Retinal alterations in patients with Lafora disease

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\textbf{ABSTRACT}

Purpose: Lafora disease is a genetic neurodegenerative metabolic disorder caused by insoluble polyglucosan aggregate accumulation throughout the central nervous system and body. The retina is an accessible neural tissue, which may offer alternative methods to assess neurological diseases quickly and noninvasively. In this way, noninvasive imaging may provide a means to characterize neurodegenerative disease, which enables earlier identification and diagnosis of disease and the ability to monitor disease progression. In this study, we sought to characterize the retina of individuals with Lafora disease using non-invasive retinal imaging.

Methods: One eye of three individuals with genetically confirmed Lafora disease were imaged with optical coherence tomography (OCT) and adaptive optics scanning light ophthalmoscopy (AOSLO). When possible, OCT volume and line scans were acquired to assess total retinal thickness, ganglion cell-inner plexiform layer thickness, and outer nuclear layer + Henle fiber layer thickness. OCT angiography (OCTA) scans were acquired in one subject at the macula and optic nerve head (ONH). AOSLO was used to characterize the photoreceptor mosaic and examine the retinal nerve fiber layer (RNFL).

Results: Two subjects with previous seizure activity demonstrated reduced retinal thickness, while one subject with no apparent symptoms had normal retinal thickness. All other clinical measures, as well as parafoveal cone density, were within normal range. Nummular reflectivity at the level of the RNFL was observed using AOSLO in the macula and near the ONH in all three subjects.

Conclusions: This multimodal retinal imaging approach allowed us to observe a number of retinal structural features in all three individuals. Most notably, AOSLO revealed nummular reflectivity within the inner retina of each subject. This phenotype has not been reported previously and may represent a characteristic change produced by the neurodegenerative process.

1. Introduction

Lafora progressive myoclonus epilepsy is a neurodegenerative metabolic disease caused by accumulation of insoluble polyglucosan aggregates.\textsuperscript{1,2} These aggregates, termed Lafora bodies, build up throughout the body and central nervous system, including the neurosensory retina, and lead to progressive symptoms early in adolescence.\textsuperscript{3}

Post-mortem investigations have found retinal cell loss and Lafora bodies within and around inner retinal neurons, predominantly in the ganglion cell and inner nuclear layers.\textsuperscript{4,5} Standard fundus examination and visual acuity testing is often unremarkable, but electroretinographic findings reveal retinal dysfunction.\textsuperscript{6} The neural retina is accessible to noninvasive imaging and provides a window into the central nervous system. Noninvasive retinal imaging has been used to demonstrate...
retinal alterations in a number of neurodegenerative diseases. Here we used spectral domain optical coherence tomography (SD-OCT) and adaptive optics scanning light ophthalmoscopy (AOSLO) in three genetically confirmed individuals with Lafora disease to look for similar retinal structural changes.

2. Materials and methods

This study followed the tenets of the Declaration of Helsinki and was approved by the Institutional Review Board at the Medical College of Wisconsin. Informed consent was obtained from all subjects, or from their legally authorized representative. Axial length was measured in each eye using an IOL Master (Carl Zeiss Meditec, Dublin, CA, USA) for scaling retinal images. One drop of phenylephrine hydrochloride (2.5%) and tropicamide (1%) was administered to one eye to dilate the pupil and suspend accommodation for imaging.

High-resolution SD-OCT volumetric (nominal 6 × 6 mm scans, 512 A-scans/B-scan, 128 B-scans) and horizontal line (nominal 6 mm) scans of the macula were acquired using a Cirrus HD-OCT device (Carl Zeiss Meditec) and/or the Bioptigen Envisu R2200 SD-OCT system (nominal 7 mm scan; Leica Microsystems, Wetzlar, Germany). Mean retinal thickness within a 6 mm radius of the fovea was calculated using Cirrus’ built-in macular analysis software. Bioptigen line scans were registered and averaged as previously described. The OCT line scans were then semi-automatically segmented using the Duke Optical Coherence Tomography Retinal Analysis Program (DOCTRAP) software to obtain retinal thickness measurements of the ganglion cell-inner plexiform layer (GCIPL) [boundaries: bottom of the retinal nerve fiber layer (RNFL) to top of the inner nuclear layer (INL)] and the outer nuclear layer + Henle fiber layer (ONL+) [boundaries: bottom of outer plexiform layer (OPL) to top of the external limiting membrane] using custom MATLAB software (MathWorks, Natick, MA).

The foveal avascular zone (FAZ) and the vasculature surrounding the optic nerve head (ONH) were assessed in Case 3 using the AngioVue OCT-Angiography system (Optovue Inc. Fremont, CA). For each eye, multiple foveal (nominal 3 × 3 mm, 304 B-scans at 304 A-scans/B-scan) and ONH (nominal 4.5 × 4.5 mm; 400 B-scans at 400 A-scans/B-scan) scans were acquired and averaged together as previously described. For FAZ measures, data were extracted from the whole retinal vasculature slab (boundaries: internal limiting membrane (ILM) to 9 μm below the OPL). For ONH scans, data were extracted from the radial peripapillary capillaries slab (boundaries: ILM to bottom of RNFL). Area and acircularity of the FAZ, segmented by a single observer (R.E.L.), as well as ONH capillary density was measured and assessed as previously described.

Using previously described AOSLO systems, confocal and non-confocal split-detection videos focusing on the photoreceptor layer or the RNFL were acquired at the macula and along the superior and nasal meridians. Individual videos were registered and averaged to produce single high-resolution images. The images were montaged together semi-automatically and regions of interest (150 × 150 μm) at 1° and 2° from the fovea were extracted. Cones were semi-automatically counted by a single observer (H.H.) and cone density was calculated using custom software (Translational Imaging Innovations, Inc., Hickory NC). GraphPad Prism (La Jolla, CA, USA) was used for statistical analysis.

3. Results

3.1. Case 1

A 24-year-old female was diagnosed with molecularly confirmed Lafora disease after presenting with typical symptoms (compound heterozygous EPM2A mutations: p.Asn163Asp & p.Ala254Mets*33). Clinical ophthalmological findings have been previously published for this patient. There was overall retinal thinning at 3 and 6 mm from the fovea in both eyes (Fig. 1A). GCIPL and ONL+ thickness measurements across the macula were within the lower limits of the previously published normative range. For each eye, optic nerve head (ONH) were assessed in Case 3 using the AngioVue using custom MATLAB software (MathWorks, Natick, MA). Mean retinal thickness within a 6 mm radius of the fovea was calculated using Cirrus’ built-in macular analysis software. Bioptigen line scans were registered and averaged as previously described. The OCT line scans were then semi-automatically segmented using the Duke Optical Coherence Tomography Retinal Analysis Program (DOCTRAP) software to obtain retinal thickness measurements of the ganglion cell-inner plexiform layer (GCIPL) [boundaries: bottom of the retinal nerve fiber layer (RNFL) to top of the inner nuclear layer (INL)] and the outer nuclear layer + Henle fiber layer (ONL+) [boundaries: bottom of outer plexiform layer (OPL) to top of the external limiting membrane] using custom MATLAB software.

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3.2. Case 2

A 20-year-old female was diagnosed with molecularly confirmed Lafora disease after presenting with symptoms (compound heterozygous EPM2A mutations: p.Arg241Ter & p.Asn163Asp). Overall retinal thinning was observed across the macula (Fig. 2A). In both eyes, six of the nine regions of the Early Treatment Diabetic Retinopathy Study (ETDRS) grid were in the lower 5th percentile for the normal distribution of retinal thickness, while one region (6 mm temporal) was in the lower 1st percentile. We were unable to acquire OCT line scans to assess GCIPL and ONL+ thickness in this patient. Parafoveal cone density estimates (Fig. 2B: 1°: ~32,800 cones/mm²; 2°: ~27,000 cones/mm²) were significantly lower than normal and showed a general trend of increased capillary density across the whole area imaged (Fig. 4B).

3.3. Case 3

A 12-year-old male, sibling of Case 2, had molecularly confirmed Lafora disease, but had not manifested clinical signs or symptoms at the time of imaging. The age of this patient was adjusted to 18 years to allow comparison to normative data on Cirrus HD-OCT device, which showed no signs of retinal thinning (Fig. 3A). GCIPL and ONL+ thickness measurements, and parafoveal cone density estimates (1°: ~50,600 cones/mm²; 2°: ~36,200 cones/mm²) were within normal limits (Fig. 3B and C). Nummular reflectivity was observed within the RNFL approximately 5° temporal to the macula (Fig. 3D). The FAZ area (right: 0.33 mm²; left: 0.36 mm²; Fig. 4A) and acircularity (right: 1.12, left: 1.19) were not significantly different in comparison to a normative database (area: 0.278 ± 0.101 mm²; acircularity: 1.19 ± 0.095). Additionally, the radial peripapillary capillary density surrounding the ONH was not significantly different from a previously published dataset, though there was a general trend of increased capillary density across the whole area imaged (Fig. 3D).

4. Discussion

The retina is an extension of the central nervous system, is amenable to noninvasive imaging, and is affected by neurodegenerative processes. Previous work has demonstrated that visual acuity often remains good, while electrotetrographic findings reveal generalized retinal dysfunction. Therefore, we sought to characterize the retinal structure of three individuals with Lafora disease using noninvasive retinal imaging (OCT and AOSLO) beyond standard fundoscopic examination. OCT is widely accessible and commonly used to evaluate RNFL and ganglion cell layer thickness to detect neurodegenerative pathology in the retina, which is used for diagnosis, assessing disease severity, and monitoring progression. AOSLO provides high resolution en-face images of the retina, where individual cells in the outer retina and hyperreflective structures within the inner retina can be identified. There were noteworthy findings observed with both imaging modalities. First, the two symptomatic patients in this series had notable retinal thinning across the macula viewed with OCT: an abnormal finding in young adults. This finding appears to be due to ganglion and bipolar cell loss, as previously suggested in histopathological reports. Additionally, ONL+ thickness (measured on OCT) and cone density (measured on
AOSLO) were unaffected in all three cases. These findings suggest that the outer retina is normal, or unaffected by Lafora disease, which corroborate findings from previous studies that suggest Lafora disease alters inner retinal neurons exclusively and coincides with inner retinal degeneration. Although Lafora bodies are not visualized with the imaging modalities, it can be suggested that the changes observed in the inner retina of the affected individuals in this study are related to the neurodegenerative effects of this disease.

Most notably, nummular reflectivity – colloquially known as Gunn’s dots – was observed in all three cases on AOSLO, including Case 3 (asymptomatic). The observed diameter in our patients (mean ± SD = 11.68 ± 2.4 μm) was similar to that reported in two prior studies in individuals with normal vision. Gunn’s dots can appear on the RNFL when imaged with AOSLO, but they reside somewhere within the ILM. They are believed to be either a by-product of microglial phagocytosis or dead microglia that fulfilled their scavenger role in retinal pathology. While nummular reflectivity can be benign, observed in younger eyes near the optic disc and in the macula of adults, nummular reflectivity has been reported across several retinal and neurological conditions in areas of nerve fiber disease or loss, and may be indicative of a nonspecific neurodegenerative process. The origin, function, and pathological nature of Gunn’s dots remains an area of active study. Further studies should be completed to determine how the presentation of Gunn’s dots varies between neurodegenerative and healthy populations, as well as the relationship between the presence of Gunn’s dots and inner retinal thinning.

There are certain limitations of this study. Lafora disease is a rare disorder, limiting the number of subjects available for imaging. The
features of this disease (e.g. seizure activity and cognitive impairment) present challenges for these individuals to perform the tasks required for imaging (i.e., fixating on a target and remaining still). This can reduce the likelihood of collecting useable images, which contributed to our inability to obtain certain modalities on each individual in this study.

This also presented practical limitations to the amount of time available for research testing, and we did not collect visual acuity or other visual function measures in this study. Further imaging and longitudinal follow-up within this population may clarify whether the imaging findings reported here are of clinical use in progressive
neurodegeneration disorders like Lafora disease. Nevertheless, our findings, in corroboration with previous reports, are consistent with the inner retina being preferentially affected by Lafora disease. This suggests that multimodal imaging of the inner retina may be a useful tool to monitor disease progression or treatment response in patients with Lafora disease.

5. Conclusions

Consistent with previous reports, we observed variable retinal structure in three individuals with Lafora disease when imaged with non-invasive retinal imaging techniques. A notable finding was nummular reflectivity observed within the inner retina on AOSLO in all three individuals, which may be indicative of a generalized neurodegenerative process. Noninvasive retinal imaging in neurodegenerative disorders, including Lafora disease, enables identification of early disease states and provides new opportunities to monitor disease progression.

6. Patient consent

Written consent was obtained from all subjects’ legal guardian or legally authorized representative. This report does not contain any personal information that could lead to the identification of these individuals.

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Authorship

All authors attest that they meet the current ICMJ criteria for Authorship.

CRediT authorship contribution statement

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Declaration of competing interest

Mrs. Linderman is a consultant for OptoVue. Dr. Rosen has personal financial interest in Opticology and Guardion, and is a consultant for OptoVue, Boehringer-Ingelheim, Astellas, Genentech-Roche, Nanoretina, OD-OS, Regeneron, and Bayer. Dr. Vincent is a consultant for Adverum Biotechnologies Inc. Dr. Carroll receives financial support from AGTC and OptoVue, is a consultant for MeiraGTX, and has personal financial interest in Translational Imaging Innovations. Dr. Minassian is chief medical advisor to Taysha Gene Therapies. The following authors have no financial disclosures: H.H., J.A.C., E.N.W., R.M., P.S., T.Y.P., and E.J.P.

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