Optical Coherence Tomography Angiography of Optic Disc Perfusion in Glaucoma

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Purpose: To compare optic disc perfusion between normal subjects and subjects with glaucoma using optical coherence tomography (OCT) angiography and to detect optic disc perfusion changes in glaucoma.

Design: Observational, cross-sectional study.

Participants: Twenty-four normal subjects and 11 patients with glaucoma were included.

Methods: One eye of each subject was scanned by a high-speed 1050-nm—wavelength swept-source OCT instrument. The split-spectrum amplitude-decorrelation angiography (SSADA) algorithm was used to compute 3-dimensional optic disc angiography. A disc flow index was computed from 4 registered scans. Confocal scanning laser ophthalmoscopy (cSLO) was used to measure disc rim area, and stereo photography was used to evaluate cup/disc (C/D) ratios. Wide-field OCT scans over the discs were used to measure retinal nerve fiber layer (NFL) thickness.

Main Outcome Measures: Variability was assessed by coefficient of variation (CV). Diagnostic accuracy was assessed by sensitivity and specificity. Comparisons between glaucoma and normal groups were analyzed by Wilcoxon rank-sum test. Correlations among disc flow index, structural assessments, and visual field (VF) parameters were assessed by linear regression.

Results: In normal discs, a dense microvascular network was visible on OCT angiography. This network was visibly attenuated in subjects with glaucoma. The intra-visit repeatability, inter-visit reproducibility, and normal population variability of the optic disc flow index were 1.2%, 4.2%, and 5.0% CV, respectively. The disc flow index was reduced by 25% in the glaucoma group (P = 0.003). Sensitivity and specificity were both 100% using an optimized cutoff. The flow index was highly correlated with VF pattern standard deviation (R² = 0.752, P = 0.001). These correlations were significant even after accounting for age, C/D area ratio, NFL, and rim area.

Conclusions: Optical coherence tomography angiography, generated by the new SSADA, repeatedly measures optic disc perfusion and may be useful in the evaluation of glaucoma and glaucoma progression. Ophthalmology 2014;121:1322-1332 © 2014 by the American Academy of Ophthalmology.

Glucoma is the second leading cause of blindness in the United States. Although elevated intraocular pressure (IOP) is a risk factor, more than one half of patients with glaucoma actually have IOP levels within the normal range at their first visit. There is a growing body of evidence suggesting that glaucoma pathogenesis is related to vascular dysfunction. More recently, prospective trials have demonstrated that optic disc hemorrhage and peripapillary atrophy are both associated with accelerated glaucoma progression. These findings may support a role for focal ischemia of the optic disc as a causative factor for glaucoma at least in some patients, either by itself or in conjunction with elevated IOP.

However, the lack of measurement tools has hampered the development of a clinical test for optic disc perfusion. Optical coherence tomography (OCT) is commonly used in clinical settings for the diagnosis and management of glaucoma. Doppler OCT has been used to obtain precise measurements of total retinal blood flow calculated from the Doppler frequency shift of backscattered light. Although Doppler OCT is appropriate for large vessels around the disc, it is not sensitive enough to measure accurately the low velocities of small vessels that make up the disc microcirculation.

We recently developed a method of measuring local circulation using high-speed OCT to perform quantitative angiography. With the use of the split-spectrum amplitude-decorrelation angiography (SSADA) algorithm, flow in the optic disc can be quantified. The purpose of this study was to investigate optic disc perfusion in glaucoma with OCT angiography, determine the relationship of flow index measurements with traditional measures of function and structure, and assess whether the SSADA-based optic disc flow index can measure vascular changes associated with glaucoma.

Methods

Study Population

This study was performed at the Casey Eye Institute at Oregon Health & Science University. The research protocols were approved by the institutional review board at Oregon Health & Science University and carried out in accordance with the tenets of...
the Declaration of Helsinki. Written informed consent was obtained from each subject after an explanation of the nature of the study.

The normal subjects and subjects with early glaucoma were part of the Advanced Imaging for Glaucoma study. One eye from each subject was scanned and analyzed. Perimetric glaucoma (PG) eyes exhibited glaucomatous visual field (VF) loss and optic disc changes, such as disc rim defects or nerve fiber layer (NFL) defects, on ophthalmoscopy. Pre-perimetric glaucoma (PPG) eyes showed similar changes to the optic disc as glaucomatous eyes while maintaining normal or borderline VF tests. Standard eye examinations and VF tests were performed on both eyes of the normal subjects. Normal VFs have a normal mean deviation (MD) (P > 0.05), pattern standard deviation (PSD) (P > 0.05), and Glaucoma Hemifield Test (P > 0.03). Glaucomatous VF loss was defined as abnormal PSD (P < 0.05) or Glaucoma Hemifield Test (P < 1%) in a consistent pattern on both qualifying VF examinations. Borderline VFs met neither normal nor PG criteria.

**Optical Coherence Tomography**

A swept-source--based Fourier-domain OCT system described in previous publications was used to obtain images for the quantification of optic disc perfusion. The system used a commercially available swept laser centered at 1050 nm (Axsun Technologies, Inc, Billerica, MA) with a 100 nm tuning range and operated at a 100 kHz repetition rate. This provided an axial resolution of 5.3 μm and an imaging range of 2.9 mm in tissue. Imaging of the optic disc was performed with an average laser output power of 1.9 mW, consistent with safety limits set by the American National Standards Institute.

**Image Acquisition and Processing**

Each subject underwent pupil dilation with 1% tropicamide and 2.5% phenylephrine eye drops before examination. Once the pupils were dilated, subjects were seated in front of the OCT scanner, and their heads were stabilized with the aid of both a supporting chinrest and a forehead rest. Subjects were directed to focus their gaze on the internal fixation target, and a real-time en face view was used by the operator to visualize the imaging area on the fundus.

The scan pattern used was optimized for the SSADA. Each set of scans comprised four 3×3-mm images of the optic disc obtained from 32,000 axial line scans. Two images in each set were captured in the x-fast configuration, with B-scans acquired rapidly along the x-axis. Each pass of the scan beam captured 200 axial lines forming a single B-scan. Eight B-scans were obtained at each fixed y-axis location before the process was repeated at the next sequential y-location. A total of 200 sampling locations were covered along a 3-mm region of the y-axis. At a capture speed of 455 B-scans per second, the 1600 B-scans were acquired within 3.5 seconds. The remaining 2 images in each set were captured in the y-fast configuration. Although the data points remained the same between the y-fast and x-fast configurations, the order in which the data were collected differed. In the y-fast configuration, B-scans were acquired rapidly along the y-axis rather than the x-axis, as was performed in the x-fast configuration.

In each scan, the SSADA was used to distinguish vessels from static tissue. As seen in real-time OCT reflectance images, the amplitude of signal returning from nonstatic tissue varies rapidly over time. By calculating the decorrelation of signal amplitude from consecutive B-scans, a contrast between static and nonstatic tissue is created that allows for the visualization of blood flow. The faster the blood particles move across the laser beam, the higher is the decorrelation of the detected signals within a velocity range set by the scan parameters. Thus, decorrelation is approximately linear to flow velocity, that is, the distance traveled by red blood cells flowing across the light beam per unit time. However, beyond a saturation velocity that is defined by the time interval between consecutive OCT B-scans, the decorrelation increases more slowly with velocity and eventually reaches an upper boundary.

Eye motion causes 2 types of artifacts in SSADA. First, motion between consecutive B-scans at the same nominal position causes decorrelation that can appear as flow. Second, motion between B-scan positions distorts the transverse position of scans along the slow scan axis. To correct the first type of motion error, B-scans with very large (saccadic) bulk motion artifacts were removed. Furthermore, decorrelation due to bulk tissue motion was calculated by histogram analysis and subtracted from each cross-sectional SSADA frame. To correct the second type of motion artifact, we used an image registration algorithm that registered 4 orthogonal raster scanned volumes. Motion correction was first performed on the structural OCT data. The motion correction algorithm generated 3-dimensional (3D) displacement fields that map A-scans from the input volumes into a common motion-corrected space. The same displacement fields were applied to the decorrelation (flow) data to produce motion-corrected flow data volumes. Flow data from 4 input volumes were weighted and merged, improving the signal-to-noise ratio in the flow signal and reducing the flow measurement variation due to local flow changes caused by the cardiac cycle.

Cross-sectional registered reflectance intensity images and flow images were summarized and viewed as an en face maximum projection. The disc boundary for each subject was manually delineated along the neural canal opening using the OCT reflectance images of normal (Fig 1A1, B1) and PG (Fig 1A2, B2) eyes. The boundaries were then transferred to the OCT angiogram map (Fig 1C1, C2) for disc region segmentation. The disc flow index was defined as the average decorrelation value within the disc, given by

\[
\int \frac{D \cdot VdA}{A} = (V = 1, \text{ if vessel}; V = 0, \text{ if not}),
\]

where D was the decorrelation value acquired by SSADA. V was 1 when the decorrelation value was above background; otherwise, V was 0. Thus, the flow index is a dimensionless parameter between 0 and 1. Because of the nonlinear relationship between decorrelation and flow velocity, the flow index mainly measured the area (or caliber) of large vessels and both the area (or vessel density) and velocity of capillaries.

To clearly present the 3D nature of OCT angiography, the 3D angigram was separately projected onto en face views (Fig 1E–G) in 3 layers. The segmentation was based on the detection of the highest gradient magnitude in OCT reflectance for specific tissue interfaces (edges). The retinal layer was defined from the internal limiting membrane to the interpolated Bruch’s membrane. The choroidal layer was defined as the layer 180 μm below the Bruch’s membrane. The scleral layer was the region below the choroidal layer (Fig 1H1, H2).

**Repeatability and Reproducibility**

Intra-visit repeatability of the disc flow index was calculated from a subset of normal subjects with 3 sets of scans performed
within a single visit. Each scan set, consisting of 2 x-fast and 2 y-fast scans, was obtained within 15 seconds of each other. The coefficient of variation (CV) was calculated by comparing 3 measurements obtained at the same location by a single operator.

The same subset of normal subjects used to calculate intra-visit repeatability were used to calculate inter-visit reproducibility obtained from 3 sets of scans performed on 3 separate visits. All scans were obtained within the timeframe of 1 year. The CV was determined from measurements made by a single operator and obtained on separate visits of the subjects.

**Visual Field Testing**

Visual field tests to determine PSD and MD were performed with the Humphrey Field Analyzer II (Carl Zeiss Meditec, Inc, Dublin, CA) set for the 24-2 threshold test, size III white stimulus, and standard Swedish Interactive Threshold Algorithm.

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**Figure 1.** Disc photographs (A1, A2), optical coherence tomography (OCT) reflectance (B1, B2), whole-depth OCT angiograms (C1, C2, en face maximum projection), and cross-sectional angiograms (D1, D2, overlaying on OCT reflectance in gray scale) in the right eye of a normal subject (A1—H1) and the left eye of a perimetric glaucoma (PG) subject (A2—H2). Disc margins are marked by the black elliptical outlines (B1, B2, C1, C2). The position of the cross-section is shown by dotted blue lines (C1, C2). A dense microvascular network was visible on the OCT angiography of the normal disc (C1). This network was greatly attenuated in the glaucomatous disc (C2). To appreciate the ability of OCT angiography to detect blood flow within the various vascular beds, the 3-dimensional angiograms were separately projected into en face maximum projection in 3 layers, that is, retinal angiograms (E1, E2), choroidal angiograms (F1, F2), and scleral/lamina cribrosa angiograms (G1, G2). The boundaries used for segmentation are indicated by dotted green lines on cross-sectional OCT reflectance (H1, H2).
Structural Analysis

The disc rim area for each subject was measured by confocal scanning laser ophthalmoscopy (cSLO) (HRT II, Heidelberg Engineering, GmbH, Dossenheim, Germany). An experienced glaucoma specialist graded vertical and horizontal cup/disc (C/D) ratio on stereo photography. The C/D area ratio was calculated by multiplying vertical and horizontal ratios. Peripapillary retinal NFL thickness was measured from a 3D volumetric scan of 8×8 mm on the same swept-source OCT system used for OCT angiography. The NFL thickness was then averaged from a circular profile of 3.4-mm diameter centered on the disc.

Statistical Analysis

Linear regression analysis was used in the normal group to investigate whether the measurement of disc flow index was affected by age, body mass index (BMI), mean ocular perfusion pressure (MOPP), or IOP. Wilcoxon rank-sum tests were used to compare average values of measurements between normal and glaucoma eyes. Univariate regression analysis was then used to determine the relationships between disc flow index and traditional measures of function and structure, such as the VF MD, VF PSD, C/D area ratio, rim area, NFL thickness, and IOP in the glaucoma group. Because VF values for MD and PSD were reported in logarithmic decibel scale, values for disc flow index, C/D area ratio, rim area, and NFL thickness were converted to logarithmic decibel scale by 10 × log10 [value/average value of the normal group)] to improve the correlation linearity and strengthen the correlation coefficient values. Multivariate linear regression was performed after log transformation of the variable into decibel units relative to the normal reference average. This model was used to analyze the effect of optic disc perfusion on the VF PSD test and vice versa while controlling for several other independent variables, including age, C/D area ratio, rim area, and NFL thickness. All statistical analyses were performed with MATLAB version R2010b (The MathWorks Inc, Natick, MA).

Results

Disc perfusion was studied in 24 normal subjects and 11 subjects with glaucoma (Table 1). The mean age in the normal group, 52±10 years, was 16 years less than in the glaucoma group. The glaucoma group consisted of 8 PG and 3 PPG eyes. Most of the glaucoma group had mild disease, in which 6 of the 11 subjects had stage 0 to 1 according to Glaucoma Staging System 2.23 There were no significant differences in BMI, diabetes mellitus, systemic hypertension, or the use of systemic antihypertensive medications between the 2 groups. There were also no significant differences in IOP, diastolic blood pressure, systolic blood pressure, or MOPP between the 2 groups. Data collection was complete except for cSLO and disc photography from 1 PPG subject.

The CV for intra-visit repeatability and inter-visit reproducibility of the disc flow index was 1.2% and 4.2%, respectively, based on measurements from 4 normal subjects. The inter-subject variability among the 24 normal subjects was 5.0% CV. There were no correlations of age, BMI, MOPP, or IOP with the disc flow index in the normal group. Three normal subjects (12%) were taking systemic antihypertensive medications: 1 (4%) taking a diuretic and 2 (8%) taking an angiotensin-converting enzyme inhibitor. The use of any systemic antihypertensive medication or any subclass did not have a significant effect on the disc flow index as determined by the Mann—Whitney U test. Subjects with glaucoma were receiving at least 1 ocular antihypertensive eye drop: a prostaglandin in 10 (91%), a beta-blocker in 8 (73%), a carbonic anhydrase inhibitor in 4 (36%), and an alpha-2 agonist in 1 (9%). One subject (9%) used no drops, 2 subjects (18%) used 1 drop, and 8 subjects (73%) used >1 drop. The use of an ocular antihypertensive eye drop or any subclass was not correlated with the disc flow index as determined by the Mann—Whitney U test.

The OCT angiography scan provided detailed 3D image sets for both disc structure (reflectance) and perfusion (flow). Example images of normal and PG eyes are shown in Figure 1. The color fundus photographs and OCT en face reflectance summation images both showed discs with moderate degrees of peripapillary atrophy. The maximum projection en face angiograms showed that normal discs had a denser microvascular network (Fig 1C1, note dense microvascular network temporally) compared with the glaucomatous disc (Fig 1C2). The disc flow indices were computed by averaging decorrelation values within the disc margin in the en face whole-depth OCT angiograms. The disc margin (ellipses in Fig 1) could be determined from the en face structural OCT images, with the aid of color disc photographs. Cross-sectional OCT angiograms were constructed by overlaying blood flow (color scale) on reflectance (gray scale). These cross-Sections showed that OCT angiography detected flow from all depths of the disc, from the inner surface to the lamina cribrosa. There appeared to be reduced flow in the lamina region of the glaucomatous disc compared with the normal disc (Fig 1D1, D2).
The 3D OCT angiograms also could be projected within separate layers (Fig 1E–G). In examining these layered projections, one needs to keep in mind that large retinal vessels cast shadows on the tissue below. At the same time, flow from large superficial vessels also can be projected onto highly reflective tissue below, creating a flow projection artifact. This can occur because of variation in the shadowing effect due to the transit of blood particles. Thus, large vessels in the retinal layer can be seen again in deeper layers as shadows or flow. Despite these artifacts, distinct patterns were present in the layered en face projection angiograms. In the retinal layer, the superficial disc vasculature blended seamlessly into the retinal vascular network. The retinal and superficial disc vascular networks were dense in the normal eye (Fig 1E1), but attenuated in the glaucomatous eye (Fig 1E2). The peripapillary choriocapillaries were nearly confluent and dominated the choroidal layer angiograms (Fig 1F1, F2), but the disc circulation was relatively low in this layer. The sclera had relatively low flow, and much of that was probably projected from the choroid above (Fig 1G1, G2). The lamina cribrosa had a dense vascular network in the normal eye (Fig 1G1) and a mildly attenuated network in the glaucomatous eye (Fig 1G2). On the basis of all of the 3D information, the impression was that glaucoma attenuated flow both in the microvascular network of the superficial disc and in the deeper lamina cribrosa. The large retinal vessels appeared to be relatively unchanged in caliber.

The disc flow index in the glaucoma group was 25% lower ($P = 0.003$) compared with the normal group (Fig 2, Table 2). Sensitivity and specificity were both 100% using a flow index cutoff value of 0.1515 (Fig 2). As expected, the glaucoma group also had significantly worse VF, rim area, C/D area ratio, and NFL thickness (Table 2).

In the glaucoma group, univariate regression analysis showed that the disc flow index was significantly correlated with VF PSD and rim area, but not with VF MD, C/D area ratio, or NFL thickness (Table 3). The rim area was significantly correlated with VF PSD, VF MD, C/D area ratio, and NFL thickness. The IOP was not significantly correlated with any other factors (data not shown).

In the normal group, there was no correlation between the disc flow index and age, VF PSD, rim area, C/D area ratio, or NFL thickness (Fig 3). The glaucoma group tended to be older and had larger PSD and C/D ratio, smaller rim area, and thinner NFL. In the glaucoma group, the disc flow index was significantly correlated with VF PSD and rim area (Fig 3, Table 3). Of note, there was no overlap in flow index between the glaucoma and normal groups, but there was much overlapping between the 2 groups for cSLO rim area, C/D area ratio, and NFL thickness.

In the multivariate analysis where the flow index was considered as the dependent variable (Table 4), VF PSD was the dominant explanatory variable, accounting for approximately 75% of the variance ($R^2$). Age, C/D area ratio, rim area, and NFL thickness were not significant explanatory variables when grouped with VF PSD in the multivariate models. This showed that disc perfusion was more strongly linked to VF PSD than any disc structural parameters. The PSD parameter was chosen to summarize VF function because it is a more specific diagnostic parameter than MD in early glaucoma.24,25

In the multivariate analysis where the dependent variable was VF PSD (Table 5), the flow index was the dominant explanatory variable, accounting for approximately 75% of the variance ($R^2$). Age, C/D area ratio, and rim area were not significant explanatory variables when grouped with the disc flow index in the multivariate models. Although NFL thickness was a significant explanatory variable, the flow index was the dominant one, accounting for more than 5 times the variance ($R^2$) in VF PSD as NFL thickness. This suggests that the flow index is a relatively strong indicator of glaucoma severity.

![Figure 2](image.png)

**Figure 2.** Box plot showing the disc flow index in normal and glaucoma groups. The median (line inside the box), interquartile range (box), and whole range of values (whiskers) are shown. This plot shows that the 2 groups are completely separated. The minimum of the normal group is 0.1516, and the maximum of the glaucoma group is 0.1513, indicating sensitivity and specificity are both 100% using a cutoff value of 0.1515.

### Table 2. Results of Diagnostic Testing

<table>
<thead>
<tr>
<th>Variables</th>
<th>Normal</th>
<th>Glaucoma</th>
<th>P Value*</th>
</tr>
</thead>
<tbody>
<tr>
<td>VF</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MD (dB)</td>
<td>0.20±0.87 (−1.59 to 1.87)</td>
<td>−3.28±4.12 (−13.35 to 0.08)</td>
<td>0.003</td>
</tr>
<tr>
<td>PSD (dB)</td>
<td>1.43±0.20 (1.10−1.86)</td>
<td>4.44±3.12 (1.59−9.07)</td>
<td>0.003</td>
</tr>
<tr>
<td>Structural assessments</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>cSLO rim area (mm²)</td>
<td>1.55±0.34 (0.35−2.26)</td>
<td>1.13±0.25 (0.71−1.56)</td>
<td>0.007</td>
</tr>
<tr>
<td>C/D area ratio</td>
<td>0.11±0.10 (0.01−0.44)</td>
<td>0.37±0.17 (0.08−0.59)</td>
<td>0.012</td>
</tr>
<tr>
<td>NFL thickness (μm)</td>
<td>107.9±19.9 (90.8−122.6)</td>
<td>82.0±17.0 (58.9−107.6)</td>
<td>0.000</td>
</tr>
<tr>
<td>Disc perfusion</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Flow index</td>
<td>0.161±0.008 (0.15−0.18)</td>
<td>0.121±0.026 (0.07−0.15)</td>
<td>0.003</td>
</tr>
</tbody>
</table>

C/D = cup/disc; cSLO = confocal scanning laser ophthalmoscopy; MD = mean deviation; NFL = nerve fiber layer; PSD = pattern standard deviation; VF = visual field.

Numbers displayed are mean ± population standard deviation (range).

*Wilcoxon rank sum.
In this study, we reported the first use of OCT angiography to quantify human disc perfusion in glaucoma. Optical coherence tomography angiography with SSADA has many properties that make it useful for clinical evaluation. First, it is a noninvasive technique that does not require the injection of any exogenous dye or contrast agent. Second, it provides 3D visualization of the optic nerve head vasculature from the disc surface to the lamina cribrosa. Third, it provides near-automated quantification of disc perfusion for diagnostic evaluation.

Preliminary validation of the diagnostic utility of the disc flow index was shown by the significant differences between normal and glaucoma eyes. Furthermore, there was a high correlation of the flow indices to visual functions, and low intra-visit, inter-visit, and inter-subject variability.

Optical coherence tomography angiography is not the first technique to be used to evaluate disc perfusion in glaucoma. Disc blood flow of patients with glaucoma was previously investigated by fluorescein angiography (FA). Prolonged arteriovenous passage times have been demonstrated in patients with primary open-angle glaucoma and normal pressure glaucoma. Fluorescein filling defects in the disc have been found in glaucomatous and ocular hypertensive patients. Fluorescein angiography studies have shown focal sector hypoperfusion of the optic disc in patients with low-tension glaucoma and diffuse disc hypoperfusion in patients with chronic simple glaucoma. However, FA is not commonly used to monitor glaucoma because of its invasive nature and difficulty in quantification. Unlike FA, SSADA is a noninvasive technique that relies on the decorrelation of OCT signal amplitude reflected from nonstatic tissue. This allows for the quantification of flow that can be used to monitor disc perfusion while avoiding potential side effects of nausea and anaphylaxis associated with dye injection.

Laser Doppler flowmetry (LDF) and laser speckle flowgraphy (LSFG) are 2 other noninvasive techniques that were reported to measure disc perfusion. With the use of single-point LDF, Piltz-Seymour et al and Hamard et al similarly reported decreased blood flow in the disc of glaucoma and glaucoma suspects when compared with normal subjects. With the use of scanning LDF, Michelson et al reported that both neuroretinal rim blood flow and peripapillary retinal blood flow were significantly decreased in patients with glaucoma compared with controls. Hafez et al also found significantly lower blood flow in the disc for patients with open-angle glaucoma compared with normal patients and suggested perfusion might be reduced before the manifestation of VF defects. By using LSFG, Yokoyama et al found that the mean blur rate for the entire optic disc was significantly lower in the glaucoma group than in the control group. Sugiyama et al reported that less blood flow was observed with LSFG at the superior and inferior sectors of the disc rim in patients with PPG compared with normal control subjects. Overall, our findings agree with previous results that disc perfusion is reduced in glaucomatous eyes. More important, OCT angiography with SSADA offers greater intra-visit repeatability (1.2% CV) and inter-visit reproducibility (4.2% CV) than LSFG or LDF. With LSFG, CVs for intra-visit repeatability ranged from 1.9% to 11.9%, and inter-visit reproducibility was 12.8%. Laser Doppler flowmetry was even less reliable, with CVs for intra-visit repeatability of 6.6% to 21.2% and inter-visit reproducibility of 25.2% to 30.1%. Inter-subject variability in the normal population was reduced when flow was assessed with SSADA-based OCT angiography (5.0%) compared with LSFG (23.2%–33%) or LDF (43.5%–47.1%).

A technical reason that SSADA has advantages over LDF is its relative insensitivity to variations in the average intensity of signal reflected from the tissue of interest, which could be affected by intrinsic tissue reflectance or beam attenuation due to defocus, media opacity, pigment absorption, and scattering in overlying tissue. In LDF, the variance of the speckle pattern is inherently proportional to average intensity of the reflected signal. In SSADA, the effect of signal intensity is effectively canceled because decorrelation is proportional to the variance divided by the average signal intensity. This has been demonstrated clinically: The SSADA-based average decorrelation in 3 normal subjects varied within ±0.8% CV, even though the average signal intensity in the same images varied by ±36.7% CV. Thus, on the basis of both theoretic considerations and clinical data, SSADA-based optic disc flow index is little affected by absorbance and reflectance of disc tissue, unlike LDF perfusion measurements. The reduced signal strength dependence by OCT angiography

| Table 3. Correlation Coefficient Matrix among Visual Field, Blood Flow, and Structural Variables in Subjects with Glaucoma |
|----------------|----------------|----------------|----------------|----------------|----------------|
| Variables (dB) | Disc Flow Index | VF MD | VF PSD | C/D Area Ratio | cSLO Rim Area |
| VF MD | 0.138 (0.146) | | | | |
| VF PSD | 0.752 (0.001) | 0.563 (0.006) | | | |
| C/D area ratio | 0.048 (0.273) | 0.307 (0.048) | 0.201 (0.097) | | |
| cSLO rim area | 0.397 (0.026) | 0.724 (0.001) | 0.638 (0.003) | 0.371 (0.031) | |
| NFL thickness | 0.004 (0.853) | 0.508 (0.014) | 0.121 (0.294) | 0.138 (0.289) | 0.408 (0.047) |

C/D = cup/disk; cSLO = confocal scanning laser ophthalmoscopy; MD = mean deviation; NFL = nerve fiber layer; PSD = pattern standard deviation; VF = visual field.

All variables were converted to dB scale by 10×log10 (value/average value of the normal group). Table cells display Pearson’s R² (P value to test |R| = 0). Statistically significant correlations (P < 0.05) are in bold.
allows for cleaner inter-individual comparison of flow index.

With OCT angiography, the principal finding of the present study is the correlation between disc flow index and VF PSD ($R^2 = 0.752, P = 0.001$), which is suggestive of a link between reduced perfusion of the disc and severity of glaucoma. Multiple regression models (Tables 4 and 5) suggest that there is a strong link between VF PSD and disc flow index that is not mediated by disc structural variables, such as C/D area ratio, rim area, and NFL thickness. This suggests that the flow index provides independent information on glaucoma severity that is not
All variables were converted to dB scale by 10

\[
\frac{C/D}{L} = \frac{L_3}{L_5} \quad \text{PSD}
\]

Most of the population with glaucoma in error, and dry eye, and therefore less reliable in early VF MD and PSD may be that the MD is easily affected by nonglaucomatous factors, such as cataract, refractive error, and dry eye, and therefore less reliable in early glaucoma.24,25 Most of the population with glaucoma in our study had early disease; therefore, PSD may be a more reliable measure of their disease severity than MD. Another possibility for the better correlation of the flow index with PSD is that they both measure segmental or focal glaucomatos changes. In contrast, overall NFL thickness average and rim area are global measures that may correlate better with MD. Another explanation is that the discrepant correlations with MD and PSD are due to random variation because of the small sample size in our study. Thus, our findings need confirmation from a larger independent study.

**Study Limitations**

There are several limitations associated with OCT angiography. First, unlike Doppler OCT, which provides absolute volumetric flow in microliters/minute, OCT angiography only yields a flow index in arbitrary units. However, the precision of the disc flow index measurement with OCT angiography is better compared with dual-circular Doppler OCT scanning to measure the total retinal blood flow, which has an intra-visit repeatability of 10.5%.29 Second, flow projection artifact from superficial blood vessels to deeper tissue levels prevents us from separately measuring superficial and deep optic nerve head flow. The artifact is caused by the moving shadow cast by flowing blood cells. Decorrelation is caused by both moving reflectors (e.g., blood cells) and moving shadows (e.g., projections on distal high reflectance tissue). These 2 types of decorrelation are not distinguished by SSADA, and both appear as flow in the 3D angiogram. The artifact is not problematic if our analysis is confined to the 2-dimensional maximum projection angiogram. Therefore, the study was limited to the use of 2-dimensional angiograms that measured superficial and deep vascular beds together. Third, the disc flow index includes measurements on both the local disc circulation and the large retinal blood vessels. Thus, it is a mixture of both disc and retinal circulations and not a pure measurement of a single vascular bed. However, because of the velocity saturation effect,15,21 OCT angiography, at current scan speeds, cannot measure flow, but it can measure the caliber of large vessels (Fig 1E1, E2), suggesting that the angiography-based flow index mainly detects changes in disc microvasculature. The velocity or the blood flow in large retinal vessels should be measured using Doppler techniques.15,50 Fourth, OCT angiography requires the use of a high-speed OCT system. In our study, we used a custom swept-source OCT device that operated at an axial scan rate of 100 kHz. Commercial spectral-domain OCT devices typically operate at an axial scan rate of 20 to 40 kHz. However, recent advancements have led to the production of commercial spectral OCT devices operating at speeds of up to 70 kHz (RTVue XR, Optovue, Inc, Fremont, CA; Cirrus HD-OCT 5000, Carl Zeiss Meditec, Dublin, CA), as well as a commercial 100 kHz swept-source OCT device (DRI OCT-1 Atlantis, Topcon Corp, Tokyo, Japan). In addition, laboratory

| Table 4. Multivariate Regression Models of Factors Affecting Disc Flow Index in Subjects with Glaucoma |
|---|---|---|---|---|---|---|---|---|
| Model | Variable (dB) | Slope (P Value) | R² | Model | Variable (dB) | Slope (P Value) | R² |
| 1 | PSD | -0.319 (0.001) | 0.75 | Age | -0.067 (0.828) | 0.001 | 0.751 |
| 2 | PSD | -0.312 (0.002) | 0.75 | C/D area ratio | 0.088 (0.306) | 0.037 | 0.789 |
| 3 | PSD | -0.367 (0.002) | 0.752 | cSLO rim area | 0.207 (0.585) | 0.012 | 0.764 |
| 4 | PSD | -0.383 (0.013) | 0.752 | NFL thickness | -0.388 (0.086) | 0.097 | 0.799 |
| 5 | PSD | -0.332 (0.000) | 0.702 |

C/D = cup/disc; cSLO = confocal scanning laser ophthalmoscopy; NFL = nerve fiber layer. All variables were converted to dB scale by 10\*log10 (value/average value of the normal group). Significant slope coefficients (P < 0.05) are in bold.

| Table 5. Multivariate Regression Models of Factors Affecting Visual Field Pattern Standard Deviation in Subjects with Glaucoma |
|---|---|---|---|---|---|---|---|
| Model | Variable (dB) | Slope (P Value) | R² | Model | Variable (dB) | Slope (P Value) | R² |
| 1 | Disc flow index | -2.353 (0.001) | 0.75 | Age | 0.373 (0.649) | 0.006 | 0.756 |
| 2 | Disc flow index | -2.254 (0.002) | 0.75 | C/D area ratio | 0.295 (0.138) | 0.071 | 0.823 |
| 3 | Disc flow index | -2.123 (0.002) | 0.752 | cSLO rim area | -1.303 (0.056) | 0.106 | 0.858 |
| 4 | Disc flow index | -1.567 (0.013) | 0.752 | NFL thickness | -1.167 (0.047) | 0.121 | 0.823 |
| 5 | Disc flow index | -2.405 (0.000) | 0.702 |

C/D = cup/disc; cSLO = confocal scanning laser ophthalmoscopy; NFL = nerve fiber layer. All variables were converted to dB scale by 10\*log10 (value/average value of the normal group). Significant slope coefficients (P < 0.05) are in bold.
OCT prototypes of multi-megahertz speed have been reported. With the use of these faster next-generation commercial OCT units, OCT angiography may become a feasible technique for clinical evaluation of glaucoma.

A notable limitation of our study is that we cannot rule out the effect of glaucoma and blood pressure medications on the disc flow index. Most patients in our glaucoma group were receiving multiple ocular antihypertensive eye drops. Therefore, it is not possible to determine their individual effects on disc perfusion with our small sample size, and we cannot entirely rule out the possibility that the glaucoma drops could somehow be responsible for the reduced disc perfusion. We consider this unlikely for 2 reasons. The first is the strong correlation between flow index and VF PSD that indicates perfusion is tightly linked to disease severity. The second is that previous studies have shown that select glaucoma eye drops, including prostaglandins, and carbonic anhydrase inhibitors, improved disc and retinal perfusion or had no significant effect. We plan to perform regional (i.e., superior and inferior hemispheres) correlation between VF and disc blood flow in a future study to remove entirely the confounding effect of global parameters, such as medications, from consideration.

In conclusion, we used OCT angiography based on the SSADA to measure human disc perfusion in vivo. We demonstrated that OCT angiography can detect reduced disc perfusion in a group of patients with early glaucoma with 100% sensitivity and specificity. This reduction in flow index is not a by-product of rim loss or cupping in glaucomatous eyes, and we were able to establish a strong link between the disc flow index and the VF PSD. Our data warrant further studies to determine whether a lower flow index is correlated with the rate of progression and whether disc perfusion can be used as a prognostic indicator for disease course. Flow index values might be used to help determine which ocular hypertensive subjects and other glaucoma suspects require treatment. We anticipate that future investigations will lead to an enhanced understanding of this disease and improved treatment strategies.

References


Footnotes and Financial Disclosures

Originally received: June 5, 2013.
Final revision: January 14, 2014.
Accepted: January 16, 2014.

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Financial Disclosure(s):
The author(s) have made the following disclosure(s): Oregon Health & Science University, Y.J., and D.H. have a significant financial interest in Optovue, Inc, a company that may have a commercial interest in the results of this research and technology. These potential conflicts of interest have been reviewed and managed by Oregon Health & Science University. J.G.F. and D.H. receive royalties on an OCT patent licensed by the Massachusetts Institute of Technology to Carl Zeiss Meditec and LightLab Imaging. M.F.K. and J.G.F. receive royalties from intellectual property owned by the Massachusetts Institute of Technology and licensed to Optovue, Inc. The other authors have no proprietary or commercial interest in any materials discussed in this article.

Supported by National Institutes of Health Grant 1R01 EY023285-01, Rosenbaum’s P30EY010572, Clinical and Translational Science Award Grant UL1TR000128, an unrestricted grant from Research to Prevent Blindness, R01-EY11289-26 and Air Force Office of Scientific Research (AFOSR) FA9550-10-1-0551, German Research Foundation DFG-HO-1791/11-1, DFG-GSC80-SAOt, and DFG Training Group 1773.

Abbreviations and Acronyms:
BMI = body mass index; C/D = cup/disc; cSLO = confocal scanning laser ophthalmoscopy; CV = coefficient of variation; FA = fluorescein angiography; IOP = intraocular pressure; LDF = laser Doppler flowmetry; LSFG = laser speckle flowgraphy; MD = mean deviation; MOPP = mean ocular perfusion pressure; NFL = nerve fiber layer; OCT = optical coherence tomography; PG = perimetric glaucoma; PPG = pre-perimetric glaucoma; PSD = pattern standard deviation; SSADA = split-spectrum amplitude-decorrelation angiography; 3D = 3-dimensional; VF = visual field.

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