

Diameter-dependent classification of dermal vasculature using optical coherence tomography angiography

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Background: Dermal blood vessels beneath the epidermis play critical roles in epidermal homeostasis and are functionally divided into several types, such as capillaries. Optical coherence tomography angiography (OCTA) is a powerful tool for the non-invasive assessment of dermal vasculature. However, the classification of vessel types has been limited. To address this issue, we proposed an algorithm for diameter-dependent classification that preserves three-dimensional (3D) information using OCTA.

Methods: OCTA data were acquired by a prototype swept-source-type optical coherence tomography (OCT) system, which was processed through several imaging filters: an optical microangiography (OMAG) imaging filter, a vesselness imaging filter, and a diameter map filter. All vessels were visually classified into three types based on their diameters, as micro-vessels, intermediate vessels, and thick vessels. Aging-related alterations and their association with the epidermis were investigated for each vessel type. The measurements were conducted on the cheeks of 124 female subjects aged 20–79 years.

Results: The 3D vascular structure was visualized by applying our proposed post-processing filters. Based on visual assessment, the thresholds for the diameters of the micro, intermediate and thick vessels were set at 80 and 160 µm. It was found that micro-vessels were predominantly located in the upper layer of the dermis and thick vessels in the deeper layer. Analysis of vessel metrics revealed that the volume density of the micro-vessels decreased significantly with age (r=-0.36, P<0.001) and was positively correlated with epidermal thickness (r=0.50, P<0.001). In contrast, the volume density of thick vessels significantly increased with age (r=-0.2, P<0.05) and was not significantly correlated with epidermal thickness (r=0.13, P ≥ 0.05).

Conclusions: In this study, we proposed a 3D quantification method using OCTA for dermal blood vessels and various vessel metrics, such as vessel volume density. This proposed classification will be beneficial for determining the function of the dermal vasculature and its diagnostic applications.

Keywords: Aging alteration; epidermal homeostasis; micro-vessels; optical coherence tomography angiography (OCTA); vascular classification

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Introduction

Cutaneous alteration with aging has been extensively studied for decades, with increasing interest in the medical and beauty industries. Given that the skin tissue is the outermost layer of the human body, extrinsic aging (photoaging) occurs due to exposure to the sun. Extrinsically aged skin is characterized by rough skin textures, uneven pigmentation, and deep wrinkles (1-4). Superficial vascular alterations, such as telangiectasias (5,6) and a decrease in capillary vessels (7,8) can also occur. These capillary alterations may be associated with changes in epidermal homeostasis (9). Non-invasive measurements of dermal vasculature would be beneficial to assess various skin conditions.

Based on optical technology, non-invasive assessment systems for assessing dermal vasculature have been developed. Representative measurement techniques include photoacoustic microscopy (10) and optical coherence tomography angiography (OCTA) (11). In the dermatological field, the OCTA system is considered to be beneficial for evaluating the capillaries beneath the epidermis with capillary-level resolution (11,12). Using OCTA, it has been suggested that skin vascularity is associated with various skin conditions in which dermalepidermal interactions play an important role, for example, solar lentigo, psoriasis, and wound healing (13-15). Furthermore, OCTA could evaluate vascular alteration with photoaging (16,17). Using OCTA and OCT, our previous study indicated decreasing vessel density and epidermal thinning with photoaging (16). One of the limitations in the prior OCTA studies, including ours, is the use of twodimensional (2D) vessel metrics (e.g., area density and diameter) to analyze 2D vascular images generated by the maximum intensity projection (MIP) of the 3D OCTA dataset. MIP analysis is particularly useful when tail-like artifacts emerge beneath blood vessels because these artifacts can prevent OCTA from visualizing the three-dimensional (3D) structural images of vascular networks (18,19). As Untracht et al. (20) demonstrated, 3D-based vessel analysis can potentially visualize the details of vascular geometry that cannot be observed using 2D analysis. Several attempts have been made to explore the possibility of 3D-based vascular analysis in OCTA measurements, showing improved vessel density assessments (21-23). Because 3D analysis is considered to retain more information about the actual vascular architecture than 2D analysis, it is beneficial for classifying the characteristics of each vessel's functional type, such as capillary and post-capillary venules. For instance,

capillaries can be recognized as thin vessels comprising a limited number of pericyte layers, implying a potential connection between their structures and the maintenance of epidermal homeostasis via nutrient supply (24). These vessel types could be roughly characterized based on their diameters in the dermatological field (25); however, this has not been proposed.

This study aimed to develop a diameter-dependent analysis method for dermal blood vessels exhibiting the complex structure shown in *Figure 1*. Various types of vessel metrics that preserve 3D information were proposed. The effects of photoaging and the correlation with epidermal thickness of diameter-dependent vascular types were investigated using the proposed method. We present this article in accordance with the STROBE reporting checklist (available at https://qims.amegroups.com/article/ view/10.21037/qims-23-1790/rc).

Methods

Optical coherence tomography (OCT) system

In this study, we employed a prototype swept-source type OCT (SS-OCT) system equipped with a swept-source laser with a central wavelength of 1,310 nm and a scanning rate of 200 kHz (SL1310 V1-20048, Thorlabs, Inc., NJ, USA). The SS-OCT system was designed with a handheld measurement probe, including a galvanometric mirror (GV S002, Thorlabs, Inc.) and an objective lens (LSM03, Thorlabs, Inc.) (26). The system delivers a dynamic range of ~105 dB with an axial resolution of ~6 µm in the tissues and a lateral resolution of ~15 µm. The field of view (FoV) was designed to be 8 mm × 8 mm. To achieve OCTA imaging, the volume scanning pattern for this FoV included 800 A-scans per B-scan and 800 B-scans per volume, with each B-scan repeated four times. This scanning pattern provided a pixel spacing between adjacent A-scans of 10 µm (pixel size of 10 μ m in x and y dimensions). In the axial direction (z dimension), the pixel size was 3.75 µm. The penetration depth was approximately 2.0 mm at maximum; however, due to the attenuation of the OCT signal, it can be visualized up to a depth of 140 pixels (500 µm) from the surface of the skin. To further improve the image quality for quantitative analysis, the volume scan was repeated twice at the targeted region of interest and then registered and averaged to give rise to a final single volume for later quantification (see below). The total scanning time to obtain the final single volume was approximately 26 s.

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Figure 1 Illustration of dermal skin vasculature and vessel metrics except vessel density in skin tissue. Epidermal thickness (μ m) is determined by detecting the dermal-epidermal interface. DEJ-vessel distance (μ m) is defined as the distance from the position of the DEJ to the depth where the vascular density in the MIP image exceeded 5%. Vessel branches (mm^{-3}) indicate the number of branching points divided by the volume of the analysis range. Vessel angle (°) denotes the mean angle of the core line in the superficial layer. DEJ, dermal-epidermal junction; MIP, maximum intensity projection.

Post-processing

After data acquisition using the SS-OCT system, the raw OCT data were processed using several imaging filters: an optical microangiography (OMAG) imaging filter, a vesselness imaging filter, and a diameter map filter, as shown in the flowchart in *Figure 2A*. The OMAG technique was used to convert the OCT signal into an angiographic image with high accuracy (11,27). In this method, the OCT signal C can be described as follows:

$$C(\mathbf{r},t) = A(\mathbf{r},t)e^{-i\Phi(\mathbf{r},t)}$$
^[1]

where $A(\mathbf{r}, t)$ is the amplitude component of the OCTA signal, $F(\mathbf{r}, t)$ is the phase component, and \mathbf{r} is the spatial position of the OCTA signal. In the OMAG imaging, the intensity (I_{OMAG}) of the image due to blood flow was calculated using Eq. [2].

$$I_{\text{OMAG}}(\mathbf{r}) = \frac{1}{N-1} \sum_{i=0}^{N-1} |C_{i+1}(\mathbf{r}) - C_i(\mathbf{r})|$$
[2]

where N is the number of repeats of the B-scans and $C_i(r)$ is the OCT signal at the spatial position r during the *i*-th repetition of the B-scan. A representative example of an OCTA image of human forearm skin is shown in *Figure 2B,2C*. As we know, an OCTA image naturally contains noise in the z-direction, called a tail-like artifact (28), which can be easily appreciated in each B-scan OCTA image (*Figure 2D*).

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While there are several methods available in the literature for tail artifact removal [e.g., (28)], here we adopted a method using an exponentially decaying function in the z-direction (29) to remove the tailing artifacts in the OCTA datasets. In this case, the tailing artifact removed from the OMAG image $I_{\text{TailRemoved}}$ can be expressed as:

$$I_{\text{TailRemoved}} = I_{\text{OMAG}} \left(z + \Delta z \right) - I_{\text{OMAG}} \left(z \right) \cdot a_0 \exp\left(-k\Delta z \right)$$
[3]

where Δz represents the distance in the z-direction (depth) from the superficial vessel, a_0 is the decay coefficient, and k is the decay rate that determines the degree of decay. The operation of this tail-removal filter would result in tail artifact-free OCTA images (*Figure 2E, 2F*).

Next, the tubular structure (i.e., vesselness) of the vessels was extracted by assuming that the shape of the dermal vessels in OCTA is tubular. A multiscale Hessian filter was applied to the tail-removed image to obtain a vesselness image as follows:

$$I_{\text{Vesselnsee}}\left(\boldsymbol{r}\right) = \max_{\{s\}} \left(\mathbf{H}\left(\boldsymbol{r},s\right) I_{\text{TailRemoved}}\left(\boldsymbol{r}\right)\right)$$
[4]

where H represents the 3D Hessian matrix, defined as $\mathbf{H} = \partial^2 / \partial \mathbf{r}^2$. The tubular structure was extracted by calculating the eigenvalues of the Hessian matrix (Figure 2G, 2H). In the vesselness imaging filter, the tubular structures were set to a non-zero value, whereas other structure types, such as planes or blobs, were set to zero. While calculating the Hessian matrix, the maximum value of vesselness for each position was determined by varying the vesselness scale s from 0.1 to 10 with logarithmic step size (30,31). Lastly, the tubular vessel structures derived through the above process were skeletonized to shrink all vessel types into one pixel in size (i.e., vessel cross-section only represented by a one-pixel width). In doing so, a centerline extraction filter was passed through the vessel image. The decision-tree method was used for this process (32). In this method, pixels were removed from the outermost layer of the tube, resulting in a skeleton of the vesselness image. The diameter-based information within the vessel was colormapped onto the skeleton using a method developed using the Euclidean distance transform (33) to preserve the actual 3D morphology of the blood vessels (Figure 21,27). Finally, diameter-dependent vessel classification was achieved based on a visual observation of the OMAG images and diameter maps.

Vessel metrics

In this study, various vessel metrics were derived from



Figure 2 Flowchart and examples of OCTA imaging and processing in the cheek. (A) Flowchart of vessel imaging and extraction. (B-L) Examples of the OCTA image processing to enhance the image quality for later quantitative assessment. (B) MIP imaging of an 8 mm by 8 mm OMAG image filter, where the scale bar is 1 mm. MIP manipulation was employed from the skin's surface at a depth of 20 pixels (75 µm) to 140 pixels (525 µm). The red rectangle shows the manually chosen ROI, and the size of ROIs was 2 mm by 2 mm, which was used to show the images (C-L). (C,E,G,I,K) are the OCTA MIP images of the ROI, and (D,F,H,J,L) are representative OCTA B-scans, i.e., cross-sectional OCTA images. (C,E,G,I,K) Representative micro- and thick vessels are denoted by red and green arrows, respectively. The scale bar (both vertical and lateral) is 200 µm. (C,D) Original OMAG images containing tailing artifacts. (E,F) Corresponding tail-removed images after applying the tail artifact removal algorithm. (G,H) Corresponding vesselness images after applying a Hessian filter to extract vessel tubular structures. (I,J) Skeletonized vessel maps with diameter information coded with color. The color bar indicates the thickness of the vessel diameter, whereas red and yellow indicate thin and thick vessels, respectively. (K,L) Skeletonized vessel maps applying the diameter-dependent classification of three vessel types: micro- (red), intermediate (gray), and thick (green) vessels. The dashed line indicates the vessels from the depth of the DEJ to 80-pixel depth (300 µm) for calculation of vessel metrics. OMAG, optical microangiography; OCTA, optical coherence tomography angiography; MIP, maximum intensity projection; ROI, region of interest; DEJ, dermal-epidermal junction.

the vessel images generated through post-processing and are summarized in *Table 1. Figure 1* illustrates the vessel metrics for skin tissue. The epidermal thickness (ET) was determined by detecting the dermal-epidermal interface, that is, the dermal-epidermal junction (DEJ), using intensity profiles of OCT signals (34). All the vessel-related indicators, except ET, were calculated using the depth obtained from ET detection as the starting depth. The

Table 1 Summary of metrics evaluated in th	his study
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Metric	Abbrv.	Unit	Description				
Epidermal thickness	ET	μm	ET of a skin tissue				
DEJ-vessel distance	-	μm	Distance between dermal-epidermal junction and peripheral blood vessels				
Vessel areal fraction	VAF	%	MIP density of the vesselness image				
Vessel volume density	VVD	%	Vessel density calculated by the vessel diameter map (all vessels)				
Micro-vessel volume density	mVVD	%	Vessel density of micro-vessels calculated by the vessel diameter map (a diameter of ${\leq}80~\mu\text{m})$				
Intermediate-vessel volume density	iVVD	%	Vessel density of intermediate-vessels calculated by the vessel diameter map (80 μm < a diameter <160 $\mu m)$				
Thick vessel volume density	tVVD	%	Vessel density of thick vessels calculated by the vessel diameter map (a diameter of ${\geq}160~\mu\text{m})$				
Vessel branch	-	m ⁻³	Number of branching points of blood vessels per volume				
Vessel angle	mVA	0	Mean angle of peripheral micro-vessels (horizontal =0°, vertical =90°)				

DEJ, dermal-epidermal junction; MIP, maximum intensity projection.

DEJ-vessel distance was defined as the distance from the location of the DEJ to the depth where the vascular density in the MIP image exceeded 5%, which is the estimated threshold excluding the noise such as dot-like structures through vesselness filters. This metric indicates that the smaller the value, the greater the number of blood vessels present beneath the epidermis. The vessel area fraction (VAF) was calculated from the MIP image relative to the vessel from the depth of the DEJ to 80-pixel depth (300 µm) by considering the individual variations in the epidermal thickness and the degree of attenuation. The VAF was calculated by counting the pixel number of the MIP of the vesselness image. The equation can be written as follows:

$$VAF = \frac{1}{a} \sum_{x,y} COUNT \left(MIP \left(I_{Vesselness} \left(\boldsymbol{r} \right) \right) \right)$$
[5]

where *a* represents the area of the ROI, MIP is an operation function taking the max value along the *z*-direction at position (x,y), and COUNT is an operation function for counting the number of pixels at which the MIP value was non-zero, namely, tubular structure.

The volume density of the vessels from the depth of the DEJ to the 80-pixel depth was denoted as the vessel volume density (VVD). The VVD was calculated by converting the diameter of the skeleton into the effective area of the tubular section and integrating it over the skeleton. VVD can be written as:

$$VVD = \frac{1}{\upsilon} \sum_{R} \pi R^2$$
 [6]

where π represents a circular constant, R represents the radius of the voxel in the skeleton, and the integrated volume is divided by the total voxel volume n within the analysis range. The integration process was diameter selective because VVD was calculated using a diameter map image. Here, VVDs comprising micro-vessels, intermediate vessels and thick vessels were defined as micro-vessel volume density (mVVD), intermediate-vessel volume density (iVVD) and thick vessel volume density (tVVD), respectively. Correlations between ET and the diameterselective VVDs were investigated to validate the diameter thresholds by setting up with two VVDs, i.e., VVDs of vessels with diameters larger or smaller than a given threshold.

The number of vessel branches was derived by calculating the number of branching points from the skeletonized vessels and dividing it by the volume of the analysis, ranging from the depth of the DEJ to 80-pixel depth. The micro-vessel angle (mVA) was defined as the mean angle of the core line from the depth of the DEJ to an 11-pixel depth (approximately 40 µm) corresponding to the thickness of the papillary dermis (35), focusing on the angle in the micro-vessels of the superficial layer. The mVA was derived by calculating the arctangent from the slope of the least-square regression line. This operation was conducted

Group	Age range (years)	Subject number		Fitzpa	atrick ski	n type		Orealizer hakita	Destruction
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G0	20–39	37	0	2	14	19	2	5	0
G1	40–59	44	0	1	19	21	3	2	17
G2	60–79	43	1	0	20	21	1	3	43

Table 2 Summary of subjects

along the skeleton, and the box size of the least-square regression was set to be (7,7,7) voxels. The mVA can be expressed as:

$$mVA = |\arctan(l)|$$
[7]

where l is the slope derived from the least-square regression analysis. The absolute value was taken to be in the range of the resulting angle from 0° to 90°.

Subjects

The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013) and approved by the Ethics Committee of Shiseido Co., Ltd. (No. C02109). The subjects were 129 healthy Asian females aged 20-79 years, they were enrolled after obtaining informed consent. The measurement positions were on the right sides of their cheeks, in the sun-exposed area. The measurements began after a moderate acclimatization period at approximately 23.0 °C and 45% RH. Five subjects were excluded due to the large body motion during OCTA measurements. The number of subjects, identified on the basis of the Fitzpatrick classification of skin type, were I (n=1), II (n=3), III (n=53), IV (n=61), and V (n=6), as determined using a questionnaire. The subjects were divided into three groups based on age, G0 (20-39 years old), G1 (40-59 years old) and G2 (60-79 years old), to examine the relationship between age and vessel metrics. Table 2 summarizes the population, smoking habits, and hormone status for each age group.

Statistical analysis

The Wilcoxon rank correlation test, in which P values were adjusted using the Bonferroni method, was employed to investigate whether there were statistically significant differences between the groups. Spearman's rank correlation coefficients were calculated for the vessel metrics.

Results

Figure 2E-2L show vascular images passing through our post-processing filters after being processed using the OMAG algorithm. The tail-like artifacts in the OMAG image (Figure 2E, 2F) were eliminated by applying the tail removal filter. Additionally, the multi-scaled Hessian filter enhanced the visibility of the vessel structures (Figure 2G,2H). Skeletonization and Euclidean distance transformation of the vesselness image allowed for a resulting diameter map in 3D (Figure 2I,27). Visual observations of the OMAG images and the diameter maps (e.g., Figure 2C, 2I) allowed for a diameter-dependent vessel classification, resulting in micro-vessels, estimated as having a diameter less than 80 µm, intermediate-vessels, estimated having a diameter between 80 and 160 µm, and thick vessels, estimated having a diameter greater than 160 µm (Figure 2K, 2L). Following these post-processing steps, a 3D vascular structure was visualized, with micro-vessels predominantly in the upper layer of the dermis and thicker vessels in the subsequent layers.

Figure 3 illustrates the aging alteration for the age groups of G0, G1 and G2. Aging alterations in ET and VVD were plotted as boxplots in *Figure 3A,3B. Figure 3C,3D* present a typical vesselness image for G0, G1, and G2. ET shows a decreasing trend with age. Conversely, no statistically significant difference was found in VVD between groups G0 and G1, although a significant difference was observed for groups G0 and G2.

In *Figure 4*, the scatter plots of the ET and vessel volume densities, mVVD, and tVVD are presented. The residuals between the vessel metrics and age were calculated to exclude the influence of aging alterations. There was a positive correlation between mVVD_{resid} and ET_{resid} (r=0.50, P<0.001), whereas there was no significant correlation between tVVD_{resid} and ET_{resid}. All correlations between vessel metrics are summarized in *Table 3*. Significant correlations with ET_{resid} were also observed for iVVD_{resid}.



Figure 3 Age-related alterations of ET and VVD and the vesselness images for each age group: G0 (20–39 y.o.), G1 (40–59 y.o.), and G2 (60–79 y.o.). (A) Boxplot of ET for age groups. (B) Boxplot of VVD for age groups. ***, P<0.001; **, P<0.01; *, P<0.05; n.s., P≥0.05. (C) Representative vesselness images (top row) and corresponding B-scan images (bottom row) for G0 (25 y.o.), G1 (50 y.o.), and G2 (74 y.o.), respectively. (D) Representative images of skeletonized vessel maps applying the diameter-dependent classification (top row) and corresponding B-scan images (bottom row) for G0 (25 y.o.), G1 (50 y.o.), and G2 (74 y.o.), respectively. Color represents three vessel types: micro- (red), intermediate (gray), and thick (green) vessels. The dashed line (bottom row) indicates the vessels from the depth of the DEJ to 80-pixel depth (300 µm) for calculation of vessel metrics. ET, epidermal thickness; VVD, vessel volume density; y.o., years old; DEJ, dermal-epidermal junction.

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Figure 4 Scatter plots between ET and vessel metrics; mVVD and tVVD. The data were normalized, and its residual against age was plotted. (A) ET_{resid} vs. mVVD_{resid}. (B) ET_{resid} vs. tVVD_{resid}. In the plots, a correlation coefficient based on Spearman's rank correlation test was calculated. ***, P<0.001; n.s., P≥0.05. ET, epidermal thickness; mVVD, micro-vessel volume density; tVVD, thick vessel volume density.

Table 3 Correlation coefficients between ET and vessel metrics ¹											
Metric	ET (µm)	DEJ-vessel distance (µm)	VAF (%)	VVD (%)	mVVD (%)	iVVD (%)	tVVD (%)	Vessel branch (m ⁻³)	mVA (°)		
Age	-0.56**	0.23*	-0.22*	-0.24**	-0.36***	-0.34***	0.20*	-0.37***	-0.21*		
ET	-	-0.54***	0.42***	0.48***	0.56***	0.47***	-0.01 ^(n.s.)	0.58***	0.17 ^(n.s.)		
ET _{resid}	-	-0.53***	0.37***	0.43***	0.50***	0.36***	0.13 ^(n.s.)	0.49***	0.05 ^(n.s.)		

[↑], Spearman's rank correlation test: ***, P<0.001; **, P<0.01; *, P<0.05; n.s., P≥0.05. ET, epidermal thickness; DEJ, dermal-epidermal junction; VAF, vessel areal fraction; VVD, vessel volume density; mVVD, micro-vessel volume density; iVVD, intermediate-vessel volume density; tVVD, thick vessel volume density; mVA, vessel angle.

DEJ-vessel distance $_{\rm resid}$, and vessel branch $_{\rm resid}$, although not for mVA $_{\rm resid}$.

Figure 5 shows a plot of the correlation coefficient between ET_{resid} and the diameter-selective vessel volume densities as a function of the diameter threshold. A significant correlation was observed between ET_{resid} and VVD_{resid} composed of vessels thinner than the diameter threshold of 80 µm, while there is no significant correlation between ET_{resid} and VVD_{resid} composed of vessels thicker than the diameter threshold of 160 µm.

Discussion

In this study, we proposed an analysis method for OCTA that preserved the 3D information of blood vessels in the skin and investigated both aging-related alterations and their association with the epidermis. Although several OCTA studies have examined the contribution of vessels to aging alterations in the skin (16,17,36), these studies were

primarily limited to two-dimensional analyses, that is, MIP vessel density analysis. This limitation is largely attributed to the tail artifacts in the OCTA signals. Our post-processing method mitigated this limitation and successfully visualized vesselness in 3D with high accuracy by applying a tail-removal imaging filter (11) to an OMAG image and passing it through a multiscale Hessian filter. Consequently, we could evaluate the vessel densities on the basis of vessel diameters, namely on the basis of vessels being micro, intermediate and thick.

Regarding photoaging alteration, this study shows that the VVD in the cheek area decreased with age. A similar aging trend was observed in the histological section images (37) and the MIP images of OCTA (16). As the VVD retains non-invasive 3D information, it represents a more accurate vessel structure of the organism. The aging trend of VVD was not monotonic and was found to decrease significantly in the G0–G2 group. However, there was no significant difference in the G0–G1 group, possibly due to a significant



Figure 5 Correlation coefficient between ET_{resid} and the diameterselective vessel volume densities as a function of the diameter threshold. In the plot, two types of VVDs are considered; VVD composed of vessels thinner than the diameter threshold and VVD composed of vessels thicker than the diameter threshold. A correlation coefficient of ET_{resid} for each VVD_{resid} was calculated based on Spearman's rank correlation test. ET, epidermal thickness; VVD, vessel volume density.

change in the secretion of female hormones caused by menopause (38). Classifying VVD on the basis of vessel diameter, mVVD decreased with age, but tVVD increased. In ovariectomized rats, mimicking human menopause, it has been reported that the capillary diameter decreases, suggesting a decrease in capillaries (38). Based on past knowledge, mVVD is suggested as being representative of the density of capillaries. Herein, it is known that there is no change in the level of skin redness with age (39,40). This unchanged skin redness may be associated with increased noncapillary, larger vessel types, such as post-capillary venules, with increased tVVD. Whether these alterations in mVVD and tVVD are specific to photoaging needs further investigation.

Dermal blood vessels visualized by OCTA may also play a crucial role in maintaining epidermal homeostasis. The correlation between vessel metrics and the epidermis was investigated to clarify the effect. Considering that both vessel metrics and ET are influenced by aging, we conducted a correlation analysis after accounting for the effect of age. A significant positive correlation with ET_{resid} for mVVD_{resid}, not for tVVD_{resid}, was observed here, suggesting that the number of micro-vessels, mVVD, may

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play an important role in securing epidermal thickness rather than the number of thick vessel, tVVD. Capillaries beneath the epidermis maintain homeostasis in the epidermis by exchanging nutrients and waste products, regulated by Starling's force (25,41). The results of this study suggested that blood vessels with diameters of 80 µm or less, as visualized by OCTA, contributed to epidermal homeostasis as capillaries, which in turn may be associated with ET.

To further investigate the association of micro- and thick vessels with epidermis, the correlation of ET for diameter-selective vessel volume densities was calculated by systematically changing the diameter threshold. The highest correlation was observed between $\mathrm{ET}_{\!\mathrm{resid}}$ and VVD_{resid} composed of vessels thinner than the diameter threshold of 80 µm, suggesting that the diameter threshold of micro-vessels estimated by the visual observation was reasonable. A positive correlation was observed between ET_{resid} and the vessel angle composed of vessels thinner than the diameter threshold of 40 µm (data not shown), implying that vessels whose diameters were approximately 40 µm can be dedicated to efficient nutrition and other aspects, for example, supporting mechanical stiffness (42), through their vertical alignment. It was also found that the correlation coefficient between ET_{resid} and VVD_{resid} for vessels thinner than the those having the diameter threshold of 40 µm was less than that of vessels of diameter 80 µm, suggesting that the number of the vessels having a diameter of less than 40 µm was relatively small even though the association with epidermis is high. In contrast, no significant correlation was found between ET_{resid} and VVD_{resid} for vessels having a diameter greater than the diameter threshold of 160 µm, which may be due to their dissociation from epidermal homeostasis.

Regarding the intermediate vessels, a similar trend against ET_{resid} was found for iVVD_{resid} (r=0.36, P<0.001). It is suggested that vessels with intermediate diameter may contribute to maintaining epidermal homeostasis by supporting capillaries. Our study is the first to focus on diameter-dependent vascular classification and examine its relationship with the epidermis, non-invasively. In addition to mVVD and iVVD, DEJ-vessel distance and vessel branch correlate with ET as shown in *Table 2*. Because capillaries are densely distributed beneath the epidermis, these metrics could also represent the condition of the capillaries as a separate aspect from mVVD.

In this study, we proposed 3D quantitative vessel parameters, represented by VVD, among the skin and vessel metrics obtained by OCTA, to investigate the aging alteration of dermal blood vessels and their contribution to the epidermis. The proposed diameter-dependent classification using OCTA is expected to be beneficial in assessing the degree of photoaging and epidermal conditions.

One limitation is that the skin types are imbalanced for Types III and IV, and it is not known whether the results obtained in this study are similar for other skin types. Another limitation is that the shape of a single vessel (e.g., dots, curved, or coiled) is not reflected in this study (43,44). The association of dermal vasculature with aging and the epidermis may become stronger by assessing the 3D geometry of a single vessel. Finally, statistical analysis revealed that the behavior of micro and intermediate vessels is similar. The interpretation of the role of intermediate vessels is still ambiguous. Further investigation regarding every vessel type is necessary to clarify the process of photoaging or intrinsic aging of the vessel in the skin.

Conclusions

In this study, we proposed a 3D quantification method for dermal blood vessels and various vessel metrics, such as VVD. Our results revealed that the mVVD decreased with age and was positively correlated with ET. These suggest that the micro-vessels observed using OCTA may play a vital role in maintaining the thickness of the epidermis. The proposed diameter-dependent vascular analysis using OCTA could be beneficial for understanding the roles of dermal blood vessels and assessing photoaging.

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Footnote

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Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). The study was approved by the Ethics Committee of Shiseido Co., Ltd. (No. C02109) and informed consent was obtained from all individual participants.

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