Structure/Psychophysical Relationships in X-Linked Retinoschisis

Lea D. Bennett,1 Yi-Zhong Wang,1,2 Martin Klein,1 Mark E. Pennesi,3 Thiran Jayasundera,4 and David G. Birch1,2

1Retina Foundation of the Southwest, Dallas, Texas, United States
2Department of Ophthalmology, University of Texas Southwestern Medical Center, Dallas, Texas, United States
3Oregon Health & Science University, Casey Eye Institute, Portland, Oregon, United States
4University of Michigan, Kellogg Eye Center, Ann Arbor, Michigan, United States

Correspondence: Lea D. Bennett, Retina Foundation of the Southwest, Suite 200, 9600 North Central Expressway, Dallas, TX 75231, USA; lbennett@retinafoundation.org.
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PURPOSE. To compare structural properties from spectral-domain optical coherence tomography (SDOCT) and psychophysical measures from a subset of patients enrolled in a larger multicenter natural history study of X-linked retinoschisis (XLRs).

METHODS. A subset of males (n = 24) participating in a larger natural history study of XLRs underwent high-resolution SDOCT. Total retina (TR) thickness and outer segment (OS) thickness were measured manually. Shape discrimination hyperacuity (SDH) and contour integration perimeter (CIP) were performed on an iPad with the myVisionTrack application. Sensitivity was measured with fundus-guided perimeter (+2 threshold testing strategy; 10-2 grid, spot size 3, 68 points). Correlation was determined with Pearson’s r correlation. Values are presented as the mean ± SD.

RESULTS. Mean macular OS thickness was less in XLRs patients (17.2 ± 8.1 μm) than in controls (37.1 ± 5.7 μm; P < 0.0001) but mean TR thickness was comparable (P = 0.5884). For patients, total sensitivity was lower (13.2 ± 6.6 dB) than for controls (24.2 ± 2.4 dB; P = 0.0008) and had a strong correlation with photoreceptor OS (R² = 0.55, P = 0.0001) and a weak correlation with TR thickness (R² = 0.22, P = 0.0158). The XLRs subjects had a logMAR best corrected visual acuity (BCVA) of 0.5 ± 0.3 that was associated with OS (R² = 0.79, P < 0.0001) but not TR thickness (R² = 0.01, P = 0.6166). Shape DH and CIP inner ring correlated with OS (R² = 0.33, P = 0.0085 and R² = 0.47, P = 0.0001, respectively) but not TR thickness (R² = 0.0004, P = 0.93; R² = 0.0043, P = 0.75, respectively).

CONCLUSIONS. When considered from a single visit, OS thickness within the macula is more closely associated with macular function than TR thickness within the macula in patients with XLRs. Keywords: retinoschisis, SDOCT, microperimetry

Juvenile X-linked retinoschisis (XLRs) is a congenital macular degeneration affecting 1/5000 to 1/25,000 worldwide.1–3 The gene associated with XLRs, Retinoschisin (RS1),4 translates to a retinoschisin protein (RS1), which assists in maintaining retinal structure by binding to the photoreceptors and bipolar cells. Patients are diagnosed in their primary school years with clinical characteristics of bilateral retinal splitting5–8 and an electronegative electroretinography (ERG) response with preserved a-wave.9 Best corrected visual acuity (BCVA) typically ranges from 20/50 to 20/120 (0.5–0.8 log minimum angle of resolution [logMAR]) and remains stable until the fifth or sixth decade of life when the cavities resolve and visual acuity decreases.10

Shape discrimination hyperacuity (SDH) and contour integration perimeter (CIP) in patients with intermediate AMD show significant deficits, with macular edema exacerbating the loss of the ability to detect distortions in circular shapes.11,12 These tests assess the global integration of visual stimuli over a large retinal area. The SDH tests parafoveal acuity, whereas the CIP determines retinal acuity outside of the central 3°. Due to the foveal edema in XLRs we hypothesize that the global integration measured by SDH/CIP may be affected, although some patients retain a relatively good BCVA. Spectral-domain optical coherence tomography (SDOCT) studies in XLRs have been reported,5,7,13,14 but rarely correlated with fundus-guided perimeter15 or shape discrimination. Clinical attributes of XLRs have been characterized, but concise relationships with structure and psychophysical function need further exploration.

A multicenter natural history study of XLRs was designed to understand disease progression and determine suitable outcome measures for future gene therapy trials. The results reported here were obtained from a single visit. A subset of patients were tested with additional measures so that we could determine whether photoreceptor outer segment (OS) and/or total retina (TR) thickness could predict performance on visual tasks such as BCVA, fundus-guided perimeter, SDH, and CIP in patients with XLRs.

METHODS

Study Population

Measures were obtained from a cohort of 24 subjects (age 32.2 years ± 17.7 SD; range, 9–79 years) from a larger group (n =
78) enrolled in a multicenter natural history study of XLRS, ClinicalTrials.gov NCT0233117. Included here are results from a single visit for patients receiving supplementary evaluations that were not part of the natural history study. These patients were enrolled consecutively after Institutional Review Board approval of protocols. Age-similar normal values were derived from contemporaneous subjects for each test. Not all control subjects performed all tests, so the controls are unique individuals for each analysis. Controls had normal eye exams and normal visual acuity. Analyses of within-patient variability were used to determine whether there were significant differences between eyes for each test. Since there were no significant differences between eyes, we averaged both eyes for $n = 1$ per test. Research adhered to the tenets of the Declaration of Helsinki and was approved by the Western Institutional Review Board.

**Spectral-Domain OCT**

High-resolution horizontal line scans of the macula were obtained using Spectralis Heidelberg retina angiography + OCT (Heidelberg Engineering, Inc., Heidelberg, Baden-Württemberg, Germany). Horizontal line scans had a mean of 100 scans over $30^\circ$ including the fovea. The average thicknesses of TR and OS were measured with manual segmentation (Igor Pro 6.03A; Wave Metrics, Inc., Tigard, OR, USA). The OS thickness was determined as the distance between the ellipsoid zone (EZ), otherwise known as the inner segment (IS)/OS junction, and the apical retinal pigment epithelium (RPE) border. The TR thickness was measured from the inner limiting membrane (ILM) to the basement membrane (BM).

**Microperimetry**

Macular sensitivity was determined under mesopic conditions on a microperimeter (MP1-S; NAVIS software, ver. 1.7; Nidek Technologies, Padova, Italy) with spot size 3 (0.43° diameter) and a 10-2 protocol. Perimetric sensitivity (with infrared illumination of the fundus) was determined as the mean of 68 points spanning 20° of the retina. The MP1-S microperimeter tests sensitivity up to 20 dB, but normal subjects and some XLRS patients need a higher dynamic range of stimuli intensity to get their true sensitivity (>20 dB). To circumvent the ceiling effect of the MP1-S, a 1.0 log neutral density filter was used when the patient exhibited maximum sensitivity (20 dB) for the majority of the individual test points. One patient had one eye with no light perception. The mean sensitivity for this eye was set to 0 dB, averaged with the fellow eye, and included in the analysis.

**Visual Acuity**

After refraction, BCVA was assessed by Electronic Visual Acuity Tester (Jaeb Center for Health Research, Tampa, FL, USA). Results for each subject were represented by the Snellen equivalent or as the logMAR.
Shape Discrimination Hyperacuity and Contour Integration Perimetry

Shape DH and CIP tests to evaluate central vision were performed on an iPad using the myVisionTrack visual function application. The SDH test displays three smooth and one distorted circle on the iPad. The subject was instructed to touch the distorted circle. The test continues as a 4-alternative, forced choice (4AFC) test algorithm with a 2-down, 1-up adaptive staircase procedure for the amount of distortion presented in each trial until the SDH is determined. The CIP test showed smooth and distorted circular contour segments spatially distributed in an “inner” or “outer” ring using a 4AFC staircase paradigm with a stimulus duration of 0.25 seconds (Wang Y-Z, Mitchel G. IOVS 2013;54:ARVO E-Abstract 5019). As with the SDH test, the subject was instructed to choose the distorted line segment. A maximum likelihood fitting procedure was implemented to estimate detecting the distortion of contour segments of inner or outer rings.

Differences between sample means were analyzed with Student’s 2-tailed t-test and the Pearson coefficient test for the correlation studies. All values are presented as the mean ± SD.

RESULTS

Patient Information

Each of the 24 patients had an identified mutation in the R51 gene. The most common type of mutation was a missense mutation (80%). Other mutations that occurred in our patient population were small frameshifting insertions/deletions (10%), intronic splice site mutations (5%), and exon deletions (5%). Of the patients presented here, two individuals are...
Fig. 1E) was absent in the SDOCT scans from eyes devoid of (Fig. 1F). Of interest, the photoreceptor layer (PRL, enlarged in cavities (six eyes, 12.5%) were also noted in our population and ONL (2.1%, not shown). Eyes without detectable schisis in the GCL only (2.1%), and one eye had cavities in nine eyes (14.6%) of our patients (Fig. 1E). One eye had cavities (12 eyes each; 25%; Figs. 1B–D, respectively).

Figures 1B through 1D, which involved the INL, ONL, and the highest incidence for cavity localizations are shown in eyes (14.6%) with cavities intruding into the inner nuclear layers (INL) and outer nuclear layer (ONL; Fig. 1A). Examples for the highest incidence for cavity localizations are shown in Figures 1B through 1D, which involved the INL, ONL, and ganglion cell layer (GCL) or had an INL-only pattern of schisis cavities (12 eyes each; 25%; Figs. 1B–D, respectively). Localization of cavities within the INL and GCL occurred in nine eyes (14.6%) of our patients (Fig. 1E). One eye had cavities in the GCL only (2.1%), and one eye had cavities in the GCL and ONL (2.1%, not shown). Eyes without detectable schisis cavities (six eyes, 12.5%) were also noted in our population (Fig. 1F). Of interest, the photoreceptor layer (PRL, enlarged in Fig. 1E) was absent in the SDOCT scans from eyes devoid of cavities (Fig. 1F) and from those with extrafoveal schisis (Fig. 1C).

Since cavity size differs among subjects, we wanted to know if age was a contributing factor to TR thickness in the macula. Figure 2A depicts how the TR and OS thicknesses were measured. The TR, determined as the distance between the ILM and the BM in the central 10°, was 335.6 ± 97.8 µm for XLRS subjects, which was not different than controls (318.1 ± 17.7 µm; P = 0.5884; Fig. 2B). However, the OS thickness was smaller in XLRS patients (17.2 ± 8.1 µm) compared to controls (37.1 ± 5.7 µm; P < 0.0001; Fig. 2C). When age was considered as a potential factor for either central TR or central OS thickness, we found that there was a weak relationship between XLRS patient age and TR (Fig. 2D; R² = 0.24, P = 0.0158) or OS (R² = 0.18, P = 0.0579; Fig. 2E) thickness.

Spectral-Domain OCT

High-resolution SDOCT scans showed different clinical features in our population of XLRS patients. There were seven eyes (14.6%) with cavities intruding into the inner nuclear layer (INL) and outer nuclear layer (ONL; Fig. 1A). Examples for the highest incidence for cavity localizations are shown in Figures 1B through 1D, which involved the INL, ONL, and ganglion cell layer (GCL) or had an INL-only pattern of schisis cavities (12 eyes each; 25%; Figs. 1B–D, respectively). Localization of cavities within the INL and GCL occurred in nine eyes (14.6%) of our patients (Fig. 1E). One eye had cavities in the GCL only (2.1%), and one eye had cavities in the GCL and ONL (2.1%, not shown). Eyes without detectable schisis cavities (six eyes, 12.5%) were also noted in our population (Fig. 1F). Of interest, the photoreceptor layer (PRL, enlarged in Fig. 1E) was absent in the SDOCT scans from eyes devoid of cavities (Fig. 1F) and from those with extrafoveal schisis (Fig. 1C).

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Microperimetry

To assess central retina function, psychophysical sensitivity was measured with fundus-guided microperimetry. The black arrow on the fundus/microperimetry grid overlay (Fig. 3A) defines the position of the horizontal line scan (Fig. 3B). Subject XLRS-001-RFS-325, a 10-year-old boy, had BCVA of 20/80 (0.6 logMAR) and a lower than normal (24.2 ± 2.4 dB) total mean sensitivity (12.8 dB) in the left eye (Fig. 3A). This subject had a similar central TR thickness (353 µm) and smaller OS thicknesses (8.4 µm) compared to normal (318.1 ± 17.7 µm and 37.1 ± 5.7 µm, respectively). An area highlighting where the EZ line was absent from the central retina is enlarged. The red arrows point to where the EZ line stops (Fig. 3B), which explains why the average of the central 10° OS for this subject was decreased. Conversely, the EZ line (red arrow) can be seen clearly across the macular region in a 19-year-old subject, XLRS-001-RFS-319 (Fig. 3C). This individual had BCVA of 20/40 (0.4 logMAR), a near-normal mean sensitivity of 20.7 dB, and a comparable TR thickness (326 ± 37.1 µm) compared to normal (318.1 ± 17.7 µm, respectively). This subject’s OS thickness was 22 µm, which was larger than the mean OS thickness from all XLRS subjects (13.2 ± 6.6 µm; Fig. 3C). Overall, the mean sensitivity in XLRS patients was lower than the control mean sensitivity (XLRS: 13.2 ± 6.6 dB; control: 24.2 ± 2.4 dB; P = 0.0008).

To evaluate the relationship between anatomical features and psychophysical functional measures, the sensitivity or BCVA was analyzed against central TR and macular OS thickness. The TR thickness had a weak relationship (R² = 0.22, P = 0.0158; Fig. 3D) circular (SDH) and (CIP inner) contour distortion did not correlate with age. Correlation was determined with Pearson’s r correlation.

Shape DH and Contour IP

Shape DH was worse in XLRS subjects (−0.4 ± 0.2 logMAR) than in controls (−0.7 ± 0.1 logMAR, P < 0.001; Fig. 4A). Detection of contour lines in a ring, designated inner CIP, was higher (worse) for XLRS subjects (−0.5 ± 0.3 logMAR) than for control subjects (−0.9 ± 0.2 logMAR, P = 0.002, Fig. 4B).
However, thresholds for contour lines in a wider ring (outer CIP) were not different between the groups (XLRS: −0.7 ± 0.2 logMAR; control: −0.8 ± 0.1 logMAR; Fig. 4C). No effect of age was found on SDH and CIP thresholds for patients with XLRS (Fig. 4).

Correlation analysis with TR or OS thickness revealed no association of TR thickness with SDH ($R^2 = 0.0004$, $P = 0.9270$), inner CIP ($R^2 = 0.0043$, $P = 0.7546$), or outer CIP ($R^2 = 0.0039$, $P = 0.7576$) (Figs. 5A–C). However, OS thickness was highly correlated with SDH ($R^2 = 0.3297$, $P = 0.0085$) and inner CIP ($R^2 = 0.4667$, $P = 0.0001$), but weakly associated with outer CIP ($R^2 = 0.2104$, $P = 0.0307$; Figs. 5D–F). Of note, the correlation was similar whether controls were included in the analysis or not.

DISCUSSION

The purpose of the present study was to compare structural properties from SD-OCT to psychophysical measures in a subset of patients enrolled in a larger multicenter natural history study of XLRS. Here we showed that the OS length was highly correlated with BCVA (Fig. 3G), fundus-guided perimetry (Fig. 3E), SDH (Fig. 5D), and CIP (Fig. 5E) but that total thickness of the retina had weak association with these measures (Figs. 3D, 3F, 5A–C). The TR thickness failed to show a negative correlation with age in our patients tested with XLRS, unlike previous reports showing that younger patients had large foveal schisis cavities and older patients had thinner retinas with minimal cavities. While this may be common in progression of the disease, it is certainly not seen in all patients. The cavity size can vary according to the individual regardless of age, which could be the result of the specific mutation, other eye diseases, or medication. For example, two patients (ages 53 and 40) in this group, using ocular carbonic anhydrase inhibitors (CAIs) to reduce the swelling in the retina, had resolution of foveal schisis. Of note, CAI use was not prohibited in this study; not all patients respond to CAIs, and other patients in this study had cystic changes in the macula without resolution while using this medication. Further, this study was not powered to detect group differences between those using and those not using the CAIs.

Although the mean sensitivity was variable in these XLRS subjects, it was still below control sensitivity. Interestingly, sensitivity, a psychophysical examination of macular function, was better correlated with macular OS thickness than central TR thickness in XLRS subjects. This suggests that a defect in the photoreceptors, not maculoschisis, contributes to macular sensitivity loss in patients with XLRS.

Similar to patients with macular edema in AMD, the XLRS patients displayed defects in SDH and CIP validating our hypothesis that these patients would have a deficit in the global integration visual acuity. This could be due to the cystic cavities distorting straight lines when maculoschisis is present. However, after further analysis, TR thickness did not correlate with SDH or CIP outcomes in XLRS subjects (Figs. 5A–C). Interestingly, it was OS thickness that correlated with the results from SDH and CIP tests (Figs. 5D–F). Thus, the outer retina is the major limitation to the altered SDH/CIP results.
shown here. However, it cannot be dismissed that the schisis could have exacerbated the loss of visual integration as found in AMD. This is the first report of shape and contour line discrimination deficits in XLRS. This supports the hypothesis that a photoreceptor defect, rather than maculopathy, is most responsible for the functional deficit in XLRS. It will be interesting to repeat these tests to determine if the SDH and CIP change over time in these patients. In particular, if age is not a contributing factor and younger patients do not differ from older patients, this would suggest that the shape discrimination defect is present at the earliest stage of disease.

Since it is believed that in the majority of patients the disease shows either no or minimal progression, accurate baseline results need to be documented from each subject when considering outcomes for a treatment trial. Furthermore, test-retest variability will also be important when determining significant change in disease progression. Test-retest variability has been obtained for microperimetry and BCVA in seven patients with XLRS, with the authors evaluating the coefficients of repeatability and associated confidence intervals so that they would know the minimum level of change required in a parameter to be considered statistically different from baseline. Test-retest statistics have yet to be determined within our patient population. These measures will be assessed to see how the data vary among these particular XLRS subjects in order to define significant change from baseline for either treatment or longitudinal studies.

Data presented here are consistent with previous measures of schisis cavities and decreased photoreceptor sensitivity in patients with XLRS. New findings include measures of OS length and the relationship between OS length and macular function based on microperimetry, SDH, CIP, and BCVA. Psychophysical outcome measures in these patients will be imperative when deciphering the effectiveness of therapies in future clinical trials for XLRS.

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References