Age-Dependent Changes in Total and Free Water Content of In Vivo Human Lenses Measured by Magnetic Resonance Imaging

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Purpose. To use magnetic resonance imaging (MRI) to measure age-dependent changes in total and free water in human lenses in vivo.

Methods. Sixty-four healthy adults aged 18 to 86 years were recruited, fitted with a 32-channel head receiver coil, and placed in a 3 Tesla clinical MR scanner. Scans of the crystalline lens were obtained using a volumetric interpolated breath-hold examination sequence with dual flip angles, which were corrected for field inhomogeneity post-acquisition using a B1-map obtained using a turbo-FLASH sequence. The spatial distribution and content of corrected total ($\rho_{\text{tot}}$) and free (T1) water along the lens optical axis were extracted using custom-written code.

Results. Lens total water distribution and content did not change with age (all $P > 0.05$). In contrast to total water, a gradient in free water content that was highest in the periphery relative to the center was present in lenses across all ages. However, this initially parabolic free water gradient gradually developed an enhanced central plateau, as indicated by increasing profile shape parameter values (anterior: 0.067/y, $P = 0.004$; posterior: 0.050/y, $P = 0.020$) and central free water content (1.932 ms/y, $P = 0.022$) with age.

Conclusions. MRI can obtain repeatable total and free water measurements of in vivo human lenses. The observation that the lens steady-state free, but not total, water gradient is abolished with age raises the possibility that alterations in protein-water interactions are an underlying cause of the degradation in lens optics and overall vision observed with aging.

Keywords: magnetic resonance imaging, crystalline lens, T1 mapping, water content, in vivo, aging

Throughout adulthood, the human eye undergoes multiple changes in overall visual function, the majority of which are associated with changes to the refractive and transparent properties of the crystalline lens.1 Perhaps the most striking of these age-related changes is presbyopia in middle age2,3 and cataract in the elderly,4 which are manifestations of a loss of lens elasticity and transparency, respectively. A more subtle phenomenon that occurs to overall vision is the gradual shift in refractive status of the adult eye toward hyperopia,5–11 which has been attributed to a decline in the refractive power of the lens with age, known as the “lens paradox” because of the counterintuitive nature in which the lens loses power.12,13 Continual lifelong growth of the lens leads to a thicker and rounder shape with age,14–16 so if one assumes the internal refractive index gradient (GRIN) remains constant, these geometric changes should over time result in a more powerful lens. The fact that the opposite occurs,3,17–19 suggests that aging processes alters the lens GRIN to counter the effects of lens growth on its geometry to produce a less-powerful lens.

Extensive work has demonstrated the lens GRIN does indeed change with age,18–25 but there remains considerable debate regarding the underlying physiological mechanisms responsible for this change. Most recently, we used magnetic resonance imaging (MRI) to noninvasively obtain the GRINs of a wide age range of human lenses in vivo26 and incorporated these values into an optical modeling platform that attempted to predict the clinically-measured visual performance of each subject. Our findings showed that the incorporation of an age-dependent factor into the MRI transverse relaxation time T2 versus refractive index (T2-n) calibration18 was necessary to accurately model the measured refractive error. The need for the age-dependent factor indicates that with advancing age the contribution of lens proteins to the refractive index changes. Because T2 measures the water-to-protein ratio, this implies that aging...
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Supporting Information

Methods

Subjects

MRI Procedure and Post-Processing

All subjects were fitted with a 32-channel head receiver coil (Siemens, Munich, Germany) before undergoing MRI using a 3T clinical scanner (MAGNETOM Skyra; Siemens) located in the Centre for Advanced MRI at the University of Auckland. Inside the scanner, subjects laid supine on a table with their heads stabilized by foam pads. A Falcon tube filled with room temperature water that served as an external water reference to obtain the relative fraction of the lens total water content was strapped next to each subject’s eye (Fig. 1A). During the scan, subjects used a 45° tilted mirror attached to the head coil to view a fixation crosshair combined with pictures that randomly changed every five seconds, giving a viewing distance of about 2.1 m.

In this study, PD- and T1-mapping of the lens were performed using a volumetric interpolated breath-hold examination sequence with dual flip angles (field of view = 159 mm; matrix size = 768 × 768; slice thickness = 3 mm; echo time [TE] = 2.7 ms; repetition time [TR] = 15 ms; flip angle [α] = 4° and 23°; parallel imaging acceleration factor = 2; total imaging time = 4.5 minutes). Post hoc field inhomogeneity correction of the raw MR images acquired at the two flip angles was then performed using a B1-map obtained by a turbo-FLASH sequence with pre-saturated preparation (field of view = 220 mm; matrix size = 160 × 160; slice thickness = 3 mm; TE = 2.23 ms; TR = 1349 ms; parallel imaging acceleration factor = 2; total imaging time = 30 s). The B1-map was resliced and coregistered with one volumetric set of images (α = 23°) for pixel-wise signal corrections.

Corrected PD and T1 maps were then calculated using the following MRI signal equation

\[
\frac{S}{\sin(ab_1)} = \frac{S}{\cos(ab_1)} e^{\frac{\alpha}{T_1}} + PD \left(1 - e^{\frac{\alpha}{T_1}}\right)
\]

where \(S\) is the signal intensity, \(a\) is the flip angle, \(PD\) is the resting spin density before excitation (i.e., at time zero), and \(b_1\) is a multiplier that denotes the ratio of actual flip angle (biased by field inhomogeneity) and ideal flip angle that is calculated from the acquired B1-map.
Corrected PD and T1 values of the lens were derived from the coefficients of the linear fitting performed between \( \frac{\text{PD}_{\text{lens}}}{\text{PD}_{\text{water}}} \) and \( \frac{\text{T1}_{\text{lens}}}{\text{T1}_{\text{water}}} \) to obtain the slope (i.e. \( e^{-\frac{TR}{T1}} \)) and the y-intercept (i.e., \( \text{PD}(1 - e^{-\frac{TR}{T1}}) \)). The PD value of the external water reference \( \text{PD}_{\text{water}} \), which is representative of 100% water, was similarly calculated with Equation 1. The total water content of the lens \( \rho_{\text{lens}} \) was expressed as the relative fraction of the external water reference by normalizing the PD value of the lens \( \text{PD}_{\text{lens}} \) to \( \text{PD}_{\text{water}} \) as in Equation 2:

\[
\rho_{\text{lens}} = \frac{\text{PD}_{\text{lens}}}{\text{PD}_{\text{water}}} \tag{2}
\]

where \( \rho_{\text{lens}} \) is the total water content of the lens, described as a percentage unit of the external water reference (pu). Using this approach, instrumental factors such as different MRI manufacturers, field strength and radiofrequency coils are eliminated, giving the nontrivial measurement of the lens total water content.

For all MR images, the slice that contained the thickest cross-section of the lens visible was chosen for data extraction. All data processing was carried out using custom-written routines in MATLAB (MathWorks, Natick, MA, USA). A one-dimensional trend analysis was performed for each lens PD- and T1-map. For this, \( \rho_{\text{lens}} \) and T1 values over a 5-pixel-wide band along the lens optical axis were extracted, averaged and then fitted to a power function (Equation 3) going anteriorly and posteriorly from the lens center to give two independent profiles for total and free water from each subject’s lens:

\[
\rho_{\text{lens}}/T1(r) = a + b(r)^c
\]

where \( r \) is the normalized radial distance \((r/a)\) from the lens surface, \( a \) is the total water \((\rho_{\text{lens}})\) or free water (T1) value at the lens center, \( b \) is the variation in \( \rho_{\text{lens}} \) or T1 between the lens center and surface (such that it is positive when water content is higher at the surface), and exponent \( c \) characterizes the rate of change in the \( \rho_{\text{lens}} \) or T1 profile gradients (i.e., the slope). Because water distribution along the equatorial axis is of limited relevance to the optics of the lens, its analysis was not included in this part of the study. The data extraction and fitting process are demonstrated in Figure 1.

**Statistical Analysis**

Linear regression of lens water parameters using age in years as the independent variable was performed to determine whether there was an age dependence. Right eyes were used in analyses unless it did not satisfy the inclusion criteria (one subject). All results were reported as mean ± standard deviation unless stated otherwise. All statistical analyses were performed using IBM SPSS Statistics (v25.0, IBM Corp. Armonk, NY, USA) and MATLAB, with a significance level of 5% used for all tests.

**RESULTS**

We first present results for total \((\rho_{\text{lens}})\) and free (T1) water content obtained by applying our optimized in vivo MRI protocols to a small cohort of younger subjects to confirm the repeatability of our measurements. We then apply our methods to larger and older subject cohorts to assess how
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Validation of In Vivo Lens Water Measurements

To establish our protocols and test their repeatability, we first recruited a smaller cohort of seven young subjects (two males, five females; mean age: 28 ± 6 years). In this cohort, \( \rho_{\text{lens}} \) and T1 values were extracted as either total (Fig. 2A) or free (Fig. 2B) water profiles, respectively, across the optical axis or as mean values of total (Fig. 2C) and free (Fig. 2D) water obtained from three regions of interest defined as the anterior cortex (\( r/a = -0.85 \)), the nucleus (\( r/a = 0 \)), and the posterior cortex (\( r/a = 0.85 \)). The total water profiles obtained for these young lenses were mostly linear (Fig. 2A), indicating that total water distribution across the lens is relatively constant. However, a repeated measures one-way analysis of variance revealed a slight but significant decrease (\( P = 0.016 \)) in the mean total water content of the nucleus (43 ± 4 pu) relative to the anterior and the posterior cortex (both 47 ± 5 pu) (Fig. 2C). In contrast, the free water distribution across the optical axis varied considerably, exhibiting a parabolic shape (Fig. 2B). Free water content in the nucleus (970 ± 82 ms) was significantly lower relative to both the anterior (1459 ± 69 ms; \( P < 0.005 \)) and posterior (1359 ± 102 ms; \( P < 0.005 \)) cortex (Fig. 2D), with no significant difference being observed between the mean free water content of the two cortical regions (\( P = 0.163 \)). Interestingly, although the variability of total water content between subjects was relatively small, presumably in part because of the normalization to an external water standard, considerably greater intersubject variability was observed for the T1 values that represent free water (Figs. 2B, 2D).

To rule out whether this observed intersubject variability in T1 values was due to MRI measurement error, the seven young subjects were rescanned at the same time of day (± 1 hour) up to five days after their initial scan. To evaluate test-retest repeatability of the lens free water measurements, both the shape of the T1 profiles (Fig. 3A) and the mean T1 values obtained in the different regions of the lens (Fig. 3B) were compared between scans using Pearson’s correlation analysis, paired t-tests, and Bland-Altman plots. Within-subject coefficients of variation (CoV) were also calculated based on a within-subject standard deviation method. These analyses showed that the shape parameter (exponent \( c \)) of the anterior and posterior free water profiles...
FIGURE 3. Validation of lens free water (T1) measurements in vivo in a cohort of young subjects. After an initial scan (Visit 1), the seven young subjects were rescanned (Visit 2) at the same time of the day (±1 hour), but up to five days after the first scan. (A) Correlation (left) and Bland Altman (right) plots of T1 profile slopes (shape parameter exponent c) fitted anteriorly (filled circles) and posteriorly (empty circles) from the lens center using Equation 3. (B) Correlation (left) and Bland Altman (right) plots of lens free water content (T1 values) at the anterior cortex (r/a = −0.85; filled squares), nucleus (r/a = 0; shaded squares), and posterior cortex (r/a = 0.85; empty squares). On the correlation plots, the black solid line indicates the best fit line, and the dashed gray line indicates the unity line. On the Bland-Altman plots, the black solid line indicates the mean bias, whereas the dashed gray lines indicate the upper and lower 95% limits of agreement.

obtained along the optical axis did not significantly differ (mean difference: −0.43 ± 1.36, 95% confidence interval [CI] [−1.22, 0.35], P = 0.255) between scans (Fig. 3A, left panel), whereas the correlation plot showed a strong agreement (r = 0.693, P = 0.006) between T1 profiles shapes between the two visits (Fig. 3A, right panel). The within-subject CoV was 16%. The free water content of the nucleus, anterior and posterior cortices also did not significantly differ (mean difference: 16 ± 97 ms, 95% CI [−29, 60] ms, P = 0.470) between the two MRI scan sessions (Fig. 3B, left panel). There was also a strong correlation between the regional free water content obtained at the two visits (r = 0.927, P < 0.0005), and the within-subject CoV was 5% (Fig. 3B, right panel). This confirms the repeatability of MRI protocols to measure lens water content and tends to suggest that the variability we have observed in lens free water is likely to be due to inherent intersubject biological variability.

Age-Dependent Changes in Lens Total and Free Water Content and Distribution

Having established and tested the repeatability of our MRI protocols to measure the lens water content and distribution in vivo on a subcohort of young subjects we then applied these protocols to a larger cohort of subjects, who had been previously recruited for another study, to investigate how total and free water changes in the lens with age. Unfortunately, due to imaging artefacts, PD and T1 lens maps could not be extracted in four subjects (3 middle-aged and 1 older) and these subjects were excluded from further analysis. Hence, the effects of age on total and free water for 53 subjects are reported (Table).

| Characteristic | Young (18–40) | Middle-Aged (41–60) | Older (>60) |
|----------------|---------------|---------------------|-------------|
| No.            | 22            | 17                  | 14          |
| Age (y)        | 24 ± 4        | 50 ± 5              | 73 ± 6      |
| Male           | 10 (45%)      | 10 (59%)            | 5 (36%)     |
| Refraction (D) | −1.52 ± 1.70  | −0.77 ± 1.75        | 0.94 ± 1.55 |

Data are reported as either mean ± standard deviation (SD) or No. (%).

| Characteristic | Young (18–40) | Middle-Aged (41–60) | Older (>60) |
|----------------|---------------|---------------------|-------------|
| Total Water    |               |                     |             |

Consistent with the PD profiles of lenses extracted from the smaller cohort of younger subjects, total water content (ρ_{total}) remained relatively constant across the optical axis of young (Fig. 4A), middle-aged (Fig. 4B), and older (Fig. 4C) lenses, and the profiles did not appear to be influenced by aging (Fig. 4D). This was confirmed by plotting mean total water content obtained from the three regions of interest of the lens against subject age (Fig. 5). Total water content in the nucleus (Fig. 5A; P = 0.617),
expected distributions, the paucity of comparable content in the in vivo human lens. The studies most comparable to our current work are those of Patz et al. and Richdale et al., who have published nuclear T1 values of 1138 ms and 1270 ms, and cortical T1 values of 1413 ms and 1630 ms, respectively, obtained from MRI experiments. Although these studies broadly agree with the water MRI signal, which would account for much of the reported discrepancies.

Free water content in these studies was also typically represented as a percentage of lens total water, which is not directly comparable to the MRI longitudinal relaxation time T1 used in this study. Our average T1 values of the lens nucleus (996 ± 120 ms) and cortex (1107 ± 105 ms) are similar to those reported by studies employing nuclear magnetic resonance (NMR), a noninvasive technique that works on the same principles as MRI to measure free water content in tissues, although NMR does not spatially resolve the regional variations in free water across the lens. The studies most comparable to our current work are those of Patz et al. and Richdale et al., who have published nuclear T1 values of 1138 ms and 1270 ms, and cortical T1 values of 1413 ms and 1630 ms, respectively, obtained from MRI experiments. Although these studies broadly agree

DISCUSSION

In this study we have optimized dual MRI parametric PD- and T1-mapping protocols to obtain for the first time total and free water gradients simultaneously in human lenses in vivo. Our technique demonstrated excellent interday repeatability of lens free water (T1) measurements (Fig. 3) and suggested that the observed variability in T1 values was due to inherent biological variability in free water content between subjects. Overall these result show that MRI can be used as a reliable tool for quantifying water distribution and suggested that the observed variability in T1 values was due to inherent biological variability in free water content and free water gradients simultaneously in human lenses.

FIGURE 4. Differences in the distribution of total water ($\rho_{\text{lens}}$) in the human lens between age groups. Average profile plots of total water content ($\rho_{\text{lens}}$ values) against normalized distance (r/a) extracted along the optical axis from lenses in young (A, blue), middle-aged (B, green), and older (C, red) age groups. Error bars: ±2 standard error. (D) Comparison of the average total water profiles obtained for each age group to emphasize that total water distribution does not markedly change with age.

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FIGURE 5. Effect of age on regional total water (\( \rho_{\text{lens}} \)) content in the human lens. Mean total water content (\( \rho_{\text{lens}} \) values) obtained from the lens (A) nucleus (\( r/a = 0; \) shaded squares), (B) anterior cortex (\( r/a = -0.85; \) solid squares), and (C) posterior cortex (\( r/a = 0.85; \) empty squares) plotted against age. No significant correlation between total water content and age was found in any of the three regions.

with our results, differences in imaging protocols (spin-echo sequence vs. variable flip angle), MRI field strengths (1.5T and 7T vs. 3T) and definitions of the nuclear and cortical regions (manual selection vs. predefined fractions) should be carefully considered given the dependence of T1 on these factors.68

Application of our optimized protocols to a larger and older subject cohort revealed that although increasing age had no discernible effects on total lens water content (Figs. 4, 5), an increase in the free water content of the lens nucleus and the development of central plateau in the free water profile occurred with increasing age (Figs. 6, 7). In a previous study performed on the same cohort of subjects,26 similar age-related changes to the GRIN, but not T2, profile of the lens were found (Fig. 8). T2 values of the lens are dominated by proton exchange between bound water molecules and crystallin proteins; they can be used as a measure of the lens water-to-protein ratio (Fig. 8B) and for this reason should be correlated to the GRIN (Fig. 8C). However, our recent discovery that the same T2 values would yield different GRINs depending on age explains why the T2 profiles appear relatively unchanged in spite of an evident decrease in the lens central refractive index and development of a flattened central plateau in the GRIN profile between subject age groups. We previously hypothesized that this was most likely due to an age-dependent decrease in the refractive index increment of lens proteins, although it could also occur from an increase in lens total water or a loss in lens protein content. When taken together, our T1, T2, and GRIN measurements suggest that with age, the observed GRIN changes are driven by changes in the free water gradient, rather than changes in the lens total water or protein concentration. These observations therefore confirm that it is the interactions between protein and water that are altered with advancing age. Interestingly, these reciprocal age-dependent changes in the free water gradient and GRIN appear to contribute to the clinically observed lens paradox by reducing the optical power of the lens.26

Until now, it is not known whether the observed age changes in lens free and bound water proportions are a cause and/or effect of protein conformational changes in the lens. It has been postulated that the conversion of bound to free water does not occur solely as a simple consequence of the protein changes over that time period.29 Although our observations do not inform us on how age alters the interaction between water and lens proteins that affects overall lens optics, we can speculate on two possible mechanisms. In the first, we envisage that the lifelong accumulation of posttranslational modifications to lens proteins69–71 directly impairs their ability to bind water, causing an increase in free water content in the deeper areas of the lens where long-lived crystallin proteins are concentrated. In line with this, several protein modifications including truncation, oxidation, and deamidation have been identified in water-insoluble protein fractions of both clear and cataractous aged lenses.72 The second possible mechanism involves age-related changes to the internal hydrostatic pressure gradient, which has been measured in all lenses studied to date.73,74 This pressure gradient is generated by the outward movement of water through an intracellular pathway mediated by gap junction channels that is driven by the lens microcirculation system.75–77 It has been revealed that this pressure gradient is regulated by a dual feedback system, which uses the mechanosensitive channels TRPV1 and TRPV4 to sense changes in pressure at the lens surface.78 The realization that the lens pressure gradient is subject to feedback regulation78 raises the possibility that age-related changes in this internal pressure gradient affect protein hydration in the lens nucleus.

Before the discovery of the internal hydrostatic pressure gradient, it was predicted on theoretical grounds that the lens undergoes reversible “syneresis” between the protein-bound state and surrounding free water as an operative response to changes in hydrostatic pressure.79 A series of experiments by Bettelheim et al.79–81 showed that applying external pressure to the lens surface causes a “reversed” syneresis, where free water enters the hydration shell around
polar protein domains to become “bound,” decreasing the lens free-to-bound water ratio. When the pressure was alleviated, this movement of water molecules is reversed and the free-to-bound water ratio increased. However, the extent to which water molecules in the lens undergo reversed syneresis in response to increasing pressure was found to diminish with age, and is actually reversed in older lenses (i.e., converts from bound to free). The relevance of this finding to aging was not fully recognized at the time, because it was thought that the pressure values tested were larger than those experienced by the lens in vivo. Consistent with this notion, an age-dependent increase in endogenous hydrostatic pressure gradient has been reported in mice lenses, but whether this also occurs in humans is yet to be determined. Of course, because protein hydration depends on protein structure (and vice versa), age-dependent modifications of lens proteins may also affect the ability of proteins to engage in syneresis. Hence, to fully understand the mechanisms responsible for the disruption of steady-state free water distribution and content and their implications on lens optics, the relative contributions of the lens physiology (pressure gradient) and protein structure to syneresis need to be further delineated.

Although free water in all lenses had the same parabolic distribution (Fig. 2B), large intersubject variability in absolute values were observed (Fig. 2D). Having shown the repeatability of our MRI measurements (Fig. 3), we concluded that this variability was due to inherent biological variability in free water content that in turn reflects individual differences in the protein content and composition of the lens nucleus. Because they are first established during embryonic development, we propose that the observed subject-specific lens free water content reflect differences in lens water transport mechanisms designed to compensate for these potential differences to achieve a consistent and appropriate nuclear refractive index and lens GRIN overall, obtained by either the active removal of free water from the nucleus or via the imposition of a pressure gradient that modulates the amount of water bound to proteins via syneresis. Furthermore, the awareness that pharmacological modulation of ciliary muscle tone to alter the zonular tension applied to the mouse lens, also alters the magnitude of the lens hydrostatic pressure gradient, suggests that the GRIN can be subjected to external regulation presumably to modify lens power and therefore overall visual performance. In the nonaccommodating mouse lens, we posit this mechanism is used to adjust the
steady-state power of the lens as it grows throughout the lifetime of the animal. In humans, we anticipate a similar modulation of lens pressure would account for the age-dependent changes in its refractive power\textsuperscript{3,17–19,26}; however, because the young human lens can accommodate, we also need to consider whether changes in zonular tension may be dynamically altering lens pressure and ergo, via syneresis, the free water distribution or content of the lens to drive the shape changes associated with accommodation.\textsuperscript{85–87} It is possible that presbyopia occurs as a consequence of the diminished ability of the aging human lens to compensate for changes in zonular tension, and therefore hydrostatic pressure, during accommodation. This would not only explain the observed age-dependent increase in free water proportion within the lens, but also be expected to have implications on the stiffness and accommodative ability of the lens. The protocols developed in this study have the potential to test this hypothesis and will be the subject of future work.

In conclusion, by using optimized MRI protocols to spatially map the total and free water distribution and content of the lens, which act as in vivo physiological biomarkers of lens water transport, we have shown that changes to the free water population of the human lens are primarily responsible for the clinically observed changes in lens power and overall visual function that occur with age. Having established and tested the utility of MRI to study water content in the human lens in vivo, we expect that these protocols can be adapted to determine whether changes in lens water transport precede the onset of the age-related lens pathologies, presbyopia, and cataract.

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