Microstructural changes of photoreceptor layers detected by ultrahigh-resolution SD-OCT in patients with autosomal recessive bestrophinopathy

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ABSTRACT
Purpose: To determine the changes in the microstructures of the photoreceptors in patients with autosomal recessive bestrophinopathy (ARB) by ultrahigh-resolution spectral-domain optical coherence tomography (UHR-SD-OCT).
Methods: Five eyes of 4 patients with ARB were studied. Cross-sectional images of the fovea were recorded by the UHR-SD-OCT system with a depth resolution of <2.0 μm.
Results: The UHR-SD-OCT images revealed changes in the outer retinal structures that were dependent on the severity of the photoreceptor atrophy. There was an increase in the reflectivity and appearance of small hyperreflective dots (HRDs) in the outer segments, followed by an irregularity and decrease in the length of the outer segments, then a disruption of the ellipsoid zone (EZ) band, and appearance of large HRDs corresponding to the segmented ellipsoids. Finally, there was a disappearance of the large HRDs followed by a localized thinning of the outer nuclear layer and appearance of hyperreflective foci above the region of the disrupted EZ.
Conclusions: UHR-SD-OCT can record images that show detailed changes of the microstructures of the photoreceptors at different stages of ARB. These observations should help in determining the mechanisms involved in retinal pathology and should provide important information on the effectiveness of treatments.

1. Introduction
Autosomal recessive bestrophinopathy (ARB; OMIM-611809) is one of the BEST1-related retinopathies that is caused by biallelic pathogenic variants of the BEST1 gene.1–2 ARB shares common clinical features with Best vitelliform macular dystrophy (VMD; OMIM-153700), which is caused by a single pathogenic variant of the BEST1 gene.1–3 The retinal structural changes of ARB have been described from the spectral-domain optical coherence tomographic (SD-OCT) images.4–6, and these features are essentially the same as those in cases of VMD.7–9 However, due to the limited axial resolution of the conventional SD-OCT, there is a significant gap in what is observed in the diseased retina by the conventional OCT and what is happening structurally during the process of neuronal atrophy of the retinas. The findings made from the conventional SD-OCT images from eyes with both VMD and ARB partially corresponded with the histological studies of postmortem eyes with VMD.10–17 However, histological studies cannot determine the structural changes of the photoreceptor layer in VMD and ARB in detail because of the fragility of the photoreceptor cells to histological processing.

An ultrahigh-resolution SD-OCT (UHR-SD-OCT; KOWA OCT Bi-μ; Kowa Company, Ltd., Japan) device has become available which has an axial resolution of <2.0 μm which is about three-times better than the conventional SD-OCT. It can obtain clearer images of the outer retinal structures both in normal18,19 and diseased eyes.20–22 Matsui et al. presented a B-scan image of a patient with ARB obtained by the UHR-SD OCT and showed detailed structural changes which could not be observed by the conventional SD-OCT devices.17 We have examined the microstructures of the photoreceptor layer in eyes with occult macular dystrophy (OMD, Miyake’s disease) by UHR-SD-OCT and were able to determine the characteristics of the photoreceptor microstructures at different stages of OMD.23

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The purpose of this study was to determine the changes in the microstructures of the photoreceptors in eyes with ARB. To accomplish this, we examined the B-scan images obtained by the UHR-SD-OCT of 5 eyes of 4 patients with ARB of the same genotypes but of different severities. Our findings will fill the gap between the OCT images and the histological findings in eyes with ARB which should help in determining the mechanisms for the degeneration of the photoreceptors.

2. Patients and methods

The procedures used in this study adhered to the tenets of the Declaration of Helsinki and were approved by the Ethics Committee of the National Institute of Sensory Organs, National Hospital Organization Tokyo Medical Center (Reference, R18-029). A signed informed consent was received from all subjects for the examinations after an explanation of the nature and possible consequences of the study.

2.1. Participants

We examined 5 eyes of 4 unrelated patients with ARB. The phenotype of these patients was determined by comprehensive ophthalmic examinations including ophthalmoscopy, fundus photography, fundus autofluorescence (FAF; HRA 2; excitation light, 488 nm; barrier filter, 500 nm; Heidelberg Engineering, Heidelberg, Germany), conventional SD-OCT (Cirrus HD OCT, version 6.5; Carl Zeiss Meditec, Dublin, CA, USA), electrooculography (SG-2002, LKC Technologies, Gaithersburg, MD, USA), and full-field electroretinography (ERG; LE4000, Tomey Corporation, Aichi, Japan). For the recording by the conventional OCT, we used the ‘HD 5 Line Raster’ mode, and averaged four 9.0 mm B-scan images to obtain one averaged B-scan image. Genetic analysis was performed on all the patients. To search for the causative genes, whole exome sequencing with targeted analysis for retinal disease-causing genes on RetNET (https://sph.uth.edu/retnet/home.html) and inheritance filtration were performed as described in detail. The details of the whole exome sequencing and in silico molecular genetic analyses are described as supplemental information (Supplementary Methods for the Genetic Analysis). All 4 cases of ARB had the same compound heterozygous pathogenic variants of the BEST1 gene (c.584C>T, p.Ala195Val; c.763C>T, p.Arg255Trp: mRNA reference sequence, NM_004183.3).

2.2. UHR-SD-OCT imaging

An UHR-SD-OCT system (KOWA OCT Bi-μ; Kowa Company, Ltd., Japan) with a broadband superluminescent diode was used on all subjects. The center wavelength of the light source was 860 nm, and the bandwidth was 135 nm. Each A-scan had a depth of 2.6 mm which consisted of 1800 A-scans. For a 9.0 mm width image, the digital width sampling was 5 μm/pixel. Each B-scan spanned 30° and consisted of 1800 A-scans. A detailed technical description of this imaging device has been published elsewhere. Cross-sectional horizontal and vertical images with a 9.0 mm width across the fovea were recorded in all the subjects with dilated pupils. The original averaging technique of ‘A-scan matching algorithm’ was taken to improve the clarity of the OCT images, and the procedures were described in detail elsewhere. Briefly, twenty B-scans images which had higher correlation coefficients to the base image were chosen from the fifty B-scan images, and averaged for the final B scan image. The examination required a fixation of 1.5 s to obtain a horizontal B-scan image from either a normal or diseased eye.

3. Results

3.1. UHR-SD-OCT images of normal eyes

The phenotypic and genotypic findings of the four cases are
presented in Table 1. Representative horizontal B-scan images across the fovea of the normal eye of a 41-year-old woman taken with the UHR-SDR-OCT device are presented in Fig. 1A and B together with the longitudinal reflectivity profiles (LRP; Fig. 1C). Hyperreflective regions were present which were not mentioned in the International Nomenclature for Optical Coherence Tomography (IN-OCT), viz., a rod IZ and Bruch’s membrane.

3.2. UHR SD-OCT images of eyes with ARB

Fundus photographs and FAF images of the 5 eyes of 4 cases with ARB are shown in Fig. 2. All of the cases were bilateral, and the fundus changes corresponded approximately with those of eyes with VMD at the vitelliform or scrambled-egg stage.3 The OCT images showed hyper-autofluorescence (AF) spots in Cases A, B, D, and E, and a hyper AF ring at the border of the affected region in Case C. The macular region showed hypo AF in all cases. In Cases B, C, and D, the affected region extended beyond the macula which is typical of cases with ARB.4-5

The conventional SD-OCT images showed these eyes had characteristic structural changes that are typical of that between the pseudo-dhyopopyon and vitelliform stages, viz., subretinal fluid (SRF) in the macula, absence of the IZ, elongation of the photoreceptor outer segments, disruption of the EZ, and focal choroidal excavation (Fig. 3F).4,5,6,8,10,11

The UHR-SD-OCT B-scan images of the five eyes with ARB (Fig. 2) are shown in Fig. 3. Cases A to E are aligned according to the severity of the photoreceptor damage from mild to severe. One B-scan image of the same eye as in Fig. 3A taken by a conventional SD-OCT is shown for comparison (Fig. 3F). Further expanded images (x 2.5) of the outer segment regions in the right column in Figures A, F, and B are shown in Fig. 3G.

The OCT images of the right eye of a 33-year-old woman whose best-corrected visual acuity (BCVA) was 20/20 are shown in Fig. 3A. There is a shallow retinal detachment in the perimacular region (Fig. 3A, white arrow), where the cone IZ is not present and the outer segments are elongated and hyperreflective. Part of the outer segments in the detached region resembled icicles hanging on the EZ (Fig. 3A and G left, asterisk).

The OCT images of the left eye of a 20-year-old woman whose BCVA was 20/20 are shown in Fig. 3B. Similar to the earlier case, the outer segments are elongated and hyperreflective material that resembled icicles can be seen hanging on the EZ (Fig. 3B and G right, asterisk). The peripheral edge of the outer segments was irregularly aligned, and the cone IZ was not observed in the detached region. There were debris-like hyperreflective spots between the outer segments and the RPE (Fig. 3B, double asterisks).

The OCT images of the left eye of a 36-year-old woman whose BCVA was 20/20 are shown in Fig. 3C. The outer segments are shortened and the EZ band is disrupted and composed of individual large hyperreflective dots (HRDs; Fig. 3C, yellow arrowheads). These HRDs were connected to the ELM with a less highly reflective bridge-like structures. The outer segments could not be clearly seen. Instead, thin icicle-like structures could be seen hanging on the ELM (Fig. 3C, yellow asterisks).

The OCT images of the right eye of a 55-year-old man whose BCVA was 20/125 due to the loss of photoreceptors at the fovea are shown in Fig. 3D. The EZ band was disrupted and a small number of large HRDs was seen connected to the ELM (Fig. 3D, yellow arrowheads). Thin icicle-like structures can be seen hanging directly on the ELM (Fig. 3D, yellow asterisks).

The OCT images of the right eye of the patient shown in Fig. 3A whose BCVA was 20/20 are shown in Fig. 3E. The expanded image shows a widely disrupted EZ with preserved ELM inferior to the fovea (Fig. 3E, white arrowhead). At the location of the disrupted EZ, the outer nuclear layer (ONL) is thin (Fig. 3E, white arrowhead). There are hyperreflective spots scattered in the ONL above the disrupted EZ (Fig. 3E, white arrowhead), which are known as hyperreflective foci (HRF).3,5

4. Discussion

4.1. Process of photoreceptor atrophy in eyes with ARB

This was an observational study of 5 eyes of 4 unrelated individuals with ARB, and we assumed that there was a longitudinal progression of the disease process that could be determined by the cross-sectional observations of the cases with different severities but with the same genotype. The early abnormality of the photoreceptor layer was observed in the perimacular region as a shallow SRF lesion in the UHR-SD-OCT images (Fig. 3A, white arrow). The cone IZ became indistinct and disappeared, and the outer segments became hyperreflective and elongated with small HRDs aligned in a row like icicles or string of pearls hanging on the EZ (Fig. 3A, asterisk and Fig. 3G, left). In the macular region with a larger SRF, the peripheral edges of the outer segments were irregularly aligned (Fig. 3B, asterisks and Fig. 3G, right) and some parts of the icicle-like outer segments appeared to be shortened with debris-like hyperreflective spots in the SRF (Fig. 3B, double asterisks). In addition, the EZ band became disrupted and was seen as large HRDs connected to the ELM with less highly reflective bridge-like structures (Fig. 3C and D, yellow arrowheads). In the next stage, the disrupted EZ was more atrophic, and the large HRDs were replaced by thin string-like icicles hanging directly on the ELM (Fig. 3D, yellow asterisks). During these processes, the ONL became thinner (Fig. 3C and D), and the outer plexiform layer appeared to be closer to the ELM with scattered HRF above the disrupted EZ in the region where icicle-like formation was completely lost (Fig. 3E, white arrowhead). These HRF could also be
cases with ARB are shown. The images in fundus photographs (left column) and FAF images (right column) of the five eyes of four retina of eyes with autosomal recessive bestrophinopathy (ARB). Fundus photographs and fundus autofluorescence (FAF) images of the eyes with ARB, showing the left eye of the patient in 9, 27. These characteristic features in eyes with ARB have been reported by the conventional SD-OCT, however detailed structural changes of the photoreceptor layer in eyes with ARB were not described. Instead, there are many studies that due to the limited resolution of the conventional SD-OCT, the results of the previous studies did not show the disease-associated microstructures of the photoreceptors such as the string-of-pearls appearance with small HRDs of the outer segments (Fig. 3A, B and 3G, asterisks) and disrupted EZ which was composed of large HRDs (Fig. 3C and D, yellow arrowheads). Thus, our UHR-SD-OCT findings not only confirmed the earlier observations but also gave new insights about the possible pathological mechanism(s) involved in the retinal damages in the ARB.

The horizontal diameter of the large HRDs in the disrupted EZ was approximately 33 μm and that of the small HRDs in the outer segment were approximately 13 μm (Fig. 3). The size of small HRDs observed in the ARB was almost the same as that in OMD.25 Because the horizontal diameter of HRDs was much larger than that of the normal cone inner segments in the paramacular regions (approx. 7 μm) or the cone outer segments at the fovea (approx. 1–2 μm),26,27 both the HRDs in the disrupted EZ and outer segment observed in the UHR-SD-OCT images most likely do not represent individual photoreceptors but clusters of cone elements or abnormally expanded individual cone photoreceptors. The small HRDs in the outer segments may be packets of abnormal outer segment disks that changed their orientation in the process of phagocytosis, or they may be phagosomes being phagocytosed by the apical processes of the RPE just below the outer segment tips.22,30,31 The abnormal phagocytosis of the outer segments during the disease process produced the cluster of rotated packets of outer segment disks which had abnormal hyperreflectivity in the outer segment region. The small HRDs may also be due to the increased reflectivity of the outer segments alone due to mechanical stress22 or degenerative changes.33 One possible source of the larger HRDs could be abnormal photoreceptor inner segment ellipsoids that have lost their connection with the normal outer segments. These ellipsoid particles may not contain normally functioning mitochondria and will lead to degeneration by apoptosis. The region with degenerated ellipsoids will lead to a further loss of nuclei and a thinning of ONL (Fig. 3E).

4.2. Common and distinct features in cases with ARB and OMD

In an earlier study with UHR-SD-OCT, we showed that during the early stages of OMD, the outer segments were hyperreflective with small HRDs aligned like icicles or string of pearls.22 In addition, there was a disruption of the EZ which then appeared as clusters of larger HRDs22 similar to the structural changes during the photoreceptor atrophy in ARB. However, the processes were distinctly different between the two diseases in other aspects. For example, the changes in the cone IZ and outer segments preceded that of the EZ in ARB, but the changes of the EZ and RPE were preserved in the later stages of OMD although the EZ was more blurred than in normal eyes. And third, the photoreceptor atrophy was followed by RPE disruption and inner retinal atrophy in ARB, but both the RPE and inner retinal structures were normally preserved in the later stages of OMD.

These differences can be explained by the pathological mechanisms of these diseases. In ARB, there is a dysfunction of the RPE leading to a
A horizontal B-scan image of the right eye of a 33-year-old woman. In the expanded image, the EZ, cone IZ, and rod IZ are preserved in the perimacular region (yellow arrows), but in the region with shallow retinal detachment, the cone IZ is absent (white arrow), and the outer segments are hyperreflective and elongated with small hyperreflective dots (HRDs) appearing like icicles or string of pearls hanging on the EZ (asterisks). The peripheral edge of the outer segments is irregularly aligned and there are debris-like hyperreflective spots between the outer segments and the RPE (double asterisks).

Another difference is that both cone and rod photoreceptors are affected in ARB due to a generalized dysfunction of the RPE, whereas only the cone photoreceptors in the macular region are affected and rod photoreceptors are normally preserved until the late stages in OMD.22 The preservation of the rod photoreceptors in OMD could partially explain the preserved EZ band and ONL thickness and sequential changes in the outer retinal structures that were dependent on the severity of the photoreceptor atrophy. Further expanded images of the outer segment regions (asterisks) in the right column in A, F, and B. Left (A x 2.5, UHR-SD-OCT) and central (F x 2.5, Conventional OCT) images show the same region of the same eye for comparison. String-of-pearls appearance of the outer segment was apparent only in the UHR-SD-OCT images (left and right, asterisks). EZ = external limiting membrane; ELM = ellipsoid zone of photoreceptor; IZ = interdigitation zone of photoreceptor. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

In ARB, the cone and rod photoreceptors are affected in ARB due to a generalized dysfunction of the RPE, whereas only the cone photoreceptors in the macular region are affected and rod photoreceptors are normally preserved until the late stages in OMD. The preservation of the rod photoreceptors in OMD could partially explain the preserved EZ band and ONL thickness and normal RPE in the late stage which differs from the eyes with ARB. These suggestions are in accordance with the results of the observations of the UHR-SD-OCT images in cases of ARB that clearly indicated sequential changes in the outer retinal structures that were dependent on the severity of the photoreceptor atrophy. First, there is an increased reflectivity and presence of the small HRDs in the outer segments (Fig. 3A and G left). Then, there were irregularities and a decrease in the
length of the outer segments which leads to a complete loss of the outer segments (Fig. 3B and G right, and 3C). These changes were followed by the disruption of the EZ and the appearance of the large HRDs, i.e., segmented ellipsoid regions, connected to the ELM (Fig. 3C and D). Then, there was a disappearance of the large HRDs and appearance of thin string-like icicles hanging directly from the ELM (Fig. 3D). Finally, there was a local thinning of the ONL and appearance of HRF above the disrupted EZ (Fig. 3E). These reflect very slowly progressive natural course in the BEST1-related maculopathies, and we assume that the serous SRF by itself may not be so harmful to the photoreceptor cells than the hyperreflective materials containing lipofuscin that were often observed in the early stage of VMD and ARB.

4.3. Limitations

Our study has several limitations. First, our study was retrospective and cross-sectional, and longitudinal changes were not observed in individual patients. Data of longitudinal follow-up examinations for individual patients will provide more precise information of the disease progression. Second, we selected patients with the same genetic abnormality because genotypic differences in the same gene may have different courses and severities in ARB. However, to investigate the general mechanism of photoreceptor atrophy in more detail, data from a wider variety of genotypes and diseases of known etiology should be made.

In conclusion, the UHR-SD-OCT device can obtain images of the photoreceptor microstructures that show the structural changes in detail during the progression of ARB. The detailed observation will help clinicians and researchers to determine the mechanism of the retinal pathology and provide important information for the effectiveness of therapeutic strategies.

5. Patient consent

Consent to publish this case report has been obtained from the patients in writing. This report does not contain any personal identifying information.

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The authors have no proprietary or commercial interest in any materials discussed in this article.

Authorship

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

Summary statement

Ultrahigh-resolution SD-OCT images revealed changes of the microstructures of the photoreceptors in eyes with autosomal recessive bestrophinopathy.

Author declaration

We wish to confirm that there are no known conflicts of interest associated with this publication and there has been no significant financial support for this work that could have influenced its outcome. We confirm that the manuscript has been read and approved by all named authors and that there are no other persons who satisfied the criteria for authorship but are not listed. We further confirm that the order of authors listed in the manuscript has been approved by all of us.

We confirm that we have given due consideration to the protection of intellectual property associated with this work and that there are no impediments to publication, including the timing of publication, with respect to intellectual property. In so doing we confirm that we have followed the regulations of our institutions concerning intellectual property.

We further confirm that any aspect of the work covered in this manuscript that has involved human patients has been conducted with the ethical approval of all relevant bodies and that such approvals are acknowledged within the manuscript.

We understand that the Corresponding Author is the sole contact for the Editorial process (including Editorial Manager and direct communications with the office). He is responsible for communicating with the other authors about progress, submissions of revisions and final approval of proofs. We confirm that we have provided a current, correct email address which is accessible by the Corresponding Author.

Declaration of competing interest

K.T.: Payment for lectures - Santen Pharmaceutical Co., Ltd., Novartis Pharma Co., Ltd., Kowa Company, Ltd., and Senju Pharmaceutical Co., Ltd. Receipt of equipment - Kowa Company, Ltd.

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Appendix A. Supplementary data

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