A New Approach for the Segmentation of Three Distinct Retinal Capillary Plexuses Using Optical Coherence Tomography Angiography

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Purpose: To segment three distinct retinal capillary plexuses by using optical coherence tomography angiography (OCTA).

Methods: This prospective study included 30 eyes of 15 healthy subjects. En face OCTA images generated by the AngioPlex platform were manually segmented by the “progressive matching” method to the superficial, middle, and deep capillary plexuses (SCP, MCP, and DCP, respectively). The estimated position of each plexus relative to the reference line was calculated. Vascular density (VD) and skeleton density (SD) analyses, as well as the interclass correlation coefficient and relative standard deviation, were performed on each capillary plexus. We also measured central retinal thickness (CRT) and ganglion cell layer thickness (GCT).

Results: Thirty eyes of 15 healthy subjects (9 females; average age of 28.33 ± 3.07 years) were included in the analysis. We defined the relative estimated positions of the outer boundary MCP to the RPEfit as MCP = 14.491 ± 0.307 CRT/C0 /1.443 GCT/C0, while the outer boundary of DCP was 37.63 ± 7.04 μm below the IPL. The VDs of SCP, MCP, and DCP were 32.97% ± 3.90%, 45.05% ± 5.34%, and 37.34% ± 4.96%, respectively, while the SDs of SCP, MCP, and DCP were 14.45 ± 1.51 mm/C0 /1, 19.80 ± 1.92 mm/C0 /1, and 17.38 ± 1.97 mm/C0 /1, respectively.

Conclusions: With the progressive matching method, we segmented three capillary plexuses and defined the relative estimated positions of each capillary plexus to the reference line and calculated the VD and SD of three capillary plexuses in healthy subjects, providing controls for future studies.

Translational Relevance: Our study provides a visual method for OCTA image vascular segmentation and provides reference and control for future studies on retinal three capillary plexuses.

Introduction

Optical coherence tomography angiography (OCTA) is an advanced, noninvasive, dye-less imaging technology that allows for direct evaluation of the retinal microvasculature with depth-resolved capability.¹⁻⁴ The technique is based on the principle of identifying the temporal evolution of the optical coherence tomography (OCT) signal caused by the motion of scattering particles, such as erythrocytes, within the vessels. This technique can be used to generate information on three-dimensional blood flow for visualization of the retinal and choroidal vasculature.⁵

OCTA studies have largely concentrated on the superficial capillary plexus (SCP) and deep capillary plexus (DCP).²⁻⁵,⁷ However, significant evidence from histopathological and anatomical studies indicates the existence of a third capillary plexus, the middle capillary plexus (MCP), which is controlled by
distinct developmental cues. Moreover, according to other studies in rodent and human retina, each capillary layer of the retina has different regulatory units, suggesting an independent neurovascular unit at each capillary plexus, favorably supporting the existence of the MCP. Given that the MCP is partially incorporated into other plexuses by standard commercial software, some pioneers began to identify the MCP by manual segmentation of OCTA volumes and came to realize the importance of MCP. Nesper et al. even described the three-dimensional image of the three capillary plexuses in their study. However, the position of the MCP is a matter of opinion. Onishi et al. and Park et al. considered the MCP a thin slab between the inner plexiform layer (IPL) and inner nuclear layer (INL) (0 μm offset from IPL and 30 μm beneath the IPL). Hagag and colleagues argued that the MCP is in the outer 20% of the ganglion cell complex and inner 50% of the INL. The approach of Garrity et al. is to segment the MCP with an inner boundary set at the IPL-INL junction and an outer boundary set at 20 μm below the IPL-INL junction. By contrast, Nesper et al. considered that the MCP was segmented from 55 to 60 μm above the IPL. In most previous studies on MCP were performed using the RTVue XR Avanti OCTA instrument (Optovue, Fremont, CA) based on split-spectrum amplitude-decorrelation angiography software, and there are few studies performed using other devices.

In this study, we used the Cirrus high-definition OCT AngioPlex instrument (Zeiss Meditec, Inc., Dublin, CA) by using the Optical Micro Angiography (OMAG) algorithm to segment the three distinct retinal capillary plexuses. A manual segmentation method called “progressive matching” was adopted to determine the locations of MCP and DCP and then calculated their approximate distribution in healthy human subjects by recording their specific locations. In addition, vascular density (VD) and skeleton density (SD) analyses were performed on each capillary plexus. Our study is an important extension of previous studies on three distinct retinal capillary plexuses and provides a basis for future research.

**Methods**

This prospective study was approved by the Lixiang Eye Hospital of Soochow University Institutional Review Board and adhered to the tenets of the Declaration of Helsinki and was Health Insurance Portability and Accountability Act compliant. Informed consent was obtained from each subject before OCTA imaging.

**Study Sample**

A total of 15 volunteers were recruited for this study, and a total of 30 normal eyes were evaluated. Each subject underwent a standard refractive diopter (D) measurement (AR-310A; Nidek, Aichi, Japan), slit-lamp biomicroscopy, and dilated fundus examination. The inclusion criteria were no evidence of ocular media opacity, retinal disease, or significant refractive error (myopia of three Ds or more or hyperopia of one D or more) in the study eye. Exclusion criteria included ocular surgery, poor-quality images with a signal strength less than 9 (maximum of 10), significant motion artifact, or inability to abstain from blinking or movement during image acquisition. No eyes were excluded in this study.

**OCTA Imaging**

Images were obtained using the commercially available instrument Zeiss Cirrus HD-OCT 5000-2328 (Zeiss Meditec, Inc.) with OMAG AngioPlex (software version 9.5.2.19038, copyright 2016). Because the OMAG algorithm does not split the OCT spectrum into subbands, it theoretically provides a higher axial resolution of approximately 5 μm in tissue. The device, with a central wavelength of 840 nm and a scan speed of 68,000 A-scans per second, minimized the effect of eye motion-related artifacts by the use of Fast Trace, as previously described. At least one 3 × 3-mm (759 × 759 pixels) scan centered on the fovea was taken in each eye of interest.

**Segmentation of the Capillary Plexuses**

The data were segmented using commercially available AngioPlex software inner-limiting membrane (ILM) and retinal pigment epithelium (RPE-fit) segmentation algorithms, and the position of the IPL was calculated by the two algorithms. According to the user manual of the device, ZIPL = ZILM + 70% * (TILM-OPL), while ZOPL = ZRPEfit – 110 μm; ZIPL is the estimated IPL boundary, ZILM is the ILM boundary, TILM-OPL is the thickness between the ILM and outer plexiform layer (OPL), and ZOPL is the estimated OPL boundary. Because the ILM line encroached upon choroidal vessels in the fovea area when moving down because of the concave area in the fovea, RPEfit and IPL were selected for manual segmentation simultaneously. A
A manual segmentation method called progressive matching was adopted to determine the locations of MCP and DCP. First, a 20-μm thin slab was set with RPEfit as the reference line and gradually moved upward from the nonvascular area of the outer nuclear layer at an interval of 6 μm until the continuous capillary plexus began to disconnect (as shown in Fig. 1). The most integrated image was selected as the DCP, and the relative estimated position from RPEfit was recorded (the outer boundary, same as follows). Then, we adjusted the thin slab to 30 μm and continued to obtain the MCP (as shown in Fig. 2). We repeated the above steps with the IPL as the reference line. The selection of the most integrated image was interpreted by the two independent researchers together, and if the two researchers’ interpretation results were inconsistent, the final interpretation was conducted by the third researcher. We further activated the manual “Remove Projections” option in AngioPlex software for MCP and DCP analyses, which uses additive mixing of overlying vessels to reduce projection artifacts in lower layers. In this way, we obtained the MCP and DCP after removal of the decorrelation tail artifact (MCP-re and DCP-re).

**Image Analysis**

En face OCTA images were exported to an external software program (ImageJ, version 1.50i; http://imagej.nih.gov/ij/; provided in the public domain by the National Institutes of Health, Bethesda, MD, USA) and opened in image analysis. To calculate the VD and SD, we applied a thresholding algorithm to en face images of each capillary plexus to create a binarized black-and-white slab. VD was calculated as a proportion of the total area of pixels with a detected OCTA signal (white pixels in the binarized image) compared to the total area of the image. SD was defined as the ratio of the skeletal length of vessels to the total area of the image. A similar approach has been described in detail in many previous reports.
A Cirrus OCT scan of the macula consisting of 128 B-scans with 512 A-scans each (Macular Cube 512 × 128 scan) was taken of each eye, and the central retinal thickness (CRT) and ganglion cell inner plexiform thickness (GCT) were measured using the existing algorithm available on the Cirrus device.

One week later, we randomly selected 20 eyes for resegmentation and analysis to calculate the interclass correlation coefficient (ICC). Moreover, one eye was selected randomly from the segmentation procedure for analysis repeated 10 times to calculate the relative standard deviation (RSD).

**Statistical Analysis**

Statistical analysis was performed using SPSS 18.0 (IBM Corp., Armonk, NY). For all tests, \( P \) values <0.05 were considered statistically significant. All data are reported as the means and standard deviations (SD). Pearson and Spearman correlations were used to study the association between the estimated location of each capillary plexus and the CRT and GCT. A linear regression model (method = stepwise, criteria = pin (0.05) pout (0.10)) was used to calculate the estimated location of the MCP and DCP outer boundaries.
Results

Of the 15 subjects, 9 (60%) were female with an overall average age of 28.33 ± 3.07 (range, 24 to 34) years, and average spherical equivalent of −0.54 ± 0.77 (range, +0.5 to −2.5) D. All images met the entry criteria. Out of a total of 60 interpretations, 7 (11.67%) required a third-party determination (6 for MCP and 1 for DCP).

The outer boundaries of the MCP were 183.30 ± 13.22 μm from the RPEfit and 15.70 ± 8.03 μm from the IPL, while the outer boundaries of the DCP were 137.60 ± 11.42 μm above the RPEfit and 37.63 ± 7.04 μm beneath the IPL (Table 1). The inner boundaries of the MCP were approximately located at the junction of the IPL with the ganglion cell layer (Fig. 2), while the GCT was approximately 85.97 ± 3.32 μm. Because the three networks around the foveal avascular zone (FAZ) would be anastomosed postnatally, we defined the SCP as a slab from 0 μm beneath the ILM to 70 μm beneath the ILM.9,22 This definition preserves the integrity of the vessels surrounding the FAZ without encroaching upon the MCP in the parafoveal region.

In the Pearson and Spearman correlation analysis, the estimated position of the MCP was strongly correlated with the CRT and GCT relative to RPEfit and IPL, while the estimated position of the DCP was weakly correlated only with the CRT and GCT relative to RPEfit (Table 2).

Linear regression indicated that the estimated position of the MCP relative to RPEfit was correlated with the CRT and GCT, while the estimated position relative to the IPL was affected by the CRT. The position of DCP relative to RPEfit was related to the GCT, while the estimated position relative to IPL was unaffected by either area (Table 3). The estimated

Table 1. Relative Estimated Position of Each Capillary Plexus With Each Reference Line and CRT and GCT

| MCP | RPEfit (μm) | IPL (μm) | DCP | RPEfit (μm) | IPL (μm) | CRT (μm) | GCT (μm) |
|-----|-------------|----------|-----|-------------|----------|----------|----------|
|     | −183.30 ± 13.22 | −15.70 ± 8.03 |     | −137.60 ± 11.42 | 37.63 ± 7.04 | 240.33 ± 21.26 | 85.97 ± 3.32 |

a Data are mean ± standard deviation, and the minus sign means above the reference line.
position of MCP relative to RPEfit had a high degree of fitting with the CRT and GCT (adjusted $R^2 = 0.494$) and had a better prediction accuracy (Fig. 5). Therefore, the outer boundary of MCP relative to the estimated position of RPEfit was set as $Y$, CRT as $X_1$, and GCT as $X_2$, and the regression equation $Y = 14.491 - 0.307X_1 - 1.443X_2$ was given.

The SCP was mainly composed of arteries, veins, arterioles, and venules arranged around the center, whereas the MCP and DCP were mainly composed of arterioles, venules, and a large number of capillaries (Figs. 3, 4). In the DCP, we also found some “vortex” systems (red arrow), which drain into central conduits that directly connect to venules in the SCP, as previously described. Among the three retinal capillary plexuses, the MCP had the highest VD (45.05 ± 5.34%), followed by the DCP (37.34 ± 4.96%), and the lowest was SCP (32.97 ± 3.90%). After the artifact was removed, the VD of the DCP-re decreased (33.42 ± 4.80%), but the VD of the MCP-re increased (57.52 ± 7.11%). Similar results were found for SD, as shown in Table 4.

VD showed high reproducibility with ICCs more than 0.941 for each capillary plexus, and SD showed high reproducibility with ICCs more than 0.931 for each capillary plexus. Both VD and SD showed high precision, with RSDs less than 2.26% for each capillary plexus. Table 5 showed the detailed results of the ICCs and RSDs.

**Discussion**

In this study, we used a progressive matching method for segmentation, which has good reproducibility and precision according to the ICC and RSD tests. To ensure the accuracy of the results, we used two judges to interpret together and introduced a third judge when the interpretation was inconsistent. The interpretation of the DCP was more consistent, whereas the MCP was more prone to divergence in interpretation due to the influence of SCP decorrelation tail artifacts.

For the thickness of the DCP, Onishi et al. and Park et al. selected a thin slab of 15 μm, but we found that a slab of this thickness could not display an integrated DCP; thus, we selected a thin slab of 20 μm to segment the DCP following the method of Bonnin et al. DCP is presumably located on the lateral side of the INL and most of the OPL, whereas the MCP is presumably located at the inner edge of the INL and most of the IPL. There was a distinct disruption of the vascular layer inside the INL. According to develop-

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**Table 2.** Pearson and Spearman Correlations between the Relative Estimated Position, CRT, and GCT

|       | RPEfit (μm) |         | IPL (μm) |         |
|-------|-------------|---------|----------|---------|
|       | MCP         | DCP     | MCP      | DCP     |
|       | Pearson     | Spearman| Pearson   | Spearman|
|       | CRT         | GCT     | CRT      | GCT     |
| $R$   | -0.650      | -0.640  | -0.478   | -0.475  |
| $P$   | <0.001      | <0.001  | 0.008$^a$| 0.008$^a$|

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**Table 3.** Linear Regression Analysis With the Relative Estimated Position, CRT, and GCT$^a$

|       | Unstandardized Coefficient | Standardized Coefficient | $P$ Value |
|-------|---------------------------|--------------------------|-----------|
| MCP-RPEfit | Adjusted $R^2 = 0.494$, Constant = 14.491 | CRT: -0.307, GCT: -1.443 | 0.002, 0.020 |
| MCP-IPL | Adjusted $R^2 = 0.201$, Constant = 27.696 | CRT: -0.181, GCT: -0.478 | 0.008, - |
| DCP-RPEfit | Adjusted $R^2 = 0.178$, Constant = -2.950 | CRT: -1.566, GCT: -0.455 | 0.012, 0.0012 |
| DCP-IPL | Adjusted $R^2 = 0.043$ | CRT: -9.75, GCT: -5.34 | - |

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$^a$ Method = stepwise, criteria = pin (0.05) pout (0.10).
mental studies, at 25 to 26 weeks of gestation, deeper plexus components form by angiogenic sprouting from the primary plexus veins, and these plexus sprouts penetrate the retina and establish two laminar networks on either side of the INL.\textsuperscript{10,22,26–28} Additionally, Tan and his colleagues\textsuperscript{8} found four layers of capillary plexuses in the human retina through anatomical studies of cadaveric eyes, which are located on the nerve fiber layer, retinal ganglion cell, border of the IPL and superficial boundary of the INL, and boundary of the deep INL and OPL. These findings are in perfect agreement with our study.

In our study, the estimated position of MCP was affected by CRT and GCT regardless of whether we used RPEfit or IPL as the reference line. According to linear regression analysis, to achieve the best fitting

**Figure 5.** Normal P-P plot of regression standardized residual. (A) The estimated DCP position with RPEfit as the reference line is the dependent variable. (B) The estimated MCP position with RPEfit as the reference line is the dependent variable. (C) The estimated DCP position with IPL as the reference line is the dependent variable. (D) The estimated MCP position with IPL as the reference line is the dependent variable. (B) Shows a better prediction accuracy, follow by (A) and (D).

**Table 4.** VD and SD of the Three Capillary Plexuses\textsuperscript{a}

|       | SCP       | MCP       | MCP-re    | DCP       | DCP-re    |
|-------|-----------|-----------|-----------|-----------|-----------|
| VD (%)| 32.97 ± 3.90 | 45.05 ± 5.34 | 57.52 ± 7.11 | 37.34 ± 4.96 | 33.42 ± 4.80 |
| SD (mm\textsuperscript{-1}) | 14.45 ± 1.51 | 19.80 ± 1.92 | 23.52 ± 2.10 | 17.38 ± 1.97 | 15.61 ± 1.93 |

\textsuperscript{a} Data are the mean ± standard deviation, MCP-re means the MCP after removal of the decorrelation tail artifact, and DCP-re means the DCP after removal of the decorrelation tail artifact.
Onishi et al.,15 the VD of the SCP was 48.19% ± 2.95%, and the VDs of the MCP and DCP without decorrelation tail artifact removal were 56.71% ± 2.73% and 59.87% ± 3.32%, respectively. In the RTVue XR Avanti OCTA system, the calculation area of VD was in the parafoveal region, which was defined as a ring around the fovea with an inner ring diameter of 1 mm and an outer ring diameter of 3 mm. Therefore, the partial or whole FAZ was excluded from the total area, and the results were relatively high.15,34–36

In the study by Garrity et al.,17 the SD values of the SCP, MCP, and DCP were 15.48 ± 2.04 mm⁻¹, 15.28 ± 1.82 mm⁻¹, and 16.33 ± 2.32 mm⁻¹, respectively, while projection artifacts were removed from en face OCTA images of the MCP and DCP with three-dimensional PAR-enabled software developed by Optovue.37,38 In our study, the values of the SCP and DCP after removal of decorrelation tail artifacts were very close to the results of Garrity et al. (14.45 ± 1.51 mm⁻¹ vs. 15.48 ± 2.04 mm⁻¹ and 15.61 ± 1.93 mm⁻¹ vs. 16.33 ± 2.32 mm⁻¹), but the results of the MCP were very different (23.52 ± 2.10 mm⁻¹ vs. 15.28 ± 1.82 mm⁻¹). As shown in Figure 4, the commercial software on the AngioPlex platform can remove the artifacts in the DCP well, whereas in the MCP, it reduces only the grey value of the decorrelation tail artifacts of the upper layer. Because the artifact was not completely removed, the artifact was still considered an OCTA blood flow signal when performing the VD calculation. The method reduces the artifacts by comparing and reconstructing two slab images generated from two or more retinal layers.20,39 When the signal of the upper layer is too strong or signals affect adjacent multiple layers, the artifact cannot be completely eliminated. Interestingly, this approach can improve the signal-to-noise ratio of the FAZ in the MCP, resulting in a lower threshold value and even a higher VD (Table 4). The projection-resolution algorithm of the AngioVue platform is compared along each axial scan in the proximal to distal direction, identifying successive higher peaks as real vessels and removing smaller peaks.40,41

The present study has limitations. First, we report the values for only a small population of healthy subjects. Further studies are needed to establish a normative database of retinal segmentation. Second, the analysis of diseased eyes is beyond this report. In some diseases, such as diabetes, hypertension, glaucoma, high myopia, and age-related macular degeneration, the structure of the retina and vessels may change, leading to different segmentation results. Moreover, as mentioned above, the artifact in the

### Table 5. ICCs and RSDs Analyzed for VD and SD

|        | SCP | MCP | MCP-re | DCP | DCP-re |
|--------|-----|-----|--------|-----|--------|
| ICC    |     |     |        |     |        |
| VD     | 0.996 | 0.971 | 0.984 | 0.950 | 0.941 |
| SD     | 0.996 | 0.963 | 0.971 | 0.948 | 0.931 |
| RSD    |     |     |        |     |        |
| VD (%) | -   | 1.36 | 1.49  | 1.27 | 2.26  |
| SD (%) | -   | 1.34 | 0.61  | 1.22 | 1.46  |

* MCP-re means the MCP after removal of the decorrelation tail artifact, and DCP-re means the DCP after removal of the decorrelation tail artifact.

Here, the MCP had the highest VD, followed by the DCP and SCP. Based on our results and other studies of human histology, the MCP lies at the boundary of the IPL and inner portion of the INL, colocalizing with the bipolar cell processes and amacrine cells in close proximity to the high oxygen demand of the IPL synapses,11 whereas the DCP lies at the boundary of the deep INL and OPL, colocalizing with horizontal cells and close to the OPL synapses.29 Retinal oxygen measurement experiments in animal models have determined that the primary oxygen consumers of the inner retinal layer are located in the plexiform layers (IPL and OPL), which may be located in mitochondria-rich synapses.30,31 Areas with high oxygen consumption tend to have thinner blood vessels and higher blood flow density to provide a higher surface area-to-blood volume ratio for material exchange.32,33 In a report by Onishi et al.,15 the VD of the SCP was 48.19% ±
MCP cannot be completely removed, so the value of the VD and SD in the MCP may be affected. We expect the emergence of a more advanced method of artifact removal by the instrument. Finally, although we determined the general position of the three layers of capillary plexuses in the retina, our results could not be directly applied to other platforms due to the different definitions of the reference line by different OCT platforms.

In conclusion, by using the progressive matching method, we segmented three capillary plexuses visually and defined the relative positions of each capillary plexus to the reference line, which provided convenience for future studies. Moreover, we calculated the VD and SD of three capillary plexuses in some normal subjects, providing controls for future studies.

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