Morphologic Choroidal and Scleral Changes at the Macula in Tilted Disc Syndrome with Staphyloma Using Optical Coherence Tomography

Ichiro Maruko, Tomohiro Iida, Yukinori Sugano, Hiroshi Oyamada, and Tetsuju Sekiryu

PURPOSE. To evaluate the macular choroidal and scleral changes in tilted disc syndrome (TDS) with staphyloma using optical coherence tomography (OCT) to determine the mechanism of serous retinal detachment (SRD) formation.

METHODS. All eyes underwent fluorescein (FA) and indocyanine green angiography (ICGA) in this retrospective, observational study. Enhanced-depth imaging (EDI) OCT and prototype high-penetration (HP) OCT were used to examine the choroid and sclera, respectively, at the upper and lower optical areas and the subfovea on vertical OCT sections.

RESULTS. Twenty-four eyes with TDS and inferior staphyloma were included. FA showed the macular area with the superior edge of staphyloma had a granular hyperfluorescent pattern and ICGA showed belt-like hypofluorescence. OCT showed SRDs in seven eyes. The mean EDI-OCT choroidal thicknesses in 19 eyes were: upper area, 211 ± 79 µm; subfovea, 153 ± 70 µm; and lower area, 158 ± 42 µm. The mean subfoveal and lower choroid were significantly (P < 0.01 for both) thinner than the upper area. The mean HP-OCT scleral thicknesses in 14 eyes were: upper area, 414 ± 36 µm; subfovea, 493 ± 40 µm; and lower area, 398 ± 83 µm. The subfoveal sclera was significantly (P < 0.01) thicker than the others.

CONCLUSIONS. The subfoveal choroid was relatively thin and the subfoveal sclera thickened in TDS with a staphyloma edge at the macula. The area with retinal pigment epithelial (RPE) atrophy was hyperfluorescent on FA; choriocapillaris occlusion was hypofluorescent on ICGA. Characteristic anatomic subfoveal scleral alterations might lead to a thinner choroid and inhibit choriocapillaris outflow; a secondary RPE disorder subsequently could cause SRDs. (Invest Ophthalmol Vis Sci. 2011; 52:8763–8768) DOI:10.1167/iovs.11-8195

Although the tilted disc syndrome (TDS) usually is associated with good visual prognosis, foveal complications, such as serous retinal detachment (SRD), have been reported.1–7 Staphyloma in TDS often is observed in the inferior ocular area, and the superior edge of the staphyloma sometimes involves the macula.6,9–12 The superior edge of the inferior staphyloma at the fovea in TDS is characterized by a window defect on fluorescein angiography (FA) due to retinal pigment epithelial (RPE) atrophy and hypofluorescence due to choriocapillaris occlusion on indocyanine green angiography (ICGA).6 These anatomic changes at the superior edge of the staphyloma might lead to foveal weakness and subsequent complications. Choroidal neovascularization (CNV) or polypoidal choroidal vasculopathy (PCV) in TDS was also reported as one of the complications due to anatomic changes at the macula.6,13

Optical coherence tomography (OCT) is a noninvasive, advanced imaging technique for viewing the fovea that is critical for diagnosis and follow-up. In fact, SRDs at the fovea in TDS were clearly visualized by OCT for the first time.5,6 Spectral-domain OCT (SD-OCT) is a high-speed, high-resolution technology that provides detailed images of the retinal structures in a short time. However, no study has investigated the choroidal and scleral changes in TDS using SD-OCT.

A new method for visualizing the choroid, enhanced-depth imaging OCT (EDI-OCT), was reported.14 EDI-OCT can visualize the sclera in cases with a thinner retina and choroid, such as in pathologic myopia.15 High-penetration OCT (HP-OCT) is another way to observe the choroid using a 1-µm wavelength.16–20 Although the device is not commercially available, it is expected to visualize both the choroid and sclera.

The present study evaluated the choroidal and scleral changes using EDI-OCT or HP-OCT to elucidate the mechanism of SRD development in TDS with the superior edge of staphyloma at the fovea.

METHODS. This retrospective study followed the tenets of the Declaration of Helsinki. The institutional review board at Fukushima Medical University School of Medicine approved this study that included OCT observation of eyes with macular and retinal disorders, observational study of age-related macular degeneration and similar disorders (including TDS), and use of the prototype HP-OCT not commercially available.

The present study involved the characteristic inferonasal tilting of the oval optic disc with a congenital inferonasal crescent with the superior edge of staphyloma at the fovea. The clinical examinations to diagnose TDS included measurement of the best-corrected visual acuity (BCVA), slit-lamp biomicroscopy with a contact or noncontact lens, indirect ophthalmoscopy, and digital FA and ICGA (TRC-50IX/IMAGEnet H1024 system, Topcon, Tokyo, Japan). The BCVA was measured with a Japanese standard decimal visual chart, and the logarithm of the minimum angle of resolution (logMAR) scale was used for statistical analysis. The spherical equivalent (SE) refractive error using an auto refractometer (Nidek, Gamagori, Japan) and the axial length using a biometer (IOL-Master; Carl Zeiss Meditec, Dublin, CA) were measured. Most eyes were examined with a commercially-available optical coherence tomograph (Heidelberg Spectralis; Heidelberg Engineering, Heidelberg, Germany) and the prototype HP-OCT with the 1060-nm wavelength (Topcon, Tokyo, Japan).

EDI-OCT. We observed the choroid, defined as the area between the outer RPE surface and the inner scleral surface, on vertical sections using EDI-
OCT, in which the OCT device (Heidelberg Spectralis) is positioned close to the eye to obtain an inverted image. Each section was obtained using eye tracking, and 100 scans were averaged to improve the signal-to-noise ratio. The standard scanning length can be 9 mm. We measured the choroidal thicknesses at the subfovea and the upper and lower points 1.5 mm from the foveal depression on the vertical OCT lines passing through the fovea (EDI-OCT) (Figs. 1, 2).

Prototype HP-OCT

We observed the choroid on vertical sections and the sclera, defined as the hyperreflective area from the inner scleral surface, using HP-OCT. This instrument can average up to 50 images to improve the signal-to-noise ratio and enhance the choroid and sclera by movement of the reference mirror to change the focus similar to EDI-OCT when positioned close to the eye. When the full-thickness sclera could not be observed, the deepest hyperreflective point was used as the measurement value. No-reflection as the dark area behind the scleral hyperreflection was recognized as the existence of the different structures including the connecting tissues, vessels, muscles, or orbital fat. The standard scanning length can be 12 mm, which is longer than the scanning length with EDI-OCT. We measured the choroidal and scleral thicknesses at the subfovea and the upper and lower points 1.5 mm.

**Figure 1.** A 67-year-old woman (patient 3) has TDS in the right eye. The BCVA is 0.50 (20/40 Snellen; 0.30 logMAR), and the spherical equivalence is −15.25 diopters. The axial length is 27.33 mm. (A) A grayscale fundus photograph of the right eye shows the tilted optic disc with a crescent border and staphyloma from the inferior disc. (B) FA in the right eye shows the granular pattern of the hyperfluorescence at the macular area with a superior edge of staphyloma. (C) Late-phase ICGA shows belt-like hypofluorescence larger than the hyperfluorescence on FA. (D) EDI-OCT images show that the choroidal thickness on vertical section is 119 μm at the upper area (U), 93 μm at the subfovea (F), and 144 μm at the lower area (L). (E) The prototype HP-OCT shows that the choroidal thicknesses on vertical section are 116 μm at the upper area (U), 73 μm at the subfovea (F), and 139 μm at the lower area (L). The dotted line indicates the posterior edge of the sclera defined as the hyperreflective area.

**Figure 2.** A 45-year-old man (patient 7) with TDS in the right eye. The BCVA is 0.30 (20/67 Snellen; 0.52 logMAR), and the spherical equivalence is −3.875 diopters. The axial length is 24.56 mm. (A) A grayscale fundus photograph of the right eye shows the tilted optic disc with a crescent border and staphyloma from the inferior disc. (B) FA in the right eye shows slight hyperfluorescence at the macular area with the superior edge of staphyloma. (C) Late-phase ICGA shows belt-like hypofluorescence larger than the hyperfluorescence on FA. (D) An EDI-OCT image shows that the choroidal thicknesses on vertical section are 151 μm at the upper area (U), 98 μm at the subfovea (F), and 124 μm at the lower area (L). (E) A prototype HP-OCT image shows that the choroidal thicknesses on vertical section are 146 μm at the upper area (U), 89 μm at the subfovea (F), and 122 μm at the lower area (L). The dotted line indicates the posterior edge of the sclera defined as the hyperreflective area.
from the foveal depression on the vertical OCT lines passing through the fovea (HP-OCT) (Figs. 1, 2).

The reported measurements obtained from the OCT images represented the average measurements obtained by three observers (IM, YS, HO). The visual acuities (VAs) are expressed as the decimal and logMAR equivalents, the standard Snellen VA values also were recorded. The results of the measurement of the choroidal and scleral thicknesses were analyzed using the Wilcoxon signed rank test. \( P < 0.05 \) was considered statistically significant.

**RESULTS**

Twenty-four eyes of 15 patients (2 men, 11 women; mean age, 56.2 years) were diagnosed with TDS with the superior edge of staphyloma at the fovea. The optic disc in all cases was elevated to the upper (superotemporal) area, with a crescent at the inferior or inferonasal margin. The mean BCVA was 0.54 (20/37 Snellen; 0.27 logMAR), and the mean spherical equivalence (SE) was \(-4.47 \) dipters (D). All eyes underwent FA and ICGA. FA showed band-shaped granular hyperfluorescence corresponding to the atrophic band with the superior edge of staphyloma in all eyes. Early phase of ICGA showed an asymmetric choroidal vascular pattern between the upper and the lower fundus across the superior edge of staphyloma. In the inferior staphyloma, less number and smaller caliber of the choroidal vessels were delineated in ICGA. The superior border of the inferior staphyloma showed band-shaped hypofluorescence during ICGA throughout the angiographic phase. The band-shaped hypofluorescent area on late-phase of ICGA was larger than that of the hyperfluorescent seen on FA. There was no lesion suggestive of CNV or PCV in the present study. Figures 1 and 2 show representative cases of TDS with the superior edge of staphyloma at the fovea. Table 1 shows the patient data including the choroidal and scleral thicknesses.

Among all cases, 19 eyes of 10 patients were examined using EDI-OCT. Seven eyes had an SRD. The mean axial length was 25.05 ± 1.26 mm. The mean choroidal thickness on the vertical EDI-OCT images was 211 ± 79 \( \mu \)m at the upper area, 153 ± 70 \( \mu \)m at the subfovea, and 158 ± 42 \( \mu \)m at the lower area. The mean choroidal thicknesses at the subfovea and the lower area were significantly \(( P < 0.01, \) for both comparisons) thinner than the upper area. The mean subfoveal choroidal thickness in eyes with an SRD was slightly thicker than in eyes without an SRD; however, the difference was not significant \(( P = 0.61 \) at the upper area; \( P = 0.49 \) at the subfovea; \( P = 0.88 \) at the lower area). The mean subfoveal choroidal thicknesses in eyes with an SRD were slightly thicker than in eyes without an SRD on EDI-OCT \((176 ± 80 \, \mu m; \, 140 ± 65 \, \mu m; \, P = 0.27)\).

Fourteen eyes of nine patients were observed with EDI-OCT and HP-OCT. Seven eyes had an SRD. The mean axial length was 24.89 ± 1.05 mm. The mean choroidal thicknesses on EDI-OCT vertical sections were 212 ± 87 \( \mu \)m at the upper area, 148 ± 76 \( \mu \)m at the subfovea, and 154 ± 43 \( \mu \)m at the lower area. The mean choroidal thicknesses at the subfovea and the lower area were significantly \(( P < 0.01 \) for both comparisons) thinner than the upper area. The mean subfoveal choroidal thickness in eyes with an SRD was slightly thicker than in eyes without an SRD; however, the difference was not significant \((176 ± 80 \, \mu m; \, 140 ± 65 \, \mu m; \, P = 0.27)\). The mean subfoveal choroidal thicknesses in eyes with an SRD were slightly thicker than in eyes without an SRD on EDI-OCT \((176 ± 80 \, \mu m; \, 119 ± 65 \, \mu m; \, P = 0.08)\). The mean subfoveal choroidal thicknesses in eyes with an SRD were slightly thicker than in eyes without an SRD on HP-OCT \((174 ± 92 \, \mu m; \, 115 ± 62 \, \mu m; \, P = 0.27)\). The mean scleral thicknesses on vertical HP-OCT images were 414 ± 36 \( \mu \)m at the upper area, 493 ± 40 \( \mu \)m at the subfovea, and 398 ± 83 \( \mu \)m at the lower area. The subfoveal sclera was significantly thicker than the other structures \(( P < 0.01, \) for both comparisons). Full thickness sclera was observed at the outside of the foveal area in all 14 eyes, and measurement values defined as the deepest hyperreflective point at the subfovea were thicker than at the outside of the foveal area even in eyes with invisible full thickness sclera at the foveal area. There was no significant difference between the mean subfoveal scleral thickness on HP-OCT in eyes with and without an SRD \((481 ± 45 \, \mu m \, \text{and} \, 505 ± 33 \, \mu m, \text{respectively}; \, P = 0.34)\).

**DISCUSSION**

In the present study, the subfoveal choroid was thinner than that in the outside area of staphyloma, and the subfoveal sclera was thicker than that in the other areas. The characteristic anatomic changes including the subfoveal scleral thickening in TDS might induce choroidal thinning and abnormal choroidal circulation at the fovea; secondary RPE atrophy could cause breakdown of the blood-retinal barrier and a subsequent SRD.

In the present study for TDS, HP-OCT showed almost the same choroidal measurements as EDI-OCT. Ikuno et al. reported the reproducibility of choroidal thickness using both EDI-OCT and HP-OCT devices. Intersystem intraclass correlation coefficient showed the high correlation values of 0.921. Thus, both OCT systems can evaluate the choroidal thickness measurements as the same values, and these are also proved in eyes with TDS.

In patients with TDS, moderate myopia is common.22 The present study supported this (e.g., the mean spherical equivalence was \(-4.5 \) dipters and the mean axial length was 25 mm). Lacquer cracks, representing breaks in Bruch’s membrane, are sometimes seen in the posterior pole in eyes with pathologic myopia.23,24 FA showed hyperfluorescence and ICGA showed well-delineated hypofluorescence corresponding to the lacquer cracks.23 These angiographic features of the lacquer cracks in eyes with pathologic myopia are similar to those of the superior edge of staphyloma in TDS. In the present study, FA showed the hyperfluorescence and ICGA showed the hypofluorescence at the superior border of the inferior staphyloma in all cases. These may indicate less number and smaller caliber of choroidal vessels and subsequent ischemic changes at the staphyloma edge provide RPE atrophy. Thus, the choroidal expansion of the staphyloma may lead to RPE atrophy in TDS. However, a macular hole retinal detachment, retinal schisis, simple subretinal hemorrhage, and even lacquer cracks were not commonly observed in TDS. This may indicate that another mechanism of myopia is associated with the pathophysiology of TDS.

Recently, a new method to visualize the choroid, EDI-OCT, was described.14 Margolis and Spaida5 reported that the subfoveal choroidal thickness in normal subjects was 287 \( \mu m \). We reported a subfoveal choroidal thickness of 250 \( \mu m \).26 Fujiwara et al.27 reported that the subfoveal choroidal thickness in highly myopic eyes was 93.2 \( \mu m \). In the present study, the subfoveal choroidal thickness in TDS was 153 \( \mu m \) on EDI-OCT, which might indicate that the subfoveal choroidal thickness in TDS is thinner than in normal eyes and thicker than in highly myopic eyes. Subfoveal choroidal thickness in typical central serous chorioretinopathy (CSC) has been reported to be thicker than normal in recent EDI-OCT studies.26,28 In the present study, the subfoveal choroid in TDS was not thick in eyes with TDS; however, subfoveal choroid in seven eyes with an SRD was slightly, but not significantly, thicker than in eyes without a detachment in TDS. Although FA did not show focal leakage in TDS, the choroidal fluid may flow from the choroid to the subretina through the damaged RPE. Thus, the pathophysiology of SRD in TDS may not be identical with that in CSC, but there can be some similarities.
TABLE 1. Clinical Characteristics of Patients with Tilted Disc Syndrome in the Current Study

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<th>SRD</th>
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Mean* M 2:F 11 56.2 0.69 (0.16) -4.47 25.05 (1.26) 7 211 (79) 153 (70) 158 (42) 211 (87) 144 (81) 156 (49) 414 (36) 493 (40) 398 (83)

Choroidal and scleral thicknesses are measured at the subfovea, upper, and lower points of 1.5 mm from the foveal depression. F, female; M, male; NA, not available.

*Values are mean (SD) except for BCVA (logMAR), and Sex, and SRD, which are totals.
A dome-shaped macula is characterized by a convex protrusion of the macula within the staphyloma in highly myopic eyes seen on OCT. Gaucher et al. reported that an SRD was present in 67% (10/15 eyes) of eyes with a dome-shaped macula using conventional (Stratus) OCT (Carl Zeiss Meditec, Inc, Dublin, CA). Imamura et al. evaluated the posterior anatomic structure of dome-shaped maculas using EDI-OCT and reported subfoveal choroidal thinning and subfoveal scleral thickening. Although the sclera can be observed by EDI-OCT in eyes with a thinner retina and choroid in patients with a dome-shaped macula associated with pathologic myopia, it is difficult to evaluate the sclera in cases with TDS with relative myopia on EDI-OCT.

Because HP-OCT has a wavelength that is 1 μm longer than the commercially-available device, HP-OCT is expected to visualize not only the choroid but also the sclera in non-myopic eyes. Using the prototype HP-OCT, the subfoveal sclera in cases of TDS was significantly thicker than the other areas. These results may be similar to scleral thickening in patients with a dome-shaped macula even though the pathogenic mechanisms and clinical conditions differ.

We recognize there are two types of diseases with SRD at the fovea; one is the disease with choroidal thickening such as CSC and Vogt-Koyanagi-Harada disease, and the other is the disease with choroidal thinning such as a dome-shaped macula associated with highly myopic eyes. The former is definitely associated with choroidal abnormalities of choroidal vascular hyperpermeability or inflammatory infiltration. Because the choroid in TDS was thin and choroidal vascular hyperpermeability was not observed on ICGA, the choroid might not be contributed to SRD in TDS. The latter is not fully understood. Imamura et al. reported that subretinal fluid in patients with a dome-shaped macula might accumulate because of impaired choroidal outflow resulting from scleral thickening. We think the similar mechanism to the dome-shaped macula occurs around the foveal area in TDS, thus the choroidal fluid in TDS might not pass through a thickened sclera and could leak into the subretina through the degenerated RPE seen on angiography. Although the subfoveal choroid in eyes with SRD was relatively thinner than in eyes without SRD, the choroidal fluid in TDS with SRD might be partially stored in the choroid because of the obstruction of choroidal outflow. The mechanism of SRD development in TDS might be completely different from CSC.

In the present study, EDI-OCT and HP-OCT showed that the subfoveal choroid was thinner than in the outside area of staphyloma and the subfoveal sclera was thicker than in other areas. Choroidal outflow obstruction and RPE damage seen on OCT and angiography might induce subsequent SRDs. Several limitations of the present study included its retrospective design and the small number of patients. CNV or PCV in TDS is sometimes observed as a complication. Although we do not have such a case with CNV or PCV in the present study, it is important enough to cause the visual loss in TDS. Further study will need to elucidate the pathogenesis of CNV or PCV in TDS. Nevertheless, no previous study has visualized the choroid and sclera in TDS using EDI-OCT and HP-OCT.

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References


