**Thickness of the Meniscal Lamellar Layer:** Correlation with Indentation Stiffness and Comparison of Normal and Abnormally Thick Layers by Using Multiparametric Ultrashort Echo Time MR Imaging

**Purpose:**
To determine the relationship between lamellar layer thickness on ultrashort echo time (UTE) magnetic resonance (MR) images and indentation stiffness of human menisci and to compare quantitative MR imaging values between two groups with normal and abnormally thick lamellar layers.

**Materials and Methods:**
This was a HIPAA-compliant, institutional review board-approved study. Nine meniscal pieces were obtained from seven donors without gross meniscal pathologic results (mean age, 57.4 years ± 14.5 [standard deviation]). UTE MR imaging and T2, UTE T2*, and UTE T1r mapping were performed. The presence of abnormal lamellar layer thickening was determined and thicknesses were measured. Indentation testing was performed. Correlation between the thickness and indentation stiffness was assessed, and mean quantitative MR imaging values were compared between the groups.

**Results:**
Thirteen normal lamellar layers had mean thickness of 232 μm ± 85 and indentation peak force of 1.37 g ± 0.87. Four abnormally thick lamellar layers showed mean thickness of 353.14 μm ± 98.36 and peak force 0.72 g ± 0.31. In most cases, normal thicknesses showed highly positive correlation with the indentation peak force ($r = 0.493–0.912; P < .001$ to $.05$). However, the thickness in two abnormal lamellar layers showed highly negative correlation ($r = -0.90, P < .001$; and $r = -0.23, P = .042$) and no significant correlation in the others. T2, UTE T2*, and UTE T1p values in abnormally thick lamellar layers were increased compared with values in normal lamellar layers, although only the UTE T2* value showed significant difference ($P = .010$).

**Conclusion:**
Variation of lamellar layer thickness in normal human menisci was evident on two-dimensional UTE images. In normal lamellar layers, thickness is highly and positively correlated with surface indentation stiffness. UTE T2* values may be used to differentiate between normal and abnormally thickened lamellar layers.

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The meniscus is a crucial structure that preserves articular cartilage integrity in the knee and contributes to healthy knee function (1–3). Given the important roles of the menisci, it is not surprising that torn menisci and/or surgical removal of the menisci are well known to result in early articular cartilage damage, and, eventually, early osteoarthritis. Early detection of subclinical meniscal abnormalities with the development and implementation of strategies to protect against progression to meniscal tear would be a valuable initial step for prevention of knee osteoarthritis, one of the most important burdens in the current health care system.

The healthy menisci are considered to be tissues that have short T2 (T2 values around 4 msec), which makes them “invisible” (ie, they have low signal intensity on conventional magnetic resonance [MR] images). However, the advent of ultrashort echo time (UTE) MR imaging enabled the selective imaging of tissues with short T2, which allows for acquisition of signal from these tissues and further provides high spatial and contrast resolution. This combination of imaging features allowed the anatomic structure of the menisci to be viewed in a noninvasive fashion (4–8). Additionally, quantitative MR imaging relaxation measurements have been used to interrogate changes in collagen matrix and water content and early proteoglycan depletion, including T2 and T2*, and T1p (6,9). When acquired with a UTE sequence, these techniques can be applied to tissues with short T2, such as the menisci (9–13).

Electron microscopy imaging studies (14) revealed three distinct layers in meniscal cross section (Fig 1): a superficial network that covers the femoral and tibial surfaces by a meshwork of very thin fibrils (30 nm); a lamellar layer beneath the superficial network, represented by a layer of lamellae of collagen fibrils (150–200 μm); and a central main portion, composed of predominantly circular-oriented bundles of collagen fibrils with occasional radial-tie fibers (15). By using UTE MR images, the anatomic structure of the menisci, including the lamellar layers and radial-tie fibers, were made visible (4). Because, to our knowledge, this degree of structural characterization is not possible by using conventional MR imaging, variations in structure can be explored by emphasizing potential clinical importance. Furthermore, little information is currently available regarding the relationship between lamellar layer thickness and local biomechanical properties in the human meniscus. Clearly, the lamellar layer becomes of paramount importance in the MR and arthroscopic diagnostic criteria of meniscal tearing because this layer must be violated in the setting of a tear.

We hypothesize that (a) the thickness of normal lamellar layer positively correlates with the biomechanical stiffness or resistance to the compressive load and (b) an abnormally thickened lamellar layer will show changes in previously established multiparametric quantitative MR imaging relaxation measurements (T2, UTE T2*, and UTE T1p) compared with the values in normal lamellar layers. The purpose of our study was to determine the relationship between lamellar layer thickness on UTE MR images and indentation stiffness of human menisci and to compare T2, UTE T2*, and UTE T1p values between two groups with normal and abnormally thick lamellar layers.

Materials and Methods

Our Health Insurance Portability and Accountability Act–compliant cadaveric study was exempted by the institutional review board, and informed consent was not required. Our study was performed between July 2012 and June 2014.

Specimen Preparation

Nine menisci were removed from eight fresh-frozen cadaveric human knees (five men, two women; mean age, 57.4 years ± 14.5 [standard deviation], and two samples with unknown age and sex) without gross meniscal tear and were included. Seven menisci were not included because of grossly visible meniscal tears. They were cut in a sagittal manner into 5-mm-thick triangular pieces.

MR Imaging

MR hardware.—A 3-T MR imager (Signa HDx; GE Healthcare, Milwaukee, Wisconsin) was used. A 16-channel UTE coil was used for imaging. A 2D UTE sequence was used with repetition time 4.5 msec, echo time 0.06–2.00 msec, bandwidth 14 kHz, and a field of view 14 x 14 cm². A UTE T1p sequence was also used with repetition time 1000 msec, echo time 2.52 msec, bandwidth 14 kHz, and a field of view 14 x 14 cm². The UTE T2* and T1p images were postprocessed on a clinical workstation (Leonardo; GE Healthcare, Milwaukee, Wisconsin) using a dedicated software application (UltraLow Echo Time; GE Healthcare, Milwaukee, Wisconsin). T2, UTE T2*, and UTE T1p images were compared with T2, T1, T1p, T2*, T1*, and T2 images that were obtained from a separate MR session with a 3-T MR imager (Signa HDx; GE Healthcare, Milwaukee, Wisconsin).

Specimen Preparation

Five menisci were removed from each of eight fresh-frozen cadaveric human knees (women, four; men, four; mean age, 57.4 years ± 14.5 [standard deviation], and two samples with unknown age and sex) without gross meniscal tear and were included. Seven menisci were not included because of grossly visible meniscal tears. They were cut in a sagittal manner into 5-mm-thick triangular pieces.

MR Imaging

MR hardware.—A 3-T MR imager (Signa HDx; GE Healthcare, Milwaukee, Wisconsin) was used. A 16-channel UTE coil was used for imaging. A 2D UTE sequence was used with repetition time 4.5 msec, echo time 0.06–2.00 msec, bandwidth 14 kHz, and a field of view 14 x 14 cm². A UTE T1p sequence was also used with repetition time 1000 msec, echo time 2.52 msec, bandwidth 14 kHz, and a field of view 14 x 14 cm². The UTE T2* and T1p images were postprocessed on a clinical workstation (Leonardo; GE Healthcare, Milwaukee, Wisconsin) using a dedicated software application (UltraLow Echo Time; GE Healthcare, Milwaukee, Wisconsin). T2, UTE T2*, and UTE T1p images were compared with T2, T1, T1p, T2*, T1*, and T2 images that were obtained from a separate MR session with a 3-T MR imager (Signa HDx; GE Healthcare, Milwaukee, Wisconsin).

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Abbreviation:

TE = echo time
TR = repetition time
UTE = ultrashort echo time

Author contributions:

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Conflicts of interest are listed at the end of this article.
Wis) with gradient capabilities of 150 (T · m−1)/sec slew rate and 40 mT/m amplitude on each axis was used in conjunction with a home-built 1.5-cm solenoid transmit-receiver coil. The tissue samples were placed in a plastic sample container filled with perfluorooctyl bromide solution (Solvay America, Houston, Tex) to minimize susceptibility artifacts from air-tissue interface during MR imaging, which was performed at room temperature (16). The container was positioned inside the solenoid coil where the main circumferential collagen fibers of the meniscus were oriented perpendicular to B₀ to minimize so-called magic angle effects. All the images were acquired in the coronal plane.

**MR pulse sequences.**—The two-dimensional UTE sequence used half-pulse radiofrequency excitation together with radial ramp sampling and fast transmit and receive switching to allow a minimal nominal echo time (TE) of 8 μsec (5,9,17–19). The approach to measurement of T2* was similar to the conventional strategy of varying the TE while keeping the repetition time (TR) constant. T1ρ of the meniscus was measured with spin-lock–prepared UTE T1ρ acquisitions at progressively increasing spin-locking times (5). T1, which was required to calculate T1ρ, was measured for each meniscus sample with a saturation recovery sequence by using a short hard 90° pulse followed by a UTE acquisition with a series of saturation recovery times (10,17). T2 of the meniscus was measured with a conventional Carr-Purcell-Meiboom-Gill spin-echo sequence. Typical acquisition parameters are summarized in Table 1.

**Indentation Stiffness Assessment**

Indentation stiffness testing was performed by using a biomechanical testing system (MACH-1; Biomomentum, Quebec, Canada). The samples were placed into a custom mold that served to inhibit sample movement and lateral

![Figure 1](image-url)  
**Figure 1:** Drawing of the microanatomy of the meniscus. 1, A superficial network; 2, a lamellar layer beneath the superficial network; 3, predominantly circular-oriented bundles of collagen fibrils; and 4, radial-tie fibers.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>MR Imaging Sequence Parameters</th>
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<tbody>
<tr>
<td>Sequence</td>
<td>TE</td>
</tr>
<tr>
<td>Spin-echo T2 mapping</td>
<td>2000</td>
</tr>
<tr>
<td>Two-dimensional UTE</td>
<td>300</td>
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<tr>
<td>UTE T2* mapping</td>
<td>100</td>
</tr>
<tr>
<td>UTE T1ρ mapping†</td>
<td>400</td>
</tr>
<tr>
<td>UTE T1 saturation recovery time</td>
<td>Various*</td>
</tr>
</tbody>
</table>

Note.—FOV = field of view.  
* 10, 20, 50, 100, 200, 400, 800, and 1200 msec.  
† Spin-locking times (saturation recovery times) = 0.02, 2, 5, 10, and 20 msec.
expansion and to hydrate during indentation testing. The samples were positioned so that either femoral or tibial surface faced up, to expose an articular surface for indentation measurement. A 1.0-mm diameter plane-ended cylinder indenter (Fig 2) was positioned orthogonal to lamellar surface, and it was lowered down until it touched the sample by using a load threshold criteria of 0.1 g. Indentation protocol consisted of 100-μm compressive displacement over 1 second, a hold for 1 second, then release at the same rate. The peak force (in grams) was determined from measured force-time data. Indentation was performed at multiple sites 1.0-mm apart along the entire length of the sample surface. Photographs were taken and compared with MR images to spatially register indentation sites to location of lamellar thickness measurement.

**MR Image Analysis**

**Morphologic assessment.**—The images of all samples obtained with all pulse sequences were reviewed in consensus by two board-certified radiologists (J.Y.C. and M.I., with 11 and 1 years of experience of musculoskeletal radiology, respectively). Samples were divided into two groups: the normal lamellar layer group and abnormally thickened lamellar layer group according to MR findings on UTE images. Normal lamellar layer was defined as a linear or mild crescent-shaped thickening pattern of the lamellar layer with smooth tapering both inward and outward (Fig 3a), and abnormally thickened lamellar layer was defined as a localized thickening of the lamellar layer with uneven articular surface or abrupt transition and without normal smooth tapering pattern (Fig 3b). A total of 17 lamellar layers including 13 normal and four abnormally thick layers were analyzed. One femoral surface with marked susceptibility artifacts was excluded. There was no other gross abnormality such as tear, signal intensity, or morphologic alteration in other meniscal structures including deep circumferential fibers and radial-tie fibers in all samples except one, which showed localized hyperintensity in the central main portion on UTE MR images.

**Thickness of the lamellar layer.**—Lamellar layer thicknesses were measured at sites that were one-to-one correspondence with the sites at which indentation stiffness testing was performed by using the open-source software (Osirix) (Fig 2). Measurements for all samples were performed independently by two radiologists.

**Spin-echo T2, UTE T2*, and UTE T1ρ measurements.**—On the images obtained with the Carr-Purcell-Meiboom-Gill and UTE sequences, regions of interest were placed on whole lamellar layer on each surface to determine spin-echo T2, UTE T2*, and UTE T1ρ values. Regions of interest were drawn freehand along the border of the entire lamellar layer (Fig 4a). T2*, T1, and T1ρ values were obtained by using a Levenberg-Marquardt fitting algorithm developed in house on the basis of equations previously described (5). The analysis algorithms were written by using software (Matlab version 7.9; Mathworks, Natick, Mass) and were executed offline. The program allowed placement of regions of interest on the first image of the series that were then copied to the corresponding position on each of the subsequent images. The mean intensity within each of the regions of interest was used for subsequent curve fitting.

**Statistical Analyses**

Interobserver reliability for measurement of lamellar layer thicknesses between two radiologists was examined by using an intraclass correlation coefficient. For each
surface, correlation analyses between the thicknesses of the lamellar layer and the peak forces from indentation test were determined by using the Spearman correlation analysis. The mean quantitative MR values (spin-echo T2, UTE T2*, and UTE T1p) and indentation peak forces were compared between normal and abnormal samples by using Mann-Whitney test. Nonparametric tests were used because of a relatively small number of samples. All the statistical analyses were performed with software (SPSS version 18.0; SPSS, Chicago, Ill). For all tests, \( P \) values less than .05 were considered to indicate statistical significance.

**Results**

**Comparison of Mean Values of Normal and Abnormal Samples**

Interobserver reliability for measurement of the lamellar layer thicknesses was excellent (intraclass correlation coefficient, 0.942) between two radiologists. Thirteen normal lamellar layers had a mean thickness of 232 \( \mu \text{m} \) ± 86 (range, 78–582 \( \mu \text{m} \)) and mean indentation peak force of 1.37 g ± 0.87 (range, 0.07–4.93 g); for four femoral lamellar layers, mean thickness was 278 \( \mu \text{m} \) ± 98 (range, 100–425 \( \mu \text{m} \)) and mean peak force was 1.30 g ± 0.92 (range, 0.11–3.72 g); and for nine tibial lamellar layers, mean thickness was 210 \( \mu \text{m} \) ± 75 (range, 132–582 \( \mu \text{m} \)) and mean peak force was 1.40 g ± 0.90 (range, 0.10–4.93 g). There was no significant difference in the thickness and peak force between femoral and tibial surfaces. Four abnormally thickened lamellar layers showed mean thickness of 353 \( \mu \text{m} \) ± 98 (range, 110–735 \( \mu \text{m} \)) and mean peak force of 0.72 g ± 0.31 (range, 0.14–2.08 g). The mean peak force tended to be lower in the abnormal surfaces \((P = .100)\) but the difference was not statistically significant.

The mean spin-echo T2, UTE T2*, and UTE T1p values of normal lamellar layers were 23.99 msec ± 11.98, 4.93 msec ± 2.17, and 9.75 msec ± 2.43, respectively, whereas those of abnormally thickened lamellar layers were higher, as follows: 47.20 msec ± 35.15, 12.66 msec ± 7.42, and 22.61 msec ± 19.43, respectively (Fig 4). In particular, UTE T2* values were significantly higher \((P = .010)\) in the abnormal surfaces, while the other values were not.

**Correlation between Lamellar Layer Thickness and Indentation Stiffness**

In normal surfaces, lamellar layer thicknesses correlated positively with the indentation peak force \((\text{Spearman} \quad \rho = 0.493–0.912; \quad P < .05\) except two layers with \(P = .094\) and \(P = .090)\) (Table 2; Fig 3c). However, in abnormal surfaces, the lamellar layer thickness showed significant negative correlation in some \((\text{Fig 3d})\), and no significant correlation in the others.

**Discussion**

The lamellar, circumferential, and radial fibers form a complex architectural network within the meniscus that help to withstand the varied forces (eg, shear, tension, and compression) to which it is exposed. The lamellar layer is known to serve as...
The results of our study indicated a substantial positive correlation between the lamellar layer thickness and indentation peak force in the normal lamellar layer of the knee meniscus, and a significant difference in UTE T2* values between the two groups with normal and abnormally thick lamellar layers. Our results strongly support our proposed hypotheses.

Figure 4: Representative MR images show meniscus with (a–d) normal lamellar layer and with (e–g) abnormally thick lamellar layer. (a) Placement of region of interest by freehand drawing. Corresponding spin-echo T2 (in milliseconds) (b, e), UTE T2* (in milliseconds) (c, f), and UTE T1p (in milliseconds) (d, g) overlay color maps apparently demonstrate differences in the lamellar layer matrix between the two groups. The spin-echo T2, UTE T2*, and UTE T1p values are generally higher in the abnormal surfaces compared with the values in the normal surfaces.

An envelope for the circumferentially oriented fiber bundles in the central main portion of the meniscus (20) and be well suited to facilitate surface to surface motion (21).
Our results suggest that normal lamellar layer plays a considerable role in the resistance to compressive load, especially at the contact surfaces between the articular cartilage and meniscus. Lai et al (22) reported that the menisci were either substantially stiffer near the surface or had comparable compressive stiffness through the depth of the meniscus through compression testing. By assuming that a normal lamellar layer is relatively homogeneous and somewhat stiffer than underlying layers, it would be logical that a thicker layer would result in higher indentation stiffness. Also, we presume that, even within an individual, the lamellar layer thicknesses could vary depending on meniscal regions such as anterior, middle, and posterior horn because the compressive loading is different depending on meniscal regions (23). Therefore, the lamellar layer thicknesses could vary depending on meniscal regions such as anterior, middle, and posterior horn because the compressive loading is different depending on meniscal regions (23).

In addition, we observed that femoral lamellar layers tended to be thicker than tibial ones in normal samples. In daily practice, we encounter horizontal tears that extend to the tibial surface more commonly than those extending to the femoral surface (24). The reason for this finding is unclear. Probably, several factors influence on this phenomenon. We guess that our observation (ie, thicker lamellar layer in the femoral surface) may be one of the factors. It would be worth researching to consider our finding as a possible factor to follow up in further studies on tear pattern of the meniscus.

Compared with normal surfaces, the abnormal surfaces had a somewhat lowered stiffness and significant increase in thickness. It is conceivable that even though abnormal lamellar layers are thicker than normal ones, they might lose their stiffness, possibly because of degenerative changes. The degenerative process may be gradual and inhomogeneous, which would lead to varying degrees of lamellar thickening and associated focal softening. In our study, this was apparent in quantitative MR imaging maps, which showed a more heterogeneous distribution of T2, T2*, and T1p values in abnormal surfaces. The mechanism of indentation softening in a heterogeneous tissue is more complex than that in a homogeneous tissue. Consideration must be given to interaction between the heterogeneous lamellar layer and underlying meniscal tissue, and it may require rigorous assessment of intratissue deformation at micron resolution (25) to fully address this issue.

We also compared T2, UTE T2*, and UTE T1p values between the normal and abnormally thick lamellar layers. On the basis of our results, abnormally thick lamellar layers showed increase of all spin-echo T2, UTE T2*, and UTE T1p values compared with normal lamellar layers, though only UTE T2* values were significantly different. These results suggest sensitivity of UTE T2* properties to pathologic and biomechanical changes of human meniscus. UTE T2* sequence may be useful for early evaluation of human meniscus.

There are several limitations in our study. First, there is a minimal discrepancy in one-to-one correlation of lamellar layer thickness and indentation force, the registration involved visual correspondence between registration photographs and during the indentation testing and MR images, which may have introduced an approximately 1-mm inaccuracy, even though care was taken to minimize this. Second, the underlying main central portion fibers were possibly engaged because of the indentation depth of approximately 100 μm and a macro-scale indenter even if this indentation test was conducted on the lamellar layers. As mentioned earlier, the mechanics of indentation is such that the biomechanical property of deeper layer meniscal tissue could influence the results. To reduce the influence of deeper layer, we used a small indenter (1 mm) and a shallow indentation depth (100 μm).

In conclusion, variation of lamellar layer thickness in healthy human menisci was evident on two-dimensional UTE MR images. In normal lamellar layers, thickness is highly and positively correlated with surface indentation stiffness.
In addition, UTE T2\* values potentially can be used to differentiate between normal and abnormally thickened lamellar layers.

Disclosures of Conflicts of Interest: J.Y.C. disclosed no relevant relationships. R.B. disclosed no relevant relationships. W.C.B. disclosed no relevant relationships. M.I. disclosed no relevant relationships. S.S. disclosed no relevant relationships. E.Y.C. disclosed no relevant relationships. J.D. disclosed no relevant relationships. G.M.B. disclosed no relevant relationships. D.D. Activities related to the present article: disclosed no relevant relationships. Activities not related to the present article: author disclosed a pending patent for electrosprining of meniscal tissue. Other relationships: disclosed no relevant relationships.

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