Purpose: Retinal vascular networks are observed as a layered structure residing in a nerve fiber layer and an inner nuclear layer of the retina. This study aimed to evaluate reflectance confocal adaptive optics scanning laser ophthalmoscopy (AO-SLO) for imaging of the layered retinal vascular networks.

Methods: This study included 16 eyes of 16 healthy cases. On the fovea, 2.8- and 3.0 mm²-areas were imaged using a prototype AO-SLO and optical coherence tomography angiography (OCTA), respectively. AO-SLO images focused on the nerve fiber and photoreceptor layers were recorded in the area. Two different vessel images (capillary networks in the superficial layer and in all layers) were generated to examine if the deep capillary network could be distinguished. We compared AO-SLO with OCTA in imaging of the layered retinal vascular networks.

Results: Sufficient images of capillary networks for analysis could be generated when the motion contrast was enhanced with AO-SLO movies in seven cases (43.8%). The deep capillary network could be distinguished in the merged image. Vascular depiction performance in AO-SLO was significantly better than in OCTA at both 0.5- and 1.0-mm areas from the fovea (P < 0.05).

Conclusions: Retinal vascular imaging using AO-SLO might be a useful adjunct to OCTA as a supportive method to evaluate the retina in healthy patients and patients with disease.

Translational Relevance: In cases requiring accurate and detailed retinal vasculature observation, AO-SLO might be useful for evaluating retinal vascular lesions as a supportive imaging method of OCTA.
be useful to examine the retinal vasculatures and its hierarchical structure. Spaide et al. also reported its superiority compared with FA in visualization of the retinal capillary layer. However, the horizontal resolution of OCT is known to be around 20 μm, which could be larger than the diameter of retinal capillaries.

Adaptive optics scanning laser ophthalmoscopy (AO-SLO) is a technique that enables observation of the retina at the cellular level in patients. Although optical fundus imaging equipment applying adaptive optics including AO-SLO have been mostly employed to visualize photoreceptors, it has also been applied to the visualization of vascular features in both normal retina and retina in vascular disorders. Furthermore, because the depth of focus is approximately 60 μm, which is calculated from the incident light wavelength (840 nm) and the numerical aperture (0.12), AO-SLO could be useful in imaging a specific layer of the retinal capillary network. Recently, Mo et al. published a study on the difference between OCTA and AO-SLO FA in imaging foveal microvasculatures. However, the utility of AO-SLO in imaging the hierarchical structure of the retinal capillary network has not been well studied. Furthermore, the difference between OCTA and AO-SLO vascular imaging with motion contrast enhancement has not yet been examined.

In this study, we evaluated the ability of reflectance confocal AO-SLO to visualize the retinal capillary layer in healthy eyes. We also compared reflectance confocal AO-SLO with OCTA in imaging retinal layered capillaries.

**Methods**

This study was approved by the Institutional Ethics Committees of the Kyushu University Hospital (Protocol No. 25233, UMIN000016858), and was performed in accordance with the ethical standards laid down by the Declaration of Helsinki. Written informed consent was obtained from all patients after a detailed explanation of the study.

**Participants**

The study included 16 healthy subjects (10 eyes of 10 females and 6 eyes of 6 males) with clear ocular media and no history of prior ocular or systemic disease.

**Adaptive Optics Scanning Laser Ophthalmoscopy (AO-SLO)**

Patients underwent full ophthalmologic examination including slit-lamp examination, dilated fundus examination, and color fundus photography to confirm no apparent retinal disorder after pupil dilatation with an eye drop of 0.5 % toropicamide and 0.5 % phenylephrine (Mydrin-P; Santen, Osaka, Japan). Based on a previous report, axial length was measured in each patient using IOL master (Carl Zeiss Meditec, Jena, Germany) to calculate the AO-SLO image scale. In addition, all patients underwent imaging using a prototype AO-SLO system (depth of focus: ~60 μm; Canon, Inc., Tokyo, Japan) as described previously. The AO-SLO device consists of an adaptive optics system that can measure and compensate optical aberrations produced by ocular media, a high-resolution confocal SLO imaging system, and a wide-field imaging subsystem. The wavelength of AO-SLO is 845 nm, and the wavelength of the beacon light for measurement of wavefront aberrations is 760 nm. The imaging light and the beacon light power are set at 400 and 100 μW, respectively, in accordance with the safety limits set by the American National Standards Institute. The optical resolution is 5 μm.

**Image Acquisition with AO-SLO**

We imaged retinal vascular networks in the macular area (2.8 × 2.8 mm) of all subjects using a prototype AO-SLO system. The movies were recorded for 3 seconds per scan area with a field size of 2.8° × 2.8° at 25 regions. The frame rate was 32 frames per second. The AO-SLO device can focus on the photoreceptor layer as well as capillaries in the nerve fiber layer while compensating for aberrations of the target eye, and automatically create an en face image in each movie. Each area was imaged in succession. If the imaging conditions were poor (poor fixation or blinking), these areas were imaged again. The layer adjustment range in the AO-SLO device is ±2 diopters (D) from the photoreceptor layer. Therefore, the AO-SLO device allows the creation of an en face image of the photoreceptor and nerve fiber layers.

**Motion Contrast Enhancement in AO-SLO Imaging**

The capillary images were constructed by AO-SLO Retinal Image Analyzer software (ARIA; Canon, Inc.) dedicated to the prototype AO-SLO. The ARIA
software adopts the same methods that was published in a paper by Tam et al.\(^2\)

**Imaging of the Hierarchical Structure of Retinal Vessel with AO-SLO**

When the AO-SLO device was focused on the photoreceptor layer and on the capillaries at the nerve fiber layer, flickering could be observed in the vessel. The flickering at the nerve fiber layer could have originated from red blood cells in the superficial vessels (superficial image), whereas the flickering when focusing on the photoreceptor could have originated from the whole vessels in the retina (deep image; Figs. 1A, 1B). As reported previously,\(^{26}\) when red blood cells are present in the light path, the light cannot reach the photoreceptor because of the hemoglobin. On the other hand, when leukocytes are present in the light path, the light can reach the photoreceptor because of the transparency. As both red blood cells and leukocytes flow through blood vessels, flickering occurs in the AO-SLO image (Figs. 1A, 2B). Twenty-five pictures were merged to make a 2.8-mm square vessel map in each of the two different layers using ARIA. Yellow dotted squares indicate an AO-SLO single image.

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**Figure 1.** Imaging of the hierarchical structure of retinal vessels with AO-SLO. (A) The vessels are shown in pink. The scanning lights are shown in blue. The photoreceptor layer is shown in yellow. Red and white circles indicate red blood cells and leukocytes, respectively. (B) When focusing on the retinal nerve fiber layer, superficial retinal vasculatures could be imaged as a superficial image with AO-SLO. When focusing on the photoreceptor layer, we could also image all retinal vasculatures as a deep image with AO-SLO. Twenty-five single images were merged to make a 2.8-mm square vessel map in each of the two different layers using ARIA. Yellow dotted squares indicate an AO-SLO single image.
2.8-mm² vessel map in each of the two different layers with ARIA (Fig. 1B).

Vessel Depth of AO-SLO Imaging

To distinguish the two different layered vessel structures, the superficial and deep capillary images, which were constructed by ARIA, were converted to green and red, respectively (Figs. 2A, 2B). These two images were merged into a single image using Image J 1.48 software (http://imagej.nih.gov/ij/; provided in the public domain by the National Institutes of Health, Bethesda, MD, USA) (Fig. 2B).

Optical Coherence Tomography Angiography (OCTA)

We imaged retinal vascular networks of all subjects using OCTA (Optovue RTVue XR Avanti: Optovue, Inc., Fremont, CA) and obtained retinal vascular mapping images of the macular area (3.0 × 3.0 mm) on the same day of AO-SLO imaging. The instrument was used to obtain split-spectrum amplitude decorrelation angiography (SSADA) images. The instrument has an A-scan rate of 70,000 scans per second, using a light source centered on 840 nm and a bandwidth of 45 nm. The tissue resolution is 5-µm axially, and there is a 22-µm beam width. Each B-scan contained 216 A-scans. Five consecutive B-scans (M-B frames) were captured at a fixed position before proceeding to the next sampling location. A total of 216 locations (B-scans) along the slow transverse direction were sampled to form a three-dimensional data cube. With a B-scan frame rate of 270 frames per second, the 1080 B-scans in each scan sequence were acquired in approximately 4 seconds. Four volumetric raster scans, including two horizontal priority fast transverse (x-fast) scans and two vertical priority fast transverse (y-fast) scans, were obtained consecutively in one session. The best x-fast and y-fast scans were registered using the contained software (ReVue; Optovue, Inc.), which has the ability to correct some motion artifacts, including residual axial motion and transverse saccadic motion. After processing of the volume scans, the decorrelation in the images, which is essentially one minus the correlation, was calculated. Stationary tissue shows a high correlation in imaging characteristics from one frame to the next. Blood flowing through vessels causes a changing reflectance over time and localized areas of low correlation between frames (or conversely a high decorrelation). This method does not use phase information from the OCTA signal. The correlated frames were evaluated and statistical outliers were removed from the averaging process to reduce the possibility of tissue motion being present. The spectrum of the light source was split into four component parts to decrease the noise present in the image; each was used to perform the decorrelation step, and the results of all four were averaged. This split-spectrum strategy trades axial resolution for decreased noise. After this step, a block of information exists that contains levels of decorrelation that range from 0 to 1. In any given region of tissue, the maximal projection image can be viewed to obtain an image of the contained blood flow. As the retina is a laminar structure with a corresponding stratification of the blood supply, segmentation of the retina in specific layers allows simple en-face visualization of the corresponding vascular supply for that layer. While we could obtain four types of vascular mapping images
of each layer of retina using OCTA (as superficial, deep, outer retina, and choriocapillaris layers), we used the superficial (upper: inner limiting membrane with offset of 3 μm, lower: inner plexiform layer [IPL] with offset of 15 μm) and deep layer images (upper: IPL with offset of 15 μm, lower: IPL with offset of 70 μm) for the current investigation.

Foveal Avascular Zone (FAZ) Quantification

The foveal avascular zone (FAZ) boundary was defined as the inner edge of the bordering vessels. With one side of the OCTA image being 3 mm, FAZ in OCTA was manually measured using ImageJ 1.48 software. Two independent researchers (YK and SN) assessed if the AO-SLO images were merged correctly for the measurement of the FAZ area. The FAZ in the merge image of AO-SLO was calculated using the distance calculated from one side (3 mm) of OCTA image. The statistical analyses (paired t-test) were performed using the software, JMP v10.0 (SAS Institute, Cary, NC).

Estimation of AO-SLO Images at the 0.5- and 1.0-mm Area From the Fovea

We investigated whether the quality of 64 AO-SLO images (820 × 820 μm) at the distance of 0.5 and 1.0 mm from the fovea in the superior, inferior, nasal, and lateral location was sufficient for image analysis. The quality of retinal vascular mapping images of AO-SLO was evaluated independently by two observers (YK and IW). In cases where the evaluation of the two observers was different, a third observer (MY) evaluated the image quality. The kappa coefficient was 0.64 and 0.78 (P < 0.001 and P < 0.0001) for estimation of the image quality at the distance of 0.5 and 1.0 mm from the fovea as defined above, respectively.

Comparison of AO-SLO and OCTA

The superficial and deep vascular images of OCTA in the same area where good quality AO-SLO images were chosen were cropped with Keynote (Apple Corp., Cupertino, CA). We evaluated the two images as follows: two researchers (YK and IW) independently compared AO-SLO and OCTA images, and counted the number of different vascular structures in the images; and each image was scored as follows: (1) in cases whereby the vessel was detectable only on either an AO-SLO image or an OCTA image but not both, the image was scored “+1” if the number of superior quality points was greater in an AO-SLO image than in an OCTA image, and scored “−1” if it was greater in an OCTA image than in an AO-SLO image; and (2) regardless of a difference in the number of superior quality points, the image was scored “0” if there was at least one superior quality point in both AO-SLO and OCTA images. Because eight pairs of corresponding images obtained by AO-SLO and OCTA were evaluated in each case, the highest and lowest score were +8 and −8, respectively. Data were examined by the Wilcoxon one-sample signed rank test and the median, standard deviation (SD), and P value were calculated. The kappa coefficient was 0.93 and 0.75 (P < 0.0001 and P < 0.0001) at the distance of 0.5 and 1.0 mm from the fovea for estimation of the two images as defined above, respectively. This result indicated good interobserver agreement. The superiority was analyzed using the Wilcoxon signed-rank test.

Estimation of Imaged Vascular Layer in AO-SLO

In AO-SLO merged images (as shown in Fig. 1B) at 1.0 mm from the fovea, three superficial and three deep vessels were randomly chosen. Two researchers (YK and IW) estimated the AO-SLO-imaged vascular layer with the corresponding OCTA images. The kappa coefficient was 1.0. The concordance rate was calculated for estimation of both superficial and deep vessels. Furthermore, the width of the vessels that AO-SLO but not OCTA could image was measured with ARIA. The lumen diameter at the next branch that both AO-SLO and OCTA could image was also measured with ARIA in AO-SLO vessel maps extracted from the photoreceptor videos.

Statistical Analysis

Statistical analyses were performed using the software, JMP v10.0 (SAS Institute). Continuous variables were analyzed using the Wilcoxon signed-rank test or two-sample t-test as appropriate. Categoric variables were analyzed using the Wilcoxon signed-rank test or two-sample t-test as appropriate. Categoric variables were analyzed using Fisher’s exact test. P values less than 0.05 were considered statistically significant.

Results

Subject Characteristics

The average subject age was 26.7 years with a SD of 2.2 years. Ten subjects were females (62.5%). The average axial length was 25.4 ± 1.7 mm. Participants characteristics in this study are summarized in Table 1.
When focusing on the retinal nerve fiber layer, superficial retinal vasculatures could be imaged (superficial image; Figs. 1A, 1B). When focusing on the photoreceptor layer, we could also image all retinal vasculatures (deep image; Figs. 1A, 1B). Thicker vessels of the superficial layer of the retina were imaged clearly in the superficial image. Part of capillary network in the inner vascular plexus could not be imaged well in the superficial image. Capillaries as well as thicker vessels could also be imaged in the deep image. Retinal vessels could be imaged thinner and brighter in the superficial image than in the deep image (Fig. 1B). These data indicate that AO-SLO could image retinal vessels in the superficial layer and in all layers by using a different focusing method.

**Distinguishing Different Layered Retinal Vessels Using AO-SLO Images**

To distinguish the two different layered vasculatures, the superficial and the deep images were converted to green and red, respectively, and these two images were merged (Fig. 2). In the merged image, yellow indicates superficial vessels, whereas red indicates deep vessels (Fig. 2B). This data suggest retinal vessels in different layers could be distinguished using AO-SLO images (Table 2).

**Quantitative Comparison between AO-SLO and OCTA**

Among 16 cases (each comprising 64 images taken at the distance of 0.5 and 1.0 mm from the fovea), we assessed that seven cases (43.8%: five eyes of five females and two eyes of two males) included images of sufficient quality for analysis while the AO-SLO images from the nine cases that were not analyzed contained insufficient image quality for the analysis (Table 2). Seventeen superficial images (26.6%) taken at 0.5-mm distance were good enough for analysis (Fig. 3A), and 14 taken at the 1.0-mm distance (21.9%) were judged to be good enough for analysis (Figs. 3B, 3C). The AO-SLO merged image was able to show retinal vessel structures similar to OCTA (Figs. 3D–G).

**Quantitative Comparison of Vessel Structure Imaging between AO-SLO and OCTA**

Retinal vasculatures at the distance of 0.5 mm from the fovea form the foveal vascular zone (Fig. 4), whereas retinal vasculatures at the distance of 1.0 mm from the fovea consist of a two-layered structure (Fig. 5). Next, we compared AO-SLO images with OCTA at distances of 0.5 and 1.0 mm from the fovea. As reported previously, retinal vessels in OCTA images...
tend to be imaged thicker than those in our prototype AO-SLO (Figs. 4 and 5). Although the differences in the width of each capillary were clearly distinguished in AO-SLO, those differences were unclear in OCTA at 0.5 as well as 1.0 mm from the fovea (Figs. 4 and 5). Close-running capillaries could also be distinguished from each other in AO-SLO images, whereas these capillaries were imaged as single-thick capillaries in OCTA images (Figs. 4C–F, and 5C, 5D). Some capillary structures, which were simply seen as branching structures in OCTA images, could be imaged as doughnut-shape structures in AO-SLO images (Figs. 5E, 5F). Furthermore, some capillaries could not be imaged by OCTA while they could be imaged by AO-SLO (Figs. 4G, 4H). However, a few capillaries could be imaged by OCTA but not by AO-SLO (Fig. 6). Our quantitative analysis showed that the ability of AO-SLO to visualize vessels was significantly superior compared with OCTA at both 0.5 and 1.0 mm from the fovea (median ± SD; +2.5 ± 2.7 points and +2.5 ± 1.9 points, P = 0.03 and 0.02, respectively; Table 2).
Lumen Diameter of AO-SLO Imaged Vessels

The lateral optical resolution of our prototype AO-SLO is higher than the optical resolution of commercial OCTA. To examine if the higher resolution could explain the superior vessel plot-out ability of AO-SLO compared with OCTA, the lumen diameter was measured with our AO-SLO image analyzer. The lumen diameter of the vessels that AO-SLO, but not OCTA, could image (10.2 ± 2.8 μm, n = 15) was similar to one of the vessels at the next branch, which both AO-SLO and OCTA could image (10.1 ± 2.4 μm, n = 15) on the vessel maps extracted from the photoreceptor (P = 0.95).

Quantitation of Foveal Avascular Zone with AO-SLO and OCTA

Next, we compared the area of the FAZ among AO-SLO and OCT images. Our quantitative analysis showed that there was no significant difference between the FAZ area in AO-SLO (0.36 ± 0.12 mm²) and in OCTA (0.34 ± 0.11 mm²; P = 0.08, Fig. 7, Table 2).

Comparison of Vessel Layer between AO-SLO and OCTA

Lastly, we investigated differences in the imaged vascular layer between AO-SLO and OCTA. All vessels that were recognized as the superficial layer in AO-SLO, were imaged in the superficial vessel layer in OCTA. However, 40.5% of vessels, which AO-SLO recognized as the deep layer was imaged in the superficial vessel layer in OCTA (Table 2). These data indicate that the superficial vascular layers could be consistent between AO-SLO and OCTA, whereas the deep vascular layer in AO-SLO included the OCTA superficial vascular image.

Discussion

Recently, there have been various reports demonstrating the usefulness of OCTA as a tool for clinical evaluations in retinal vascular disorders. OCTA provides both structural and functional (i.e., blood flow) information of retinal vasculatures. A distin-
The distinguishing feature of OCTA that differs from existing imaging equipment is its ability to image retinal vascular layers. In this study, we investigated the potential of AO-SLO for imaging the retinal vascular layer in healthy eyes. As the depth of focus of our prototype AO-SLO is approximately 60 μm, AO-SLO has excellent representation of a specific layer structure, such as the photoreceptor layer. To distinguish the retinal vascular layers, we performed a method that subtracts superficial vascular images from images that include both vascular layers. Using this method, the distinction of the superficial retinal vessels and deep retinal vessels was possible in high resolution. However, it was more difficult to image the hierarchical structure compared with OCTA. Only 43.8% AO-SLO images were of sufficient quality for use in this current study (Table 2). This could be due to eye motion or an unstable tear film. Furthermore, disorder of the retinal layer structure in an image area could also influence the method. Although the healthy eye could be investigated in this study, retinal diseases such as edema and high myopia might pose difficulty in imaging the layered retinal vascular networks.

A clinical gold standard in imaging the retinal vessels has still been FA. However, it is known that angiography using an injected dye sometimes can lead to adverse events including anaphylaxis. Therefore, a novel method to image retinal capillaries is desirable. The best feature of OCTA could be its ability to visualize the flow without using an injected dye. A recent study that compared AO-SLO with OCTA also used fluorescein for the AO-SLO. Two methods to image retinal vessels using AO-SLO have been reported. One method was AO-SLO FA using injected dye and the other method was AO-SLO using motion contrast, which were introduced by Pinhas and Tam, respectively. In this study, we investigated the ability to image retinal vessels by observing the retinal vessels using motion contrast in AO-SLO instead of injected dyes. We confirmed that retinal vessels that could not be visualized with the OCTA could be observed with our prototype AO-SLO. Specifically, close-running capillaries as well as doughnut-shaped vessel structures could not be imaged well in OCTA. Our observation is consistent with a recent study using AO-SLO FA. Therefore, similar to AO-SLO FA, AO-SLO with motion contrast is also useful in imaging retinal capillaries without any dye. Furthermore, although the utility of AO-SLO FA in imaging the hierarchical structure of retinal capillaries has not yet been examined, the method introduced in this study might be also available for AO-SLO FA.

As the lateral optical resolution of AO-SLO (5 μm) is around four times higher than OCTA (20 μm; Table 2), the superiority of the vessel plot-out ability of AO-SLO could be due to the resolution. However, the diameter of vessels that OCTA was unable to image was not smaller than the diameter of the vessels that were observed in this study. Another possible reason is the difference in imaging frame rate. The frame rate of AO-SLO is known to be 32 frames per second (96 frames in three seconds imaging), whereas the OCTA frame rate could be 1.3 to 1.4 frames per second (4 frames in 3-second imaging) that is lower than AO-SLO.

Diameter differences of each retinal vessel could be observed with AO-SLO while OCTA tends to equalize the vessel thickness. The lateral image resolution is 2 μm in AO-SLO, whereas it is 13.9 μm in OCTA (Table 2). Therefore, it was thought that there is a tendency for OCTA to equalize without depicting differences in vessel diameter. However, it
may be difficult to simply compare the diameter of the retinal vessel between the AO-SLO images and OCTA images. As the AO-SLO device creates en face images, retinal vessels could be imaged thicker in the deep image than in the superficial image. Thus, the retinal vessel diameter is dependent on the depth of the imaged vessels. Therefore, measurement of vessel diameter using AO-SLO might be of limited use. It is expected that a method to measure the diameter that is corrected for the depth would be developed. However, because the imaged vessels in AO-SLO are derived from the flow using motion contrast, the imaged lumen diameter cannot be equal to the actual anatomic diameter.

Based on previous reports, FAZ was clearly recognized and quantifiable in this study. A recent report showed that the area of FAZ on the OCTA is larger than that on AO-SLO when quantified. However, our quantification did not show this difference (Table 2). This might be due to the AO-SLO FA in the previous reports. Due to leakage in AO-SLO FA, it is likely to be smaller than the real image. In addition, diseased eyes were included in the previous reports. Although the presence or absence of axis length correction, difference in analysis software, and image merging might affect the FAZ measurement, the difference in FAZ area may also occur in AO-SLO due to difference in blood vessel imaging method (FA and motion contrast).

This study has the limitations inherent in any study of limited sample size. A dedicated adaptive optics instrument is used to acquire a large number of images, and this is followed by image averaging, which in turn is followed by montaging separate images to obtain a field of view large enough to have clinical utility. Furthermore, this study evaluated the ability of AO-SLO to visualize the retinal vascular layer only in healthy eyes. Despite this, we could only analyze half of all images (Table 2). Therefore, it would likely be even more difficult in diseased eyes.

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