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Retina

Inter-Individual Topographic Variation of Choroidal Thickness in Healthy Eyes on Swept-Source Optical Coherence Tomography

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Methods: This was a retrospective study. Macular CT data for 900 0.2 \times 0.2-mm grids from 410 healthy eyes were collected from swept-source optical coherence tomography. Following the analysis of factors associated with mean CT, the β -coefficients of the included associated factors in each grid were summarized for choroidal thickness changes analysis. Additionally, the coefficient of variance (CoV), coefficient of determination (CoD), and coefficient of variance unexplained (CoVU) for CT were calculated in each individual grid to investigate the inter-individual choroidal variations pattern.

Results: Sex ($\beta = -17.26$, female vs. male), age ($\beta = -1.61$, per 1 year), and axial length ($\beta = -18.62$, per 1 mm) were associated with mean macular CT. Females had a thinner choroid in all 900 grids (0.5–26.9 µm). As age increased, the CT noticeably decreased (8.74–19.87 µm per 10 years) in the temporal regions. With axial length elongation, the thinning (7.94–24.91 µm per 1 mm) was more evident in subfoveal and nasal regions. Both the CoV (34.69%–58.00%) and CoVU (23.05%–40.78%) were lower in the temporal regions, whereas the CoD (18.41%–39.66%) was higher in the temporal regions.

Conclusions: Choroidal thinning is more predominant in the subfoveal and nasal regions with axial length elongation, but in the temporal region with aging. The interindividual variation of CT is higher and less determined by sex, age, or axial length in the nasal regions.

Translational Relevance: Topographic variation should be considered when interpreting choroidal thickness.

Introduction

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The choroid, a vascular structure ranging from Bruch's membrane to the choroidal–scleral interface, plays an important role in supporting the structure and function of the retina.¹ The development of enhanced depth imaging² and swept-source optical coherence tomography (SS-OCT)³ makes it possible for in vivo choroidal structure visualization and accurate thickness measurement. Previous studies have reported that changes in choroidal thickness (CT) are associated with physiological processes such as aging,⁴ axial length elongation,⁵ and the development of various eye diseases,⁶ including age-related macular degeneration,⁷ myopia,⁸ and pachychoroid diseases.⁹

However, most of these studies focused only on the mean and subfoveal CT^{10} or, in the case of the Early Treatment Diabetic Retinopathy Study (ETDRS), regional $CT.^{11,12}$ Indeed, the choroid is a three-dimensional structure, and the distribution of CT in healthy eyes remains inconclusive.¹³ In widefield monitoring of the choroidal thickness and

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vascular densities, the thickness values, symmetrical/asymmetrical patterns, and topography of the choroid are diverse.^{14–16} Various studies have reported that the thickest areas are the superior,¹⁷ temporal,^{8,18} and subfoveal regions.¹⁹ Also, studies have reported varied choroidal thickness distribution patterns,¹⁴ suggesting the inter-individual CT variations are too complex to be evaluated based on several subregions.

Recently, a 900-grid analysis algorithm for CT has been developed for Triton Multimodal SS-OCT (Topcon Healthcare, Tokyo, Japan), allowing for nuanced evaluation of the topographic features of macular CT.^{20,21} In this study, we investigated the relationship between CT in each of the 900 grids and various CT-associated factors to map CT variations. Additionally, we constructed distributions for the coefficient of variance (CoV), coefficient of determination (CoD), and coefficient of variance unexplained (CoVU) across the 900 grids. Incorporating all of these parameters resulted in comprehensive representations of the nuanced topographic patterns of interindividual choroidal variation.

Methods

Data Collection

Our research adhered to the tenets of the Declaration of Helsinki. Institutional Review Board (IRB) approval was obtained from the Joint Shantou International Eye Center (JSIEC) of Shantou University and The Chinese University of Hong Kong. Informed consent was not required by the IRB as this was a retrospective study.

Outpatient records with OCT examinations from January 2018 to November 2020 at JSIEC were reviewed to identify healthy individuals with no systemic diseases. OCT images with a resolution of 512 \times 256 pixels were obtained from the Triton Multimodal SS-OCT using a three-dimensional 7 \times 7-mm macular scanning mode. Patients with chorioretinal diseases, glaucoma, history of ocular surgery or treatment, or OCT image quality scores below 40 were excluded. Also, given the interocular symmetry pattern,^{22,23} only one eye of each included individual was enrolled randomly for analysis.

For every included eye, segmentation of the choroid (from Bruch's membrane to the choroidal–scleral junction) in each B-scan image was manually checked by a trained clinician who was masked to the subjects' clinical information to rectify any potential segmentation errors. Axial length and refractive error were input to correct magnification errors using the inbuilt algorithm of the Triton Multimodal SS-OCT.²⁴ Finally, CT data for 900 0.2×0.2 -mm grids in the 6 \times 6-mm macular areas were collected.

Also included for further correlation analysis were general information regarding sex and age; basic ophthalmic parameters of uncorrected visual acuity on a logMAR scale; diopter spherical and cylindrical refraction; axis of astigmatism; and spherical equivalents obtained from a Topcon full automatic optometer (KP8800). Other data collected included intraocular pressure (Haag-Streit AG, Köniz, Switzerland) and biological parameters including axial length, anterior chamber depth, lens thickness, and central corneal thickness obtained from ocular biometry (OA-2000; Tomey Corporation, Nagoya, Japan).

Study Design

Figure 1 shows a flowchart of the study protocol and data analysis. First, the association of all demographic and clinical parameters with the mean CT in the whole macular region was analyzed by univariable regression. The parameters with P < 0.05 in univariable regression were then subjected to multivariable regression analysis for the mean macular CT. Statistically significant variants of all parameters from the multivariable regression were included in the further analysis of each grid. Then, each 0.2×0.2 -mm grid was regarded as an independent area in the following data analysis.

The associated factors identified in the previous step were included in multivariable regression analyses to observe their relationship of the CT in each grid ($y_1 \sim y_{900}$) by R Studio 4.1.3 (R Foundation for Statistical Computing, Vienna, Austria). The β -coefficients of each factor were collected to analyze changes in the thickness of the choroid for every 1 unit of change in the corresponding factor for each grid.

The inter-individual variation of CT was evaluated using our self-defined CoV, CoD, and CoVU. The CoV, calculated as the ratio of the CT standard deviation to its mean thickness value in the 410 eyes, was used to describe the inter-individual variation in each grid. The CoD, the R^2 values of the associated factors from multivariable regression, was collected to evaluate the extent to which each associated factor accounted for the inter-individual variation within each grid. The CoVU values were computed as CoV × (1 – CoD) to determine the residual inter-individual variations that remained unexplained by the associated factors.

Finally, the β -coefficients of the 900 grids for each associated factor (CoV, CoD, and CoVU)



Figure 1. Flowchart of the study protocol.

were summarized as topographic maps constructed using Excel (Microsoft Corporation, Redmond, WA). For better visualization, left-to-right reversal was performed in all left-eye samples, aligning the temporal-to-nasal direction with the left-to-right.

Results

General Information and Screening of Influence Factors

A total of 410 samples were collected, encompassing individuals 6 to 87 years old (mean age, 33.59 \pm 25.21 years), including 198 males (48.29%) and 212 females (51.71%), and the mean CT of the whole macular area was 195.20 \pm 70.53 µm (Table). The mean macular CT was found to be associated with various factors in the univariable regression analyses, including age, sex, image quality, uncorrected visual acuity, intraocular pressure, spherical diopter, spherical equivalent, axial length, anterior chamber depth, and lens thickness. But, in multivariable regression, only sex (standardized $\beta = -17.26$, female vs. male), age $(\beta = -1.61 \text{ per } 1 \text{ year})$, and axial length $(\beta = -18.62 \text{ per } 1 \text{ year})$ per 1 mm) were statistically associated with the mean macular CT (Table) and were used in the analysis of the CT in each grid.

CT Variation Related to Sex, Age, and Axial Length in 900 Grids

Compared to males, females exhibited thinner choroid in all 900 grids, ranging from 0.5 to 26.9 μ m. The difference was minimal in the inferior nasal, superior nasal, and middle temporal regions but was more pronounced in the central regions (Fig. 2A). Both aging and axial length elongation were associated with choroid thinning in all 900 grids. For every 10-year increase in age, the decrease in CT ranged from 8.74 to 19.87 μ m, which was more prominent in the temporal choroid than nasal (Fig. 2B). With a 1-mm increase in axial length, CT thinning ranged from 7.94 to 24.91 μ m and was more prominent in subfoveal and nasal areas (Fig. 2C).

Inter-Individual Variation in CT in the 900 Grids

Inter-individual variation of CT is reflected by the distribution of CoV, CoD, and CoVU. Figure 3A illustrates the inter-individual CoV of CT distribution, which ranged from 34.69% to 58.00% (mean $41.90\% \pm 5.28\%$) and increased from the temporal to the nasal. The CoD distribution pattern is shown in Figure 3B; values ranged from 18.41% to 39.66% and decreased from the temporal to nasal regions. Figure 3C shows a map of CoVU values (range, 23.05%–40.78%) to

Table.Univariable and Multivariable Regression Associations of Demographic and Ophthalmic ParametersWith Mean Choroidal Thickness

		Univariable Regression		Multivariable Regression	
Parameter		β -Coefficient	Р	β -Coefficient	Р
Age (y), mean \pm SD	33.59 ± 25.21	-1.439	<0.001	-1.607	<0.001
Female, <i>n</i> (%)	212 (51.71)	-23.634	0.001	-17.261	0.011
Right eye, <i>n</i> (%)	207 (50.49)	10.639	0.127	NI ^a	NI ^a
TopQ image quality, mean \pm SD	56.90 \pm 9.12	1.823	<0.001	-0.275	0.497
Uncorrected visual acuity (logMAR), mean \pm SD	0.78 ± 0.39	-25.774	0.004	-17.253	0.076
Intraocular pressure (mmHg), mean \pm SD	15.36 ± 3.55	2.656	0.007	0.473	0.603
Spherical diopter (D), mean \pm SD	-1.96 ± 3.53	4.082	<0.001	-3.647	0.519
Cylindrical diopter (D), mean \pm SD	-0.70 ± 1.29	0.202	0.074	NI ^a	NI ^a
Axis of astigmatism, mean \pm SD	82.28 ± 70.47	-0.025	0.620	NI ^a	NI ^a
Spherical equivalent (D), mean \pm SD	-2.31 ± 3.77	3.598	<0.001	1.72	0.745
Axial length (mm), mean \pm SD	24.17 ± 1.47	-10.213	<0.001	-18.619	<0.001
Anterior chamber depth (mm), mean \pm SD	3.46 ± 0.43	37.407	<0.001	16.533	0.154
Lens thickness (mm), mean \pm SD	3.89 ± 0.55	-48.65	<0.001	8.925	0.515
Central corneal thickness (µm), mean \pm SD	531.79 \pm 33.30	0.079	0.499	NI ^a	NI ^a
Mean choroidal thickness (μ m), mean \pm SD	$195.20~\pm~70.53$	NI ^a	NI ^a	NI ^a	NI ^a

Bold value means statistically significant (P < 0.05).

^aNot included in the regression model.



Figure 2. Topographic maps of CT attenuation with sex, age, and, axial length in 900 grids. (A) Differences in choroidal thickness between males and females. (B) Choroidal thickness thinning per 10 years of increasing age. (C) Choroidal thickness thinning per 1-mm increase in axial length. The left-to-right direction in all maps reflects temporal-to-nasal regions. The central *black dots* indicate the fovea. All numbers in the distribution maps are micrometers.

highlight the unexplained effect of factors other than sex, age, and axial length that were more pronounced in the nasal regions compared to the temporal region.

Discussion

Our study has found that the thinning pattern of the choroid with aging is notably more pronounced in the temporal area, whereas it becomes more evident in the subfoveal and nasal areas with axial length elongation. Margolis and Spaide¹⁹ observed a more pronounced decrease in CT with aging in normal eyes at positions 2.5 mm and 3.0 mm temporal to the fovea, but their investigation was limited to the CT on a horizontal line at 500-µm intervals crossing the fovea. Our map of thickness changes in the 900 grids effectively demonstrates that aging has more effect on the choroidal thinning process in the temporal area, which can be seen in samples 1 and 2 in Figure 4. Previ-



Figure 3. Topographic maps of inter-individual variation of macular CT. (**A**) Coefficient of variation. (**B**) Coefficient of determination, which represents the percentage of variation that can be explained by gender, age, and axial length. (**C**) Coefficient of variance unexplained, which represents the residual variation. The left-to-right direction in all maps reflects temporal-to-nasal areas around the fovea. The central *black dots* indicate the fovea. All numbers in the distribution maps are percentages.

ous studies have shown decreases in vascular density, overall luminal area, and choriocapillaris diameter with age,²⁵ whereas the temporal parafoveal and perifoveal areas exhibit higher choriocapillaris and choroidal vascular density.²⁶ Thus, the patterns of choroidal thinning associated with aging may be attributed to the greater impact of vascular density decreasing in the temporal regions.

Elongation of the axial length has a greater impact on the subfoveal and nasal areas, which are critical areas for ophthalmologists to evaluate to monitor the progression of myopia, particularly the development of myopic maculopathy. Deng et al.²⁷ reported that myopic individuals exhibit pronounced thinning of the choroid at the central foveal region, as well as in the temporal and inferior sectors of the peripapillary region, which can be explained by the uneven expansion of the posterior pole during the development of myopia. Fang et al.²⁸ demonstrated that, in eyes with myopia, choroidal thinning initially manifests as tessellation and subsequently progresses from the peripapillary to macular regions, resulting in diffused choroidal atrophy and the development of myopic maculopathy. Our choroidal changes map based on axial length elongation provides support for the observed choroidal thinning pattern in myopia eyes. Samples 1 and 3 in Figure 4 are vivid examples of the choroidal attenuation in the nasal region. The detailed distribution in the 900 grids provides us with a more intuitive way to monitor the progression of myopia, as compared to the sectors of the ETDRS and 3-mm areas toward the fovea.

Previous studies have found that the mean CT,²⁹ subfoveal CT,^{10,30} and CT at 1 or 3 mm from the fovea³¹ were significantly thicker in men compared to women. Our study initially found that in 900 subregions the CT values of males were consistently thicker than those of females, particularly in the subfoveal and inferior areas. Samples 1 and 4 in Figure 4 illustrate the difference in CT values between males and females. The variation in choroidal thickness between men and women may be explained by the elevated basal sympathetic tone in women,^{25,32} which in turn may lead to relatively narrower choroidal blood vessels. Also, the difference in hormone levels may be another factor, given that women experience a significant decrease in choroidal thickness during the mid-luteal phase of the menstrual cycle.^{25,29} In our study, the map of CT differences between males and females provides additional evidence of the physiological distinctions in different sexes, and this may explain the sex differences in the prevalence of some macular diseases. For example, central serous chorioretinopathy is more prevalent among males than females.³³

To our knowledge, no comparable studies have been conducted that would provide a comparison for our results of the distributions of CoV, CoD, and CoVU for the choroid. The CoD map (Fig. 3B) showed greater CT variation attributable to sex, age, and axial length in the temporal region. Additionally, the CoV and CoVU maps indicate a substantial decrease from the temporal regions to the nasal regions, signifying that the inter-individual choroidal variation in the tempo-



Figure 4. Examples of CT maps and B-scans of the fovea. (**A**, **B**) A 10-year-old male with an axial length of 23.08 mm. (**C**, **D**) A 73-year-old male with an axial length of 22.99 mm. (**E**, **F**) An 11-year-old male with an axial length of 26.13 mm. (**G**, **H**.) A 13-year-old female with an axial length of 22.84 mm.

ral area is less pronounced compared to the nasal region. This is the first attempt to describe the interindividual CT variances within precise areas of 0.2×0.2 mm. Several studies that have also utilized the 900grid algorithm to analyze CT distribution divided the macular region into five zones according to the CT in 900 grids but did not investigate the inter-individual variation.^{20,21} Therefore, the potential mechanisms of inter-individual choroidal variation remain uncertain and deserve further exploration in the future.

We recognize that the current study had some limitations. First, the retrospective design might lead to biased results due to not considering choroidal diurnal variation or water intake. Second, all samples were obtained from eyes without macular disorders, and the changing pattern of CT in diseased eyes requires further investigation. Finally, the analysis was limited to the macular region of the choroid, but other areas of the fundus should also be investigated in future studies.

In conclusion, our study innovatively regarded each small choroidal subregion $(0.2 \times 0.2 \text{ mm})$ as an independent area to develop a precise topographic map for better understanding of the inter-individual choroidal variations in healthy eyes. Such an approach could be helpful for identifying the choroidal physiology and distinguishing pathological alterations from physiological variations.

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