Optical Detection of Early Damage in Retinal Ganglion Cells in a Mouse Model of Partial Optic Nerve Crush Injury

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PURPOSE. Elastic light backscattering spectroscopy (ELBS) has exquisite sensitivity to the ultrastructural properties of tissue and thus has been applied to detect various diseases associated with ultrastructural alterations in their early stages. This study aims to test whether ELBS can detect early damage in retinal ganglion cells (RGCs).

METHODS. We used a mouse model of partial optic nerve crush (pONC) to induce rapid RGC death. We confirmed RGC loss by axon counting and characterized the changes in retinal morphology by optical coherence tomography (OCT) and in retinal function by full-field electroretinogram (ERG), respectively. To quantify the ultrastructural properties, elastic backscattering spectroscopic analysis was implemented in the wavelength-dependent images recorded by reflectance confocal microscopy.

RESULTS. At 3 days post-pONC injury, no significant change was found in the thickness of the RGC layer or in the mean amplitude of the oscillatory potentials measured by OCT and ERG, respectively; however, we did observe a significantly decreased number of axons compared with the controls. At 3 days post-pONC, we used ELBS to calculate the ultrasound marker (D), the shape factor quantifying the shape of the local mass density correlation functions. It was significantly reduced in the crushed eyes compared with the controls, indicating the ultrastructural fragmentation in the crushed eyes.

CONCLUSIONS. Elastic light backscattering spectroscopy detected ultrastructural neuronal damage in RGCs following the pONC injury when OCT and ERG tests appeared normal. Our study suggests a potential clinical method for detecting early neuronal damage prior to anatomical alterations in the nerve fiber and ganglion cell layers.

Keywords: ganglion cells, spectroscopy, optical coherence tomography, electroretinography, axonal degeneration

Optic neuropathy (or retinal neuropathy) is often characterized by the loss of or damage to the optic nerves that leads to retinal ganglion cell (RGC) death and vision impairments.1,2 One of the earliest recognized manifestations of optic neuropathy is the ultrastructural alteration in the axon or the body of RGCs.3 It has been reported that a myriad of molecular alterations observed in early-stage RGC/axon damage result in ultrastructural changes, including DNA fragmentation, axon disruption, and mitochondria swelling, among others.3,4 Although beyond the optical diffractive limit, these ultrastructural alterations collectively change the optical-scattering properties in tissue. Thus, optical detection that is sensitive to ultrastructural alterations in the nerve fiber layer (NFL)/RGC layer can be an effective method for identifying the early onset of optic neuropathy.

Elastic light backscattering spectroscopy (ELBS) has demonstrated extraordinary sensitivity to ultrastructural alterations at a length scale of tens of nanometers in tissue without actually resolving the nanoarchitecture.5–8 Elastic light backscattering spectroscopy has been used to detect various cancers at early stages, even in “histologically normal” precancerous lesions.9,10 The measured spectroscopic biomarkers were found to be associated with some ubiquitous structural hallmarks in cancers such as enlarged nuclear sizes, chromatin compaction, and extracellular matrix crosslinking.11,12 Using backscattering light to detect structural abnormalities in retina, Huang et al.13,14 pioneered the work showing that wavelength-dependent scattering changes occur in NFL in elevated intraocular pressure (IOP) rat and murine models, and revealed that the optical changes were linked to cytoskeleton distortion.

In this study we investigated whether ELBS can be used to detect earlier biomarkers of NFL/RGC damage compared with other morphologic and functional methods/techniques. We induced rapid RGC death by partial optic nerve crush injury in mice.15 The damage to the optic nerve was verified by the gradual decrease of axon density. The consequential functional and morphologic changes were quantified by ERG and optical coherence tomography (OCT). The spectroscopic markers were quantified by analyzing the wavelength-dependent scattering signals measured by confocal reflectance microsco-
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py. We found that the spectroscopic changes in the backscattered light can be observed at the very early stage of RGC death, prior to the detectable functional and morphologic alterations.

**Materials and Methods**

All animal procedures were approved by the institutional animal care and use committee at Northwestern University and conformed with the guidelines on the Use of Animals from the National Institutes of Health and the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research.

**Partial Optic Nerve Crush (pONC) Procedure**

First, C57BL/6j mice (Jackson laboratory, Bar Harbor, ME, USA) were anesthetized with an intraperitoneal injection of ketamine (100 mg/kg; Lloyd, Inc., Shenandoah, IA, USA) and xylazine (10 mg/kg; Butler Schein Animal Health, Dublin, OH, USA). The surgery of the partial optic nerve crush was as described in Puyang et al.15 In brief, a small incision was made in the superior and lateral conjunctiva, and a gentle blunt dissection was performed to expose the optic nerve using forceps (Dumont #5B, WPI, Sarasota, FL, USA). The optic nerve was partially clamped with self-closing forceps (WPD) for 3 s/site approximately 0.5 to 1 mm behind the globe. The superior and temporal area was thus defined as the crush-operated (CO) side, and the inferior and nasal area as the nonoperated (NO) side.

**Optical Coherence Tomography**

The system has been described in detail in previous publications.16,17 Briefly, we used a super continuum light source (SuperK EXW-6; NKT Photonics, Birkerød, Denmark) to provide visible-light illumination from 520 to 620 nm, which permitted a higher axial resolution than OCT systems operating within near-infrared light spectral range. The illuminating light was delivered by an optical fiber, collimated into a cube beam splitter (BS, CM1-BS013; Thorlabs, Inc., Newton, NJ, USA), by which the light was divided into a sample arm and a reference arm. The sample arm consisted of a two-dimensional scanning mirror and a 1:5 telescope system. A line scan CCD camera (sp2k; Basler, Wetzlar, Germany) mounted on a glass slide with the same condition. The normalized spectra were fitted with a power law function based on a spectral model of \( J(D-1) \), as illustrated in Figure 1C. The modeling is described in detail in the next section.

**Confocal Reflectance Microscopy**

Mice were perfused with 4% paraformaldehyde fixation and eyes were then enucleated. A blue mark was made on the superior quadrant to indicate the orientation. The anterior chamber, ocular lens, and vitreous were carefully removed. The exposed eye cup was immersed in PBS for 15 minutes. The retina was then carefully removed from the choroid and flat-mounted on a glass slide with PBS as the mounting medium. Confocal reflectance microscopic images were taken by a microscope system (Leica SP5; Leica Microsystems, Inc., Wetzlar, Germany). The focus was set on to the NFL with the depth of focus \( \sim 6 \) μm, which is determined by the pinhole size (Fig. 1A). The illumination wavelength swept from 470 to 670 nm by 10-nm intervals, resulting in 21 wavelength-dependent images (Fig. 1B). The spectral fluctuation in the scattering intensity was compensated by taking a reflectance image from a clear glass surface with the same condition. The normalized spectra were fitted with a power law function based on a spectral model of \( J(D-1) \), as illustrated in Figure 1C. The modeling is described in detail in the next section.

**Analytical Model for ELBS Analysis**

Macromolecules are the fundamental building blocks of living tissue based on the Gladstone-Dale equation, the tissue’s refractive index (RI) is proportional to the local macromolecular mass density.21,22

\[
\beta = n_B + \rho \gamma
\]

where \( \rho (\text{g/mL}) \) is the local mass density of the solid material (e.g., macromolecules), \( \gamma (\text{mL/g}) \) is the RI increment, usually approximately 0.17 mL/g for biological materials23,24; and \( n_B \) is the RI of water. The heterogeneous distribution of mass density causes RI fluctuation, which leads to detectable backscattered light due to elastic scattering.

Because of the heterogeneous nature of the mass density distribution, we modeled tissue as a random medium with continuously fluctuating mass density. The most comprehensive way to describe such a medium is by its correlation function (i.e., mass density correlation function).25-27 We used a three-parameter functional family, the Whittle-Matérn (WM) function, to quantify the correlation functional form.28 This versatile functional family covers essentially all of the commonly used correlation functions, including the power law, exponential, stretched exponential, and the Gaussian function. The functional family of WM is formulated as11:

\[
B_n(r_d) = A_n \left( \frac{r_d}{L_n} \right)^{(D-3)/2} K_{(D-3)/2} \left( \frac{r_d}{L_n} \right),
\]

where \( K_{(D-3)/2} \) is a modified Bessel function of the second type, \( A_n \) is the amplitude of the RI fluctuation, \( D \) is the “shape factor” determining the type of the function,27 and \( L_n \) is the length scale of the correlation function whose exact meaning depends on \( D \). When \( 0 < D < 3 \), the correlation function can be described by a power law and the tissue is organized as a fractal.25 Given the linear relationship between RI and mass
density, \(D\) is the mass density fractal dimension \(D_m\). When \(3 < D < 4\), the functional form becomes a stretched exponential function. For \(D = 4\), \(B_n(r_n)\) is an exponential function, where \(B_n = B_0(0)\). When \(D\) approaches infinity, the function turns into a Gaussian form.

The first-order Born approximation can be used to predict the scattering power spectral density \(\Phi\) by the Fourier transform of \(B_n(r_n)\), \(\Phi(\kappa) = \text{FT}(B_n)\). Then, the differential cross-section per-unit volume \(\sigma(\theta, \phi)\) can be analytically expressed, where \(\theta\) and \(\phi\) denote the scattering angle in a spherical coordinate with respect to the incident direction. Since confocal reflectance detects the backscattered light, the light intensity is proportional to the backscattering coefficient \(\mu_b\). The conditioned function of \(\mu_b\) is \(\Phi_0\):

\[
    \mu_b = A_0 \sigma \left( \frac{\lambda}{\kappa} \right) \Gamma \left( \frac{D}{2} \right) L^3_{\text{D}} k^{4-D}, \quad kL_\mu \gg 1, \quad (3)
\]

where \(\Gamma\) is the gamma function. According to Equation 3, the backscattering spectrum follows \(\mu_b \Phi_0\) where the exponent becomes \(4 - D\) with respect to wavelength. By fitting the spectrum with a power law function, \(D\) can be inversely calculated.

**Axon Counting**

A cross-section of the proximal optic nerve was prepared as described previously. In brief, the optic nerves were fixed in glutaraldehyde in resin (EOPN-Araldite; Electron Microscopy Sciences, Hatfield, PA, USA) and cut into 1-\(\mu\)m sections. The sections were then stained with p-phenylenediamine (Sigma-Aldrich Corp., St. Louis, MO, USA) and imaged under an inverted microscope (Carl Zeiss Microscopy LLC, Thornwood, NY, USA). The axon numbers per image were counted using ImageJ, averaged, and then normalized to the total area of the optic nerve.

**Statistics**

We used two sample student’s \(t\)-tests to calculate the statistical significance. For electroretinogram, 2-way ANOVA was used to examine the influence of two independent variables (time point and light intensity). For spectroscopic analysis from the confocal measurements, the paired \(t\)-test was used to calculate the statistical difference between the operated eye and the fellow eye from the same subjects.

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**Results**

We used a pONC model to induce rapid RGC and axon loss. First, we evaluated the retinal morphological changes by in vivo OCT (Figs. 2A–C). We took OCT images from the mouse retinas before crush, and at 3 days and 1 week postcrush (\(n = 5\) for each group). The thickness of NFL + GCL was measured separately in selected regions from the inferior and nasal side (referred as the nonoperated side) and the crush-operated (CO) side in high-resolution cross-sectional OCT images (Figs. 2A, 2B). At 3 days post-pONC injury, no significant decrease in NFL + GCL thickness was observed compared with the control mice (\(n = 5, P > 0.05\) in student’s \(t\)-test; Fig. 2C). At 1 week postcrush injury, the thickness of NFL + GCL decreased in both areas (NO: 15 ± 1 \(\mu\)m, CO: 14 ± 1 \(\mu\)m, \(n = 5, P < 0.001\) in student’s \(t\)-test; Fig. 2C). Interestingly, damage was found in the entire retina (i.e., both CO and NO areas). No significant difference between NO and CO was observed in NFL + GCL thickness (\(P > 0.05\) in student’s \(t\)-test, Fig. 2C).

We performed scotopic full-field ERG recordings to examine functional changes during the first week following the crush injury. Our data suggested the mean amplitude of OPs, which reflects the inner retinal function, was not significantly reduced (\(n = 5\) in each group, \(P < 0.001\) in 2-way ANOVA with Tukey’s posttest; Fig. 2D) until 1 week postcrush. At 3 days postcrush, we only observed the trend of decrease in the OPs, but the change was not significant (\(n = 5, P > 0.05\) in 2-way ANOVA with posttest; Fig. 2D). These results demonstrated a robust reduction in visual responses from the inner retina at 1 week postcrush injury, consistent with our morphologic findings.

We next examined the degeneration of optic nerve at 3 days postcrush in fixed tissues. The optic nerve samples from the controls and the mice at 3 days postcrush were dissected, fixed, and cryosectioned (see “Methods” section). In the cross-sections of optic nerves from the control mice, axons appeared as dark circles of myelin surrounding unstained axoplasm (Fig. 2E). We observed the disruption of the myelin sheath, hyperdense axoplasm, and vacuolization in the images taken from 3 days postcrush groups. In addition, glial reaction was activated and resulted in numerous large, lipid-containing glial scars (Fig. 2E). We counted the total number of axons from both groups. For the control eyes, the mean axon number was 46,465 ± 1358 (\(n = 9\)). For the 3 days postinjury group, the total axon number was substantially reduced to 28,990 ± 1930 (\(n = 4, P < 0.001\) in student’s \(t\)-test; Fig. 2F). Together our data suggest there is a significant reduction in axon number, though...
the change may not be detected by in vivo OCT and ERG at 3 days postinjury. Having characterized the morphologic and functional changes in retina, we then investigated whether the optical spectroscopic marker can be a more sensitive indicator of NFL/RGC damage. We calculated the values of \( D \) (i.e., the “shape factor” defining the functional type of the mass density autocorrelation function) within 3 days postinjury, when neither NFL/RGC thickness nor ERG showed significant differences between injured and control eyes. Figure 3A shows an example of the averaged spectra from the injured (left) and control (right) eyes. The injured retina has a reflectance spectrum that decreases faster at longer wavelengths, indicating a lower \( D \), and also has more fragmented ultrastructures. When we overlaid the \( D \) values onto the grayscale confocal reflectance images, as shown in Figure 3B, it exhibited overall lower \( D \) 3 days postcrush compared with the control retina. It also appeared that \( D \) was lower near the optic nerve head, indicating that the damage was propagating from the optic disk. Statistically, the reduction of \( D \) is significant (control: \( D = 4.0 \pm 0.2 \); CO: \( D = 3.56 \pm 0.3 \), \( n = 10 \) animals, \( P = 0.001 \) paired student’s \( t \)-tests), as shown in Figure 3C. We also

FIGURE 2. Characterization of retinal damage by OCT, ERG, and axon counting post-pONC injury. (A–C) Imaging of OCT detected a significant reduction in the NFL+GCL thickness at 1 week but not 3 days post-pONC injury. (A) Example of en face projection of 3D OCT from a mouse retina in vivo. Areas of CO (red) and NO (blue) were labeled. I, inferior; N, nasal; S, superior; T, temporal. Scale bar: 0.4 mm. (B) Representative cross-sectional OCT images of mouse retinas. ELM, external limiting membrane; INL, inner nuclear layer; IPL, inner plexiform layer; IS/OS, inner segment/outer segment; ONL, outer nuclear layer; OPL, outer plexiform layer. (C) The thickness of NFL + GCL before pONC, 3 days and 1 week post-pONC. Areas of NO and CO were measured separately post-pONC; \( n = 5 \) mice for each group. *** \( P < 0.001 \) in student’s \( t \)-tests. (D) Mean amplitudes of OPs before crush, at 3 days and 1 week post-pONC. Example of a filtered OP was shown as an inset. *** \( P < 0.001 \) in student’s \( t \)-tests. (E) Representative micrographs of optic nerve sections in a control and a 3-day postcrush eye. (F) Total axon numbers were quantified (\( n = 9 \) in control, \( n = 4 \) in 3 days). *** \( P < 0.001 \) in Student’s \( t \)-tests.

FIGURE 3. Ex vivo ultrastructural quantification of early NFL damage. (A) Examples of reflectance spectrum from a left (ONC) and right (control) eye. The shaded areas showed the standard errors. (B) Representative confocal reflectance images of flat-mounted retinas from left (ONC) and right (control) eye from the same animal. \( D \) was pseudocolor-encoded on the images. (C–D) Paired comparison of \( D \) between left (ONC) and right (control) eyes, and between NO quadrants and CO quadrants; \( n = 10 \) mice.
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In this study, we tested whether the ultrastructural marker $D$ obtained by ELBS can be an early indicator for NFL/RGC damage. We introduced acute axon damage by pONC in mice, which propagated to the retina and caused RGC death. We found that neither NFL/RGL thickness nor ERG OP amplitude showed significant changes within 3 days post-pONC, while there is significant reduction of ultrastructural marker $D$ from the injured eye compared with the fellow eye from the same animals, as well as from the crushed site compared with the noncrushed site.

The spectroscopic analysis that was used to calculate $D$ is rooted from the tissue’s ultrastructural properties. The fundamental origin of this elastic optical backscattering is the RI fluctuation within tissue. Based on the Gladstone-Dale equation, tissue RI is linearly proportional to the local macromolecular mass density; thus, the measurement of backscattered light allows us to quantify the native tissue structures. Because of the heterogeneous nature of the tissue, we can use the mass density autocorrelation function to describe the tissue structures and establish a model where $D$ quantifies the shape of the autocorrelation function. Conceptually, a more condensed structure will yield higher $D$ than a more fragmented structure. More importantly, it has been theoretically and experimentally shown that the length-scale sensitivity by measuring $D$ can reach as low as 40 nm, beyond the diffraction-limited resolution of any ophthalmic imaging modality. In this study, we observed lower $D$ values from NFL/GCL in crushed eye, indicating more fragmented ultrastructures. This result aligns well with the previous finding that DNA fragmentation and cell disintegration occurs in the early stage of RGC damage, as well as the disruption of cytoskeleton.

Interestingly, we observed that the overall change of $D$ across all quadrants compared with the control was greater than that of the crushed quadrant compared with the nonoperated quadrants. This suggests that secondary effects may exist in the ultrastructure to the entire retina in response to the nerve crush. This could also be evident by the propagation of the optic nerve damage. Secondary degeneration of RGCs, which has been found in pONC or transection models of different animals, could be caused by the deleterious molecules released from the RGCs or glial cells damaged by the primary direct insults. First, oxidative stress, calcium overload, or mitochondrial dysfunction in the primarily injured RGCs could produce or regulate proapoptotic signals to neighboring RGCs, causing the DNA fragmentation and cell disintegration. Second, retinal glial cells such as microglia, astrocytes, and oligodendrocytes also played a role in the secondary degeneration after pONC, which may also contribute to the $D$ change observed here; however, the specific mechanisms of these complex pathologies need further study.

The reflectance spectroscopic contrast in NFL has been previously studied by Huang et al. In their study, an elevated IOP rat model was used to introduce global damage to the NFL. Using a multispectral imaging microreflectometer, the spectroscopic contrast was found between the glaucomatous and the control eyes. A neural fiber bundle model was used to elucidate the structural mechanism, where the loss of the thin neural fiber bundles from the simulation underpinned two experimental observations. In our study, we focused on investigating longitudinally whether an ultrastructural marker can be used to detect early NFL damage by comparing it with the morphological marker (NFL thickness) and the functional markers (OP amplitude) measured at the same time point. We used confocal gating by confocal reflectance microscopy to collect the reflected light from a ~6-μm-thick tissue layer, which was mostly composed of NFLs reflection and partially reflected from soma toward periphery. We adopted a continuous RI fluctuating model to interpret our observation. The model suggests the ultrastructural fragmentation within NFL/RGCs. This fragmentation also may contribute to the loss of neural fiber bundles previously found by Huang et al.

This paper presents a proof-of-concept study for using ELBS for early detection of optic neuropathy. For in vivo measurements, the ultrastructural marker $D$ can be measured and derived using other ophthalmologic imaging modalities that provide depth discrimination on the retina. For example, hyperspectral adaptive optics scanning laser ophthalmoscopy potentially could be used to provide spectroscopic measurement from NFL/RGCs layers. Optical coherence tomography, which is the standard of care in morphologic ophthalmic imaging, could also provide depth-resolved spectroscopic measurements. We recently demonstrated a novel OCT method to quantify $D$ values locally through spectroscopic analysis, which could potentially translate this study into clinical and preclinical in vivo applications.

The limitation of this study is the possible influence of the fixation process on the tissue’s ultrastructures. The fixatives denature proteins and introduce crosslinking to preserve the structure while at the same time could in principle increase the $D$ values. However, the advantage of fixation is more rigorous control over the confocal microscopic measurements between the crushed and normal retinas, and the significant difference of the ultrastructural marker still can be observed in the current study. Future in vivo measurement of retinal ultrastructure from intact retinas is work in progress.

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