The IS/OS Junction Layer in the Natural History of Type 2 Idiopathic Macular Telangiectasia

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PURPOSE. To document the progression of a break in the photoreceptor inner segment/outer segment (IS/OS) junction layer and its functional correlates over time in the natural history of type 2 idiopathic macular telangiectasia (type 2 MacTel).

METHODS. Patients with at least 1 year of follow-up were selected from the MacTel Study. En face images were created by manual segmentation of the IS/OS junctional line in volume scans acquired using a spatial-domain optical coherence tomography retinal imaging unit. Retinal sensitivity thresholds were determined using a retinal microperimeter unit. Aggregate retinal sensitivity loss within IS/OS lesions was calculated. Changes over time in an area of IS/OS defects and retinal sensitivity were analyzed.

RESULTS. Thirty-nine eyes of 23 patients (mean age: 62.3 ± 9.2 years) were analyzed. Mean follow-up time was 1.9 years (range: 1–3 years). Mean IS/OS break area at baseline was 0.575 mm² (SE = 0.092, 95% confidence interval [CI]: 0.394–0.756 mm²). The cluster-adjusted mean annual progression rate in IS/OS break area was 0.140 mm² (SE = 0.040, 95% CI: 0.062–0.218 mm², P < 0.001). Mean aggregate retinal sensitivity loss was at baseline 28.56 dB (SE = 5.43, 95% CI: 17.32–39.80 dB, n = 28), a positive correlation with IS/OS lesion area was present (P < 0.001). The mean annual rate of change in aggregate sensitivity loss was 5.14 dB (SE = 1.51, 95% CI: 2.19–8.10 dB, P < 0.001, n = 37), a significant correlation with lesion area increase was found (P = 0.006).

CONCLUSIONS. Both IS/OS break area and rate of enlargement correlate with aggregate retinal sensitivity loss in type 2 MacTel. En face OCT imaging of the IS/OS layer provides a functionally relevant method for documenting disease progression in type 2 MacTel. (Invest Ophthalmol Vis Sci. 2012; 53:7889–7895) DOI:10.1167/iovs.12-10765

Type 2 Idiopathic Macular Telangiectasia (type 2 MacTel) is a retinal disease that affects the juxtafoveal region of both eyes.1–2 The etiology and pathogenesis of the disease are unknown and no proven treatment is currently available.

The diagnosis of type 2 MacTel is based on stereoscopic biomicroscopic and fundus fluorescein angiographic (FFA) observations, including a loss of retinal transparency; dilated, blunted, and right-angled veins; dilated capillaries and telangiectatic vessels in the deep retinal plexus; hyperfluorescence in the angiogram without cystoid edema; and pigment plaques. Fibrosis accompanied by eventual neovascularization, scarring, and atrophy are associated with a loss of visual acuity. Early clinical signs may be quite subtle and challenging to quantify.

The introduction of optical coherence tomography (OCT) imaging has provided a valuable tool for identifying neurodegenerative changes in type 2 MacTel. An overall thinning of the retinal temporal to the fovea; a deformation of the foveal contour, hyporeflective spaces in the inner and outer retina, and a break in the line considered to represent the junctions between the photoreceptor inner and outer segments (IS/OS line)3–6 are commonly found. Disruption of the IS/OS line has been demonstrated to correlate with retinal function loss in several retinal disorders5–7,25 and is considered a useful indicator of photoreceptor integrity and predictor of visual function.

In a previous study we demonstrated that imaging the IS/OS layer en face (in the coronal plane) provides a functionally relevant method for quantifying the extent of outer retinal abnormalities as well as for assessing their topographic relationships in type 2 MacTel.26 The aim of the present study was to document the progression of the abnormalities of the IS/OS layer and retinal function over time using en face spectral domain (SD)-OCT imaging and mesopic microperimetry.

METHODS

Patients

Thirty-nine eyes of 23 patients, ranging in age at baseline from 46 to 76 years (mean = 62.3 years, SD = 9.2 years; 11 males, 12 females), were
density. RPE, retinal pigment epithelium.

B-scans within a 6 × 6-mm retinal area, with a resolution of 512 A-scans per B-scan, were acquired using an SD-OCT retinal imaging unit (Topcon 3D-OCT1000; Topcon Medical Systems, Inc., Oakland, NJ). A total of 105 OCT volume scans were processed. The Topcon Q factor (reflecting signal strength) ranged from 30.12 to 83.54 (mean = 57.50, SE = 1.44 95% confidence interval [CI]: 54.68–60.32). Motion artifacts due to microsaccades and/or drift parallel to the B-scans present in 11 volume scans were corrected. Six scans with incorrigible motion artifacts (e.g., where disparate retinal areas were scanned into a volume due to microsaccades with a vector at an angle to the direction of the B-scans) were discarded. Uneven field illumination was noted in 7 scans.

**SD-OCT Image Processing**

The procedure used in this study for creating and processing en face images has been described previously. Briefly, the IS/OS line was segmented manually, using dedicated 3D image-analysis software (Visage Imaging Amira v5.3.3; Pro Medicus Ltd., Richmond, Victoria, Australia); en face images exported in grayscale using orthogonal projection were resampled using an image editing program (Adobe Photoshop CS5 Extended; Adobe Systems Inc., San Jose, CA); a 3 × 3 pixel normal filter was applied to reduce noise emanating from high-frequency random variation and from slight misalignments and variation in reflectivity between individual B-scans within the SD-OCT volume. Delineation of the IS/OS lesions was performed manually. The IS/OS break area and the radial distance of the nearest lesion edge from the anatomic center of the fovea were measured, expressed in pixels.

**Functional Testing**

Monocular best-corrected visual acuities (BCVAs) were determined according to a standardized protocol, using Early Treatment Diabetic Retinopathy Study visual acuity charts at a distance of 4 meters. Scoring of the test was based on the number of letters read correctly. Possible scores ranged from 0 (Snellen equivalent < 20/800) to 100 (Snellen equivalent 20/12). Fundus-correlated automated microperimetry was performed using a retinal micrometer unit (Nidek MP1 and Navis software version 1.7.3; Nidek Technologies, Albignasego, Italy), following pupil dilatation with 1.0% tropicamide (Mydriacyl Eye Drops; Tocris Bioscience, Bristol, UK) and 2.5% phenylephrine hydrochloride and 5 minutes of visual dark adaptation. The technique has been described previously. Results are reported in decibels.

**Data Analysis**

An approximate calibration of distances within the en face OCT image to metric units was performed based on the uniform 6-mm width of the scan raster. En face SD-OCT images and micropimetric retinal sensitivity threshold data were superimposed over images of the fundus and adjusted to attain exact correspondence. To reduce bias from conditions unrelated to MacTel, only micropimetric data from test points within the central 10° of the MP1 grid were considered in calculations. For assessing change over time in retinal function, aggregate sensitivity loss was calculated: The mean of retinal sensitivity values measured within the central 10° of the grid at test points not within the area of the IS/OS break was calculated and considered the background sensitivity. Aggregate loss was defined as the sum of deviations from the background sensitivity of values measured at test points within the area of the IS/OS break. Annual rate of change in the lesion area and aggregate loss were calculated and compared.

**Statistical Methods**

A value of $P < 0.05$ was accepted as statistically significant. Since some participants in this cohort have both eyes included (clustered data), analytic methods accounting for the correlation between eyes
(marginal generalized estimating equations [GEE]) were used. In the GEE, dependence within clusters is treated as a nuisance parameter and inferences are predicted for population average effects. The least-square means, SEs, and 95% CI values from the models are provided. All analyses were conducted using commercially available statistical software (SAS version 9.02; SAS Institute, Cary, NC).

**RESULTS**

**Phenotype**

Mean area of the IS/OS break accounting for correlation between eyes for those participants with two eyes at baseline was 0.575 mm$^2$ (SE = 0.092, 95% CI: 0.394–0.756 mm$^2$). The cluster-adjusted mean annual progression rate in the area of IS/OS break was 0.140 mm$^2$ (SE = 0.040, 95% CI: 0.062–0.218 mm$^2$, $P < 0.001$). Concordance of the rate of expansion of the area of disruption between fellow eyes was low ($\rho = -0.20$, $P = 0.46$, concordance correlation coefficient = -0.14, $n = 16$). Expansion was observed both along distinct edges as well as through gradual thinning of the IS/OS (Fig. 1).

Within the area of the lesion, a decrease in size of hyporeflective spaces in the outer retina was observed with a simultaneous increase in thickness and reflectivity of the "collapsed layers," as described in Figures 2B–D. Topographically, the IS/OS break initially located on the temporal side of the foveal center, and islands of preserved IS/OS may also present as areas with high reflectivity. Images on the left were taken in 2008, on the right in 2010. Orange lines mark the position of respective B-scan within the en face image. (B, D) In the en face images, a significant increase in break area is accompanied by a reduction in the area of the cross-sections of the outer retinal empty spaces (nearblack areas) and the enlargement of a highly reflective area corresponding in B-scans (A, C) to outer retinal atrophy and a pathologic vertical restructuring of the retina. (F, H) Although the overall increase in the break area is minor, the progression of retinal restructuring is detectable in the en face image, the shape of the outer empty space changes, and the area and optical density of the collapsed layers increase (compare with [E] and [G]).

**FIGURE 2.** Progression of the IS/OS break in 2 years. Within the area of the IS/OS break, insular variations of backscatter may be present. Low optical density corresponds to the cross-sections of outer retinal atrophic cavities, a high reflectivity to cross-sections of an abnormal retinal tissue with a vertical orientation. In these, retinal layers between the outer plexiform layer and the RPE seem to be absent and the structurally disorganized remaining retinal layers give the impression of “collapsing” onto the RPE. This structure was always seen initially on the temporal side of the foveal center. Islands of preserved IS/OS may also present as areas with high reflectivity. Images on the left were taken in 2008, on the right in 2010. Orange lines mark the position of respective B-scan within the en face image. (B, D) In the en face images, a significant increase in break area is accompanied by a reduction in the area of the cross-sections of the outer retinal empty spaces (nearblack areas) and the enlargement of a highly reflective area corresponding in B-scans (A, C) to outer retinal atrophy and a pathologic vertical restructuring of the retina. (F, H) Although the overall increase in the break area is minor, the progression of retinal restructuring is detectable in the en face image, the shape of the outer empty space changes, and the area and optical density of the collapsed layers increase (compare with [E] and [G]).
yearly loss was 0.267 letters (SE = 0.079 letters, 95% CI: 0.143 to 0.39 letters, \( P = 0.65 \)). From a generalized linear model (GLM) accounting for clustering, the relationship with the annual rate of change in the area of disruption did not meet statistical significance (regression model \( \beta = 0.115 + 0.032 \times \text{[annual rate of change in BCVA]} \); \( P \) value for slope = 0.19). The slope, interpreted as an increase for each unit annual rate of change in BCVA, is associated with an estimated increase of 0.032 mm\(^2\) in the mean annual rate of change in area of disruption.

Mean aggregate MP1 retinal sensitivity loss at baseline was 28.56 dB (SE = 5.43, 95% CI: 17.52–39.80 dB, \( n = 28 \)). From a GLM the model was computed as \( 0.183 + 0.013 \times \text{[MP1 retinal sensitivity]} \), with a \( P \) value for the slope of <0.001. The slope, interpreted as an increase of 1 dB in baseline MP1 retinal sensitivity loss, is associated with an estimated increase of 0.013 mm\(^2\) in the mean baseline IS/OS break measurement. The mean annual rate of change in aggregate sensitivity loss was 5.14 dB (SE = 1.51, 95% CI: 2.19–8.10 dB, \( P < 0.001 \), \( n = 37 \)). From a GLM accounting for clustering, the relationship with the annual rate of change in the area of disruption met statistical significance (regression model \( \beta = 0.079 + 0.006 \times \text{[MP1 aggregate retinal sensitivity loss annual rate of change]} \); \( P \) value for slope = 0.006). The slope, interpreted as an increase for each unit annual rate of change in MP1 aggregate retinal sensitivity loss annual rate of change, is associated with an estimated increase of 0.006 mm\(^2\) in the mean annual rate of change in the area of disruption.

**DISCUSSION**

We undertook this study to identify a sensitive, functionally relevant outcome to monitor progression of type 2 MacTel. Visual acuity is a poor measure since the disease may become advanced in the perifoveal region without affecting visual acuity. Microperimetry is a functional measure, but it is subjective. We have demonstrated in this study that en face imaging of the area of IS/OS disruption correlates strongly with loss of macular sensitivity measured by microperimetry, and that expansion of the area of photoreceptor dysfunction correlates with progressive loss of macular sensitivity. En face imaging of the area of the IS/OS break may therefore be considered as an outcome measure for clinical trials of interventions for type 2 MacTel.

The main change over time in the morphology of a manifest break in the IS/OS junction layer is an increase in its area. This was observed both by progression of distinct edges as well as a gradual diffuse fading over a larger area (Fig. 1). The mean annual increase in area of 0.140 mm\(^2\) in our cohort was easily detectable at the currently available resolution of the en face images. An outright recovery of a clear IS/OS break was not seen, although a variation between scans in break shape was noted, possibly due to the optical properties of the layer or the OCT system used. We previously found variation in break area size between fellow eyes to be smaller than that between patients.\(^{26}\) However, a similar significant symmetry between left and right eyes of the increase in IS/OS area over time could not be demonstrated in this cohort. This may be attributable to the relatively short follow-up time in this study.

We found a significant and close correlation between area size of an IS/OS break and aggregate mesopic retinal sensitivity loss. Furthermore, an enlargement of the break area over time was associated with an increase in aggregate sensitivity loss, although the correlation of break area change with BCVA was not significant.

Within the area of the break, cross-sections of outer retinal cavities, islands of preserved IS/OS, or cross-sections of an abnormal retinal tissue with a vertical orientation may be present ("collapsed layers"; see Fig. 2). These features are
smaller and some are scattered and indistinct, such that their
time, not amenable to accurate quantification.

However, as a trend it was noted that an increase in break area
was often accompanied by a progression of retinal restructuring
(“collapsed layers”), with a simultaneous overall decrease
of the adjacent outer atrophic spaces.

Based on our own and previous observations by other
authors3,6,25 a hypothetical sequence of neurodegenerative
signs in MacTel may be outlined: Inner hyporeflective spaces
near the foveal center appear, initially convex both anteriorly
and posteriorly, distending the retina, the IS/OS junction layer
deviates toward the retinal pigment epithelium (RPE) (Fig.
3A). Subsequently, the convexity disappears, leaving the
impression of an inner atrophic cavity (Figs. 2G, 5A). Minor
vertically oriented oblong spaces along the border of the
outer nuclear and outer plexiform layers (OPL and ONL) may
also be present. Temporal to the foveal center, the IS/OS
signal attenuates and breaks and outer retinal spaces appear,
with a shape suggesting atrophy (lateral boundaries in B-
scans near the RPE appear vertical, in line with photorecep-
tor morphology). Focally, the ONL thickness decreases, layers
internal to the ONL deviate toward the RPE/choroid, become
disorganized, and “collapse” onto the RPE. As the collapse
widens, outer empty spaces shrink. Some authors character-
ize the “collapse” as a “contraction” of retinal layers.5
Indeed, a contraction in the plane of the retina centered on
these foci is frequently seen in fundus images (see Fig. 4),
along with apparent anastomoses between branches of the
supero- and inferotemporal venous systems. A vertical
component is also conceivable. We noted in en face images
that the cross-section of the collapsed tissue often colocalized
with the tips of blunted veins in the fundus image (Fig. 5).

We were unable to clearly demonstrate blood vessels within
the collapsing layers, possibly due to the relatively low
resolution of the en face images. However, aberrant blood
vessels within the outer retina, near the foveal center, have
been reported previously,25,36 and vascular involvement in the
“collapse” is possible. Although the pigment plaques
characteristic of type 2 MacTel are in the mid layers of the
retina, smaller foci were observed in the deeper layers of the
retina also in this cohort. Vessels with a vertical disposition
combined with the propensity of pigment for propagating
along vessels37 may offer an explanation for pigment plaque
genesis. Progression of the phenotype, however, may not
necessarily always pass through the same sequence of events
in all cases.

We acknowledge some limitations of this study. Sample size
and follow-up time were limited. For an accurate calibration of
measurements within OCT images, the axial length and the
refractive power of the eye must be taken into consideration.
In our study these data were not collected; “typical” values
were used as provided by the manufacturer. However, we
compared each individual eye over time in terms of structure
and function. Unless there is a significant change in the main

Figure 4. Focal lateral contraction of the retina temporal of the foveal center in type 2 MacTel. (A) In 2008, in an early-phase FA image, dilated veins from the superior and inferior temporal branches and a dilated deep capillary plexus are evident. (B) By 2010, pigment appears, the retina around it contracts, radial vessels straighten, and point toward the tissue surrounding the pigment deposit. (C) The tissue corresponds in the en face OCT image to the cross-section of “collapsed” layers.

Figure 5. “Collapsed” retinal layers with indications of vascular involvement. (A) B-scan OCT image. (B) OCT en face image of the IS/OS junction layer, an orange line marks the location of the B-scan in image (A); a white arrow marks a dilated vein. (C) Red-free image of the same retinal location. The cross-section of the collapsing layers colocalizes with the tip of a dilated vein (arrow) that changes caliber abruptly. (D) 3D perspective rendering of the OCT data; note the sunken appearance of the IS/OS layer around the break in the temporal part of the fovea externa and the vertical, optically dense tissue extending between the inner and outer layers.
parameters (due to, e.g., cataract or refractive surgery, a change in the refractive index of the lens due to maturing cataract, or an increase in axial length in progressive myopia), differences in area measured are expected to emanate from the progression of the lesion alone. The low test point density of the MP1 grid used did not permit a detailed analysis of whether function loss is associated with the IS/OS break itself or lesions within. It was noted, however, that high loss was consistently present over “collapsed layers.” A progression of the collapsing layers may have a functional relevance that would not be fully reflected in increasing lesion area size alone. Furthermore, we used volume scans from a commercially available SD-OCT machine without a real-time eye-tracking system. Fixation stability in type 2 MacTel patients is affected early and motion artifacts may be a source of error in 3D analysis. In en face images, near the foveal center, in the absence of vascular landmarks, it may not always be possible to detect these. OCT devices with active eye tracking have a significant advantage.

SD-OCT and other recent imaging techniques offer possibilities for characterizing the neurodegenerative aspects of the disease3,5,6,25,38,39 and may provide new morphologic landmarks for refining the staging system, especially in early disease.40 Break area in en face images of the IS/OS layer is a quantifiable morphologic sign that correlates with function also in its progression over time, even at stages of the disease where vascular signs may be less sensitive indicators. The IS/OS break and “collapsing tissue” may potentially be new landmarks for following disease progression in the natural history as well as in interventional studies of MacTel.

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References


APPENDIX

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