Parafoveal Microvascular Alterations in Ocular and Non-Ocular Behçet’s Disease Evaluated With Optical Coherence Tomography Angiography

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Purpose. To compare quantitative optical coherence tomography angiography (OCT-A) measurements of the parafoveal microvasculature in retinal capillary plexuses among Behçet uveitis (BU) patients, non-ocular Behçet’s disease (NOBD) patients, and healthy volunteers (HVs).

Methods. Sixty-eight subjects were enrolled in this prospective observational cross-sectional study. OCT-A imaging was performed using the Heidelberg Engineering Spectralis OCT. A custom algorithm was developed to calculate the vessel density (VD) in three retinal vascular layers: deep capillary plexus, intermediate capillary plexus, and superficial vascular plexus. The foveal avascular zone (FAZ) and acircularity index were calculated for the whole retinal vascular complex.

Results. We analyzed one eye from 21 BU patients (age, 51 ± 10 years), 23 NOBD patients (age, 48 ± 14 years), and 22 HVs (age, 44 ± 13 years). One-way multivariate analysis of covariance showed a statistically significant difference in VD among the three groups when combining the layers after controlling for scan quality (P < 0.001). The VD was lowest in the BU group and highest in the HV group in all layers. The FAZ area was also statistically significant different among the groups (P < 0.005), with the largest FAZ areas in BU patients and smallest FAZ areas in the HV group. However, no statistically significant difference was found for the acircularity index.

Conclusions. The parafoveal microvasculature is affected not only in BU patients but also in NOBD patients. Most deviations in the retinal microcirculation in Behçet patients were found in the deeper layers of the retina by using the quantitative VD measurement.

Keywords: Behçet’s disease, OCT angiography, vessel density, foveal avascular zone, acircularity index

Behçet’s disease is a systemic autoinflammatory vasculitis affecting multiple organs.1,2 Ocular involvement occurs in approximately 50% to 90% of Behçet patients.3 Uveitis in Behçet is characterized by recurrent episodes of intraocular inflammation, most commonly presenting as bilateral panuveitis with retinal vasculitis.1,4,5 Retinal vasculitis can cause vascular occlusions leading to visual deterioration and often requires intensive immunosuppressive treatment.2,5,6,7

Fluorescein angiography (FA) is the golden standard for detecting and monitoring retinal vasculitis in Behçet uveitis (BU).5,8, however, FA has some limitations with regard to visualization of the retinal microvascular circulation. First, due to dye leakage, the microvascular circulation is only visible in the early phases.5 Furthermore, FA has a very poor depth resolution and is unable to distinguish the retinal capillary plexuses separately.5,9 The functional extension of conventional optical coherence tomography (OCT), OCT angiography (OCT-A), enables visualization of the various microvascular retinal plexuses without the disturbance of dye leakage.10-12 OCT-A is even considered superior to conventional FA for visualization of microvascular changes in active BU that involve the posterior segment.13

Previous studies have demonstrated that patients suffering from Behçet’s disease with ocular involvement show hypo- or non-perfused areas on OCT-A in the superficial vascular plexus (SVP) and deep capillary plexus (DCP),13,14 in which the DCP seems most severely affected.13,15 Also, in comparison to healthy eyes, reduced vessel density (VD) and an enlarged foveal avascular zone (FAZ) are seen in Behçet uveitis and are considered indicators for microvascular alterations.6,16,17 Furthermore, it is suggested that FAZ irregularity could be a marker of ocular involvement of Behçet’s disease.16,18 Because FAZ irregularity has only been assessed subjectively, further exploration for quantitative measurement of FAZ irregularity is warranted.
Because Behçet's disease is a systemic vasculitis, we hypothesize that preclinical retinal microvascular changes may occur in patients with non-ocular Behçet's disease (NOBD). Some studies have investigated VD and FAZ area changes in NOBD patients, but have shown inconsistent results.\textsuperscript{5,18–20} VD was reduced in NOBD patients compared to healthy control subjects in three out of the four studies,\textsuperscript{8,10,20} and FAZ area was larger in the NOBD than in healthy controls in only one of the four studies.\textsuperscript{19}

Given the limited number of studies and their inconsistent results mainly due to small sample sizes, further exploration of preclinical retinal microvascular changes in NOBD patients compared to BU patients in a large group and a comparison to healthy volunteers (HVs) is warranted. Also, a quantitative measure for FAZ irregularity as a marker of ocular involvement in BD has not been explored. Therefore, the purpose of this study was to compare quantitative measurements of the parafoveal microvasculature among patients with BU, patients with NOBD, and HVs.

**Methods**

**Study Design and Population**

This prospective observational cross-sectional study was approved by the internal review board of the Rotterdam Eye Hospital and the Medical Research Ethical Committee of the Erasmus University Hospital (Rotterdam, The Netherlands; MEC-2018-050), and it adhered to the tenets of the Declaration of Helsinki. All subjects signed an informed consent prior to participation. Subjects were recruited between July 2018 and June 2019. BU patients were recruited at the uveitis service of the Rotterdam Eye Hospital and patients with NOBD were recruited at the immunology department of Erasmus Medical Center. Behçet's disease was diagnosed based on the diagnostic criteria of the International Study Group for Behçet's disease.\textsuperscript{21} With unilateral involvement, the non-uveitis eye was excluded for analysis. The NOBD group consisted of patients that have never suffered from any form of uveitis. The healthy volunteers were recruited among employees from the Rotterdam Eye Hospital and among the partners and family members of participants in the Behçet's disease groups. They were gender and age matched to Behçet's disease participants. The exclusion criterion for all subjects was a refractive error higher than 4 diopters (D).

**Study Measurements**

All subjects underwent a complete ophthalmic examination, including best-corrected visual acuity (BCVA) measurement, dilated fundus examination, enhanced depth imaging OCT, OCT-A, and slit-lamp examination performed by the ophthalmologist. All participants were checked for signs of active uveitis at the time of the study visit, and non-uveitis participants were also checked for signs of past inflammation (posterior synechiae, old vitreous cells, vascular sheathing, chorioretinal scars\textsuperscript{22}).

**OCT-A Acquisition and Processing**

OCT-A images were acquired with a Spectralis OCT2 (Heidelberg Engineering, Heidelberg, Germany), which has a wavelength of 840 nm and operates at a 40-kHz A-scan rate. The distance between B-scans was 6 μm, and the pattern size (width × height) was 10° × 10° (± 3.0 mm × 3.0 mm), resulting in 512 B-scans per OCT-A image. Scanning was performed by a single operator. Images of both eyes were acquired, but only one eye per participant was included for analysis. The study eye was chosen based on the highest scan quality or, when applicable, on the involvement of uveitis. In case of unilateral involvement in a BU patient, the affected eye was included irrespective of scan quality.

The automated segmentation by the Heidelberg software (version 6.9) of the DCP, ICP, and SVP was manually verified for each OCT-A scan and manually adjusted when needed. The DCP included the outer plexiform layer and the outer half of the inner nuclear layer. The ICP consisted of the inner half of the inner nuclear layer and the outer half of the inner plexiform layer. The SVP included the inner half of the inner plexiform layer and the whole ganglion cell layer.\textsuperscript{23} The resulting en face images of the DCP, ICP, and SVP and the whole retinal vascular complex (i.e., including the whole ganglion cell layer until the whole outer plexiform layer) were exported for analysis of OCT-A features.

A custom-made algorithm (MATLAB, The MathWorks, Inc., Natick, MA, USA) was used for calculation of VD and the FAZ. VD was the percentage of vessels within the parafoveal area (between concentric circles with a diameter of 1 mm and 2.5 mm centered on the fovea), and the FAZ area was calculated in square millimeters. VD was calculated for the DCP, ICP, and SVP, whereas the FAZ was calculated for the whole retinal vascular complex. Furthermore, we determined irregularity of the FAZ both qualitatively and quantitatively. Qualitative assessment of contour irregularity of the FAZ was performed on all images in a random order by one researcher, who looked at the image and compared the FAZ to an imaginary perfect round circle. The FAZ contour scores could be either regular or irregular. For the quantitative approach, we used the acircularity index,\textsuperscript{24} which was defined as the ratio of the perimeter (in mm) of the demarcated FAZ to the perimeter of a circle with an area equal to that of the FAZ.

**Statistical Analysis**

Statistical analysis was performed using SPSS Statistics 24 (IBM, Armonk, NY, USA). $P < 0.05$ was considered statistically significant. Normality of the distribution of the data was evaluated with the Kolmogorov–Smirnov and Shapiro–Wilk tests. A transformation of data was necessary for both VD and FAZ area to attain a reasonably normal distribution. For analysis, transformed data were used; however, to aid in clinical interpretation, the data were back transformed for comprehensive presentation in the figures. Differences in VD among the BU, NOBD, and HV groups were analyzed using a one-way multivariate analysis of covariance (MANCOVA), with scan quality as a covariate. Differences in FAZ area and acircularity index among the groups were separately analyzed using a one-way ANOVA. The qualitative score for irregularity of the FAZ area was compared among the groups using a Kruskall–Wallis test. Spearman’s $\rho$ was used to analyze whether BCVA was correlated with the vascular parameters in the BU group, and a partial correlation with age as a covariate was used to determine a correlation between disease duration and the vascular parameters in both BU and NOBD groups. Continuous data were expressed as mean ± SD, and categorical data were expressed as count and percentage.
Table 1. Demographic Data for BU Patients, NOBD Patients, and HV Control Group

|                     | BU (n = 23) | NOBD (n = 23) | HV Control Group (n = 22) | P     |
|---------------------|-------------|---------------|--------------------------|-------|
| Age (y), mean ± SD (range) | 51 ± 10 (27–72) | 48 ± 14 (23–71) | 44 ± 13 (27–74) | 0.25† |
| Gender, n (%)        |             |               |                          | 0.66‡ |
| Male                | 12 (52)     | 12 (52)       | 14 (64)                  |       |
| Female              | 11 (48)     | 11 (48)       | 8 (36)                   |       |
| Disease duration (y), mean ± SD (range) | 17 ± 9 (2–36) | 11 ± 8 (1–31) | NA                       | 0.070§ |
| BCVA study eye (letters), mean ± SD (range) | 43 ± 9 (1–63) | 59 ± 3 (51–64) | 61 ± 5 (47–70) | <0.001*† |
| Spherical equivalent, mean ± SD (range) | 0.2 ± 1.0 (–18 to 2.5) | –0.3 ± 1.3 (–3.9 to 2.0) | –0.3 ± 1.4 (–3.3 to 1.4) | 0.36† |
| Current immunosuppressive treatment, n (%) |          |               |                          | 0.82† |
| No medication (%)   | 3 (13)      | 4 (17)        | NA                       |       |
| One drug (%)        | 9 (40)      | 10 (44)       | NA                       |       |
| Two drugs (%)       | 7 (30)      | 5 (22)        | NA                       |       |
| Three or more drugs (%) | 4 (17) | 4 (17)        | NA                       |       |
| Concomitant diseases, n (%) |     |               |                          | 0.066‡ |
| Hypertension        | 4 (17)      | 1 (4)         | 0 (0)                    |       |
| Diabetes mellitus   | 2 (9)       | 1 (4)         | 0 (0)                    |       |

NA, not applicable.
† Statistically significant.
‡ One-way ANOVA.
§ χ² test.
* Independent samples t-test.

RESULTS

Demographic Data

Sixty-eight subjects were enrolled in the study, 46 patients with Behçet’s disease and 22 control subjects (HVs). The patients with Behçet’s disease included 23 patients without ocular involvement (NOBD) and 23 patients diagnosed with concomitant uveitis (BU). Two BU patients were excluded for the study due to an inability to acquire OCT-A images. One eye per subject was included for analysis. None of the participants had active ocular inflammation at the time of investigation, and none of the NOBD patients or HVs had any sign of past inflammation. Demographic data are presented in Table 1, and additional information on immunosuppressive medication and ophthalmological features in the BU group is presented in Supplementary Tables S1 and S2. Examples of the OCT-A en face images of BU, NOBD, and HV subjects are shown in Figure 1.

Vessel Density

Figure 2 shows the VD for each group in all three layers. One-way MANCOVA, with DCP, ICP, and SVP VD as dependent variables and OCT-A scan quality as the covariate, showed a statistically significant difference among the groups (P < 0.001). The VD was also statistically significantly different among the groups for each of the retinal vascular layers separately (Table 2). Bonferroni post hoc results are presented in Figure 2. A positive correlation between BCVA and VD in DCP, ICP, and SVP was found in the BU group (Table 3). In the BU and NOBD groups, no correlation was found between disease duration and VD in any of the layers (Table 4).

Foveal Avascular Zone

Figure 3 shows the FAZ area and the acircularity index for all three groups. The area of the FAZ was statistically significantly different among the groups (Table 2). A Bonferroni
The distribution of VD (%) in the three groups (BU, NOBD, and HV) in the DCP, ICP, and SVP. The dotted lines represent the median and quartiles.

Table 2. Quantitative OCT-A Features for VD, FAZ Area, and Acircularity Index for BU Patients, NOBD Patients, and HV Control Group

|          | BU    | NOBD  | HV    | P     |
|----------|-------|-------|-------|-------|
| VD       |       |       |       |       |
| DCP      | 25 ± 7| 30 ± 4| 33.5 ± 1.9| <0.001* |
| ICP      | 25 ± 5| 30 ± 3| 31.8 ± 1.1| <0.001* |
| SVP      | 30 ± 9| 36 ± 4| 38.9 ± 1.6| <0.001* |
| FAZ area (mm²) | 0.5 ± 0.3 | 0.4 ± 0.1 | 0.2 ± 0.1 | 0.003* |
| Acircularity index | 1.3 ± 0.2 | 1.2 ± 0.1 | 1.2 ± 0.1 | 0.27 |

*Statistically significant.

Table 3. Spearman’s ρ Between BCVA and Vascular Parameters in BU Patients

|          | BU     | NOBD   | HV     | P     |
|----------|--------|--------|--------|-------|
| VD       | 0.76   | <0.001*|        |       |
| DCP      | 0.80   | <0.001*|        |       |
| ICP      | 0.52   | 0.02   |        |       |
| SVP      | -0.37  | 0.10   |        |       |
| FAZ area | -0.31  | 0.18   |        |       |
| Acircularity index | -0.62  | 0.003* |        |       |
| Qualitative FAZ irregularity | -0.62  | 0.003* |        |       |

*Statistically significant.

Table 4. Partial Correlation Between Disease Duration and Vascular Parameters With a Correction for Age in BU and NOBD Patients

|          | BU     | NOBD   | P     |
|----------|--------|--------|-------|
| VD       |        |        |       |
| DCP      | -0.05  | 0.85   | -0.40  | 0.06  |
| ICP      | -0.03  | 0.91   | -0.28  | 0.20  |
| SVP      | 0.02   | 0.92   | -0.23  | 0.30  |
| FAZ area | 0.13   | 0.60   | 0.27   | 0.22  |
| Acircularity index | 0.23   | 0.33   | 0.02   | 0.93  |
| Qualitative FAZ irregularity | 0.18   | 0.46   | 0.17   | 0.46  |

post hoc test showed that there were statistically significant differences in FAZ area between BU and NOBD and between NOBD and HVs (Fig. 3). No statistically significant difference between any of the groups was found for the quantitative FAZ irregularity measurement (i.e., the acircularity index of the FAZ) (Table 2). In contrast, the qualitative FAZ irregularity was significantly different among the groups (P = 0.02). Pairwise comparisons showed that BU differed significantly from HVs (P = 0.004) and from NOBD (P = 0.011). BCVA was negatively correlated with the qualitative FAZ irregularity in BU patients (Table 3), but no significant correlation was found with the acircularity index nor the FAZ area. No correlation was observed between disease duration and both qualitative and quantitative FAZ measurements (Table 4).

**DISCUSSION**

In this study, quantitative measurements of the parafoveal microvasculature demonstrated that the parafoveal microvasculature is most eminently affected in BU patients, but deviations were also observed in the eyes of NOBD patients. Most alterations in parafoveal microvasculature of Behçet’s disease patients were found using the quantitative vessel density measurement. Also, vessel density was significantly correlated with BCVA in BU patients.

The vessel density was significantly lower in the BU group than in the control group in all three retinal layers, which is in line with results of vessel density or hypoperfusion area measurements in previous studies. The VD of NOBD patients was significantly different from both the vessel density in BU patients in all three layers and from the VD in healthy control subjects in the deeper retinal layers (DCP and ICP). This suggests that the deeper layers of the retina in NOBD patients deviate more from a normal microcirculation than does the SVP. Previous studies indicate that the DCP is more involved than the SVP in BU patients, which also suggests that microcirculation in the DCP is more affected by the disease. The DCP and ICP are both thin layers of capillaries, supplied by vertical anastomoses from the SVP, which consists of a mixture of large and small vessels. Because the vasculitis in BD involves all vessel types, it is less likely that the inflammation itself explains the more extended involvement of retinal capillaries. One possible explanation is that, due to their smaller diameter, capillaries might be more vulnerable to ischemia and subsequent drop-out. On the other hand, as blood flow velocity in normal capillaries is slower than in larger vessels, blood flow velocity in diseased capillaries may be decreased toward a level under the minimal detectable blood flow velocity of OCT-A.

We demonstrated that the FAZ area was significantly larger in both uveitic and non-uveitic Behçet’s disease patients compared to the FAZ area of healthy volunteers. Our results are consistent with previous studies that compared FAZ area between BU patients and HVs. For the comparison of FAZ area between NOBD patients and HVs, other studies have reported inconsistent results. Similar to our results, Goker et al. found FAZ enlargement, whereas three
other studies\textsuperscript{8,18,20} did not find a significant difference in FAZ area between NOBD patients and HVs. These inconsistent reports on FAZ area differences between NOBD eyes and healthy eyes could be explained by the high between-subject variability of FAZ area measurements in healthy eyes.\textsuperscript{25–30} Another explanation for the discrepancies in FAZ area findings is the difference in FAZ definition. Raafat et al.\textsuperscript{26} only analyzed the FAZ area in the SVP, and Gomez et al.\textsuperscript{23} looked at the FAZ area in the superficial and deep plexus separately. However, it is more logical to analyze the FAZ area including all retinal vascular plexuses together, as the three retinal vascular layers merge at the edge of the FAZ and therefore define the borders of the FAZ together.\textsuperscript{25,31} Another bias can be found in the FAZ data reported by Koca et al.,\textsuperscript{18} who included 94 eyes from 49 Beh\c{c}et’s disease patients, of which 43 eyes had ocular involvement, and one eye of 53 healthy controls. The fellow eyes of uveitic patients were included in their NOBD group. It may be assumed that microvasculature in the fellow eyes of unilateral BU patients will be affected more severely despite the lack of clinical inflammation, compared to the eyes of NOBD subjects. Hence, these fellow eyes cannot be classified in the NOBD group. To present the most unbiased data as possible, we chose to increase the number of patients per group, included only one eye per subject in all three groups, and combined all retinal capillary layers to define the FAZ.

Koca et al.\textsuperscript{18} suggested exploring the irregularity of the FAZ, as it might be a more valuable marker than FAZ area to indicate ocular involvement. They found a significant difference in FAZ irregularity between BU patients and healthy control subjects and found FAZ irregularity to be correlated with a lower BCVA in BU patients. Our qualitative analysis of FAZ irregularity showed similar outcomes. We also performed a quantitative measurement, the acircularity index, which quantifies the deviation of the FAZ perimeter from a perfectly round circle,\textsuperscript{24} but this did not differ significantly among the three groups. Furthermore, this acircularity index was not significantly correlated with BCVA in BU patients, whereas the qualitative FAZ irregularity measurement was (Table 3). Therefore, we still consider FAZ irregularity to be a potentially valuable marker for ocular involvement, but this irregularity is not quantified adequately by this acircularity index. Table 3 shows that BCVA in BU patients is also significantly correlated with vessel density in DCP, ICP, and SVP. Therefore, this quantitative marker is superior to qualitative FAZ measurements as an indicator of ocular involvement.

Several hypotheses can be proposed to explain the occurrence of preclinical retinal vascular alterations in NOBD patients. First, most clinical manifestations in Beh\c{c}et’s disease are attributed to vascular involvement, although sensitivity is in general low for detecting those vascular abnormalities.\textsuperscript{30,32} Uveitis is in many patients one of the presenting symptoms of Beh\c{c}et’s disease; thus, we suspect that subclinical retinal vascular damage can also occur at the onset of the disease without symptomatic ophthalmic inflammation.\textsuperscript{21} With OCT-A, we were indeed able to detect subclinical deviations in the retinal microcirculation of NOBD patients. Second, we hypothesized that duration of disease activity in NOBD patients could correlate with progressive worsening of the retinal vasculature abnormalities, as this was also described in previous literature for BU patients.\textsuperscript{17} After correcting for age, however, we did not find a correlation between disease duration and any of the OCT-A variables in either uveitic or non-uveitic Beh\c{c}et patients (Table 4). Because vessel density is known to decrease with age,\textsuperscript{33} this may be an important and overlooked bias in previous reports. Another possibility is that preclinical retinal microvascular alteration in NOBD patients predict future development of uveitis; however, this hypothesis can only be tested with a longitudinal cohort study in NOBD patients to evaluate whether a first uveitis event occurs.

Because the diagnosis of Beh\c{c}et’s disease currently relies on clinical signs and symptoms, it remains uncertain in many cases. In addition, diagnosis may be delayed because clinical manifestations can present asynchronously.\textsuperscript{21,34} Quantitative measurements to confirm diagnosis would possibly eliminate or confirm some of those suspected Beh\c{c}et’s disease cases. Furthermore, because not all Beh\c{c}et’s disease patients have ocular involvement, quantitative measurements could perhaps be of help in predicting future ocular events. OCT-A, through quantitative parameters such as VD estimations, has the potential to fulfill a role as an additional diagnostic tool in the future. However, to get to this point, robust longitudinal follow-up studies are needed to build a reliable Beh\c{c}et’s disease database so that we can better define which OCT-A quantitative measurements are most sensitive for the detection of Beh\c{c}et’s disease and which factors (e.g., age, scan quality) most influence these quantitative outcomes measurements.

This study has several strengths. As mentioned before, we aimed to present the most unbiased data as possible by including a high number of subjects and only one eye per subject per group, in addition to combining all retinal capillary layers to define the FAZ. NOBD patients and healthy control subjects and found FAZ irregularity to be correlated with a lower BCVA in BU patients. Our qualitative analysis of FAZ irregularity showed similar outcomes. We also performed a quantitative measurement, the acircularity index, which quantifies the deviation of the FAZ perimeter from a perfectly round circle,\textsuperscript{24} but this did not differ significantly among the three groups. Furthermore, this acircularity index was not significantly correlated with BCVA in BU patients, whereas the qualitative FAZ irregularity measurement was (Table 3). Therefore, we still consider FAZ irregularity to be a potentially valuable marker for ocular involvement, but this irregularity is not quantified adequately by this acircularity index. Table 3 shows that BCVA in BU patients is also significantly correlated with vessel density in DCP, ICP, and SVP. Therefore, this quantitative marker is superior to qualitative FAZ measurements as an indicator of ocular involvement. Several hypotheses can be proposed to explain the occurrence of preclinical retinal vascular alterations in NOBD patients. First, most clinical manifestations in Beh\c{c}et’s disease are attributed to vascular involvement, although sensitivity is in general low for detecting those vascular abnormalities.\textsuperscript{30,32} Uveitis is in many patients one of the presenting symptoms of Beh\c{c}et’s disease; thus, we suspect that subclinical retinal vascular damage can also occur at the onset of the disease without symptomatic ophthalmic inflammation.\textsuperscript{21} With OCT-A, we were indeed able to detect subclinical deviations in the retinal microcirculation of NOBD patients. Second, we hypothesized that duration of disease activity in NOBD patients could correlate with progressive worsening of the retinal vasculature abnormalities, as this was also described in previous literature for BU patients.\textsuperscript{17} After correcting for age, however, we did not find a correlation between disease duration and any of the OCT-A variables in either uveitic or non-uveitic Beh\c{c}et patients (Table 4). Because vessel density is known to decrease with age,\textsuperscript{33} this may be an important and overlooked bias in previous reports. Another possibility is that preclinical retinal microvascular alteration in NOBD patients predict future development of uveitis; however, this hypothesis can only be tested with a longitudinal cohort study in NOBD patients to evaluate whether a first uveitis event occurs.

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