DIRECTIONAL OPTICAL COHERENCE TOMOGRAPHY PROVIDES ACCURATE OUTER NUCLEAR LAYER AND HENLE FIBER LAYER MEASUREMENTS

BRANDON J. LUJAN, MD,*† AUSTIN ROORDA, PhD,* JASON A. CROSKREY, MD,‡ ADAM M. DUBIS, PhD,§ ROBERT F. COOPER, BS,¶ JAN-KRISTINE BAYABO, BA,* JACQUE L. DUNCAN, MD,** BHAVNA J. ANTONY, PhD,* JOSEPH CARROLL, PhD‡§¶

Purpose: The outer nuclear layer (ONL) contains photoreceptor nuclei, and its thickness is an important biomarker for retinal degenerations. Accurate ONL thickness measurements are obscured in standard optical coherence tomography (OCT) images because of Henle fiber layer (HFL). Improved differentiation of the ONL and HFL boundary is made possible by using directional OCT, a method that purposefully varies the pupil entrance position of the OCT beam.

Methods: Fifty-seven normal eyes were imaged using multiple pupil entry positions with a commercial spectral domain OCT system. Cross-sectional image sets were registered to each other and segmented at the top of HFL, the border of HFL and the ONL and at the external limiting membrane. Thicknesses of the ONL and HFL were measured and analyzed.

Results: The true ONL and HFL thicknesses varied substantially by eccentricity and between individuals. The true macular ONL thickness comprised an average of 54.6% of measurements that also included HFL. The ONL and HFL thicknesses at specific retinal eccentricities were poorly correlated.

Conclusion: Accurate ONL and HFL thickness measurements are made possible by the optical contrast of directional OCT. Distinguishing these individual layers can improve clinical trial endpoints and assessment of disease progression.

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The outer nuclear layer (ONL) of the retina contains the nuclei of the cone and rod photoreceptors.¹ Loss of the cellular machinery found in these nuclei causes irreparable loss of the photoreceptors and the capacity for visual function. The ONL thickness, consequently, has been a critical anatomical endpoint for ex vivo and animal studies of retinal degenerations.²⁻⁵ In vivo imaging using spectral domain optical coherence tomography (SD-OCT) has delivered the ability to directly measure the ONL thickness in animal models over time to monitor the natural history of disease and the effects of therapeutic interventions.⁶⁻⁸ Although it is tempting to use SD-OCT in a similar fashion to measure the ONL thickness in the macula of human subjects, standard SD-OCT image acquisition is confounded by the presence of Henle fiber layer (HFL) and cannot reliably identify the true ONL.⁹,¹⁰

Henle fiber layer consists of the photoreceptor axons and Müller cell processes that are substantial in the human macula.¹¹ Because of the directional reflectivity properties of HFL and its oblique course in the macula, it typically appears isoreflective to the true ONL on standard SD-OCT images.⁹ Consequently, HFL has been routinely included in manual and automated segmentations of the apparent ONL, thus resulting in an artificially thick assessment of the true ONL thickness.¹²⁻¹⁸ Although the influence of HFL has been recognized in several publications, it has been grouped together as the ONL+ or ONL + HFL without independent analysis of the contribution of each layer.¹⁹,²⁰ Without independent measurements, the validity of grouping these structures together as a surrogate for the true ONL thickness cannot be known.

Directional OCT (D-OCT) is a technique that can be applied to any type of OCT system that involves purposefully altering the OCT beam entry position. By changing the orientation of light incident on the retina,
D-OCT adds optical contrast to directionally reflective structures. In the application of D-OCT presented in this study, we identified a robust boundary between the true ONL and HFL and measured the thickness of these layers using horizontal and vertical cross-sectional images of normal subjects acquired with a commercial SD-OCT system. We tested the hypothesis that D-OCT can be used to determine the independent contributions of ONL and HFL thickness and that they were symmetrically distributed about the fovea on SD-OCT scans.

Given our ability to use this technique, we will refer to the ONL to mean the layer of the retina containing only the photoreceptor nuclei and HFL to mean the layer of the retina containing only the photoreceptor axons and Müller cell processes.

Methods

The Medical College of Wisconsin Institutional Review Board approved the protocol, each subject gave written informed consent to participate in the study, and the Declaration of Helsinki guidelines were followed throughout the study. Healthy volunteers without macular pathology were recruited for the study, and pupils were dilated using 2.5% phenylephrine and 1% tropicamide.

Fifty-seven eyes of 31 subjects were imaged at the Medical College of Wisconsin by a single operator using a single Cirrus HD-OCT system (Carl Zeiss Meditec Inc, Dublin, CA). Sets of D-OCT images were acquired using a previously described protocol. Briefly, this consisted of the acquisition of a central scan using a horizontal high definition 5-line raster setting which used an average of 20 B-scans, each comprised 1,024 A-scans over 20°. The central scan was defined by the pupil entry position that resulted in a “flat” appearing cross-sectional image (Figure 1, top). While the subject remained at the chin-rest, 2 additional horizontal scans using the same parameters were obtained from off-axis pupil positions between 1.5 mm to 2 mm away from the pupil position used to obtain the central scan (Figure 1, middle and bottom). An analogous protocol was then followed for vertical scans. During each scan, the subject was asked to maintain fixation on the center of the internal fixation target. The signal strength and quality of the scans were checked immediately after each acquisition and repeated if necessary.

Off-axis “tilted” B-scans were registered to the centrally acquired “flat” image by means identification of similar points of agreement between the two images using custom software written in MATLAB (Mathworks, Natick, MA). At least 20 uniformly distributed homologous control points were identified within each image set. A second-order polynomial 2-D spatial transformation was performed to obtain registered sets, which were verified by toggling between images. Registration was graded as acceptable if motion of either the internal limiting membrane or retinal pigment epithelium contours was undetectable. Images were registered pairwise between the flat image and a single tilted image, and then the flat image and other tilted images to result in a composite stack of three images of the same part of the macula having been acquired through the three different pupil positions. Left eyes were horizontally flipped to be right-eye equivalents for purposes of nasal/temporal comparison.

Axial length measurements were acquired on each subject using the intraocular lens master (Carl Zeiss Meditec, Inc) Axial length measurements were used to correct for retinal magnification differences as previously described.

Registered D-OCT image sets were segmented using ImageJ software (NIH, Bethesda, MD). Using all images in the set, boundaries were determined and manually segmented at the vitreoretinal interface, the innermost aspects of the interface between the synaptic outer plexiform layer (OPL) and HFL, HFL and the ONL, the ONL and the external limiting membrane and the bottom of the retinal pigment epithelium. Linear interpolation was then performed in MATLAB to obtain segmentation lines from the different boundary positions and standardize distances from the center of the dataset, the foveal center, as defined by the thinnest vitreoretinal interface–retinal pigment epithelium distance.
Manual segmentations were performed on all of the data by two independent masked graders. Intergrader variability was assessed by comparing the locations and area of the ONL and HFL segmentations. Additionally, a randomly chosen subset of five eyes was selected for the same grader to segment the data twice to measure intrgrader variability. The coefficient of variation between the graders was determined as the SD of the thicknesses of retinal layers divided by the mean value in pixels.

The correlation between the ONL and HFL thickness at 18 horizontal and vertical locations across 23 right eyes from 23 subjects having well-registered bilateral scans was computed using the open-source statistical package R (http://www.r-project.org/). The thickness at each location was a mean measurement over 0.3 mm. Spearman correlation coefficient $\rho$ and the $P$-value associated with each correlation coefficient were computed between values of the ONL and HFL thicknesses. At multiple eccentricities from the foveal center, the correlation between the ONL thickness and HFL thickness immediately overlying it was computed. Additionally, because of the oblique orientation of HFL, the correlation between ONL thickness and HFL thickness in more peripheral regions of the macular scans was also analyzed.

**Results**

Fifty-seven eyes of 31 subjects were imaged with 26 subjects having both eyes imaged, 4 subjects only had the right eye imaged and 1 subject only had the left eye imaged. Subjects ranged in age from 18 years to 34
years old (mean = 23 year), and 71% were male. All subjects were found to have no retinal pathology on the horizontal or vertical cross-sectional SD-OCT images. Fifty-two datasets were registered successfully to the “flat” image. Five datasets could not be sufficiently registered for accurate segmentation and were excluded from analysis due to a change in fixation and lack of image homology. Axial length measurements were obtained on all subjects. Mean (±SD) axial length was 24.2 mm (±1.1 mm).

The average coefficient of variation between graders was 0.05 indicating excellent reproducibility where grading differences were found to be less than 1 pixel (approximately 4 μm axially) in either direction from the junction of the ONL and HFL. Intragrader reproducibility for each grader was assessed, and the average coefficient of variation was 0.01.

Henle fiber layer hyperreflectivity contralateral to the side of pupil entry and HFL hyporeflectivity ipsilateral to the side of pupil entry was apparent in unregistered images (Figure 1). After registration to the flat image, the change in appearance of HFL and the outer retinal bands became more apparent. The change in reflectivity as a function of pupil entry position for each pixel of the registered scans can be visualized in a single image using the chromatic D-OCT visualization method described by Makhijani et al (Figure 2).21 Internal to HFL, the nondirectionally reflective synaptic OPL was equally reflective from each pupil entry position and consequently is visualized as a grayscale structure. This is in sharp distinction from HFL itself, which is prominently visualized as red or green, depending on a nasal or temporal pupil entry position, respectively.

A substantial decrease in intensity of the second and third hyperreflective outer retinal bands23 was apparent on the off-axis images of all eyes as compared with the flat images due to photoreceptor waveguiding and the optical Stiles-Crawford effect.24 The inner segment/outer segment junction, or ellipsoid zone,25 was maximally reflective in the central flat scans and diminished in intensity in each of the off-axis tilted images. Even more striking, the band containing the photoreceptor outer segment tips, or interdigitation zone,25 almost entirely disappeared in the tilted scans. These effects of directional reflectivity resulted in a blue coloration of these structures in the chromatic D-OCT image shown in Figure 2 because maximal reflectivity occurred in the flat position and a decrease in reflectivity occurred in each of the off-axis scans.

We hypothesized that within an individual that the ONL and HFL thicknesses were radially symmetric around the fovea. We tested this in horizontal and vertical scans for left and right eyes. The differences between measures at equal distances were calculated, and one sample Hotelling T2 test was performed. The hypothesis was rejected in all comparisons (P < 0.0001 for each), indicating that the ONL and HFL measures were not symmetric within the individual eyes.

The ONL and HFL thicknesses were compared at each retinal eccentricity in the right and left eyes of the 23 subjects where well-registered bilateral scans were obtained. The ONL thickness values between eyes were significantly correlated at each measured eccentricity with a mean Pearson correlation coefficient \( r = 0.78 \) (SD = 0.10), \( P < 0.0001 \) for the horizontal measurements and mean \( r = 0.64 \) (SD = 0.23) \( P = 0.046 \) for the vertical. Henle fiber layer showed an even higher correlation by retinal eccentricity between eyes with a mean \( r = 0.90 \) (SD = 0.06) \( P < 0.0001 \) horizontally and \( r = 0.91 \) (SD = 0.04) \( P < 0.0001 \) vertically. Because of this high correlation between both eyes in this cohort, only one eye from each subject was chosen for detailed comparison of the ONL and HFL horizontal and vertical thickness values.

The ONL thickness was found to be substantially less than measurements possible with standard SD-OCT imaging that include HFL and varied significantly with eccentricity from the foveal center. Figure 3 illustrates the average thickness measurements and SDs for the ONL and the ONL + HFL by eccentricity in horizontal and vertical dimensions of a single eye of each subject. In the central fovea, the true ONL curve and the ONL + HFL were relatively close together, reflecting the substantial ONL contribution. The parafoveal thickness of the ONL rapidly decreased while the ONL + HFL curve actually increased initially and then slowly decreased in both horizontal and vertical meridians. With increasing eccentricity, the ONL + HFL thickness continued to decrease out to the most peripherally measured locations. However, the isolated mean ONL thickness reached a nadir at 0.8 mm temporally and 1.0 mm nasally. The ONL thickness then increased at greater horizontal eccentricities as the population of rod nuclei increased.26 The vertical ONL thickness curve differed from the horizontal ONL thickness curve in several respects. Its nadir was symmetric between the inferior and superior retina and occurred at 0.8 mm eccentricity. Distal to this, it demonstrated a more rapid rise and greater ultimate thickness at the most eccentric locations. These findings are consistent with the topographic distribution of macular rods found in histological studies.26

The relative proportion of the thickness occupied by the ONL compared with the standard measurements containing the ONL and HFL was also analyzed. Figure 4 shows the mean percentage of the total
ONL + HFL thickness comprised the ONL and HFL for all studied individuals at each eccentricity. In the center of the fovea, the ONL predominates, with HFL comprising less than 10.7% of the measured thickness on average. HFL was found to have its largest mean contribution around 0.75 mm eccentricity from the foveal center, where a mean 60.0% of the thickness was accounted for by HFL.

The axons from cones that form the HFL radiate outward, eccentric to the foveal center, to synapse with their corresponding bipolar cells. Because of this anatomical arrangement, we hypothesized that the ONL thickness would not show a positive correlation with HFL thickness directly overlying it and that ONL thickness would show a positive correlation with HFL thicknesses at greater eccentricities. To investigate this, we performed a systematic series of correlations between the ONL thickness and the HFL thickness directly above it as well as at more eccentrically displaced locations. The correlations are shown in Figure 5, where the correlation coefficients at each position and correlation coefficients with $P < 0.05$ were plotted for the horizontal and vertical scans. In the foveal center, where the ONL predominates there was a statistically significant negative correlation found between the ONL and HFL thicknesses. In the remaining macula, there were no other statistically significant correlations found between the thickness of the ONL and HFL immediately overlying it. The central ONL was found to have a statistically significant positive correlation with more peripheral HFL at 0.6 mm eccentricity in horizontal meridian and 0.9 mm eccentricity in the vertical meridian.

**Discussion**

Photoreceptor nuclei are contained in the ONL, and the thickness of this layer is a key biomarker in identifying normal aging changes and the progression of degenerative diseases. The ONL is an appealing biomarker for disease progression because photoreceptors are the direct mediators of vision, and the symmetry between eyes found in this study would allow for its...
measurement where one eye was treated and the contralateral eye was a control. Spectral domain optical coherence tomography images acquired in the standard fashion are unable to reliably distinguish between the true ONL and the overlying HFL. Consequently, we used D-OCT to optically delineate these layers. We found a dramatic difference in the thickness measurements resulting from the inclusion or exclusion of HFL.

Depending on eccentricity, the incorporation of HFL into the calculation of the ONL thickness results in a doubling of the thickness estimates. The HFL has its most substantial contribution between 0.5 mm and 1.5 mm eccentricity, where it accounts for over half of the thickness of the 2 layers combined. Furthermore, there is a real difference in the ONL thicknesses between the horizontal and vertical meridians due to differences in the cone and rod contributions that only becomes obvious on OCT when the contribution of HFL is unmasked. Because of the confounding effect of HFL thickness, both manual and automated algorithms reporting ONL thickness values must ensure that HFL is not included in the measurements.

Given the course of macular photoreceptor axons as they move from more central ONL to more peripheral synaptic OPL, our hypothesis that no fixed correlation exists between HFL and ONL at the same eccentricity was confirmed. There was a positive correlation between the ONL at the foveal center and more eccentric macular HFL. However, given the reported lengths of the longest Henle fibers is 675 μm and that these were found to originate in the parafovea, these correlations with the central ONL are likely spurious.28 Given this lack of correlation in individuals without macular pathology, the thickness of the ONL cannot be computed retrospectively after measuring the ONL and HFL together with standard OCT acquisition techniques.
Visualization of parts of the border between HFL and the ONL may be possible without D-OCT under several circumstances. These include high-quality frame averaged images in certain eyes, serendipitous off-axis scan acquisition, or secondary to the presence of deforming pathology. However, a reliable and robust method for routinely delineating these boundaries across the entire cross-section or volume scans should be implemented, particularly for use in prospective clinical trials that ideally use automated segmentation algorithms. Purposeful prospective imaging methods such as D-OCT are required if accurate ONL measurements are sought.

The ability to measure the ONL and HFL thicknesses independently may be beneficial to understanding normal foveal development, following the natural history of retinal degenerations, and monitoring the effects of therapeutic interventions. Grouping the two layers together results in conclusions that are, in the best-case scenario, imprecise and, in the worst-case scenario, erroneous. Without accounting for HFL, the ONL thickness changes have been reported in numerous disease conditions including outer retinal atrophy, geographic atrophy, and ABCA4 mutations. However, because HFL was not identified and accounted for in these measurements, the true extent of photoreceptor nuclear loss cannot be known. Furthermore, conclusions regarding the extent of retinal damage and remodeling cannot accurately be made without distinguishing the ONL from HFL as these layer thicknesses will vary with eccentricity and may respond differently to cellular injury over time.
Change in the ONL thickness has been reported in ex vivo studies in normal aging individuals as well as in age-related macular degeneration. Changes in the thickness of both the ONL and HFL have also been reported in human and rodent eyes with inherited retinal degenerations. If HFL thickness increases but the ONL is thinning with age or in association with drusen as demonstrated by Curcio et al., the combined ONL and HFL thicknesses may appear relatively constant despite wide variability in each of these components. Such ambiguity is unnecessary now that these layers can be optically differentiated.

Future studies will have to establish the effect of photoreceptor cell death on the thicknesses of the ONL and HFL and the time course over which change happens. While the abundant photoreceptor axons that comprise HFL may undergo anterograde degeneration, the surrounding Müller scaffolding may not. Histological data actually demonstrates hypertrophy of the surrounding cells in response to retinal degeneration, and it is possible that this mechanism alters the visible thickness or appearance of HFL. If focal loss of photoreceptor nuclei occurs, the angle of incidence of the OCT beam relative to HFL may change, thus revealing the true thickness of the ONL in the surrounding areas. However, if there were diffuse loss of the ONL over an area, the orientation would not be expected to change in the same focal manner, therefore, requiring the additional optical contrast that D-OCT provides. Directional OCT may allow disambiguation of optical effects of disease progression versus true anatomical changes.

Henle fiber layer has classically been considered a component of the histological OPL along with the synaptic termini of the photoreceptor cells. While this designation may have been appropriate when histology was the predominant means of assessing HFL and its characteristics, the lumping together of the different components of the OPL is no longer necessary or beneficial in the age of OCT. A plexus, by definition, is a complex intermixing of dendrites and axonal synaptic termini. Henle fiber layer, in contrast, has an oblique, regular, and radially symmetric pattern. Directional OCT allows not only HFL to be visualized as hyperreflective on the contralateral side to the pupil but also hyporeflective on the ipsilateral side, thereby permitting both the anatomical distinction between the ONL externally and the synaptic OPL internally. Given the precision that in vivo imaging can now provide in distinguishing these layers, the disambiguation of the literature by the
assignment of HFL as an independent layer in posterior segment OCT is a welcome improvement.  

Limitations of this study include the use of a cohort of only young healthy normal subjects. Given a goal of this research is the clinical application of D-OCT to patients with all forms of retinal disease, imaging eyes from a full range of ages is a critical next step. An additional limitation includes the fact that a single operator acquired all of the images in this study using a single commercially available OCT system. While it is intuitive that the thickness of the retinal layers should not change between operators, this must be confirmed. In this study, the amount of pupil displacement from the central position varied depending on where the pupil entry position resulting in a normal incidence scan was obtained, which varied between individuals. Indeed, a skilled operator using the altered reflectivity of the HFL and overall scan quality as endpoints for displacement of the pupil entry position was used in this study. Additional studies are underway to determine whether a novice operator can reliably acquire images resulting in the same measurements.

Optical coherence tomography can live up to the promise of in vivo histology only if the optical properties of the tissue are accounted for. The reliance of confocal detection of back-scattered light in serving as the signal and the contrast agent in OCT means that the appearance of images is governed not only by the presence of the tissue itself, but also by its optical properties, including directional reflectivity. Directional OCT takes advantage of this optical property to add additional contrast to the retinal anatomy. The visual display of directional reflectivity content within a single chromatic D-OCT image conveys both optical contrast between retinal layers and provides information about the orientation of cellular components. The application of D-OCT demonstrated in this article would allow more accurate measures of the ONL and HFL in clinical trials. Future studies seeking to report ONL thickness measurements should differentiate between ONL and HFL so that the most accurate anatomical outcomes can be made and key insights into pathology can be revealed.

Key words: spectral domain, Fourier domain, directional reflectivity, outer plexiform layer, Henle’s fiber layer, photoreceptors, cones, rods, biomarker.

References


