Age-Related Differences in Longitudinal Structural Change by Spectral-Domain Optical Coherence Tomography in Early Experimental Glaucoma

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PERSPECTIVE. To characterize age-related differences in the magnitude of spectral-domain optical coherence tomography (SD-OCT) structural change in early experimental glaucoma (EG).

METHODS. Both eyes from four young (1.4–2.6 years) and four old (18.6–21.9 years) rhesus monkeys were imaged at least three times at baseline, and then every 2 weeks after laser-induced, chronic, unilateral IOP elevation until the onset of EG (confocal scanning laser tomographic surface change confirmed twice). Two to 20 weeks after EG onset, animals were euthanized and optic nerve axon counts for all eyes were performed. Masked operators delineated retinal and ONH landmarks in 40 radial B-scans from each eye and imaging session to quantify change from baseline in five SD-OCT neural and connective tissue parameters. The effects of EG, age, and EG × age interactions on the magnitude, rate (magnitude per postlaser time), and IOP responsiveness (magnitude per cumulative IOP insult) of postlaser parameter change were individually assessed using general estimating equation models.

RESULTS. Presac SD-OCT RNFLT and minimum rim width change and postmortem axon loss was not significantly different in old compared with young EG eyes. The rate of change and IOP responsiveness of the parameters anterior lamina cribrosa surface depth relative to Bruch's membrane opening (BMO) and BMO depth relative to peripheral Bruch's membrane were significantly lower (P < 0.05) in the old compared with the young EG eyes.

CONCLUSIONS. At similar postlaser times, levels of cumulative IOP insult and axonal damage, SD-OCT-detected ONH connective tissue structural change is greater in young compared with old monkey EG eyes.

Keywords: glaucoma, optic nerve head, lamina cribrosa, BMO minimum rim width, optical coherence tomography

"Cupping” is a clinical term that is used to describe enlargement of the optic nerve head (ONH) cup in all forms of optic neuropathy.1–3 However, “cupping” also is used as a synonym for the pathophysiology of glaucomatous damage to the ONH neural and connective tissues.4 Because of this link to pathophysiology, it is often suggested that “cupping” is pathognomonic of glaucoma, yet there is much literature that discusses the presence, importance, and meaning of "cupping" in a variety of optic neuropathies, including those of glaucoma, compressive orbital masses, ischemia, inflammation, hereditary disorders and, most recently, primary cerebrospinal fluid pressure lowering.5–9 This is because the optic cup should enlarge if the ONH neuroretinal rim thins due to degeneration and loss of axons in all forms of optic neuropathy regardless of etiology.

What distinguishes glaucoma as an optic neuropathy is not cupping, per se, but rather the type of cupping, which is classically deep; more pronounced within the superior and inferior temporal quadrants; and is associated with excavation, nerve fiber layer (NFL) hemorrhages, and a pattern of retinal ganglion cell (RGC) axon injury and peripapillary retinal NFL (RNFL) loss that leads to characteristic patterns of visual field loss.2,4 This is in contrast to "nonglaucomatous cupping,” which manifests greater pallor and less deformation for a given amount of visual field loss.2,6

However, the clinical appearance of glaucomatous cupping is variable.10 We have previously proposed that in all optic neuropathies, regardless of the location and etiology of the primary insult to the visual system, the clinical phenomenon of cup enlargement (herein referred to as clinical cupping) has two principal pathophysiologic components: "prelaminar thinning” and "ONH connective tissue deformation.”11,12 We define "prelaminar thinning” to be that portion of cup enlargement that results from thinning of the prelaminar tissues, which consist of axonal, astroglial, and vascular...
components. We define “ONH connective tissue deformation” to be that portion of cup enlargement that results from permanent posterior lamina cribrosa deformation and/or remodeling that may be associated with enlargement of the scleral canal and posterior deformation of the scleral flange and peripapillary scleral connective tissues.11–13

We have further proposed that regardless of its contribution to the mechanisms of RGC axonal insult, it is the presence of ONH connective tissue deformation and/or remodeling that underlies and defines a “glaucomatous” form of cupping, and its character and magnitude that determine the cupping “type” (i.e., its glaucoma phenotype)11–11 in a given eye (Fig. 1). With these concepts in mind, it is important to note that with the development of spectral-domain optical coherence tomography (SD-OCT) imaging, the prelaminar and ONH connective tissue components of cupping are beginning to be separately detected and quantified.15,16

From a biomechanical standpoint, the clinical manifestation of glaucomatous damage to a given ONH should depend on a host of neural and connective tissue factors, including eye-specific differences in ONH and peripapillary scleral connective tissue structural stiffness and remodeling.12,17–19 The structural stiffness of a tissue is determined by its geometry (i.e., its thickness and curvature) and its material properties (i.e., the arrangement, amount, and character of its constituents). A growing body of direct and indirect evidence suggests that the structural stiffness of the monkey and human lamina cribrosa and peripapillary scleral connective tissues increases with age.20–22 Practically speaking, this should translate to the aged eye (or stiff eye regardless of age) deforming less for a given level of loading than a young eye (or compliant eye regardless of age).

Two recent cross-sectional studies used SD-OCT lamina cribrosa imaging to report that in human patients with glaucoma or ocular hypertension, the eyes of the younger patients demonstrated a deeper laminar position relative to Bruch’s membrane opening (BMO) when controlled for the level of visual field loss and RNFL loss.20,21 These observations are clinically important because they support the notion that age exerts an important influence on structure/structure relationships (e.g., lamina cribrosa deformation versus RNFLT change) and structure/function relationships (e.g., lamina cribrosa deformation versus visual field change).20,21 They also suggest that age-related differences in ONH connective tissue structural stiffness and/or remodeling may be two mechanisms by which this occurs.

Eye-specific differences in structural stiffness and/or remodeling, regardless of age, should therefore contribute to the glaucomatous phenotype expressed by an individual eye (Fig. 1).13 We recently reported that SD-OCT ONH change detection commonly preceded or coincided with ONH surface change detection by confocal scanning laser tomography (CSLT) in four young and four old monkeys with chronic unilateral experimental glaucoma (EG).15 In this study, SD-OCT ONH change also preceded detection of RNFL thickness (RNFLT) change by SD-OCT: RNFL retardance change by scanning laser polarimetry, and retinal functional change by multifocal electroretinogram.15 In the present report, we analyze the same longitudinal SD-OCT data sets to test the hypothesis that SD-OCT-detected ONH structural change was greater in the four young as compared with the four old monkey EG eyes at similar postlaser time intervals, similar levels of postlaser cumulative IOP insult, and at the onset of CSLT ONH surface change.

**METHODS**

**Acronyms and Abbreviations**

Please see Table 1 for definitions and descriptions of all acronyms and abbreviations.

**Animals**

All experiments were performed in accordance with an institutionally approved animal use protocol at Legacy Health and conformed to the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research. Four young (<3 years old) and four old (>18 years old) rhesus monkeys (Macaca mulatta) were used in the current study.

**Overall Experimental Design**

Both eyes of each animal underwent CSLT and SD-OCT imaging on at least three separate occasions during baseline (i.e., prelaser, when both eyes were normal). One eye of each monkey was then randomly chosen to receive 360° of argon laser photoacoagulation to the trabecular meshwork in two sessions of 180°, 2 weeks apart.32,33 Postlaser imaging sessions in both the lasered (hereafter referred to as the EG) eye and
Age-Related Longitudinal Change in Glaucoma Eyes

Table 1. Full Parameter Names, Acronyms, and Their Descriptions

<table>
<thead>
<tr>
<th>Parameters/Abbreviations</th>
<th>Full Name/Description</th>
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<tbody>
<tr>
<td>BM</td>
<td>Bruch’s membrane</td>
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<tr>
<td>BMO</td>
<td>Bruch’s membrane opening</td>
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<tr>
<td>BMO depth</td>
<td>BMO depth relative to a BM reference plane 1500 μm from BMO centroid</td>
</tr>
<tr>
<td>CSLT</td>
<td>Confocal scanning laser tomography</td>
</tr>
<tr>
<td>Cumulative IOP difference</td>
<td>Cumulative IOP difference between experimental glaucoma eye and control eye</td>
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<tr>
<td>EG</td>
<td>Experimental glaucoma</td>
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<tr>
<td>GEE</td>
<td>Generalized estimating equation</td>
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<tr>
<td>ILM</td>
<td>Internal limiting membrane</td>
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<tr>
<td>IOP</td>
<td>Intraocular pressure</td>
</tr>
<tr>
<td>Laminar surface depth</td>
<td>Anterior lamina cribrosa surface depth relative to BMO</td>
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<tr>
<td>MPD</td>
<td>Mean of all CSLT topographic values within the operator-assigned disc margin</td>
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<td>MRW</td>
<td>Minimum rim width measured from BMO</td>
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<td>ONH</td>
<td>Optic nerve head</td>
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<tr>
<td>Prelaminar tissue thickness</td>
<td>Prelaminar tissue thickness measured from anterior laminar surface to ILM surface</td>
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<tr>
<td>RPE</td>
<td>Retinal pigment epithelium</td>
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<tr>
<td>RNFIT</td>
<td>Retinal nerve fiber layer thickness measured 12° from the disc center</td>
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<tr>
<td>SD-OCT</td>
<td>Spectral-domain optical coherence tomography</td>
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<tr>
<td>TCA</td>
<td>CSLT topographic change analysis</td>
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</table>

Contralateral control eye continued in 1- to 3-week intervals. Laser photocoagulation was repeated in additional 90° increments as necessary to achieve evidence of structural change or to maintain a target of 20 to 30 mm Hg in its absence. Where additional laser was indicated, we chose to perform more rather than less laser in the old eyes so as to maintain the old EG eyes at higher IOPs than the young EG eyes throughout the study. Although our goal was equal IOP exposure in both groups, we sought to minimize risk of undermining the hypothesis that older eyes would deform less for a given IOP exposure by maintaining a bias toward slightly higher IOPs in the old group (i.e., if the older group exhibited less deformation but also lower average IOP levels, the hypothesis test would be confounded). All animals were followed to the onset of CSLT-detected ONH surface change confirmed on two subsequent occasions in the EG eye.

Confocal Scanning Laser Tomography–Detected EG Onset

A trained technician outlined the optic disc margin within the baseline image of each eye using a disc photograph for reference where necessary. This contour line was automatically transferred to all subsequent images in the longitudinal series. For the current study, the parameter mean position of the disc (MPD) was calculated for each CSLT session as described previously. Mean position of the disc was defined as the height of the surface of the ONH (i.e., average height of all pixels located within the disc margin contour line) relative to the height of a standard reference plane (a plane that runs parallel to the peripapillary retinal surface, and is set 50 μm below the retinal surface height at the papillomacular bundle, which is located in the −10° to −4° section temporal of the contour line (HRT II Operating Instructions; software version 1.6)). Mean position of the disc onset was defined as the first postlaser session in which the MPD value was outside of the eye-specific 95% confidence interval (95% CI) determined by the baseline testing sessions, which was then confirmed in the two subsequent imaging sessions. Therefore once three consecutive imaging sessions resulted in MPD outside the confidence interval, the first of those three sessions was defined as MPD onset.

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variability that was confirmed in the two subsequent sessions exceeding the magnitude and position of red TCA super-pixels within the baseline images. Tomographic change analysis onset at a given postlaser imaging session required a clinician to see similar TCA change in the two subsequent sessions. Where necessary, interobserver differences in TCA were adjudicated by the two readers through discussion. For this study, EG onset was CSLT onset determined by either MPD or TCA, whichever occurred earlier (four TCA, two MPD, and two simultaneous).

Animal Euthanization and Axon Count
Seven of the eight animals were euthanized within 2 to 8 weeks post-EG onset (i.e., the time it took to confirm CSLT onset twice). One monkey (O1) was euthanized 20 weeks after retrospective assignment of a CSLT end point that was not appreciated at the time it occurred. Post mortem, optic nerve axon counts for both eyes of each animal were performed using a previously described automated segmentation algorithm that samples 100% of the optic nerve cross-section and counts 100% of the detected axons.40

Spectral-Domain OCT Data Set Delineation
Our methodology for delineation of SD-OCT ONH data sets has been described in detail in prior studies.15,41–43 Customized “Multiview” software (built on the Visualization Toolkit [VTK; Kitware, Inc., Clifton Park, NY, USA]) was used to delineate the following anatomic features within 40 of the 80 radial B-scans (every other scan) of each SD-OCT data set by two trained technicians who were masked to the status of each eye (EG versus control) and time point (baseline or postlaser): internal limiting membrane (ILM), outer boundary of RNFL, Bruch's membrane/retinal pigment epithelium complex (BM/RPE), BMO (innermost termination of the BM/RPE complex), neural boundary (inner border of the neural canal), and anterior lamina cribrosa surface (Fig. 2).

For the ILM, RNFL, BM/RPE, and neural boundary categories, each layer was delineated using discrete points interconnected by a linear Bézier curve. The position of each point in each category was finely adjusted so that the fitted Bézier curve matched the feature of interest as closely as possible. Bruch's membrane opening was delineated using two discrete points at either side of the neural canal. The anterior lamina cribrosa surface was delineated based on previous direct comparisons between SD-OCT B-scans and matched histologic sections,41 recent comparisons to three-dimensional (3D) histomorphometric reconstructions (Zanggalli C, et al. IOVS 2014;55:ARVO E-Abstract 4747), as well as our previous publications on SD-OCT laminar visualization12 and longitudinal change detection.15,43 For each SD-OCT B-scan data set, a point cloud including all the above landmarks was generated (Figs. 3, 4). Because the Spectralis x,y transverse dimensions are calibrated for human eyes, a

FIGURE 2. Original and delineated SD-OCT ONH data sets in a normal monkey eye. Green lines/points: ILM; blue lines/points: outer boundary of the RNFL; orange lines/points: BM/RPE; red points: BMO; purple points: neural boundary; yellow points: anterior lamina cribrosa surface. One of two B-scans was delineated and thus yielded 40 delineated data sets for each monkey eye at a time.
A scaling factor of 0.857 was used to correct the difference between the ocular magnification of the average monkey eye and the human eye assumed by the instrument, as previously described.

Spectral-Domain OCT ONH Connective Tissue, Prelaminar Tissue and RNFLT Parameterization

To parameterize each SD-OCT ONH data set, two reference planes were used (Figs. 3, 4). A BMO reference plane was based...
TABLE 2.
Demographics, IOP Characteristics, and Measures of EG Eye MRW, RFNLT, and Optic Nerve Axon Loss for Each Study Animal

<table>
<thead>
<tr>
<th>Animal ID</th>
<th>Animal Sex</th>
<th>Animal Age, y</th>
<th>IOP C/EG, mm Hg</th>
<th>Prelaminar Tissue Thickness, mm</th>
<th>BMO Depth, mm</th>
<th>BMO Rim Width, mm</th>
<th>RNFLT Change, µm</th>
<th>MRW Change, µm</th>
<th>Optic Nerve Axon Loss, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>V1</td>
<td>M</td>
<td>25.5±4.6</td>
<td>10/9</td>
<td>6/11</td>
<td>12/22</td>
<td>13/28</td>
<td>15/26</td>
<td>25/35</td>
<td>10/11</td>
</tr>
<tr>
<td>V2</td>
<td>M</td>
<td>26.0±7.2</td>
<td>13/10</td>
<td>8/11</td>
<td>15/22</td>
<td>13/28</td>
<td>15/26</td>
<td>25/35</td>
<td>10/11</td>
</tr>
<tr>
<td>V3</td>
<td>M</td>
<td>26.1±4.2</td>
<td>13/10</td>
<td>8/11</td>
<td>15/22</td>
<td>13/28</td>
<td>15/26</td>
<td>25/35</td>
<td>10/11</td>
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<tr>
<td>V4</td>
<td>M</td>
<td>26.1±4.2</td>
<td>13/10</td>
<td>8/11</td>
<td>15/22</td>
<td>13/28</td>
<td>15/26</td>
<td>25/35</td>
<td>10/11</td>
</tr>
</tbody>
</table>

Data Analysis

To estimate the magnitude of IOP insult for each EG eye, we calculated the parameter cumulative IOP difference as the IOP difference between the EG and its control eye on each measurement day multiplied by the number of days from the last measurement and summed over the period of postlaser follow-up (cumIOP, in units of mm Hg × days).

The effect of age and EG on the postlaser data of the eight EG and eight control eyes were analyzed for each parameter in turn using generalized estimating equation (GEE) models. In this analysis, the effects of EG, age and EG × age interactions on IOP, MRW, and RNFLT were derived from the slope coefficients from the GEE model in original scale (days). Intraocular pressure responsiveness (change/cumulative IOP insult) for laminar surface depth, BMO depth, prelaminar tissue thickness, MRW, and RNFLT were also derived from the slope coefficients from the GEE model in original scale (mm Hg × days).

RESULTS

Animal demographics are listed in Table 2. The median age of the young animals was 1.9 years (range, 1.4–2.6) at study initiation and the median length of postlaser follow-up was 33 weeks (range, 26–34 weeks). The median age of the old animals was 20.95 years (range, 18.6–21.9) at study initiation and the median length of postlaser follow-up was 23 weeks (range, 21–39 weeks) (P = 0.34 for postlaser follow-up times, Wilcoxon rank sum test).

Age-Related RNFLT and Orbital Optic Nerve Axon Loss Differences at Euthanization

The percent change of RNFLT and MRW relative to the baseline average for each individual EG eye is shown in Table 2. Median RNFLT change when animals were euthanized was 5.7% (range, −16.8% to 9.7%) in the four young EG eyes and −1.1% on the best-fitting ellipse through the 80 delineated BMO points in a 3D space (2 points/B-scan × 40 B-scans). A BM reference plane was constructed in two steps as follows. First, to identify constituent BM points, a circle within the BMO reference plane 1500 µm from the BMO centroid was identified and perpendiculars were dropped from the circle to the BMO reference plane (Fig. 3). The ONH connective tissue parameters, anterior lamina cribrosa surface depth relative to the BMO reference plane (laminar surface depth), and BMO depth relative to the BM reference plane (BMO depth) are also described in Figure 3. The ONH prelaminar parameters, prelaminar tissue thickness, and BMO minimum rim width (MRW) are described in Figure 4. A standard RNFLT circle scan at 12° angle was also performed (not shown).
TABLE 3: Age-Related Differences in Control and EG Eyes Pre- and Postlaser IOP

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Control Eyes</th>
<th>EG Eyes</th>
<th>EG Eye Axon Loss</th>
<th>EG Eye Associated</th>
<th>Median Control Postlaser IOP</th>
<th>EG Eye Associated Postlaser IOP</th>
<th>Control EG Eye Axon Loss</th>
<th>EG Eye Axon Loss</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peak IOP in Control Eyes, mm Hg</td>
<td>15 (10-13)</td>
<td>8 (8-9)</td>
<td>15.3 (13-15)</td>
<td>15.3 (13-15)</td>
<td>8 (8-9)</td>
<td>15.3 (13-15)</td>
<td>15.3 (13-15)</td>
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<tr>
<td>Mean Postlaser IOP in EG Eyes, mm Hg</td>
<td>15.3 (13-15)</td>
<td>15.3 (13-15)</td>
<td>15.3 (13-15)</td>
<td>15.3 (13-15)</td>
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<tr>
<td>Mean Postlaser IOP in Young EG Eyes, mm Hg</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
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<tr>
<td>Mean Postlaser IOP in Old EG Eyes, mm Hg</td>
<td>15 (10-13)</td>
<td>15 (10-13)</td>
<td>15 (10-13)</td>
<td>15 (10-13)</td>
<td>15 (10-13)</td>
<td>15 (10-13)</td>
<td>15 (10-13)</td>
<td>15 (10-13)</td>
</tr>
<tr>
<td>Mean Prelaminar Tissue Thickness in Control Eyes, mm</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
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</tr>
<tr>
<td>Mean MRW in Control Eyes, mm</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
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<tr>
<td>Mean MRW in Young EG Eyes, mm</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
</tr>
<tr>
<td>Mean MRW in Old EG Eyes, mm</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
</tr>
<tr>
<td>Mean Control Eye Postlaser IOP, mm Hg</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
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</tr>
<tr>
<td>Mean EG Eye Postlaser IOP, mm Hg</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
<td>10 (10-13)</td>
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<td>10 (10-13)</td>
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<td>10 (10-13)</td>
</tr>
</tbody>
</table>

Mean IOP and SD were based on Wilcoxon rank sum test. Bold text indicates <P = 0.05, which is statistically significant.

Age-Related IOP Differences in Control and EG Eye Postlaser IOP

Median control eye postlaser IOP was slightly higher in the old (10.5 mm Hg, range, 9–14 mm Hg) compared with the young (9 mm Hg, range, 8–9 mm Hg) eyes (P = 0.07, Wilcoxon rank sum test) (Table 3). Experimental glaucoma eye peak postlaser IOP was greater in the old (median 39.5 mm Hg, range, 38–42 mm Hg) compared with the young eyes (median 30.5 mm Hg, range, 28–37 mm Hg) (P = 0.03, Wilcoxon rank sum test). Although mean and median EG eye cumulative IOP insult at the CSLT onset, and pre-euthanization sessions were greater (nearly doubled) in the old compared with young EG eyes, the difference did not achieve significance.

Age-Related Differences in the Magnitude of Parameter Change at CSLT Onset

By GEE analysis, at CSLT onset in the EG eye, the effect of EG was significant for SD-OCT laminar surface depth, prelaminar tissue thickness, and MRW in the young animals but achieved significance only for SD-OCT laminar surface depth and MRW in the old animals (Table 4). Among all parameters, an EG x age interaction effect was significant for laminar surface depth only.

Age-Related Differences in the Rate of Postlaser Parameter Change

Although the mean postlaser IOP, peak postlaser IOP, and cumulative IOP insult were considerably lower in the young EG eyes (Table 5), the rate of posterior (outward) laminar surface deformation was 4.5 times faster in young (–0.460 µm/d) compared with old (–0.102 µm/d) EG eyes (P < 0.0001, GEE). Deformation of BMO relative to a peripheral BM reference plane (BMO depth) also demonstrated a significantly faster rate of postlaser change in the young (–0.109 µm/d) compared with the old EG eyes (–0.003 µm/d, P < 0.0001, GEE). In fact, the rate in old EG eyes was not significantly different from old control eyes for this parameter. Postlaser prelaminar tissue thickness significantly increased in young eyes (0.088 µm/d) and decreased in old eyes (–0.040 µm/d, P < 0.0001, GEE).

Age-Related Differences in the IOP Responsiveness of Parameter Change

Because the rate of postlaser parameter change versus time (in days, Table 5) does not take into account the magnitude of IOP exposure, the magnitude of parameter change was also evaluated per unit of cumulative IOP insult (mm Hg × day), as reported in Table 6. For a given increase in cumulative IOP insult, the anterior laminar surface deformed posteriorly at magnitudes that were 3.64 times faster (P < 0.0001, GEE) in the young (–0.237 µm/mm Hg × day) compared with the old EG eyes (–0.065 µm/mm Hg × day). Bruch’s membrane...
Table 4. Age-Related Differences in the Magnitude of Parameter Change at CSLT-Detected EG Onset

<table>
<thead>
<tr>
<th>SD-OCT Parameters</th>
<th>Laminar Surface Depth, μm</th>
<th>BMO Depth, μm</th>
<th>Prelaminar Tissue Thickness, μm</th>
<th>MRW, μm</th>
<th>RNFLT, μm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean ± SD changea in young EG eyes (EG – Control)</td>
<td>−95 ± 36</td>
<td>−9 ± 5</td>
<td>26 ± 34</td>
<td>−39 ± 30</td>
<td>3.6 ± 7.0</td>
</tr>
<tr>
<td>Mean ± SD changea in old eyes (EG – Control)</td>
<td>−22 ± 10</td>
<td>−7 ± 6</td>
<td>−17 ± 12</td>
<td>−24 ± 10</td>
<td>−0.2 ± 4.0</td>
</tr>
<tr>
<td>P value EG: young effect†</td>
<td>0.0001</td>
<td>0.4742</td>
<td>0.0262</td>
<td>0.0007</td>
<td>0.8758</td>
</tr>
<tr>
<td>P value EG: old effect†</td>
<td>&lt;0.0001</td>
<td>0.1074</td>
<td>0.2824</td>
<td>0.0065</td>
<td>0.2424</td>
</tr>
<tr>
<td>P value EG × age interactions‡</td>
<td>0.0008</td>
<td>0.8302</td>
<td>0.0896</td>
<td>0.3252</td>
<td>0.3281</td>
</tr>
</tbody>
</table>

Positive change
- Anterior to BMO: Thickening
- Anterior to PeriBM: Thickening

Negative change
- Posterior to BMO: Thinning
- Posterior to PeriBM: Thinning

a Mean ± SD change: difference between the changes from the mean baseline value to CSLT onset of the EG eye and control eye for the four young and four old animals. Bold text indicates P < 0.05, which is statistically significant.
† GEE test if mean change above is significantly different between EG and control eyes at CLST onset in young or old monkeys, respectively.
‡ GEE test if there is interaction between eye condition (EG or control) and age (young and old animals) on changes in parameters at CSLT onset.

opening depth (3.39 times more posterior deformation, P = 0.0106) also demonstrated age-related differences in IOP responsiveness.

Change from baseline data at each postlaser imaging session is plotted for a subset of parameters relative to the imaging session cumulative IOP insult in Figure 5. In general, it is evident that the young eyes were followed to lower levels of cumulative IOP insult than the old eyes, in that the young eye data points (red dots) end at less than 600 mm Hg × day, whereas the old eye data points (blue dots) extend to more than 1200 mm Hg × day. These data again indicate that CSLT-detected ONH surface change occurred (and was confirmed on two subsequent occasions) at lower levels of cumulative IOP insult in the young EG eyes.

DISCUSSION

This is the first study to longitudinally characterize age-related differences in the character of glaucomatous “cupping” at its onset, that is, at the conversion from experimental ocular hypertension to ONH structural change detected by SD-OCT. In these four young and four old animals, the magnitude of SD-OCT ONH parameter change was greater in the young compared with the old eyes both at the point of CSLT-detected ONH surface change and also when measured as continuous variables relative to postlaser time (in days) and postlaser cumulative IOP insult (in mm Hg × days). The data also suggest that structural change was greater for deep connective tissues in the young eyes.

Regarding the ONH deep connective tissues, when compared by age group versus time or versus cumulative IOP insult, lamina cribrosa deformation relative to BMO (laminar surface depth) and BMO deformation relative to peripheral BM (BMO depth) were both greater in young compared with old monkey eyes. We believe that two mechanisms likely underlie the age-related differences in EG ONH connective tissue deformation we report: age-related differences in ONH connective tissue structural stiffness13 and age-related differences in ONH connective tissue remodeling.12 Although the two can be thought of as separate mechanisms, they are also linked.

The findings of our current study offer indirect rather than direct evidence that increased structural stiffness is a contributing mechanism to age-related differences in ONH structural change in early EG. We did not perform acute IOP

Table 5. Age-Related Differences in the Rate of Parameter Change in EG Eyes

<table>
<thead>
<tr>
<th>SD-OCT</th>
<th>Anterior Laminar Depth, μm/d</th>
<th>BMO Depth, μm/d</th>
<th>Prelaminar Tissue Thickness, μm/d</th>
<th>MRW, μm/d</th>
<th>RNFLT, μm/d</th>
</tr>
</thead>
<tbody>
<tr>
<td>Change/day† in young EG eyes</td>
<td>−0.460</td>
<td>−0.109</td>
<td>0.088</td>
<td>−0.202</td>
<td>−0.004</td>
</tr>
<tr>
<td>Change/day† in old EG eyes</td>
<td>−0.102</td>
<td>−0.003</td>
<td>−0.040</td>
<td>−0.099</td>
<td>−0.010</td>
</tr>
<tr>
<td>Change rates young vs. old EG eyes, P value‡</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
<td>0.1433</td>
<td>0.4901</td>
</tr>
<tr>
<td>Change rates in EG vs. control eyes: young animals, P value§</td>
<td>&lt;0.0001</td>
<td>0.0002</td>
<td>0.2013</td>
<td>0.0029</td>
<td>0.7957</td>
</tr>
<tr>
<td>Change rates in EG vs. control eyes: old animals, P value</td>
<td></td>
<td></td>
<td>0.0001</td>
<td>0.6175</td>
<td>0.0186</td>
</tr>
</tbody>
</table>

Positive change
- Anterior to BMO: Thickening
- Anterior to PeriBM: Thickening

Negative change
- Posterior to BMO: Thinning
- Posterior to PeriBM: Thinning

a Rates of change were derived from the slope coefficients from the GEE model for each SD-OCT parameter. Bold text indicates P < 0.05, which is statistically significant.
† The longitudinal change rate of each parameter represents how much the parameter changed for each unit of time (units in day) relative to mean baseline. The data reported here are the mean value for the n = 4 old and n = 4 young EG eyes.
‡ P value for whether the rate of parameter change is significant between young and old EG eyes.
§ P value for whether the rate of parameter change in the EG eye is significantly different from in control eyes in young animals.
|| P value for whether the rate of parameter change in the EG eye is significantly different from in control eyes in old animals.
10/30 compliance testing\(^4\) on the young and old animals of this report at baseline (in our laboratory, acute IOP compliance testing refers to obtaining SD-OCT ONH imaging 30 minutes after IOP is manometrically lowering to 10 mm Hg and again 30 minutes after increasing IOP to 30 mm Hg). Therefore, we do not have direct evidence that the old eyes of this report were structurally stiffer than the young eyes before the onset of unilateral chronic IOP elevation. However, we recently reported that SD-OCT–detected ONH structural change is greater in a separate group of young versus old normal monkey eyes following acute IOP elevation (Qin L, et al. IOVS 2013;54:ARVO E-Abstract 53).

We have previously reported that ONH connective tissue remodeling in early monkey EG includes outward migration of the lamina cribrosa insertions\(^1\) and retrolaminar septal recruitment of orbital optic nerve septa into more transversely

<table>
<thead>
<tr>
<th>SD-OCT Parameters, (\mu\text{m}/(\text{mm Hg} \times \text{day}))</th>
<th>Laminar Surface Depth(^{\ast})</th>
<th>BMO Depth</th>
<th>Prelaminar Tissue Thickness</th>
<th>MRW</th>
<th>RNFLT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Change/CumIOP(\dagger) in young EG eyes</td>
<td>-0.237</td>
<td>-0.035</td>
<td>0.0156</td>
<td>-0.152</td>
<td>-0.022</td>
</tr>
<tr>
<td>Change/CumIOP(\dagger) in old EG eyes</td>
<td>-0.065</td>
<td>-0.010</td>
<td>-0.0143</td>
<td>-0.048</td>
<td>-0.004</td>
</tr>
<tr>
<td>(P) value for change/CumIOP in young EG eyes(\ddagger)</td>
<td>&lt;0.0001</td>
<td>0.0001</td>
<td>0.5794</td>
<td>0.0216</td>
<td>0.0554</td>
</tr>
<tr>
<td>(P) value for change/CumIOP in old EG eyes(\ddagger)</td>
<td>&lt;0.0001</td>
<td>0.0015</td>
<td>0.0001</td>
<td>&lt;0.0001</td>
<td>0.0722</td>
</tr>
<tr>
<td>(P) value for difference in rates of change(\S)</td>
<td>&lt;0.0001</td>
<td>0.0106</td>
<td>0.2911</td>
<td>0.1177</td>
<td>0.1302</td>
</tr>
</tbody>
</table>

\(\ast\) Rates of change were derived from the slope coefficients from the GEE model for each SD-OCT parameter. Bold text indicates \(P < 0.05\), which is statistically significant.

\(\dagger\) The longitudinal change rate of each parameter represents how much the parameter changed for each unit of cumulative IOP insult relative to mean baseline. The data reported here are the mean value for the \(n = 4\) old and \(n = 4\) young EG eyes.

\(\ddagger\) \(P\) value for whether the rate of parameter change is significant in the young or old EG eyes.

\(\S\) \(P\) value for the comparison of the rate of parameter change within the young versus the old EG eyes.

\(10/30\) compliance testing\(^4\) on the young and old animals of this report at baseline (in our laboratory, acute IOP compliance testing refers to obtaining SD-OCT ONH imaging 30 minutes after IOP is manometrically lowering to 10 mm Hg and again 30 minutes after increasing IOP to 30 mm Hg). Therefore, we do not have direct evidence that the old eyes of this report were structurally stiffer than the young eyes before the onset of unilateral chronic IOP elevation. However, we recently reported that SD-OCT–detected ONH structural change is greater in a separate group of young versus old normal monkey eyes following acute IOP elevation (Qin L, et al. IOVS 2013;54:ARVO E-Abstract 53).

We have previously reported that ONH connective tissue remodeling in early monkey EG includes outward migration of the lamina cribrosa insertions\(^1\) and retrolaminar septal recruitment of orbital optic nerve septa into more transversely

![Figure 5](https://example.com/figure5.png)

**Figure 5.** Change from baseline for selected testing modalities and parameters at each postlaser testing session in the young (red) and old (blue) EG eyes plotted relative to cumulative IOP insult. Testing sessions are ordered by EG eye cumulative IOP insult (bottom of each column). Change from baseline for each parameter at each postlaser testing session is plotted for all four young (red dots) and all four old (blue dots) EG eyes. Note the following. First, in general, the young eyes were followed to lower levels of cumulative IOP insult than the old eyes (red dots end at less than 600 mm Hg \(\times\) day and blue dots extend to more than 1200 mm Hg \(\times\) day), reflecting the fact that ONH surface change as detected by CSLT occurred at lower levels of cumulative IOP insult in young eyes. Second, age-related differences in the overall rates of change are apparent qualitatively for most of the parameters and were confirmed as statistically significant for a subset, as reported in Tables 5 and 6.
Age-related differences in ONH remodeling, if present, may be multifactorial. Inherent differences in the ONH of young versus old eyes, such as astroglial and peripapillary scleral fibroblast responsiveness to stretch and/or less connective tissue deformation and/or basement membrane strain, in stiffer, aged eyes at similar levels of mechanical load may also contribute.

The ONH rim tissue parameter MRW showed a tendency to decrease more in young compared with old EG eyes versus time or versus cumulative IOP insult (although neither comparison achieved significance), which suggests the ONH rim tissue underwent greater compression and/or thinning in young eyes. Interestingly, even though the lamina bowed more posteriorly and the rim thinned more in young eyes, the prelaminal tissue thickness actually increased in young eyes (but thinned in old eyes) over time and cumulative IOP insult (with only the comparison versus time achieving significance). These data support our previous report of prelaminal tissue thickening within 3D histomorphometric reconstructions of the EG versus control eyes of three young-adult animals with early EG. Age-related differences in this finding and its clinical importance need to be studied in larger groups of animals. Optic nerve head rim tissue and prelaminal tissue change (whether detected by MRW measurements made relative to BMO or thickness measurements made relative to the anterior lamina cribrosa surface) may represent RGC axonal loss or degenerative changes that could include both increased and decreased axon caliber, astroglial loss, hypertrophy and/or proliferation, and vascular constriction or engorgement. Although all of these changes may represent varying degrees of pathophysiologic response, it is also possible that all (except RGC axonal loss) are potentially reversible responses to conformational change that follow anterior or posterior lamina cribrosa deformation and/or expansion or contraction of the scleral canal and BMO. Because axon loss was not greater in the young eyes, we believe that both the greater MRW thinning and prelaminal tissue thickening they demonstrate reflect either conformational changes or early degenerative changes that precede frank axonal loss. The nature and clinical significance of conformational change requires further clarification.

Not all structural change results in RGC injury lead to a change in visual function. Optic nerve head conformational change that is not due to RGC axon loss may not result in visual field progression. However, it is important to note that six of the eight EG eyes in this report (three young and three old), had 12% to 29% optic nerve axon loss by postmortem histology. Thus in those six EG eyes, although there was no detectable RNFLT (~8.3% to 9.7% change), RGC axon loss was modest to moderate in magnitude. It is also important to note that one old (O2) and one young (Y4) EG eye demonstrated similar SD-OCT ONH changes to those observed in the other EG eyes while demonstrating minimal orbital optic nerve axon loss (~2% and ~4%, respectively) by postmortem histology. Additional studies in the eight EG eyes of this report are under way to correlate SD-OCT ONH sectoral change to adjacent SD-OCT RNFL thinning and colocalized optic nerve axon loss so as to identify those forms of SD-OCT ONH and RNFL parameter change that most consistently and closely correlate to RGC axon injury.

The fact that the old animals experienced similar or higher IOPs at the point of CSLT-detected ONH surface change is important because it suggests that if our study is biased by IOP exposure, it was biased to detect more, rather than less, deformation in the old eyes. The mean postlaser IOP, the mean postlaser peak IOP, and the cumulative IOP insult at CSLT onset were either significantly higher in the old compared with the young EG eyes, or not significantly different. Where possible, we chose to maintain the old EG eyes at higher IOPs than the young EG eyes, so as to achieve a bias toward more rather than less deformation in the old EG eyes. Our IOP data suggest we achieved this bias, although measurements were made only weekly, whereas continuous telemetric IOP characterization would have been necessary to most accurately quantify IOP insult.

The fact that the final measurement of SD-OCT peripapillary RNFLT demonstrated minimal change when using standard peripapillary RNFLT circle scan data in both young and old EG eyes and that postmortem, optic nerve axon loss was not detectably greater in these four old compared with these four young EG eyes is important because it is not consistent with the concept that old eyes are more susceptible than young eyes to RGC axon loss at all levels of IOP. We have previously outlined the logic and literature that suggests aged eyes should be more susceptible to glaucomatous RGC axon damage than young eyes. This finding is unexpected and needs to be confirmed in a larger study.

Interpretation of our results should be limited by the issues we have mentioned as well as the following additional considerations. All parameters in this report were calculated on a global basis, which may diminish their ability to detect focal and regional change. In particular, because the major ONH blood vessels most commonly shadow the superior and inferior quadrants, the superior and inferior sectors of the lamina cribrosa (where early change in monkey and human glaucoma is expected) are less robustly represented in our SD-OCT ONH connective tissue parameters. This effectively gave more weight to the nasal and temporal quadrants when lamina cribrosa-associated parameters were averaged for a global calculation. Regionalization of each structural parameter relative to an anatomically consistent, foveal-BMO nasal temporal axis will be the focus of the next report in this series.

In summary, our findings suggest that at similar levels of cumulative IOP insult and at the onset of CSLT ONH surface change, SD-OCT detected ONH prelaminar and connective tissue structural change is greater in young compared with old monkey eyes. These data support the concept that age-related differences in ONH connective tissue structural stiffness and/or remodeling may contribute to age-related differences in the appearance of early glaucomatous cupping in a given eye. Our findings also suggest that although SD-OCT-detectible ONH change that precedes detectible RNFL change was associated with minimal optic nerve axon loss in two eyes, it was associated with modest to moderate (12% to 29%) axon loss in the majority of monkey eyes.

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