} = -i[H^*_Q, \rho^*]$$

(12)

The time evolution of the fluctuating quadrupolar
Hamiltonian $H^*_Q(t)$ can be treated according to the
Redfield relaxation theory (98,99). Changes in the density
operator are small on the time scale $\tau_r$ of the lattice
motions i.e. $< H^2_Q > \tau_r \ll 1$. Therefore we can justify
the use of second-order perturbation theory for our cal-
culations. In most of the biological systems this condition
is satisfied because of the rather long correlation times
associated with macromolecular motions and relatively
small values of corresponding coupling constants. We can
also include the contribution from relaxation:

$$\frac{d \rho^*}{dt} = -i[H^*_Q, \rho^*] + \Gamma(\rho^*)$$

(13)

where the relaxation superoperator is given by:

$$\Gamma(\rho^*) = \int_0^\infty \left[ H^*_Q(t), \exp(-iH^*_Q t) H^*_Q(t - \tau) \right] \exp(iH^*_Q \tau), \rho^*(t)] > \delta \tau$$

(14)

Since the static quadrupolar Hamiltonian commutes with
$H^*_Q(t)$, it vanishes in the relaxation superoperator. If
the EFG is completely averaged to zero by molecular
motion, then $H^*_Q = 0$ and the spin dynamics are governed
exclusively by relaxation. This equation can be recast into a
more familiar master equation in terms of the Redfield
relaxation matrix $R$ as:

$$\frac{d \rho^*_{\alpha \beta}}{dt} = -i[H^*_Q, \rho^*_{\alpha \beta}] + \sum_\alpha \sum_{\alpha'} R_{\alpha \beta} R_{\alpha' \beta'} \rho^*_{\alpha' \beta'}$$

(15)

where $|\alpha> \ldots$ and $|\beta> \ldots$ are the eigenfunctions of unperturbed
Hamiltonian (in the presence of nonaveraged static
quadrupolar interaction), identifies the phase coherence
between states $\alpha$ and $\alpha'$ which is the $\alpha - \alpha'$th element of
spin density operator, and is defined as:

$$\langle \alpha | \rho(t) | \alpha' \rangle = \rho_{\alpha \alpha'} (t)$$

(16)

Several authors have solved this equation and described
the spin evolution in response to RF pulses, under pure
quadrupolar relaxation and in the presence of both
quadrupolar relaxation and nonaveraged static quadrupo-
lar interaction. Expressions for single, double and triple
quantum-filtered signals have been derived under different
limiting conditions. Following an RF pulse, the density
matrix evolves in the presence of quadrupolar relaxation
and static quadrupolar interaction and can be solved using
the solutions to eqns (13) and (15).

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Energy levels of sodium nuclei

The energy level diagram shown in Figure 4 depicts the energy level pattern of a spin 3/2 nucleus in different motional regimes. The following three motional regimes are possible.

Case 1: Isotropic motion with motional narrowing ($\omega_0 \tau_c \ll 1$). Figure 4(A) illustrates the spectrum from sodium in an isotropic solution state ($\tau_c$ is the rotational correlation time). The rapid tumbling of molecules containing sodium lead to a fluctuation of the EFG more rapidly than the Larmor period ($2\pi/\omega_0$), hence the quadrupole interaction is ‘motion-narrowed’ to zero. Consequently, the Zeeman interaction alone will result in four energy levels with equal frequency separation $\omega_0$. Transitions induced by RF pulses at this frequency lead to a single resonance line at $\omega_0$. Both transverse and longitudinal relaxations are simple exponential decays (100).

Case 2: Isotropic motion without motional narrowing ($\omega_0 \tau_c \approx 1$). In biological tissues where the macromolecular motion associated with the nucleus is isotropic but $\omega_0 \tau_c \approx 1$, the quadrupolar interaction dominates the relaxation. In this case, the satellite and central transitions will have different relaxation rates $R_1^{(\pm1)}$ and $R_2^{(\pm1)}$, respectively, with $R_1^{(\pm1)} > R_2^{(\pm1)}$. These fast and slow decaying components are dynamically shifted from the Larmor frequency with shifts $K_1$ and $K_2 - K_1$, respectively. However, these shifts are much smaller than the line-widths and very difficult to detect. Hence, in the subsequent discussion we will ignore these dynamic shifts. This situation is still described by Figure 4(A) with the difference that the relaxation is now bi-exponential.

$$
\begin{align*}
R_1^{(\pm1)} &= J_0 + J_1 \pm iK_1 \\
R_2^{(\pm1)} &= J_1 + J_2 \mp i(K_1 - K_2)
\end{align*}
$$

(17)

where

$$
\begin{align*}
J_m(\text{m}) &= \frac{(2\pi)^2}{20} \chi_2^2 (1 + (\omega_0/\tau_c)^2) \\
K_m(\text{m}) &= \omega_0 J_m(\text{m})
\end{align*}
$$

(18)

where the spectral densities $J_m$ and $K_m$ are the real and imaginary parts of the Fourier transform of the EFG correlation function (83). $\chi_2$ is the root mean square coupling constant.

Even in the absence of any quadrupolar splitting, biexponential relaxation can create multiple quantum coherences (MQC), which can be detected (82,101,102). In this case, the observed double quantum coherence is solely due to odd rank, $T_3^{(1)}$, and the even-rank double quantum state ($T_2^{(1)}$) is never created. Owing to its higher sensitivity, triple quantum filtered (TQF) sodium spectroscopy is well suited for measuring bi-exponential transverse relaxation rates (96).

Case 3: Anisotropic motion ($\omega_0 \tau_c > 1$). In this case, the single quantum relaxation eigenvalues are given by

$$
\begin{align*}
R_1^{(1)} &= J_0 + J_1 + J_2 - \sqrt{(J_2^2 - \omega_0^2)} \\
R_2^{(1)} &= J_1 + J_2 \\
R_3^{(1)} &= J_0 + J_1 + J_2 + \sqrt{(J_2^2 - \omega_0^2)}
\end{align*}
$$

(19)

$R_1^{(1)}$ is present due to the formation of even rank two-quantum coherence, $T_3^2$.

The following three situations can be identified based on the magnitude of $J_2$:

1. If $\omega_Q < J_2$, the relaxation eigenvalues are real and line splitting is absent, despite the presence of a nonzero EFG. The single quantum spectrum is a sum of three Lorentzians and $\omega_Q$ influences the line-widths and amplitudes of these components but it is difficult to fit these lines to determine individual line-widths.

In ordered biological tissues where the molecular motion is slow ($\omega_0 \tau_c > 1$), there is an RQI that arises as a result of the ions interacting with structural proteins such as collagen fibers in connective tissues and proteins that are part of cell membranes. This results in a very small splitting in the spectrum of sodium, which is masked by the larger single quantum line-widths. Further, the dominant presence of sodium in surrounding fluids in isotropic motion complicates the detection of RQC. Thus, measuring the relatively small RQC from motion-restricted sodium in biological tissues using conventional single quantum NMR is difficult and alternative approaches are required. However, if the residual $\omega_Q$ is comparable to line-widths, then it does not lead to line splitting but it still creates double quantum coherence due to the even-rank coherence ($T_3^{(2)}$). In this case, the double quantum filtered (DQF) signal due to even rank coherence ($T_2^{(2)}$) can be used to detect and measure the RQC (103–105).

2. If $\omega_Q > J_2$, the satellite signals do not overlap with the central transition and the relaxation eigenvalues $R_1^{(1)}$ and $R_3^{(1)}$ corresponding to outer transitions are complex and the satellite transitions are shifted by $\pm(\omega_Q^2 - J_2^2)^{1/2}$ from the central line. The satellite signals in the single quantum spectrum will have complete amplitudes and are phase twisted with respect to the central line. In this case too, the DQF signal from $T_2^{(2)}$ can be used to measure $\omega_Q$.

3. If $\omega_Q \gg J_2$, the energy levels are all shifted by $\omega_Q$ resulting in three distinct frequency separations between the energy levels [see Figure 4(B)]. The transitions between these energy levels are governed by the allowed selection rule ($\Delta m = \pm 1$) and lead to three distinct resonance lines: a central transition and two satellite transitions separated by $\omega_Q$. The nuclear spin relaxation rates of the central line and the

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Multiple quantum coherences. Consider a spin 3/2 nucleus in biological tissues with bi-exponential relaxation and nonzero RQC in a magnetic field. Immediately following the application of a nonselective 90° pulse, the longitudinal magnetization is flipped into transverse plane and then evolves under the influence of biexponential relaxation rates and RQC during the acquisition period, τ. Dropping the factors representing temperature, the signal expression following a 90° pulse is given by (83,84,92),

\[
S(t) \sim M_0 \frac{1}{5} \left( \frac{3}{2} e^{-(R^{(1)}_s - i\omega_q)\tau} + 2e^{-R^{(1)}_s\tau} + \frac{3}{2} e^{-(R^{(1)}_s + i\omega_q)\tau} \right)
\]

where \(M_0\) is the thermal equilibrium magnetization, \(R^{(1)}_s\) and \(R^{(1)}_c\) are relaxation rates of satellite and central transitions respectively and are given above. Here, for the case of illustration, it is assumed that the nonzero quadrupole interaction is large enough to create line splittings.

Triple quantum filtered signal expression. The triple quantum filtered signal is detected by setting \(\theta_1 = \theta_2 = \theta_3 = \pi/2\), and \(\phi_1 = \phi_2 = \phi; \phi_3 = \phi + \pi/2; \phi_4 = 0^\circ\) where \(\phi\) is cycled through 30, 90, 150, 210 and 330° while alternating the receiver phase between 0 and 180°. The addition of all six acquisitions produces the TQF signal:

\[
S(\tau, t) \sim M_0 \frac{9}{80} \left[ \left( e^{-(R^{(1)}_s - i\omega_q)\tau} - 2e^{-R^{(1)}_s\tau} \right) + e^{-(R^{(1)}_c + i\omega_q)\tau} \right] \\
\times \left( e^{-(R^{(1)}_s - i\omega_q)\tau} - 2e^{-R^{(1)}_s\tau} + e^{-(R^{(1)}_c + i\omega_q)\tau} \right)
\]

A series of TQF spectra are collected (for a fixed \(\delta\), usually in few ms) as a function of preparation time, \(\tau\), and the resulting spectral amplitudes are fitted to the signal expression to compute the relaxation rates (an example is shown in the case of in vivo data). The triple quantum filtered signal has 50% higher sensitivity than the DQF signal and is preferred for measuring biexponential relaxation. It does not have complications of mixing of odd and even rank coherences associated with the DQF signal. Furthermore, it can be used with surface coils, which have inhomogeneous \(B_1\) fields, to study relaxation in biological tissues (106,107).

Double quantum filtered signal expression. A double quantum filtered magic angle (DQF-MA) signal is detected by setting \(\theta_1 = \pi/2, \theta_2 = \theta_3 = 54.7^\circ\) and \(\phi_1 = \phi_2 = \phi_3 = \phi\) and \(\phi_4 = 0^\circ\) where \(\phi\) is cycled through 0, 90, 180 and 270° while the receiver phase is alternated between 0 and 180°. This sequence exclusively detects \(T^J_2\) and is important for studying ordered structures.
However, if the flip angles of the last two pulses are set to 90° then the detected DQF signal will be a combination of both $T_2^3$ and $T_3^3$. Addition of all four acquisitions produces the DQF-MA signal:

$$S(t) = M_0 \frac{\left( e^{-(R_1^1 - i\omega_Q)t} - e^{-(R_1^1 + i\omega_Q)t} \right) \left( e^{-(R_1^3 - i\omega_Q)t} - e^{-(R_1^3 + i\omega_Q)t} \right)}{10}$$

Situations corresponding to different motional regimes can be arrived at by choosing appropriate values for $\omega_Q$ and spectral densities and the above signal expressions. The DQF-MA sequence has been used to study ordered structures in cartilage, and cytoskeleton (93,103,105,108,109). Typical simulated spectra using the above signal expressions are shown in Figure 6, and Fig. 7 shows typical SQ, DQF and TQF sodium spectra from articular cartilage. The pronounced negative lobes of DQF spectra are due to the presence of $(T_{2z}^2)$ contribution in addition to $T_{2z}^3$.

Other pulse sequences that are useful in studying ordered structures are based on the Jeener-Broekaert (108,110) and spin-locking sequences (84,111). In both these sequences, the signal is detected by filtering through the quadrupolar order $T_{2z}^0$. The efficiency of both DQF-MA and JB sequences in detecting ordered sodium depends on the precision of the flip angles and hence these methods are very susceptible to $B_1$ inhomogeneities. The spin-locking method is independent of flip angle precision and is more tolerant to $B_1$ inhomogeneities. Recently, newer methods have been introduced to study sodium in ordered systems based on suppression of the central transition by applying soft pulses and employing double frequency sweep pulses. These sequences suppress signal from sodium in the isotropic regime and detect only the satellite signals (112) or convert them to a central transition signal (113,114). These methods are robust in spite of $B_1$ inhomogeneities. Other methods that are promising in detection of ordered sodium ions include quadrupolar filter by nutation (QFN) (115). QFN exploits the dependence of quadrupolar interaction on nutation frequencies to suppress isotropic sodium and detect the central transition of ordered sodium.

Single and multiple quantum sodium NMR has been used extensively to study various biological tissues such as brain (89,90,104,116–120), breast (121) heart (122–128), muscle (129,130), tumors (131–133), blood (122) and cartilage (27,93,134–136) and reviewed in several articles (86,87,91,108,119,137–141). In this review, however, we restrict our discussion to sodium NMR of cartilage.

**Sodium NMR of cartilage**

Owing to the sparse cellular content (~2% by volume) and highly ordered nature of cartilage, most of the sodium in ECM of cartilage is in a slow motion regime and exhibits multiple quantum coherences. The sodium NMR
signal from biological tissues is governed by the relevant values of $M_{0}$, $R_{s}$, $R_{c}$ and $\omega_{Q}$ of the system. It should be noted here that 40% of total sodium is observed directly through the central transition, which decays with a longer time constant ($T_{2}^{s} = T_{2}^{fall} = 1/R_{s}^{(1)}$) compared with the 60% of sodium that contributes to the satellite transitions, which decay with a faster relaxation time ($T_{2}^{fast} = T_{2}^{rise} = 1/R_{c}^{(1)}$). Therefore, in order to quantify the absolute sodium concentration from biological tissues, one has to use ultra-short RF pulses and fast acquisition schemes. Otherwise, significant amount of the fast-decaying signal will be lost before detection and this will lead to an underestimation of the total [Na].

**Sodium MQF NMR of cartilage**

**MQ spectroscopy.** As stated above, anisotropic motion due to an averaged quadrupolar interaction in biological tissues can be detected via the even-ranked double quantum coherence $T_{2}^{s}$. In the presence of anisotropic motion, transverse relaxation times of the satellite and central transitions measured via TQF sequence are referred to as $T_{2}^{rise}$ and $T_{2}^{fall}$. The RQC constant can serve as the local order parameter for a given tissue. Although the SQC signal is sum of exponentials and influenced by RQC, it is difficult to determine the relaxation times accurately from this signal. However, as described above, MQ spectroscopy with ultrashort duration RF pulses enables the measurement of all the parameters that govern sodium spin dynamics in tissues. To this end, TQF spectroscopy has been used to measure biexponential relaxation rates of cartilage both in *vitro* (142) and in *vivo* (143). In this study, a flip-angle independent TQF pulse sequence was employed that permitted the use of a surface coil to transmit and receive. Transverse relaxation times of sodium obtained from asymptomatic volunteers were found to be: $T_{2}^{rise} = 1 \pm 0.12$ ms, $T_{2}^{fall} = 12.0 \pm 0.75$ ms (mean ± SD).

Anisotropic motion of sodium ions was also detected via $T_{2}^{z}$ in cartilage and in collagen fibers. DQF spectroscopy was used to measure the RQC constant in nasal cartilage (93). Analytical expressions for the DQF spectra were obtained by calculating the evolution of the second and third rank tensors by solving a modified Redfield equation. It was shown that anisotropy was due to local rather than macroscopic order. The observed spectra were fitted to DQF signal expressions from several models by assuming an isotropic distribution of the local directors of the locally ordered sites. They showed that a model that assumes a Gaussian-weighted distribution of $\omega_{Q}$ values provided the best fit for the MQF-NMR spectra from fresh bovine nasal cartilage. In this analysis, the local RQC was found to be 550 Hz in nasal cartilage. Using isolated models of cartilage, it was shown that the contribution to RQC in cartilage was due to the ordering of collagen bundles and that PG did not contribute to the RQC. A similar model provided the best fit for Jeener–Broekaert spectra obtained from bovine articular cartilage (144).

The effect of interleukin-1β (IL-1β), a cytokine known to induce matrix degradation, was studied in bovine cartilage explants using TQF and double quantum filtered magic angle (DQF-MA) methods (109). Changes on relaxation times and RQC are measured in bovine cartilage specimens. $T_{2}^{rise}$ ranged between 2.26 and 3.5 ms, decreasing with increased PG loss. $T_{2}^{fall}$ increased from 12.3 to 14.9 ms and $T_{1}$ increased from 16 to 21 ms while $\sigma$ (RMS $\omega_{Q}$) decreased from 180 to 120 Hz over the range of PG depletion investigated. These results indicate that the IL-1β-induced macromolecular depletion has resulted in changes in the local ordering of the tissue. The effect of mechanical compression of cartilage on the MQF sodium spectral line shapes was also analyzed (135). A nonmagnetic compression cell, which enables the NMR experiments while the tissue is compressed, is employed in this study. Bovine cartilage plugs were subjected to single and MQF sodium NMR spectroscopy during a uni-axial compression at 0.7 MPa for 1 h. Compression, although affected the signal amplitude, did not influence the lineshapes of the SQ and TQF spectra significantly. The DQF spectra showed marked line shape changes in the compressed samples, which were attributed to the reduced RQC in compressed samples.

**MQ imaging.** Feasibility of performing TQF sodium imaging of articular cartilage was first demonstrated on bovine cartilage samples (142) then TQF imaging of the human knee was performed *in vivo* (143). In the *in vivo* study, a twisted projection imaging (TPI) sequence with an ultra-short 400 μs echo time was employed. Unlike SQ, TQF signal reaches a maximum at a preparation time of $\tau \approx 3$ ms. Images were acquired with a voxel size of 0.5 cm. The total imaging time for a three-dimensional data set of 16 slices was ~20 min and provided images with a SNR of 8:1. Single quantum images were also acquired with a voxel size of 0.06 cm. Total SQ imaging time for a three-dimensional data set with SNR of 16:1 was ~10 min. The TQF signal obtained a particular $\tau$ value was fitted to the TQF signal expression to calculate the transverse relaxation times. $T_{2}^{rise}$ and $T_{2}^{fall}$ (the asterisk indicates a measurement of $T_{2}$ in the presence of static $B_{0}$ inhomogeneities) measured from these studies are 0.84 and 9.6 ms, respectively. These studies clearly demonstrate the importance of a short-echo imaging sequence like TPI in imaging SQ as well as TQF signal from sodium. It also shows that it takes almost three times the total imaging time to acquire TQF images with the same SNR (but 10 times larger voxels) than SQ images, i.e. the TQF signal is an order of magnitude weaker than the SQ signal. In spite of this, MQF sodium NMR studies are indispensable in the quantification of critical parameters (such as relaxation rates and RQC) that characterize sodium dynamics in biological tissues (137).
Sodium MRI of cartilage fixed charge density (FCD)

As discussed in the Introduction, loss of PG is the initiating event in the OA. The ability to quantify these molecular changes will provide a handle for the early diagnosis and treatment monitoring. Based on the fact that Donnan equilibrium holds for cartilage equilibrated in very dilute solutions, Maroudas et al. have shown that FCD of cartilage is correlated to the GAG content of cartilage (145). Since the FCD is counter-balanced by the Na⁺ ions, loss of PG (hence GAG and FCD) due to cartilage degeneration results in the loss of sodium ions from the tissue. The loss of the negatively charged PG lowers the FCD in the tissue, thereby releasing positively charged sodium ions. Using ideal Donnan equilibrium conditions, FCD can be related to tissue sodium concentration according to the following equation:

\[
\text{FCD (mM)} = \frac{\left[\text{Na}^{+}\right]_{\text{SF}}^2}{\left[\text{Na}^{+}\right]_{\text{tissue}}} - \left[\text{Na}^{+}\right]_{\text{tissue}} \tag{24}
\]

where \(\left[\text{Na}^{+}\right]_{\text{SF}}\) is the sodium concentration in the synovial fluid and \(\left[\text{Na}^{+}\right]_{\text{tissue}}\) is the sodium concentration in the tissue. \(\left[\text{Na}^{+}\right]_{\text{SF}}\) is typically in the range 140–150 mM, while in phosphate-buffered saline (PBS) it is 154 mM.

Healthy human cartilage FCD ranges from −50 to −250 mM, depending on the age and location in the tissue (136). The FCD from [GAG] can be calculated by the following equation by assuming 2 mol of negative charge per mole of chondroitin sulfate (one from sulfate and one from carboxylate) and a molecular weight of chondroitin sulfate of 502.5 g/mol (136):

\[
\text{FCD (mM)} = -2 \times \frac{[\text{GAG} \text{ (mg/L)}]}{502.5 \text{ (mg/mM)}} \tag{25}
\]

Because of very short relaxation time of outer (satellite) transitions, one has to perform spectroscopic and imaging experiments with ultrashort echo times. Since invariably NMR and MRI experiments involve finite pulse lengths, some of the fast decaying components are lost and the signal detected following an RF pulse underestimates the true sodium content. In spite of this, as long as the observed signal is calibrated with appropriate relaxation matched reference sodium phantoms, it is possible to measure absolute sodium content of the tissue.

In an early work, sodium NMR spectroscopy was used to study changes in sodium content and relaxation rates in cartilage. In preliminary studies on cartilage, it was found that loss of PG can cause a decrease in FCD and mobility of sodium ions in the ECM. Paul et al. (146) and Jelick et al.’s (147) measured sodium relaxation rates using single-quantum NMR in nasal cartilage subjected to proteolytic enzymes, trypsin or papain to degrade PG. In this study, it was observed that both \(T_1\) and \(T_2\) relaxation rates were increased following the trypsin or papain treatment. Sodium NMR visibility in cartilage was determined by measuring sodium content using NMR and comparing it with that measured from inductively coupled plasma emission spectroscopy and it was found that sodium in cartilage was 100% NMR visible (136). Sodium content measured using NMR was then used with ideal Donnan theory to estimate FCD and for calf articular cartilage, near physiological conditions, calculated FCD was found to be −280 ± 30 mM. In the epiphyseal cartilage, FCD varied with the position of origin of the tissue within the joint, ranging from −190 to −350 mM in a manner that correlated with tissue GAG content. Similar variations of sodium concentration were also found in a low-resolution sodium image of intact ulnar epiphyseal cartilage.

Changes in sodium content and relaxation times using single quantum sodium MRI and TQF spectroscopy were investigated (148). Over a 50% PG depletion induced by trypsin from bovine cartilage, sodium content changed almost linearly and \(T_1\) increased from 18 to 26 ms, \(T_2\) increased from 7.5 to 12 ms and \(T_2^*\) decreased from 2 to 1 ms. These results indicate that care must be taken to account for these changes in the relaxation times with PG depletion/degeneration of the tissue when sodium MRI is used to quantify tissue [Na]. Also in this study, sodium images of bovine cartilage plugs were presented that displayed reduced sodium intensity with increased PG depletion. However, no attempts were made to calculate the FCD of the tissue.

Low-resolution sodium MRI was acquired from human knee in vivo (149) at 1.5 T. Reddy et al. (27) were the first to demonstrate the feasibility of acquiring a high-resolution (voxel size of 6.25 μL) three-dimensional data set of sodium images of the knee of healthy human volunteers with excellent SNR (16:1) at 4 T (Figure 8). Sodium images were also compared with corresponding proton images to demonstrate the differences in tissue contrast. It was further demonstrated that high-resolution images obtained on bovine patellar cartilage, half of which was subjected to trypsin-induced PG depletion, clearly demarcated the intact tissue (with high sodium content) from PG-depleted tissue (which had lower sodium content). However, in these studies neither sodium nor FCD measurements were performed.

The effect of mechanical compression on sodium and proton NMR relaxation times of bovine articular cartilage specimens was measured as a function of PG depletion. Uni-axial mechanical compressions were performed with an MR-compatible pressure cell and evaluated dynamically via interleaved one-dimensional proton and sodium MR projection imaging (150). Upon full compression, in normal cartilage, sodium \(T_1\) and \(T_2\) relaxation times were decreased by 38 and 37%, respectively, whereas in PG depleted tissue, following full compression, sodium \(T_1\) and \(T_2\) were decreased by 20 and 39%, respectively.

The sensitivity of sodium and proton MRI in detecting trypsin-induced PG changes in bovine articular cartilage


---

**Equation 24**

\[
\text{FCD (mM)} = \frac{\left[\text{Na}^{+}\right]_{\text{SF}}^2}{\left[\text{Na}^{+}\right]_{\text{tissue}}} - \left[\text{Na}^{+}\right]_{\text{tissue}} \tag{24}
\]

**Equation 25**

\[
\text{FCD (mM)} = -2 \times \frac{[\text{GAG} \text{ (mg/L)}]}{502.5 \text{ (mg/mM)}} \tag{25}
\]
specimens was investigated at 4 T (151). Over a ~20% PG depletion, sodium image signal change correlated well with the observed PG loss ($r^2 = 0.85, p < 0.01$) while proton density-weighted image intensity change did not exhibit a definite trend ($r^2 = 0.10, p < 0.8$). The change in proton $T_1$ and $T_2$ between depleted and nondepleted tissue regions also did not correlate with PG loss ($r^2 = 0.07$ and $r^2 = 0.06$ respectively). Results from this study indicate that sodium MRI is both sensitive and specific in detecting small changes in PG content, whereas proton density and relaxation properties are not sensitive to small changes in cartilage PG.

Shapiro et al. described a method to quantify sodium concentration in cartilage (29). Sodium concentration in bovine patellar cartilage was measured by three different methods: NMR spectroscopy of whole cartilage, NMR spectroscopy of liquefied cartilage in concentrated HCL and inductively coupled plasma emission spectroscopy. Using a three-dimensional fast gradient-echo (FGRE) pulse sequence, intact bovine patellae were imaged along with relaxation normalized calibration phantoms to map sodium concentration in cartilage (Figure 9). It was found that sodium concentrations in intact articular cartilage ranged from ~200 mM on the articular and sub-chondral bone surfaces to 390 mM in the middle with an average of 320 mM in several patellae studied. Average measurements from these sodium maps correlated well with those obtained from the spectroscopic methods and sodium was found to be 100% NMR visible in cartilage plugs. In another study, macromolecular depletion was induced in bovine cartilage plugs using trypsin, and the change in [Na] was measured via sodium MRI using the same approach. Following the imaging experiments, the tissue and PG depletion media were subjected to standard dimethylmethylene blue PG assay. A high correlation (slope = 0.89 and $r^2 = 0.81$) between the FCD measure-

**Figure 8.** (A) Axial slice of sodium image of a healthy human knee. The high sodium content in articular cartilage is clearly visible. The signal profile across the cartilage is shown in (B) (27).

**Figure 9.** Quantitation and calibration of sodium concentration in articular cartilage. Sodium image of bovine cartilage (A). Trypsin-treated region has low [Na] due to reduced GAG. Circular sodium agarose phantoms that were used to quantify [Na] are also seen in this figure. A plot of FCD measured from sodium MR and that obtained from DMMB assay (B) shows a strong linear correlation between these measurements (29).
ments obtained by $^{23}$Na MRI and those obtained by the PG assay (Figure 9) was found. Using the above-described approach involving calibration phantoms, sodium concentration was measured from human cartilage specimens obtained from knee replacement surgery. Clear differences in sodium concentrations in healthy and osteoarthritic specimens were noted (Figure 10).

The same group measured FCD from articular cartilage in the knee of healthy volunteers using two different RF coils, a birdcage coil and a transmit/receive surface coil. High-resolution sodium imaging data (voxel size $= 14.6 \mu L$, SNR $= 12:1$ and a total imaging time of 30 min) obtained with both methods gave similar results, with an average FCD of $-158$ to $-182 \text{mmol}$. This variation in FCD has been interpreted as being due to differences in PG content across the tissue. The calibration studies from this work form the basis for computing FCD using sodium MRI. In surface coil experiments, phantom positions were adjusted to match the distance of cartilage from the plane of the surface coil (141).

Although this approach avoids the necessity of compensating for signal drop-off from the inhomogeneous $B_1$ of a surface coil, any mismatch would lead to erroneous results, a problem that was corrected in a subsequent study by Wheaton et al. (30). In this method, the signal intensity of each sodium MR image was corrected for $B_1$ inhomogeneity, as well as for $T_1$ and $T_2$ weighting, on a pixel-by-pixel basis using a factor derived from the following equation:

$$S_{\text{cor}}(x,y) = \frac{S_{\text{ori}}(x,y)}{B_{1\text{map}}(x,y)e^{-\frac{TE}{T_2^*}}(1 - e^{-\frac{TR}{T_1}})}$$

where $S_{\text{cor}}(x,y)$ and $S_{\text{ori}}(x,y)$ are the corrected and original signal intensities, respectively, and $B_{1\text{map}}$ is the value of the $B_1$ sensitivity map.

The $T_1$ and $T_2$ values of the phantoms measured in progressive saturation experiments were 22 and 8 ms, respectively. The $T_1$ and $T_2$ values used for the human patellar cartilage were 23 and 4.5 ms, respectively, as estimated from data obtained in progressive saturation experiments performed in ex vivo bovine patellae. Using this approach, sodium MRI experiments were performed on the knee cartilage of healthy as well as early stage OA patients at 4 T and demonstrated the feasibility of sodium MRI in computing PG loss in early stage OA. The sodium three-dimensional image data set was acquired with a voxel size of $5.3 \mu L$ with an SNR of 12:1 in about 20 min. In comparison, it took 30 min to obtain a three-dimensional data set using a volume coil with the same SNR but an increased voxel size of $14.6 \mu L$. With the effects of $B_1$ inhomogeneity and voxel size accounted for, the surface coil provides an improvement in SNR by a factor of 2. Alternately, an image can be obtained with a surface coil in 25% of the time that it takes to acquire one with a volume coil image with an identical SNR and voxel size. Results from this study revealed that cartilage of healthy subjects had a mean FCD of $-182 \pm 9 \text{mm}$.

Figure 10. Sodium concentration maps of human patellar cartilage specimens obtained following knee replacement surgery. The top image is from a healthy cartilage while the bottom image is that from an osteoarthritic patient. The scale bar indicates sodium concentration in mm.
Table 2. Sodium concentration and FCD of healthy human subjects measured from sodium MRI (30)

<table>
<thead>
<tr>
<th>Sodium concentration (mmol/l)</th>
<th>FCD (mmol/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>248 ± 39</td>
<td>−175 ± 50</td>
</tr>
<tr>
<td>248 ± 29</td>
<td>−175 ± 35</td>
</tr>
<tr>
<td>259 ± 24</td>
<td>−187 ± 68</td>
</tr>
<tr>
<td>262 ± 60</td>
<td>−192 ± 69</td>
</tr>
<tr>
<td>245 ± 48</td>
<td>−171 ± 80</td>
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<tr>
<td>249 ± 49</td>
<td>−176 ± 64</td>
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<td>254 ± 36</td>
<td>−182 ± 45</td>
</tr>
<tr>
<td>260 ± 60</td>
<td>−190 ± 75</td>
</tr>
<tr>
<td>265 ± 56</td>
<td>−196 ± 74</td>
</tr>
</tbody>
</table>

Table 3. FCD measured from sodium MRI of symptomatic osteoarthritic subjects. Percentage change is measured w.r.t the values obtained from healthy subjects (30)

<table>
<thead>
<tr>
<th>Subject</th>
<th>FCD (mmol/l)a</th>
<th>Percentage changeb</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>−108 ± 56</td>
<td>−41</td>
</tr>
<tr>
<td>2</td>
<td>−114 ± 30</td>
<td>−38</td>
</tr>
<tr>
<td>3</td>
<td>−144 ± 43</td>
<td>−22</td>
</tr>
</tbody>
</table>

aMean FCD± SDs.

bPercentage change in mean FCD from normal value—that is, relative to mean FCD of patellae of the healthy subjects (−182 mmol/l).

Table 4. $T_{1p}$ relaxation data from osteoarthritic subjects (239)

<table>
<thead>
<tr>
<th>Subjects</th>
<th>$T_{1p}$ (ms), mean ± SD</th>
<th>$T_{1p}$ elevation (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>63 ± 4</td>
<td>26</td>
</tr>
<tr>
<td>2</td>
<td>64 ± 3</td>
<td>28</td>
</tr>
<tr>
<td>3</td>
<td>65 ± 5</td>
<td>30</td>
</tr>
<tr>
<td>4</td>
<td>72 ± 4</td>
<td>44</td>
</tr>
<tr>
<td>5</td>
<td>75 ± 6</td>
<td>50</td>
</tr>
<tr>
<td>6</td>
<td>95 ± 12</td>
<td>90</td>
</tr>
</tbody>
</table>

Feasibility of detecting cytokine-induced arthritic changes in an animal model using sodium MRI. In this study, a biochemical condition similar to OA was created by an intra-articular injection of recombinant porcine interleukin-1β (IL-1β) into the knee joint of Yorkshire pigs. The contralateral knee joint was given a saline injection to serve as the control. Sodium MRI data was acquired at 4 T after 6 h following the injection of IL-1β. Following in vivo imaging, the tissue and joint fluid were harvested and subjected to in vitro sodium MRI and histologic and immunohistochemical analyses as independent measurements of the cytokine activity and PG loss. Sodium images obtained from treated and untreated joints of pigs are shown in Figure 12. Sodium image data was used to generate FCD maps. On average, the FCD of cytokine-treated cartilage was 49% lower than that of saline-treated cartilage, reflecting a loss of PG content. These results were supported by histologic and immunohistochemical findings, most notably a reduction in staining for PG and an increase in matrix metalloproteinases in the synovial fluid.

Summary

The sodium spectroscopy work described above was performed at varying field strengths ranging from 2 to 9.4 T. MQF spectroscopy is ideally suited for measuring all the parameters that govern sodium dynamics in cartilage. These are indispensable in obtaining accurate quantification of relaxation rates and RQC, and hence information about ordered tissue. However, the order of magnitude lower SNR of MQF signals restricts them to spectroscopy mode. Most MQF spectroscopy to date has been carried out at magnetic fields greater than 4 T.

The majority of sodium MRI experiments, however, have been performed at 3 and 4 T. In these studies, three-dimensional sodium MRI of 16 slices in the knee joint was performed with volume RF coils in less than half an hour with a voxel size of 6 µL with an SNR of 12:1. The echo times employed in these experiments was a little over 2 ms. However, by exploiting the increased sensitivity of surface coils, it was possible to improve the SNR to 16:1 and the imaging time was reduced to less than 20 min. Sodium imaging at 4 T clearly demonstrated the feasibility of measuring cartilage FCD in vivo in...
of imaging experiments reviewed here employed echo times and other hardware requirements, most of the sodium furthermore, due to the limitations of gradient strengths that enable accurate quantification of cartilage FCD. the disadvantages of sodium MRI are that it requires field strengths of \( \geq 3 \) T to obtain quality sodium images that enable accurate quantification of cartilage FCD. Furthermore, due to the limitations of gradient strengths and other hardware requirements, most of the sodium imaging experiments reviewed here employed echo times of \( \geq 2 \) ms. Since the \( T_{2f} \) of cartilage lies in the range 1–2 ms, substantial signal is lost before the acquisition. This is a major contributor to the low SNR of sodium compared with conventional proton MRI. Additionally, the sodium gyromagnetic ratio, \( \gamma_s \), is one-quarter that of protons, hence sodium MRI requires four times stronger gradients to obtain images with identical resolution to that of proton MRI. With the exception of MQF-prepared TPI imaging, sodium MQF imaging is currently not clinically feasible due to the low sensitivity of the technique and has therefore been relegated to spectroscopic methods to quantify structures of materials and specimens.

However, recent advances in the gradient technology (with a gradient strength of \( \geq 4 \) G/cm) may enable one to achieve ultrashort \( TE \) (\(<200\) \( \mu \)s) that can significantly improve resolution and SNR. Radiofrequency coil technology (multiple channel capability) and parallel imaging approaches such as SENSE (156) and SMASH (157) and tuned pre-amplifiers would further contribute to high SNR. These advances may potentially make clinical sodium MRI feasible on 3 T scanners. Further, the recent proliferation of 7 T whole-body MRI scanners in clinical research centers could have a significant impact on sodium MRI and its potential for clinical use. Since SNR scales as \( B_0^{2/3} \) (158–161) and the lack of \( B_1 \) penetration and \( B_0 \) susceptibility are issues that plague proton imaging, sodium MRI can be particularly advantageous at higher fields. Further, unlike proton \( T_1 \), which increases with field, as the \( T_1 \) of sodium is predominantly due to quadrupolar interaction, it may not change appreciably at higher field. This retains the rapid averaging capability of sodium MRI even at high fields. The low \( \gamma \) of sodium will also mean significantly lower power deposition compared with proton imaging. It is therefore very likely that, with the improved SNR, sodium MRI at 7 T and higher fields would emerge as a robust tool for quantitative imaging of cartilage integrity.

**\( T_{1P} \) MRI**

In \( T_{1P} \) MRI, a long-duration, low-power RF pulse referred to as the ‘spin-lock’ (SL) pulse is applied to the magnetization in the transverse plane. The magnetization undergoes relaxation in the presence of the applied \( B_1 \) field in the rotating frame, a situation similar to that of the longitudinal magnetization in the \( B_0 \) field. This spin-locked magnetization will relax with a time constant \( T_{1P} \), the spin-lattice relaxation in the rotating frame, during the time of the spin-lock pulse (TSL). The \( B_1 \) field attenuates the effect of dipolar relaxation, static dipolar coupling, chemical exchange and background gradients on the signal. \( T_{1P} \) is always greater than \( T_2 \). In a typical \( T_{1P} \) MRI experiment, the TSL time is incremented while the amplitude of SL pulse (\( \gamma B_1 \approx 0.1 \) to a few kHz) is fixed. Alternatively, the measurement of \( T_{1P} \) as a function of the \( B_1 \) amplitude, for a fixed spin-lock length, is also possible. The ‘\( T_{1P} \)-dispersion’ curve obtained in this case is
governed by the spectral density components of the sample that are in the neighborhood of $gB_1$.

Measurement of the MR signal following the SL pulse (Figure 13) is ideal for spectroscopic measurement of $T_1^r$ of a sample. However, for imaging applications, it is often more convenient to prepare the magnetization using a pulse cluster shown in Figure 14. The thermal equilibrium magnetization vector ($M_0$), initially along the $z$-axis (panel a), is nutated by the first hard 90° pulse which is applied along the $x$-axis in the rotating frame of reference, into the transverse plane (panel b). The magnetization, now along the $y$-axis, is immediately spin-locked by the SL pulse (panel c). For the duration of the TSL period, the magnetization decays with a time constant $T_1^r$ (panel d). The magnetization thereby becomes ‘$T_1^r$-prepared’ and results in a $T_1^r$-weighted signal as a function TSL. This $T_1^r$-prepared magnetization is then restored to the longitudinal axis by the second hard 90° pulse applied along the negative $x$-axis (panel e) and can be spatially encoded by appending any imaging pulse sequence to this $T_1^r$ pulse cluster. A pixel-by-pixel fit of the spin-locked signal to an appropriate expression provides $T_1^r$. In the above description, on-resonance condition is assumed. The $T_1^r$ experiment can also be performed in off-resonance condition to quantify $T_{1^o}^r$, where the magnetization is spin-locked at an effective field given by:

$$\omega_{\text{eff}} = \sqrt{\omega_1 + \Delta \omega}$$

(27)

where $\Delta \omega$ is the resonance offset.

In biological tissues, off-resonance RF pulse, in addition to spin-locking, also leads to ‘magnetization transfer’ effect (162) which may complicate the analysis of $T_{1^o}^r$. The details of off-resonance experiments can be found in the literature (163–165). However, for the rest of the sections we restrict ourselves to describing $T_1^r$ in the on-resonance condition. $T_1^r$ studies in solution-state NMR have been reviewed extensively (166–168). There has been a considerable amount of work on biological tissues using $T_1^r$-spectroscopy and imaging dealing with tumors, muscle, myocardium, blood flow and cartilage (26,165,169–179).

**Imaging pulse sequences**

**Two-dimensional $T_1^r$ MRI.** Initial $T_1^r$ imaging experiments were performed on articular cartilage using a $T_1^r$
sequence based on a spin-echo, or SE (24), and fast spin echo, or FSE (26). Both these sequences have been implemented on multiple clinical MRI systems and field strengths to produce a single-slice $T_{1,ρ}$-weighted image. A typical image data set containing four or five images acquired at different TSL times can be fitted to the following expression to generate a pixel-by-pixel $T_{1,ρ}$-map:

$$S_y(TSL) = S_0 \exp\left(-\frac{TSL}{T_{1,ρ}}\right) + C \quad (28)$$

### Three-dimensional $T_{1,ρ}$ MRI

Recently, a $T_{1,ρ}$-prepared three-dimensional gradient-echo (GRE) based pulse sequence was developed and implemented using either adiabatic pulses (131) or hard pulses (180) in the $T_{1,ρ}$ preparation period. In the three-dimensional sequence, the magnetization is $T_{1,ρ}$-prepared just as in the case of the two-dimensional sequences. However, in place of the 90° slice-selective pulse of the SE or FSE sequence, a small angle $α$ pulse is used instead to excite a slab; this is followed by a conventional three-dimensional GE readout.

Signal intensities from three-dimensional data sets obtained at varying spin-lock lengths are fitted to the following signal expression:

$$S(TSL) \propto \frac{e^{-\frac{TSL}{T_{1,ρ}}} \left(1 - e^{-\frac{(TR - TSL)}{T_1}}\right) \sin α}{1 - e^{-\frac{TSL}{T_{1,ρ}}} e^{-\frac{(TR - TSL)}{T_1}} \cos α} \quad (29)$$

where $S(TSL)$, $TSL$, $TR$, $T_1$ and $α$ represent signal intensity at a given spin-lock duration and frequency, spin-lock length, pulse repetition time, longitudinal relaxation time and flip angle of the slice selective pulse, respectively.

Like most three-dimensional GE-based sequences, the three-dimensional $T_{1,ρ}$ sequence enables rapid data collection but produces images with a reduced SNR compared with SE-based images. Further, to accelerate the acquisition, a short $TR$ is typically employed, which introduces significant $T_1$ weighting in the resultant image and may obscure $T_{1,ρ}$-based contrast.

Advantages of this sequence are that it provides volumetric $T_{1,ρ}$ maps and that one of the early data set can be used to compute cartilage volume.

### Multi-slice $T_{1,ρ}$ MRI with spin-echo readout

To overcome the limitations, such as low SNR and poor $T_{1,ρ}$ contrast, associated with volumetric data acquisition with a three-dimensional $T_{1,ρ}$ prepared sequence, a two-dimensional multi-slice spin-lock (MS-SL) pulse sequence was developed (181). One of the challenges in implementing the $T_{1,ρ}$ sequence in multi-slice mode is due to the nonselective nature of the spin-lock pulse, which saturates the longitudinal magnetization from nonexcited regions. In this sequence, saturation of longitudinal magnetization by the application of nonselective SL pulses is experimentally measured and theoretically modeled as $T_{2,ρ}$ decay. The saturation data was used to correct the image data as a function of the SL pulse duration to make quantitative measurements of $T_{1,ρ}$.

It was found that $T_{1,ρ}$ measured using saturation corrected MS-SL data is identical to that measured using SS-SL sequence (182). $T_{1,ρ}$ relaxation maps are computed for each slice by fitting the image data pixel-by-pixel to eqn (28). During the period between each SL pulse, $τ = TR/N$, where $N$ is the total number of slices acquired, the longitudinal magnetization recovers according to $T_1$. After each $τ$ delay, the magnetization becomes saturated by a factor of $M^{sat}$. Therefore, at the end of the $N$th growth delay period, the longitudinal magnetization, $M_{z,n}$, is given by:

$$M_{z,n} = M^{sat}(TSL) \times \left[ M_{z,n-1} \exp\left(-\frac{τ}{T_1}\right) + M_0 \left(1 - \exp\left(-\frac{τ}{T_1}\right)\right) \right] \quad (30)$$

where

$$M^{sat}(TSL) = \exp\left(-\frac{TSL}{T_{2,ρ}}\right) \quad (31)$$

and

$$\frac{1}{T_{2,ρ}} = \frac{1}{2} \left[ \frac{1}{T_1} + \frac{1}{T_2} \right] \quad (32)$$

### Multi-slice $T_{1,ρ}$ MRI with spiral readout

Recently a multi-slice $T_{1,ρ}$ mapping with spiral readout was implemented at 3 T (183,238). In this sequence, $T_{1,ρ}$ prepared magnetization was read by a multi-slice spiral readout. Acquiring a second image that has an inverted longitudinal magnetization compensated for $T_1$-weighting. The total imaging time for collecting a three-dimensional data set of 14–16 slices was $\sim 13$ min. Representative data obtained with this sequence is shown in Figure 34 and was limited to an axial orientation because spiral acquisition does not allow the use of an anti-aliasing filter that is necessary for sagittal or coronal plane acquisition with a knee coil.

### Rapid volumetric $T_{1,ρ}$ MRI with SSFP readout

In order to implement any research sequence in a clinical setting for quantitative imaging, it should provide three-dimensional coverage and have high precision and high temporal efficiency without exceeding SAR limits. Since $T_{1,ρ}$ mapping involves the collection of at least four three-dimensional data sets at varying SL times, it is inherently inefficient. Although the three-dimensional FGRE and multi-slice sequences described above have been implemented on clinical scanners, they typically require 20–25 min to gather a single $T_{1,ρ}$ map. Further, three-dimensional $T_{1,ρ}$ maps are typically collected with 2–4 mm slice thickness as it is too time-consuming to
collect three-dimensional maps with isotropic voxel sizes. Therefore, at least two views, e.g. sagittal and axial, are required to properly visualize anatomical structures in three-dimensional $T_1$ maps.

To accomplish this, we recently incorporated $T_1$ imaging with a balanced steady-state true precession (SSFP or true FISP or TRUFI) pulse sequence (192). The performance of this sequence (Figure 15) was verified by comparing the $T_1$ relaxation maps obtained with this sequence with that of a standard single slice $T_1$-TSE sequence. The error in computing $T_1$ with $T_1$-TRUFI was ~5%. At the same time, the total imaging time for collecting three-dimensional $T_1$ maps in both axial and sagittal views was ~20 min. The average $T_1$ of cartilage was insignificantly different between the $T_1$-TRUFI (39.4 ms) and $T_1$-TSE (38.4 ms) readout methods (Figure 16). Integrating this sequence with parallel imaging techniques is expected to further reduce imaging time.

**SAR considerations**

During MRI, the power of the applied RF pulses has to be monitored at all times to ensure that the energy deposited in the patient remains below Food and Drug Administration (FDA) mandated safety levels for the specific absorption rate. As a safety precaution, invariably all the clinical scanners have built in mechanism to stop the scan if the FDA-mandated power level is exceeded at any point. While the most accurate method would be to measure the power output to the RF coil in real time, a priori knowledge of the specific absorption rate (SAR) of a pulse sequence can provide additional safety and save valuable experiment time. A method to calculate SAR has been proposed by Collins et al. (193), in which the SAR was calculated from the electric field generated by a typical quadrature coil on a simulated head of known tissue electrical properties and material density. Using this method, the maximum SAR for a 3 ms long
rectangular pulse that achieves a flip angle of 90°, or SAR(90°, 3), in a simulated quadrature head coil was determined to be 1.46 W/kg in 1 g of tissue. The value was actually smaller (0.708 W/kg in 1 g of tissue) if a sphere containing only muscle tissue was used in the simulation instead. The SAR, of a single pulse of flip angle \( \alpha \) and duration \( \tau \) (in ms), may be calculated using the eqn (193):

\[
SAR(\alpha, \tau) = f \left( \frac{3}{2} \right)^2 \left( \frac{\alpha}{90°} \right)^2 SAR(90°, 3) \tag{33}
\]

where \( f \) is a shape factor and equals 1 for a hard pulse or equals the width of the central lobe at the zero crossing point for a sinc pulse.

The minimum \( TR \) for the pulse sequence was determined from the equation:

\[
TR_{\text{min}} = \frac{\sum_{n=1}^{N} SAR(\alpha_n, \tau_n) \times \tau_n}{\text{SAR}_{\text{FDA}}} \tag{34}
\]

where SAR(\( \alpha_n, \tau_n \)) was calculated from eqn (33) for all \( N = 4 \) pulses in the sequence and \( \text{SAR}_{\text{FDA}} \) is the FDA-suggested maximum SAR level and equals 12 W/kg in 1 g of tissue in the extremities (193).

SAR measurements for three-dimensional \( T_{1\rho} \) weighted MRI at 1.5 T are described. The signal expression required to fit the data is introduced and critical analysis of energy deposition of the RF pulses in the sequence was performed and validated with experimental data on a known phantom. The temperature increase in the phantom resulting from the sequence was 0.015°C, which is well below the established safety guidelines. Performance of the three-dimensional \( T_{1\rho} \) pulse sequence was shown by computing \( T_{1\rho} \) maps of bovine patellar cartilage and comparing the data with that obtained with two-dimensional \( T_{1\rho} \) mapping sequence. The calculated \( T_{1\rho} \) of articular cartilage in the specimen was similar for both three-dimensional and two-dimensional methods (84 ± 2 and 80 ± 3 ms, respectively). Three-dimensional \( T_{1\rho} \)-weighted images obtained from human knee joint \textit{in vivo} demonstrate an improved contrast between cartilage and surrounding structures.

A low SAR \( T_{1\rho} \) pulse sequence

Recently, a reduced SAR version of a \( T_{1\rho} \) mapping sequence was developed (194). This method exploits a partial k-space acquisition approach in which a full power spin-lock pulse is applied only to the central phase-encode lines of k-space, while the remaining phase encode lines receive a low-power (50% amplitude of the maximum) spin-lock pulse. Acquisitions of high- and low-power phase-encode lines are interleaved temporally to minimize average power deposition. This strategy ensures that the majority of signal energy in the central portion of the k-space was fully \( T_{1\rho} \)-weighted, while at the same time the overall SAR of the acquisition was lower, and consequently, total imaging time was reduced. It was demonstrated that, using this approach in the human brain, the SAR could be reduced by 40% while the measurements of \( T_{1\rho} \) changed by only 2%. Similarly, a ‘keyhole’ acquisition approach was exploited in reducing the overall imaging time by 40% for \( T_{1\rho} \) mapping (195). These approaches can be combined with any of the acquisition sequences described above to reduce SAR and/or improve temporal resolution.

Mechanism of \( T_{1\rho} \) relaxation in cartilage

The spin interactions with different internal Hamiltonians (\( J \)-coupling, chemical shift, dipole–dipole interaction, chemical exchange, etc.) will take place in the presence of continuous RF fields of SL pulse, thereby imparting time dependence to the interaction Hamiltonians. Thus the RF field not only introduces time dependence into the Hamiltonian, but also changes the axis of quantization of the spins and modifies other characteristics that determine the relaxation behavior. Redfield first demonstrated the spin-lock phenomenon in solid materials and presented relaxation theory in the presence of a time-dependent Hamiltonian (196,197). Later, several investigators developed the theory for applications in liquid state in different conditions (167).

The interactions that are studied using this methodology can be broadly categorized into (i) scalar-coupling, (ii) dipole–dipole and (iii) chemical exchange processes. In NMR, the spin-lattice relaxation in the laboratory frame or rotating frame, the dynamical information is contained in spectral density functions. Depending upon the method used, the experiment may be simultaneously sensitive to more than one time scale. One method of probing relaxation mechanisms is to use the so-called NMR ‘dispersion’, which measures relaxation times at different field strengths using a field cycling approach (198). Although this method has provided insights into relaxation phenomena of tissues, it is limited to measuring \( T_1 \) at different field strengths. With this method, study of slow motions on the order of kHz is difficult as they suffer from poor SNR. On the other hand, \( T_{1\rho} \) experiments can be performed at high fields and have the potential to provide information about the low frequency motions (~few kHz) in biological systems with high SNR. Although there is abundant literature on studies of \( T_1 \) and \( T_2 \) relaxation times in biological systems, there are relatively few studies on \( T_{1\rho} \) mechanisms in biological tissues at high static fields.

In biological tissues, the \( T_{1\rho} \) relaxation may have contributions from several interactions. Depending upon the tissue type, more than one mechanism may be operative simultaneously but with different relative contributions. In what follows, a brief description of individual relaxation mechanism that may be operative in biological tissues is provided.
Dipolar relaxation. Since the first introduction by Redfield, spin-lattice relaxation in the rotating frame has been studied extensively under different limiting conditions. Using the density matrix approach assuming stochastic time dependence for the lattice variables, a general equation representing the contribution of nuclear spin relaxation due to fluctuating magnetic dipoles to $T_{1 \rho}$, in weak collision limit was obtained by Kelly and Sholl (199):

$$
\frac{1}{T_{1 \rho}} = \frac{3}{8} \gamma^4 \hbar^2 I(I + 1)
$$

$$
\times \left[ \frac{\tau_r}{1 + 4 \omega_1^2 \tau_r^2} + \frac{\tau_r}{1 + \omega_0^2 \tau_r^2} + \frac{\tau_r}{1 + 4 \omega_0^2 \tau_r^2} \right]
$$

and

$$
\frac{1}{T_1} = \frac{3}{2} \gamma^4 \hbar^2 I(I + 1) \left[ \frac{\tau_r}{1 + \omega_0^2 \tau_r^2} + \frac{\tau_r}{1 + 4 \omega_0^2 \tau_r^2} \right]
$$

where $I$ is the nuclear spin number, $\omega_1 = \gamma B_1$, and $\omega_0 = \gamma B_0$, $\gamma$ is the gyromagnetic ratio of the nuclei, $B_0$ the Zeeman static magnetic field and $\tau_r$ is the molecular rotational correlation time.

From the $T_{1 \rho}$ equation, it is clear that it is sensitive to two different time scales, $\omega_1$ and $\omega_0$. Therefore, if the experiment is carried out at constant Zeeman field, the last two terms of the equation will be constant. The last two terms, representing the spectral densities at $\omega_0$ and $2\omega_0$ ($J_1$ and $J_2$) can be determined from combination of $T_1$ and $T_{1 \Omega}$ (Jeener–Broekaert) experiments at the same Zeeman field. Then $J(2\omega_0)$ can be determined by measuring $1/T_{1 \rho}$ at different $B_1$ fields. In general, dipolar relaxation is mostly governed by molecular rotational motion with rotational correlation time $\tau_r$. In biological tissues, frequency dependence on relaxation rates, relaxation-diffusion, may arise from (i) rotational motion of a fraction of water bound to proteins, (ii) exchange of protons on macromolecules with bulk water and (iii) the nonaveraged residual dipolar interaction (RDI) of spin associated with oriented macromolecules in the tissue.

Effect of diffusion. It has been shown that, in biological tissues, diffusion (rotational and translational) contributions to all relaxation rates are constant and independent of frequency (200).

Effect of chemical exchange. In proteins and biological tissues, exchange between protons in different environments is expected to contribute to $T_{1 \rho}$ relaxation. Prominent exchange mechanisms may be classified as exchange between (i) water molecules in bulk and hydration water on proteins, (ii) hydration water and -OH and -NH protons on proteins (exchange between two sites) and (iii) $H_2^{17}$O and $H_2^{18}$O molecules (scalar relaxation).

Simple exchange of protons between water molecules causes $T_{1 \rho}$, dispersion provided the exchange times ($\tau_e$) satisfy the condition that $\omega_1^2 \tau_e^2 \approx 1$. When this is valid, $\omega_0^2 \tau_e^2 \gg 1$. This type of exchange contribution to the $T_1$ relaxation rate is zero, it contributes a constant to the $T_2$ relaxation rate $[1/T_1 \approx 0 \quad 1/T_2 \approx \Delta \tau_e]$ and the $T_{1 \rho}$ relaxation rate has a frequency dependence form as shown below:

$$
\frac{1}{T_{1 \rho}} \approx A \cdot \frac{\tau_e}{1 + \omega_1^2 \tau_e^2}
$$

Several groups have investigated exchange mechanisms in a variety of cases using $T_{1 \rho}$ and Carr–Purcell–Meiboom–Gill (CPMG) pulse sequences (201–206). Hills and Virta et al. (207–213) performed $T_{1 \rho}$ studies on proteins, carbohydrates, gelatin, glucose and dextran solutions and demonstrated that the primary mechanism operative in these systems is the exchange between protons in -OH groups on carbohydrates and -OH groups on water molecules. Starting with the Bloch–McConnell equations, Trott et al. have derived new expressions for the spin-lattice relaxation rate constant in the rotating frame ($1/T_{1 \rho}$), $R_{1 \rho}$, for chemical exchange between two sites, A and B, that have distinct magnetic environments and Larmor frequencies (214,215).

$$
R_{1 \rho} = R_1 \cos \theta + R_2 \sin^2 \theta + \frac{\sin^2 \theta p_a p_b \delta^2 k}{\omega^2 \text{eff}^2 + \Delta \Omega^2 + k^2}
$$

$$
\delta = \delta_b - \delta_a = \Omega_b - \Omega_a
$$

$$
\omega^2 \text{eff} = \omega^2_a + \omega^2_b
$$

$$
\omega^2_a = \Delta \Omega^2 + \omega^2_0
$$

$$
\theta = \arctan \left( \frac{\omega_0}{\Delta \Omega} \right)
$$

$$
\Delta \Omega = \Omega_a + \omega_0
$$

where $R_1$ and $R_2$ are the intrinsic longitudinal and transverse relaxation rates respectively, resulting from processes other than chemical exchange.

If $R_1$ and $R_2$ are assumed to be population averages, then the effect of difference between the intrinsic relaxation rates for species in sites A and B on $R_{1 \rho}$ is negligible. $\Omega_a$ and $\Omega_b$ are the Larmor precession frequencies of sites A and B, respectively, and $k = k_a + k_b$ is the exchange rate and site populations are $p_a = k_a/k$ and $p_b = k_b/k$.

This equation is simplified if one of the sites is much more populated than the other. In this asymmetric populations limit, $p_a \gg p_b$; $\delta_a \approx \Delta \Omega$ and $\omega^2 \text{eff} \approx \omega^2 \text{eff}$:

$$
R_{1 \rho} = R_1 \cos \theta + R_2 \sin^2 \theta + \frac{\sin^2 \theta p_a p_b \delta^2 k}{\omega^2 \text{eff}^2 + k^2}
$$

This equation generalizes the expression previously reported for the special case $R_1 = R_2$ and $\Omega = \omega_0$. When
‘θ’ is 90°, which is the on-resonance condition, then it simplifies to more familiar expression with \( R_2 \) representing ‘exchange free’ transverse relaxation rate:

\[
R_{1\rho} = R_2 + p_1 \rho_0 \delta^2 k \frac{\omega_1^2}{\omega_1^2 + k^2}
\] (43)

These results are accurate provided that the spin relaxation decay is dominated by a single exponential damping constant, and are applicable to a wider range of conditions than existing theoretical descriptions. These \( R_{1\rho} \) expressions will be useful in analyzing experimental data when exchange is not fast and site populations are unequal. Since this expression involves a chemical-shift difference term that is field-dependent, there will be an increase in \( R_{1\rho} \) with field strength. Field-dependent \( T_{1\rho} \) relaxation studies can also be used to tease out any contribution from exchange between two chemically shifted nuclei. This exchange model has been used to analyze \( T_{1\rho} \) dispersion in cartilage (216).

Scalar relaxation (74). (a) Scalar relaxation of the first kind: this is described in the context of two spins, I and S, that are scalar coupled with a spin–spin coupling constant \( J_{IS} \). The exchange time constant between these two spins is denoted by \( \tau_e \). If the relaxation time, \( T_1 \), of each spin and \( \tau_e \) is much larger than \( 1/(2\pi J_{IS}) \) then the resonance line of each spin acquires a multiplet structure due to coupling with each other. If \( 1/T_1 \) of either spin or \( 1/\tau_e \) is much larger than \( 2\pi J \), then the multiplet structure disappears and each spin will exhibit a single line spectrum and the scalar coupling \( (J_{IS}) \) can become a mechanism of relaxation. When the coupling is modulated by (fast) exchange, this type of relaxation is referred to as ‘scalar relaxation of the first kind’.

(b) Scalar relaxation of the second kind: if the S spin’s \( T_1 \) is much shorter than \( 1/(2\pi J_{IS}) \) and \( \tau_e \) due to anything other than scalar coupling, then the splitting due to scalar coupling is absent and the I spin spectrum will have a single resonance line. This type of situation can occur if S is quadrupolar nuclei. Quadrupolar nuclei couple to the EFG within the molecule in which they are present. In solution state, the fast rotation of the molecules produces a rotating EFG, which leads to a fluctuation of the local magnetic field experienced by the nuclei and results in a short \( T_1 \). When the rate of these fluctuating local magnetic fields of quadrupolar nuclei is greater than the coupling constant, the splitting will average to zero. This type of relaxation mechanism is known as ‘scalar relaxation of the second kind’.

Contribution of both these types of scalar relaxation leads to broadening of the I spins’ spectral lines and an identical effect on the relaxation rate of the I nuclei. An example of scalar relaxation of the first kind is proton relaxation in H\(_2\)\(^{17}\)O. The \(^{17}\)O nucleus has spin 5/2 and its \( T_1 \) is dominated by quadrupolar relaxation shortening \( T_1 \) to ~4 ms. Protons on H\(_2\)\(^{17}\)O are in constant exchange with those on H\(_2\)\(^{16}\)O and at neutral pH \( \tau_e \approx 1.0 \) ms. Taking into consideration the concentration differences between H\(_2\)\(^{17}\)O and H\(_2\)\(^{16}\)O, Meiboom (201) derived equations for proton \( T_{1\rho} \) in \(^{17}\)O-enriched water under conditions of \(^{17}\)O–H] scalar coupling and chemical exchange. In the fast-proton-exchange regime (i.e. \( \delta t < 1 \), where \( 2\delta \) is the \( J \)-coupling constant and \( \tau \) is the proton exchange time), it can be shown that:

\[
\frac{1}{T_{1\rho}(\omega_1)} = \frac{1}{T_{01\rho}} + \frac{\tau}{(1 + \tau^2 \omega_1^2)} \sum_i \frac{p_i \delta_i^2}{(1 + \tau^2 \omega_1^2)}
\] (44)

where \( p_i \) is the relative intensity of the \( i \)th spectral line of an \(^{17}\)O multiplet, and \( 1/T_{01\rho} \) is the rate of all \(^{17}\)O-independent relaxation processes.

Substituting for \( \delta_1 \) and letting \( p_1 = \rho \delta(0) \), where \( f \) is the \(^{17}\)O atom fraction in H\(_2\)\(^{17}\)O, the above equation becomes:

\[
\frac{1}{T_{1\rho}(\omega_1)} = \frac{1}{T_{01\rho}} + \frac{(35/12)\pi J^2 f}{(1 + \tau^2 \omega_1^2)}
\] (45)

where \( J = 2\delta \).

An imaging method that uses \( T_{1\rho} \)-weighted images to quantify [H\(_3\)\(^{17}\)O] based on this theory has been developed (217). Scalar relaxation of the second kind has been demonstrated in the case of amide protons in farmamide (218). Since quadrupolar relaxation and scalar coupling are both independent of the static field, scalar relaxation of either kind is not field-dependent.

In biological tissue, the protons on amide and amine groups are expected to be influenced by the scalar relaxation (second kind) from the quadrupolar nucleus \(^{14}\)N \((I = 1)\) and their exchange with bulk water could influence water \( T_{1\rho} \) relaxation times.

Nonaveraged or residual dipolar coupling. In biological tissues, the presence of nonaveraged RDI can contribute to ‘\( T_{1\rho} \) dispersion’-variation of \( T_{1\rho} \) as a function of the spin-locking field. Since the RDI stems from the ordered structures in the tissues, computation of the residual dipolar coupling (RDC) provides information about the structural integrity of tissue. Previously, different types of multiple quantum (MQ) coherence transfer methods have been developed to characterize this interaction. However, due to an order of magnitude low SNR of MQ filtered methods, their applications have been largely limited to \textit{ex vivo} studies and some imaging experiments that were performed on highly ordered tissue such as tendons (86,219). Spin-locking methods to characterize this interaction offer high SNR (comparable to single quantum coherence) and can be implemented \textit{in vivo} with relative ease.

Recently it has been shown that, because of the changing ratio of the residual dipolar coupling to the B\(_I\) of the spin lock pulse, even in the absence of slow motional frequencies, RDI leads to \( T_{1\rho} \) dispersion (220).

Theoretical analysis of the \( T_{1\rho} \) relaxation, in the presence of RDI, in the simplest model of two chemically
equivalent dipolar coupled nuclear spins under the influence of spin-locking, has been described (221). During spin-locking, the presence of RDI creates anti-phase magnetization, zero and double quantum coherences and longitudinal two spin-order, some of which evolve into observable coherences with different relaxation rates. The signal expression for a dipolar-coupled spins following spin-locking can be written as (221):

\[
S(t) = X_1 \exp(-R_{1p}^\text{app} t) + X_2 \exp(-R_{1p}^\ast t) \cos(2\pi v_{\text{osc}} t + \phi)
\]  

where

\[
x_1 = \frac{(R_{1p}^2 - R_{1p}^\ast)^2 + 4\pi^2(v_{\text{osc}}^2 - v_d^2)}{(R_{1p}^2 - R_{1p}^\ast)^2 + 4\pi^2 v_{\text{osc}}^2}
\]

and

\[
x_1 + x_2 \cos \phi = 1
\]

and

\[
\tan \phi = \frac{(R_{1p} - R_{1p}^\ast) + (R_{1p}^\ast - R_{1p}^\text{app})X_1}{2\pi v_{\text{osc}}(1 - X_1)}
\]

where

\[
x_0 = (-R_{1p}^\text{app} - R_m)
\]

\[
+ \left\{-q/2 + \left[\frac{(q/2)^2 + (p/3)^2}{2}\right]^{1/2}\right\}^{1/3}
\]

\[
+ \left\{-q/2 - \left[\frac{(q/2)^2 + (p/3)^2}{2}\right]^{1/2}\right\}^{1/3}
\]

where

\[
p = -\frac{(R_{1p}^2 + R_{1p}^\ast + 2R_{1p}^\text{app} - R_{1p}R_{1p}^\ast - R_{1p}R_{1p}^\ast)}{3}
\]

\[
+ 4\pi^2 v_d^2 + 16\pi^2 v_{\text{osc}}^2
\]

and

\[
q = (R_{1p} + R_{1p}^\ast + 2R_{1p}^\text{app})
\]

\[
	imes \frac{(2R_{1p}^2 + 2R_{1p}^2 + 2R_{1p}^\ast + 5R_{1p}R_{1p}^\ast - 5R_{1p}R_{1p}^\ast - 5R_{1p}R_{1p}^\ast)}{27}
\]

\[
+ R_{1p}R_{1p}^\ast + R_{1p}^\text{app} - R_{1p}^\ast 4\pi^2 v_d^2
\]

\[
+ \left\{(R_{1p} - R_{1p}^\ast)^2 + 16\pi^2 v_{\text{osc}}^2\right\}^{1/2}
\]

\[
\times \frac{(2R_{1p} - R_{1p}^\ast - R_{1p}^\ast)16\pi^2 v_{\text{osc}}^2}{3}
\]

\[
x_{1,2} = \frac{-3R_m}{2} - \frac{x_0}{2} \pm 2i\pi v_{\text{osc}}
\]

where

\[
v_{\text{osc}} = \frac{1}{4\pi} \sqrt{4p + 3(x_0 + R_m)^2}
\]

\[R_{1p}^\ast\] is the relaxation rate of single quantum coherence \((I_s + S_s)\), \(R_{1p}^\text{app}\) is the relaxation rate of anti-phase two spin order \((2I_s - 2I_s)\) and \(R_{1p}^\ast\) is the relaxation rate of \((2I_s - 2I_s)\), which is a mixture of zero, double quantum coherences, \(2I_s = (I_sS_s + I_sS_s - I_sS_s - I_sS_s)/2\), and longitudinal two-spin order.

\[
R_m = \frac{(R_{1p} + R_{1p}^\ast + R_{1p}^\text{app})}{3}
\]

The following three parameters can be measured from the signal expression:

1. The damping factor of the slowly decaying part of the evolution curve that yields the apparent rotating frame relaxation rate, \(R_{1p}^\text{app}\).

2. The initial oscillation frequency, \(v_{\text{osc}}\), corresponding to nutation phenomena [due to the presence of two modes: \((2I_s - 2I_s)\) and \((2I_s + 2I_s)\)] with respect to the inhomogeneous spin-locking RF field and associated relaxation rate \(R_{1p}^\ast\), which is similar to the \(R_2^\ast\) when one is dealing with precession with respect to an inhomogeneous \(B_0\) field. When spin-locking field is zero, \(R_{1p}^\ast \approx R_2^\ast\).

3. The relative contributions of the fast and slow decaying components \(X_1\) and \(X_2\).

**Limiting cases.** Case 1: when a small molecule is dissolved into a liquid crystalline solvent, the relaxation rates are expected to be much smaller than \(v_1\) and \(v_d\). In this limiting situation:

\[R_{1p}^\text{app} = \frac{4v_1^2}{v_d^2 + v_d^2} R_{1p} + \frac{v_d^2}{4v_1^2 + v_d^2} R_{1p}^\ast\]

and

\[
v_{\text{osc}} = \sqrt{4v_1^2 + v_d^2}
\]

when \(v_1 \gg v_d\), \(R_{1p}^\text{app}\) is equal to the true rotating frame relaxation rate, \(R_{1p}^\ast\), whereas for \(v_1 \ll v_d\), \(R_{1p}^\text{app}\) tends toward the third-mode relaxation rate, \(R_{1p}^\ast\).

\[
x_1 = \frac{R_{1p}^2 + 16\pi^2 v_1^2}{R_{1p}^2 + 4\pi^2 v_d^2}
\]

and

\[
\tan \phi = -\frac{R_{1p}^\ast}{2\pi v_{\text{osc}}}
\]

Case 2: when \(v_1 \gg v_d\) and both are distributed over a range of values as in any polycrystalline, amorphous solid, or in ordered biological tissues. In this limiting situation:

\[R_{1p}^\text{app} = R_{1p} + \frac{v_d^2}{4v_1^2} (R_{1p}^\ast - R_{1p})\]

Several experiments have shown the feasibility of measuring the residual dipolar coupling using spin-
locking techniques in a liquid crystal (Figure 17 and Figure 18 spectra and early portion of oscillations). As shown in this example, in single and liquid crystals the dipolar coupling is manifested as resolved resonance splitting in a pulse-acquired spectrum. Hence, spin-locking or other experiments involving complex pulse sequences are not required to measure RDI. However, biological tissues exhibit a distribution of RDI and it may be difficult to resolve the resulting small splitting. Further, due to RF inhomogeneities during spin-locking, the dipolar oscillation frequency will be damped faster and thereby mask the intrinsic relaxation rates. Dominant bulk water, without any RDI, may mask these small and unresolved splittings. In these cases, the spin-locking experiment is well suited to computing the RDI and associated relaxation rates.

Depending upon the limiting case that is applicable for a given tissue, one has to include the appropriate $R_{1p}^{\text{app}}$ in the expression for observed $T_1$ in biological tissues. As long as $\omega_1$ is on the order of a few kHz, dipolar relaxation due to molecular rotational processes such as fluctuating dipolar fields and diffusion do not contribute to the $T_1$ dispersion but they add as constants to the overall value of $T_1$. The $T_1$ contribution can either be calculated as mentioned above or treated as a constant. The remaining terms are due to the exchange processes that are associated with water protons and other exchangeable protons on macromolecules (such as -OH and -NH), and residual static dipolar coupling. As discussed above, depending upon the tissue type and limiting cases, appropriate contributions due to dominant exchange interaction and static dipolar interaction terms should be included in the total expression for $T_1$. Therefore, the observed $T_1$ in biological tissues can expressed as:

$$\frac{1}{T_1/\text{tissue}} = \frac{1}{T_1/\text{D}} + \frac{1}{T_1/\text{Diff}} + \frac{1}{T_1/\text{exchange}} + \frac{1}{T_1/\text{RDI}} \quad (61)$$

**Dipolar interaction in cartilage**

Using the approach described above (221), the existence of dipolar oscillation frequency due to static dipolar interaction and its origin in the case of cartilage and model systems is demonstrated (222).

Figure 19 shows the $T_1$-weighted spectral intensity from a native bovine cartilage specimen as a function of spin-lock pulse length (for $\omega_1$ of 200 Hz). The inset shows

![Figure 18. Evolution of dipolar doublet as function of the spin-lock period duration; spin-lock amplitude: $\omega_1 = 4$ kHz. These experimental results were obtained by subtracting the anti-phase doublet signal. Note the initial rapidly damped oscillations and the mono-exponential decay for the major part of the evolution curve (221).](image18)

![Figure 17. Pulse-acquire proton NMR spectrum of CH$_2$Cl$_2$ in a nematic solvent (doublet of 4.2 kHz, arising from the dipolar interaction between the two protons). The distorted baseline is due to the broad NMR spectrum of the liquid crystal (221).](image17)

![Figure 19. $T_1$-weighted signal intensity from a bovine cartilage as a function SL length is shown. The $\omega_1$ used for this study is 250 Hz. The inset shows the expanded version of the initial portion of the signal. The oscillation pattern is characteristic of the presence of RDI in the tissue (240).](image19)
the expanded initial portion of the data that has persisting oscillation pattern for \( \sim 20 \text{ ms} \). This data, with a fast decaying component with dipolar oscillations and a slow decaying component, has a striking resemblance to the one obtained from liquid crystalline sample with resolved dipolar splitting. This clearly demonstrates the presence of RDI in cartilage. As shown in the inset, in the bovine cartilage the dipolar oscillations lasts \( \sim 20 \text{ ms} \) during the spin locking. However, since in human cartilage \( T_{1\rho} \) relaxation time is about 50% smaller than that in bovine tissue, the oscillation pattern may lasts for about \( \sim 10 \text{ ms} \). Since there is a distribution of RDI in cartilage, it should be modeled to fit the observed oscillation pattern to quantify RDI from the tissue similar to the analysis presented by Chaumette et al. (221).

Several studies have demonstrated that the static RDI between water molecules associated with fibrillar collagen is the primary source of laminar appearance (alternating bright-dark pattern of signal intensity variation across cartilage) in \( T_2 \)-weighted images (31,223–225). Given the \( \sim 20\% \) (of wet weight) collagen content in cartilage ECM, water proton \( T_2 \) relaxation is expected to be dominated by the RDI and therefore masks any small changes in the relaxation times due to alterations in other matrix properties. Further, there are several studies indicating that PG changes in cartilage either do not influence the \( T_2 \) or the changes in \( T_2 \) do not follow a particular trend (226–228). These observations have led to the argument that \( T_2 \) relaxation mapping is either more sensitive to collagen content or only sensitive to changes in collagen (11,228).

**\( T_{1\rho} \) Dispersion in cartilage**

There are only a few studies dealing with \( T_{1\rho} \) dispersion in cartilage. In one of the spectroscopic studies of \( T_{1\rho} \) dispersion in bovine cartilage, it was suggested that the exchange between protons on -OH and -NH of GAG with bulk water may be the dominant source for the low frequency (0–1.5 kHz) \( T_{1\rho} \) dispersion in cartilage (229). Low-frequency dispersion changes are correlated with loss of PG from the ECM of cartilage. Supplementing the experimental data with literature results, logical arguments were presented in support of the mechanism in model systems. However, the data in support of the exchange mechanism in cartilage was extrapolated from that of isolated protein model systems and therefore may not be translated to the case of cartilage with ordered structures. Another problem with this study is that the data was gathered using spectroscopy rather than imaging methods. This will provide an average \( T_{1\rho} \) value from the entire sample and could mask subtle regional changes in the tissue characteristics and is highly susceptible to tissue handling (e.g. un-blotted excess water). These experimental and data analysis aspects have to be taken into account when inferring anything from this study.

Recently, \( T_{1\rho} \) mapping at two different scanner field strengths was analyzed using a two-site exchange model as described above. It was demonstrated that the \( T_{1\rho} \) dispersion is primarily due to RDI and contribution due to chemical exchange between -OH and -NH protons on GAG and bulk water \( \leq 3 \text{ T} \) is only \( \sim 6\% \) (216). Their data also shows that the exchange contribution may increase to \( \sim 25\% \) at higher fields of \( \sim 7 \text{ T} \). The effect of refocusing the dipolar interaction on the measured \( T_2 \) of articular cartilage was also investigated using spectroscopic methods and demonstrated that dipolar interaction contributes significantly to \( T_{1\rho} \) dispersion in cartilage. \( T_{1\rho} \) dispersion in bovine cartilage and its effect on cartilage contrast, and the effect of spin-locking on the laminar appearance in MRI of cartilage, were also investigated at 4.7 T using orientation-dependent studies (222).

It was found that, when the normal to the surface of cartilage was parallel to \( B_0 \), a typical laminar appearance was present in \( T_2 \)-weighted images but was absent in \( T_{1\rho} \)-weighted images of the same specimen (Figure 20). At the ‘magic angle’ orientation (when the surface normal was 54.7° with respect to \( B_0 \)), neither \( T_2 \) nor \( T_{1\rho} \) images demonstrated laminae (Figure 21). However, \( T_{1\rho} \) values were greater than \( T_2 \) at both orientations throughout the cartilage layers. These studies imply that, while RDI contributes to a laminar appearance in \( T_2 \) images, its effect is attenuated by spin-locking. Further, if the dipolar interaction is the only contributor to the observed dispersion, then at the magic angle all \( T_2 \) values should be equivalent to \( T_{1\rho} \) obtained at a spin-lock field at \( \sim 2 \text{ kHz} \). They observed significantly higher \( T_{1\rho} \) values compared with \( T_2 \) even at the magic angle. This may indicate that there are other relaxation mechanisms that contribute to \( T_{1\rho} \) in addition to RDI.

Although there are conflicting reports in the literature on the contribution to \( T_{1\rho} \) dispersion on cartilage, the bulk of the existing data in the literature and data presented here support the nonaveraged dipolar interaction between water protons associated with collagen being the predominant contributor. However, the dipolar interaction alone cannot account for the complete dispersion observed. Contributions from other sources including exchange cannot be ruled out completely, especially at higher static fields. Further work in this area is required to address this issue.

Small variations in \( T_{1\rho} \) dispersion profiles were observed in spectroscopic studies of cartilage specimens (229). However, except for a constant shift, there were no significant changes observed in the \( T_{1\rho} \) dispersion curve with varying levels of GAG depletion by trypsin (25). Discrepancy between these results may be due to the measurement of global (spectroscopic) vs (imaging) regional \( T_{1\rho} \) and difference in static field strength. In imaging experiments, \( T_{1\rho} \) mapping is performed at a particular spin-lock frequency (e.g. 500 Hz) and \( T_{1\rho} \) relaxation time, not dispersion, is measured. In spin-locking experiments, as long as the \( B_1 > \Delta B_0 \), field
inhomogeneities are refocused. In most of the in vivo experiments, this condition is often fulfilled because typical $\Delta B_0$ values are between 100 and 200 Hz and $B_1$ values used are $>250$ Hz. At about 500 Hz, (i) reduced dipolar interaction by spin-locking minimizes or eliminates laminae (or relaxation effects due to oriented collagen). At this field, it is likely that scalar-relaxation due to proton exchange between $H_2^{17}O$ and $H_2^{16}O$ and or between $-NH$ and $H_2O$ may also be attenuated substantially (217). As a consequence of the attenuation of the $\Delta B_0$, RDI and exchange broadening increases the dynamic range of the $T_{1p}$ relaxation time scale by $\sim$200% in bovine cartilage and by 60% in human cartilage compared with $T_2$. This improved dynamic range (due to the attenuation of background relaxation of oriented collagen matrix) of $T_{1p}$ may facilitate the detection of small changes in relaxation times due to the loss of PG or other matrix component from the tissue. At this point, available data on measured $T_{1p}$ in CS phantoms and trypsin-treated cartilage (226) suggest that it may be predominantly due to a combination of changes in [GAG] (221) and, to a lesser extent, from exchange between protons on GAG and bulk water. Further work is required to tease out these contributions. Since the relaxation rate is measured at a fixed $B_1$ amplitude of the spin-lock pulse, the static dipolar effect adds as a constant. However, $T_{1p}$ dispersion maps measured from data obtained as a function of varying $B_1$ is expected to demonstrate the otherwise dominant dipolar effects and may provide information about the ordered collagen.

**Experimental $T_{1p}$ studies**

**Ex vivo animal specimens.** Potential of proton $T_{1p}$ imaging of cartilage and its applications to study of cartilage degeneration was first proposed by Reddy et al. (230). In this study at 2 T, $T_{1p}$ mapping and dispersion was

![Figure 20. $T_2$- and $T_{1p}$-weighted images of bovine articular cartilage. In $T_2$-weighted images, laminae of alternating high and low signal intensity pattern are evident. In $T_{1p}$-weighted images, the laminar appearance is gradually reduced with increasing $\omega_1$, making the signal more homogeneous across the cartilage (222).](image)

![Figure 21. $T_{1p}$ dispersion in bovine cartilage; the cartilage surface was oriented parallel to $B_0$ and at the magic angle $\sim$55°. The data at spin-lock frequency of zero correspond to the respective $T_2$ values (222).](image)
performed on bovine cartilage specimens and the observed dispersion was interpreted by invoking the presence of exchange-modulated dipolar coupling of water protons associated with macromolecules in the tissue (Figure 22). Later studies were performed by subjecting bovine articular cartilage specimens to trypsin- and collagenase-induced PG and collagen degradation, respectively (24). Results demonstrated that the $T_1\rho$-weighted images had improved contrast and SNR compared with $T_1$- and $T_2$-weighted images and $T_1\rho$-weighted images are sensitive to PG change in the tissue. In the same study, it was suggested that collagenase-induced collagen degradation did not exhibit changes in $T_1\rho$. However, later it was found that there were errors in the protocol of the collagenase degradation and the conclusions about effect of collagenase on $T_1\rho$ were not reliable.

In an elegant study on bovine articular cartilage patellae, sequential trypsin-induced PG depletion was performed (25). Proton $T_1\rho$ maps were generated at varying $B_1$ fields at 4 T using a home-built RF coil. Depleted PG in the medium was quantified using standard dimethyl methylene blue (DMMB) assay and histology. $T_1\rho$ maps clearly demonstrated the PG depleted regions and corresponded well with the histological analysis performed with Safranin-O staining. In another study at 4 T, $T_1\rho$ and $T_2$ relaxation rates were computed from bovine cartilage subjected to sequential PG depletion (226). $T_1\rho$ and $T_2$ maps were correlated with PG in the tissue (Figure 23) and it was found that there was an excellent correlation ($r^2 = 0.89$) between $1/T_1\rho$ and [GAG] while the correlation between $1/T_2$ and [GAG] was rather poor ($r^2 = 0.01$). The $T_1\rho$ maps reflected clear changes in PG depleted regions of the tissue (Figure 24). Later on, spectroscopic studies were also demonstrated to have the same trend, although as they are based on global relaxation numbers they underestimate the magnitude of the change in the relaxation rates (231). These studies form the basis for using the $T_1\rho$ mapping studies in studying cartilage pathology.

$T_1\rho$, $T_2$, and $T_1$ studies were also performed on osteoarthritic human specimens obtained following the knee replacement surgery (173). In these studies spin-locking frequency used was ~2.5 kHz. It was found that neither $T_2$ nor $T_1\rho$ were significantly changed from studies performed on ex vivo human cartilage. These results were in contradiction to the results from several studies described above. The primary reason for this discrepancy appeared to be the status of the osteoarthritic tissue. Prior to the surgery, the patients were intravenously administered GdDTPA$^{2-}$, a contrast agent known to dominate the relaxation mechanism of water protons. Since the tissues were imaged in the presence of the contrast agent it dominates the relaxation contribution to both $T_1\rho$ and $T_2$.
as well and masks any changes due to the loss of PG/macromolecules from the tissue. These studies clearly demand a thorough investigation of the influence of contrast agents like GdDTPA$^2$ on the magnitude of relaxation contribution to $T_1$ and $T_2$.

$T_1$ and $T_2$ measurements were made in collagen and GAG suspensions, and an exponential decrease in $T_1$ and $T_2$ was found with the increasing [collagen] and/or [GAG]. Relaxation time measurements were made in native and trypsin- and interleukin-1$\beta$-induced bovine cartilage and some human OA specimens as well (227). In these studies, it was found that $T_1$ and $T_2$ are sensitive to biologically meaningful changes in cartilage. However, they are not specific to any one inherent tissue parameter.

In this study, data from phantoms show that both PG and collagen contribute to $T_1$ rate ($R_1$) in a concentration-dependent manner (at low concentrations, <20%, this is almost linear). However, there are several differences between this study and other $T_1$ mapping studies in the literature and the following factors should be taken into account when drawing conclusions from this study:

1. The experiments were performed at a very high (8.5 T) field strength where other mechanisms, e.g. exchange from -OH and -NH to H$_2$O, will dominate and could contribute to $T_1$ [see above and ref. (173)]. Most other $T_1$ studies in the literature were performed between 1.5 and 4 T.

2. Cartilage samples subjected to IL-1$\beta$ exhibited decreased $T_2$ but unchanged $T_1$ when compared with healthy and trypsin-treated cartilage. This is completely in conflict with several results from IL-1$\beta$ studies on a larger group of samples and on an in vivo animal model where a significant increases in $T_1$ (Figure 25) were observed (54,177).

3. At the higher field (8.5 T), $T_1$ should be lower than that at 1.5 T, but the $T_1$ numbers measured in human cartilage in this study are almost 50% higher than those observed at 1.5 T. This result was not explained.

4. In one of the human specimens, the GAG-depleted region shows 19% elevated $T_1$ compared with the healthy-appearing region in the same tissue. In fact, even though a suboptimal color scaling was used and there was a possibility of slice mismatch and associated partial volume effects, the $T_1$ map almost mirrors the dGEMRIC map, indicating that there is indeed an elevated $T_1$ in regions with lower GAG. At the same time, the observed $T_1$ values in two other samples [Fig. 2(C) and (D) with more advanced OA] are not consistent with those observed in the GAG-depleted regions. Given the thin sections of the images (2 mm thick) gathered in two different imaging sessions there is a possibility that the slice mismatching and partial voluming effects may be present in the data.

5. $T_1$ imaging on human specimens was performed in multi-slice mode but there are no details of the

Figure 24. Comparison of $T_2$, and $T_1$ maps of control and 40% of PG depleted bovine patellae. (A) Control $T_2$ map; (B) control $T_1$ map; (C) 40% PG depleted $T_2$ map; (D) 40% PG depleted $T_1$ map. The color scale bar shows the relaxation numbers from 0 to 256 ms (25).
sequence. In fact, in multi-slice mode, there is a substantial $T_2$ saturation that, uncorrected, will lead to erroneous results (237).

In direct contradiction to the above study, in human OA specimens ($n = 7$) with different Mankin grades (0–3), a strong correlation between $T_1$ rate and FCD measured from sodium MRI was observed (Figure 28), indicating that $T_1$ relaxation rate may be useful in studying the OA process in humans (177). Nevertheless, a study on a large group of human osteoarthritic specimens is needed to obtain statistically significant results.

Correlation of changes in cartilage biomechanical and biochemical properties with $T_1$ relaxation rate was made in a cytokine-induced model of osteoarthritic condition (54). $T_1$ mapping was performed at 4.7 T, and PG, collagen and water content were measured via biochemical assays. Stress-relaxation biomechanical tests were conducted with confined compression apparatus to measure uni-axial aggregate modulus ($H_A$) and hydraulic permeability ($k_0$) using linear bi-phasic theory. Results from this study demonstrated that $T_1$ relaxation rate was strongly correlated with PG content ($r^2 = 0.93$), $H_A$ ($r^2 = 0.83$) and $\log_{10} k_0$ ($r^2 = 0.86$), demonstrating the potential of $T_1$ relaxation as a surrogate index for changes in cartilage biochemical and biomechanical properties as well (Figure 26). These studies clearly provide a link between $T_1$ relaxation rate and [GAG] content of the tissue and biomechanical properties. These results are significant because they suggest that non-invasive $T_1$ mapping may be used to obtain biomechanical properties of the tissue without subjecting the tissue to compression in magnet, a difficult task to begin with.

Results from this work clearly show the potential of $T_1$ relaxation mapping in providing information about biomechanical properties

**Ex vivo human specimens.** Studies on isolated chondroitin sulfate and collagen suspensions show that $T_1$ is sensitive to both the components of cartilage matrix (227). Owing to a well-ordered structure of collagen and the swelling pressure of PG in intact cartilage, the observed proton $T_1$ relaxation rates in the intact cartilage cannot be extrapolated to those obtained from isolated PG and collagen suspensions. Therefore, in order to tease out
the individual contribution of PG and collagen to the observed $T_{1\rho}$, selective depletion studies are needed. It is relatively easy to quantify PG’s exclusive contribution to $T_{1\rho}$; however, it is difficult to discern the collagen contribution alone to the $T_{1\rho}$ relaxation. This is because depletion of collagen from cartilage is not feasible without affecting PG content. To overcome this limitation, an indirect approach was used (177) in which spatial maps of proton $T_{1\rho}$ and sodium maps of FCD were computed on bovine specimens subjected to trypsin-induced PG degradation and human OA specimens with different Mankin grade (232). The loss of PG was confirmed by histology in these specimens. Since trypsin predominantly degrades cartilage PG and leaves collagen almost intact, changes induced in $T_{1\rho}$ in this model are primarily from PG loss. In human OA specimens; however, while there may be a predominant contribution from loss of PG, there may also be changes in collagen content and structure depending on the Mankin grade of the tissue. Plots of normalized $1/T_{1\rho}$ rate vs FCD (Figure 27) were found to be strongly correlated with $r^2 > 0.75$ and 0.85, respectively, in both tissue types with nearly identical slope of $\sim 0.5$ ($p > 0.51$). The strong correlation of the FCD and $T_{1\rho}$ data, with identical slopes, in both the models has been interpreted as that the contributions to $T_{1\rho}$ changes are predominantly from PG and any contribution from collagen may be rather small even in natural OA cartilage. A similar trend is shown in the sodium image and $T_{1\rho}$ map of OA specimen (Figure 28).

![Figure 27](image1)

**Figure 27.** Correlation of normalized FCD and $R_{1\rho}$ data obtained from the trypsin treated model (A). The slope of the linear regression was $0.48 \pm 0.004$ with $r^2 > 0.75$ ($p < 0.001$). The data are arranged in four clusters representing the four groups of trypsin concentration. The cluster about zero, including negative values, represents the distribution of the mean of the control data. Correlation of normalized FCD and $R_{1\rho}$ data obtained from the natural OA model (B). The slope of the linear regression was $0.51 \pm 0.06$ with $r^2 > 0.85$ ($p < 0.001$) (177).

![Figure 28](image2)

**Figure 28.** Comparison of sodium concentration and $T_{1\rho}$ maps of a representative ex vivo patella from an osteoarthritic joint. The sodium map reveals a distinct reduction in FCD on the lateral side of the patella (ROI A) indicating a reduction of PG content. Elevation of $T_{1\rho}$ in ROI A is observed as an analogous measure of PG loss (177).
Studies on animal models in vivo. Animal models serve as a bridge between tissue culture and human disease. The advantage of the animal models is that the disease can be controlled. Therefore, animal models are ideal for research in developing, optimizing and validating therapies, stimulating repair of damaged tissues and understanding the disease process, as well as developing noninvasive diagnostic tools to detect early OA.

The potential of $T_{1p}$ mapping in detecting rapidly induced cytokine-mediated cartilage degeneration in porcine animal model in vivo was investigated (54). Six Yorkshire pigs were given an intra-articular injection of recombinant porcine interleukin-1$\beta$ in the knee joints before imaging. The contra-lateral knee joints were given a saline injection to serve as an internal control. Six hours following the injection, $T_{1p}$ mapping was performed using $T_{1p}$-prepared FSE sequence and sodium imaging was performed using a previously validated method at 4 T. The effect of IL-1$\beta$, primarily loss of PG, was confirmed by histological and immunohistochemical analysis (Figure 29). It was found that the average $T_{1p}$ rate ($1/T_{1p}$) of the IL-1$\beta$-treated patella was about 25% lower than that of saline-injected patella. There was an average reduction of 49% in FCD, measured via sodium MRI. These results demonstrate that changes seen in both $T_{1p}$ and FCD indicate the loss of PG.

Studies on guinea pig model of spontaneous OA. The advantages of the guinea pig model are that OA occurs spontaneously, it is an inexpensive model compared to anterior cruciate ligament (ACL) transection model, and it has been well studied (233). Initial studies demonstrating the feasibility of performing $T_{1p}$ mapping on guinea pig cartilage (Figure 30) exhibited excellent contrast between the cartilage and the surrounding tissues. The in-plane resolution in axial image is 60 $\mu$m and that in sagittal images is 117 $\mu$m. The signal-to-noise ratio in these images is ~25:1. The color scheme and $T_{1p}$ values are shown in the bar-scale. A clear region with elevated $T_{1p}$ number (possibly a lesion) can be seen on the femoral cartilage (indicated by an oval shape). In this map, $T_{1p}$ numbers of the load-bearing (LB), nonload-bearing (NLB) and lesion regions are: 44 $\pm$ 5, 58 $\pm$ 5 and 97 $\pm$ 14 ms, respectively. The imaging parameters are: spin-lock pulse power $= 500$ Hz, $TR = 200$ ms, slice thickness $= 1$ mm. The spin-lock duration varied from 1 to 40 ms. Average $T_{1p}$ relaxation

Figure 29. $T_{1p}$ maps of saline- and IL-1$\beta$-injected patellae of a Yorkshire pig (A) in vivo (top row) and the same ex vivo (bottom row). Color-coded $T_{1p}$ maps corresponding to cartilage are overlaid on the original $T_{1p}$-weighted MR image, indicating elevated $T_{1p}$ in each IL-1 treated case (54). Also shown is a plot of $1/T_{1p}$ measured in vivo and FCD measured by post-mortem PG assay from animals (B).

Figure 30. Representative $T_{1p}$-weighted images of a guinea pig (age ~4 months) and corresponding color-coded $T_{1p}$ maps (overlaid on the $T_{1p}$-weighted images). (A) An image of the knee joint showing the $T_{1p}$ map of patellar cartilage in color. The average $T_{1p}$ of patellar cartilage was measured at $57 \pm 5$ ms. (B) A sagittal slice of the same animal’s knee joint showing both femoral and tibial cartilage.
numbers from different regions of guinea pig \( n = 4 \) cartilage are as follows: patellar cartilage \( (59.6 \pm 4.4 \text{ ms}) \), femoral cartilage load-bearing \( (43.8 \pm 4.6 \text{ ms}) \) and femoral cartilage non-load-bearing \( (65.2 \pm 5.4 \text{ ms}) \). These results, although preliminary, demonstrate the feasibility of obtaining \( T_{1p} \) relaxation maps in guinea pig cartilage.

**Clinical \( T_{1p} \) MRI.** Proton \( T_{2} \) and \( T_{1p} \)-weighted images of healthy human knee joint acquired on a 1.5 T clinical scanner with a spin-locking amplitude of 500 Hz are shown in Figure 31. In the \( T_{2} \)-weighted image, the laminar appearance due to different signal intensity patterns corresponding to collagen fiber orientation, although not clearly observed owing to the lack of adequate resolution, is manifested as low signal intensity across the cartilage, whereas in the \( T_{1p} \)-weighted image the laminar appearance is substantially attenuated and results in a more homogeneous appearing and elevated signal intensity across cartilage (222).

In vivo \( T_{1p} \) relaxation studies on healthy human knee joints \( n = 6 \) were performed with a \( T_{1p} \)-prepared FSE sequence at 1.5 T (26). Significant \( T_{1p} \) dispersion \( (34–50 \text{ ms}; T_{1p} \) changes with spin-lock frequency) in the frequency range 0–375 Hz was observed. Quantitative comparison of \( T_{2} \) and \( T_{1p} \)-weighted images has demonstrated a 25% higher signal-difference-to-noise ratio in a chondral lesion, in \( T_{1p} \)-weighted image than in comparable \( T_{2} \)-weighted images in vivo.

Clinical \( T_{1p} \) relaxation mapping studies were also performed in femoral–tibial cartilage and wrist joint \( \textit{in vivo} \) at 1.5 T, using home-built bird-cage coils (234,235). In both these studies a single-slice \( T_{1p} \) prepared FSE sequence was used with a spin locking frequency ranging from 0 to 500 Hz. It was found that the average \( T_{1p} \) \( (\text{at } B_1 = 500 \text{ Hz}) \) in the weight-bearing and non-weight-bearing regions of the femoral condyle were 42.2 ± 3.6 and 55.7 ± 2.3 ms, respectively. In wrist joint cartilage, at 500 Hz spin-locking amplitude, \( T_{1p} \) ranged from 40.5 ± 0.9 to 56.6 ± 4.8 ms. Both studies demonstrated 20–30% higher SNR of cartilage in \( T_{1p} \)-weighted images compared with \( T_{2} \)-weighted images. It was suggested that the primary purpose of these studies was to demonstrate the feasibility of quantifying relaxation maps in these complicated structures.

Three-dimensional \( T_{1p} \) mapping of human knee cartilage was performed at 1.5 T on both healthy and symptomatic osteoarthritic subjects (180,236). Average \( T_{1p} \) values \( (\text{at spin locking amplitude of } 440 \text{ Hz}) \) obtained from eight healthy volunteers \( (\text{all male with age range } 22–45 \text{ years}) \) with three-dimensional \( T_{1p} \) mapping was 49.7 ± 3.2 ms \( (\text{mean ± SD}) \) and that obtained from two-dimensional \( T_{1p} \) mapping was 48.5 ± 2.6 ms \( (\text{mean ± SD}) \). In all the symptomatic subjects \( (n = 6, \text{ four men and two women, age range } 28–63 \text{ years}) \) none had radiographic OA except one. \( T_{1p} \) relaxation times were elevated compared with healthy subjects. In symptomatic subjects, \( T_{1p} \) values \( (\text{at } 440 \text{ Hz spin lock amplitude}) \) varied from 63 ± 4 to 95 ± 12 ms \( (\text{mean ± SD}) \) depending on the degree of cartilage degeneration (Table 4). Although this study was performed on a limited number of subjects, it still demonstrates the potential of \( T_{1p} \) as a surrogate marker for pre-radiographic OA. Study of a larger group of subjects is required to further evaluate the sensitivity and specificity issue of the method.

**\( T_{1p} \) MRI of osteoarthritic subjects.** Figure 32 shows a \( T_{1p} \)-weighted image \( (\text{in gray-scale}) \) of the knee from a clinically-diagnosed OA subject \( (38\text{-year-old female}) \). These images were read by a radiologist and confirmed the presence of minor irregularities on cartilage surface. In this \( T_{1p} \)-weighted image, \( TSL + TE = 60 \text{ ms} \) and the spin-lock power employed is 500 Hz, with an in-plane resolution of \( \sim 390 \mu \text{m} \) and a 3 mm slice. The lateral side of the \( T_{1p} \) map \( (\text{overlaid in color}) \) shows larger region of elevated \( T_{1p} \) values \( (\text{indicated with an oval}) \). Although, there is an increase in \( T_{1p} \) across the cartilage compared with the values from healthy controls, in the elliptical ROI, \( T_{1p} \) is elevated by \( \sim 44\% \) compared with the healthy-appearing region.

Figure 33 shows \( T_{1p} \)-weighted images from a clinically diagnosed OA patient \( (40\text{-year-old female}) \) with an absence of cartilage abnormalities on radiographs. A rheumatologist and an orthopedist diagnosed the subject with osteoarthritis and the subject has been on medication.
for about one year. A practicing radiologist read these images and diagnosed that the patient has cartilage softening on the lateral side without any surface irregularities. Without the knowledge of imaging sequence used, he found that the $T_{1\rho}$ image had superior contrast and excellent delineation of the lesion. Figure 32 (A) and (B) shows $T_{1\rho}$ images acquired with an effective echo time ($TSL + TE$) of 45 and 60 ms, respectively. In both $T_{1\rho}$ images, the lateral side of cartilage (indicated by thin arrow) shows elevated signal (in these images, white, yellow and red represent high, medium and low signal intensities, respectively).

$T_{1\rho}$ MRI of a chondromalacia subject. In Figure 34, a three-dimensional surface-rendered $T_{1\rho}$ map and a cross-sectional image of patella of an osteoarthritic human subject who has undergone arthroscopy is presented. At arthroscopy, grade I chondromalacia was identified on the lateral facet of the patellar cartilage while the medial facet was identified with grade 0. The lateral facet of the $T_{1\rho}$ relaxation map (dashed oval indicates the arthroscopically identified chondromalacia region) clearly shows the elevated $T_{1\rho}$ numbers compared with that of medial facet. Red and yellow voxels in this region indicate substantial increase in the $T_{1\rho}$. These data clearly demonstrate that, in cartilage regions of chondromalacia, $T_{1\rho}$ is increased by $\sim 46\%$ compared with healthy cartilage. The cross sectional image shows the $T_{1\rho}$ map of one of the slices overlaid on the $T_{1\rho}$-weighted image.
image. In this map, elevated $T_{1,ρ}$ across the cartilage depth can be visualized. These results show the feasibility of computing $T_{1,ρ}$ maps of grade I chondromalacia of cartilage in vivo. Although arthroscopy is a semi-quantitative technique which provides information about cartilage softening, it is a subjective and invasive procedure and not suitable for longitudinal follow-up studies. However, the relaxation maps presented here are quantitative and noninvasive.

A multi-slice spin-lock (MS-SL) pulse sequence was developed and implemented on clinical scanner. Its utility in a clinical setting was demonstrated by measuring $T_{1,ρ}$ maps of the knee joints of six healthy human subjects (237). Since MS-SL $T_{1,ρ}$ measurements are confounded by the saturation effects due to the $T_{2,ρ}$ process, strategies to compensate for these saturation effects are presented. $T_{2,ρ}$-compensated $T_{1,ρ}$ maps were computed from multi-slice data from human cartilage tissue. These values differed by 5% when compared with the ‘gold-standard’ single-slice $T_{1,ρ}$ values. Although this sequence is superior in SNR and contrast, when compared with three-dimensional FGRE based sequence, it needs a separate saturation and compute the actual $T_{1,ρ}$ values. Although early $T_{1,ρ}$ experiments on cartilage were spectroscopic, later on these were extended to single-slice imaging and this was followed by the development of three-dimensional and multi-slice imaging methods. Studies on phantoms and isolated matrix components showed that both PG and collagen contribute to $T_{1,ρ}$. Ex vivo studies on bovine specimens subjected to sequential PG depletion have shown a strong correlation between $1/T_{1,ρ}$ and $[\text{GAG}]$, although correlation between $1/T_{2}$ and $[\text{GAG}]$ was rather poor. Ex vivo bovine studies and in vivo studies on animal models have shown that treatment with IL-1β elevated the $T_{1,ρ}$. In osteoarthritic human specimens, there was a strong correlation between percentage change in FCD measured from sodium MR imaging was unchanged. In some specimens elevated $T_{1,ρ}$ was observed in GAG-depleted region and in some other specimens it remained almost unchanged. Similarly, in studies (at 8.5 T) on a small group of bovine specimens subjected to IL-1β, $T_{1,ρ}$ was found to be unchanged.

![Figure 35. Colour-coded $T_{1,ρ}$ maps overlaid on SPGR images. Left: patellar cartilage; middle: anterior femoral (trochlea) cartilage; right: posterior femoral cartilage. (a) a healthy volunteer, male, 30; (b) a patient with early OA, female, 27. The $T_{1,ρ}$ values were 40.05 ± 11.43 ms in the volunteer and 50.56 ms ± 19.26 ms in the patient, respectively.](image-url)
Existing results to date suggest that contributions to $T_1^*$ dispersion are predominantly due to dipolar interaction of water molecules associated with collagen and they are influenced by the orientation of cartilage. Although the proton exchange between amide and hydroxyl protons on GAG molecules and bulk water may also contribute to the dispersion, its contribution is relatively small at lower fields.

Based on the current data, it appears that the strong correlation between [GAG] and $1/T_1^*$ may be predominantly due to concentration effect in bovine cartilage. The fact that IL-1b-treated specimens, and in human OA specimens, in which some collagen changes are expected, suggests that, in addition to PG contribution, there may be a (small) contribution from collagen as well to the observed $T_1^*$.

When compared with healthy subjects, OA patients presented elevated $T_1^*$ even when their radiographic images did not indicate OA. Measured in vivo, $T_1^*$ was found to have a stronger correlation with severity of OA, while differences in $T_2$ were not statistically significant. $T_1^*$ was also elevated in a patient with arthroscopically confirmed grade I chondromalacia but without radiographic OA. These results clearly indicate that $T_1^*$ may serve as a surrogate marker for molecular changes (pre-radiographic) in cartilage and may be useful in longitudinal studies. So far the $T_1^*$ studies have been mostly ‘demonstration of feasibility’ in nature, involving a very limited number of healthy and OA subjects. $T_1^*$ studies on a large group of OA specimens with varying degree of degeneration are needed to determine the correlation between $T_1^*$ rate and OA disease severity as determined by histopathology. Similarly, studies on a large group of OA subjects with mild to moderate OA (with a different degree of radiographic OA) and age-matched healthy subjects are needed to address the issues regarding the precision of $T_1^*$ measurement, the effect of age and disease severity on these measurements and the capability to longitudinally measure OA disease progression.

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