Combined wide-field optical coherence tomography angiography density map for high myopic glaucoma detection

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The present study aimed to evaluate the diagnostic ability of wide-field optical coherence tomography angiography (OCTA) density map for detection of glaucomatous damage in high myopic (HM) eyes and to further compare the diagnostic ability of OCTA with that of conventional imaging approaches including red-free photography and swept-source OCT (SS-OCT) wide-field maps. A total of 77 healthy HM eyes and 72 HM eyes with open angle glaucoma (OAG) participated in this retrospective observational study. Patients underwent a comprehensive ocular examination, including wide-field SS-OCT scan and peripapillary area and macular OCTA scans. An integrated OCTA density map thereafter was merged by vascular landmark-guided superimposition of peripapillary and macular superficial vascular density maps onto the red-free photography (resulting in the OCTA-PanoMap). Glaucoma specialists then determined the presence of glaucomatous damage in HM eyes by reading the OCTA-PanoMap and compared its sensitivity and specificity with those of conventional images. Sensitivity and specificity of OCTA-PanoMap for HM-OAG diagnosis was 94.4% and 96.1%, respectively. Compared with other imaging methods, the sensitivity of OCTA-PanoMap was significantly higher than that of red-free photography (P = 0.022) and comparable to that of wide-field SS-OCT maps. Specificity of OCTA-PanoMap was significantly higher than those of other conventional imaging methods (except for wide-field thickness map). The OCTA-PanoMap showed good diagnostic ability for discrimination of HM-OAG eyes from healthy HM eyes. As a complementary method of an alternative imaging modality, OCTA-PanoMap can be a useful tool for detection of HM-OAG.

Myopia has been considered as an independent risk factor for glaucoma, and moderately to highly myopic (HM) eyes have a higher risk for developing this condition than low myopic or hyperopic eyes do1-3. The prevalence of myopia has been increasing dramatically, especially among East Asians4,5. Therefore, diagnosis and monitoring glaucoma in myopic eyes are becoming very important; however, it is challenging to diagnose this condition, especially in HM eyes6. The reasons for the difficult diagnosis of glaucoma in HM include optic nerve head tilt, increased ovality/rotation/torsion, larger peripapillary atrophy7-9, and the temporal shift of Bruch's membrane opening10 and the retinal nerve fiber layer (RNFL) peaks11,12.

Various imaging modalities are used for evaluating the glaucomatous structural damages in HM eyes. Due to structural distortion of the optic disc in these eyes (including disc tilt or large peripapillary atrophy), evaluation of the optic disc through the stereoscopic photographs is very difficult. Further, the visibility of the RNFL is poor in HM eyes on red-free photography. Optical coherence tomography (OCT) is used as a relatively objective imaging method13; however, image analysis is not easy due to the absence of normative database for HM.

There have been many attempts to overcome these difficulties when using OCT. The macular parameters (ganglion cell complex)14, its asymmetry analysis15, and asymmetrical differences across the horizontal raphe16 have shown improved diagnostic power or been adjunctive diagnostic tools for glaucoma in HM eyes. Some studies suggested approaches to improve the diagnostic power by modifying the RNFL temporalization or magnification by using the self-constructing myopic normative database in peripapillary17,18 or macular area19. Recently,
Kim et al. reported that wide-field thickness and deviation map of swept-source OCT (SS-OCT) exhibited better accuracy and diagnostic power for detection of glaucomatous defects in myopic eyes than the conventional maps from spectral domain OCT did.\(^\text{10}\)

OCT angiography (OCTA) is a relatively new imaging modality that enables us to noninvasively assess the peripapillary and macular microvasculature. Studies have shown that OCTA is useful for detecting and quantifying glaucomatous damage and its progression. Peripapillary retinal vessel density is lower in glaucomatous eyes than in healthy eyes, and the decreased vessel density in OCTA images shows an exact topographical correlation with the RNFL defect.\(^\text{14}\) The location of focal perfusion loss in peripapillary retina showed good agreement with visual field defect.\(^\text{22}\) There are many studies on the association between peripapillary choroidal microvascular dropout (MVD), which is visible in the deeper layer of OCTA, and glaucoma.\(^\text{15–30}\) There are also studies on changes in the macular OCTA in glaucomatous eyes. Furthermore, many OCTA studies have been conducted on HM eyes, the relationship between MVD and healthy HM eyes and HM glaucoma, and the diagnostic abilities of several superficial vessel density parameters.\(^\text{30}\)

Recently, many studies have attempted to integrate the information of peripapillary and macular areas analysis together in the field of glaucoma.\(^\text{31–37}\) Nowadays, with the development of OCT for diagnosing and monitoring glaucoma, it is becoming more common to analyze the optic nerve and the macular area together (both in software and hardware) rather than each area separately. However, there were not many studies that have incorporated this concept into OCTA and furthermore, as yet, there have been no studies using wide-field OCTA Map itself.

The purpose of the present study was to evaluate the diagnostic ability of the wide-field OCTA density map for the detection of glaucomatous damage in HM eyes and further compare the diagnostic ability of OCTA with that of conventional imaging approaches including red-free photography and SS-OCT wide-field maps.

**Results**

**Clinical demographics.** Table 1 shows the clinical demographics in all patients at the time of enrollment. This retrospective cross-sectional study enrolled 149 eyes: 77 healthy HM eyes and 72 HM eyes with open-angle glaucoma (OAG). Four healthy HM eyes and five HM eyes with OAG were excluded due to poor image quality. The average age was 45.7 ± 14.4 years for HM-OAG eyes and 42.3 ± 15.8 years for healthy HM eyes (the controls) (p = 0.182). The differences in age, gender, IOP, spherical equivalent, AXL, and previous refractive or cataract surgery were not significant between the two groups. However, the MD, PSD, visual field index, peripapillary RNFL, macular ganglion cell—inner plexiform layer (GCIP), and macular ganglion cell complex (GCC) thickness differed between the healthy HM eyes and those with HM-OAG (p < 0.001).

| Age (years) | High myopic OAG (N=72) | Healthy high myopia (N=77) | P value |
|------------|------------------------|-----------------------------|---------|
| 45.7 ± 14.4 | 42.3 ± 15.8            | 0.182                       |
| Gender/female (%) | 24 (33.3) | 35 (45.5) | 0.136 |
| IOP (mmHg) | 15.8 ± 3.5             | 16.1 ± 4.0                  | 0.724   |
| SE (Diopter) | −8.79 ± 2.81           | −8.21 ± 2.47                | 0.273   |
| AX (mm) | 27.46 ± 1.31           | 27.03 ± 1.07                | 0.086   |
| Refractive surgery (%) | 13 (16.9) | 9 (12.5) | 0.495 |
| Cataract surgery (%) | 7 (9.7) | 11 (14.3) | 0.457 |
| RNFL thickness (μm) | 69.3 ± 18.5 | 89.6 ± 17.0 | <0.001 |
| GCC thickness (μm) | 86.2 ± 12.5 | 103.1 ± 8.7 | <0.001 |
| GCIP thickness (μm) | 54.9 ± 8.8 | 64.5 ± 5.4 | <0.001 |
| MD (dB) | −7.40 ± 6.35           | −2.47 ± 2.02                | <0.001 |
| PSD (dB) | 6.07 ± 4.02            | 2.32 ± 1.48                 | <0.001 |
| VFI (%) | 82.7 ± 19.5            | 96.8 ± 3.5                  | <0.001 |

Table 1. Clinical demographic characteristics of the participants. The data are expressed as means ± standard deviation or no. (%). Comparisons are performed using the chi-square test for categorical variables and the independent t-test for continuous variables. OAG open angle glaucoma, IOP intraocular pressure, SE spherical equivalence, AX axial length, RNFL retinal nerve fiber layer, GCC ganglion cell complex, GCIP ganglion cell inner plexiform layer, MD mean deviation of visual field, PSD pattern standard deviation of visual field, VFI visual field index.

Inter-rater agreement between two glaucoma specialists for detection of glaucomatous defect. Two independent glaucoma specialists (WJL and KIN) rated the presence of RNFL defects from red-free photography, wide-field thickness, and deviation map from SS-OCT and OCTA-PanoMap. The agreement obtained was moderate for red-free photography (Kappa = 0.596) and substantial for other wide-field maps (including OCTA-PanoMap) from SS-OCT (Kappa 0.741–0.771) between the two specialists.

Diagnostic ability for discriminating between HM healthy eyes and HM glaucomatous eyes. The sensitivity and specificity of the categorical variables including OCTA-PanoMap, red-free photography, wide-field RNFL thickness map, wide-field deviation map (according to manual rating and criteria),
and peripapillary RNFL, GCIPL, and GCC thicknesses using the abnormal criteria (<5% level or <1% level) provided by the built-in normative database for discriminating between HM healthy and HM-OAG eyes are presented in Table 2. Sensitivity and specificity of OCTA-PanoMap was 94.4% and 96.1%, respectively. Compared with other categorical variables, the sensitivity of OCTA-PanoMap was significantly higher than that of the red-free photography (p = 0.022). Other variables sensitivities showed no significant difference compared to that of OCTA-PanoMap. Among the categorical variables, the two highest specificity values were those for the OCTA-PanoMap (96.1%) and the wide-field RNFL thickness map (97.4%). Specificity of OCTA-PanoMap was significantly higher than those of other categorical variables (except for those of wide-field RNFL thickness map). Representative cases showing usefulness OCTA-PanoMap in detection of HM glaucomatous defect are provided in Figs. 1 and 2.

The AUCs in terms of quantitative measurement are presented in Fig. 3. AUC values were 0.800, 0.889 and 0.852 for peripapillary RNFL, GCC and GCIPL thicknesses, respectively. The diagnostic ability of categorical variables was also plotted on the graph as a point.

Table 2. Sensitivities and specificities for discrimination between high myopic open angle glaucoma and healthy high myopia. To qualify as “yellow” or “red” was at least 1 sector for the analysis being yellow or red. P value index compared with OCTA-PanoMap (McNemar’s test). Bold values indicate statistical significance with p-value less than 0.05. OCT-A optical coherence tomography—angiography, RNFL retinal nerve fiber layer, GCC ganglion cell complex, GCIPL ganglion cell inner plexiform layer.

|                        | Sensitivity | P-value | Specificity | P-value |
|------------------------|-------------|---------|-------------|---------|
| OCTA-PanoMap           | 94.4%       |         | 96.1%       |         |
| Red-free RNFL photography | 81.9%   | 0.022   | 85.7%       | 0.039   |
| Wide-field RNFL thickness map | 84.7%   | 0.065   | 97.4%       | 1.000   |
| Wide-field deviation map (manual) | 97.2% | 0.687   | 77.9%       | 0.001   |
| Wide-field deviation map (criteria) | 100.0% | NA      | 9.1%        | <0.001  |
| Peripapillary RNFL thickness |         |         |             |         |
| Quadrant < 5% (yellow) | 93.1%       | 1.000   | 23.4%       | <0.001  |
| Quadrant < 1% (red)    | 86.1%       | 0.146   | 48.1%       | <0.001  |
| Clock hours < 5% (yellow) | 98.6%  | 0.250   | 18.2%       | <0.001  |
| Clock hours < 1% (red) | 91.7%       | 0.727   | 40.3%       | <0.001  |
| Macular parameters |         |         |             |         |
| GCC < 5% (yellow)      | 95.8%       | 1.000   | 57.1%       | <0.001  |
| GCC < 1% (red)         | 91.7%       | 0.727   | 76.6%       | <0.001  |
| GCIPL < 5% (yellow)    | 98.6%       | 0.250   | 42.9%       | <0.001  |
| GCIPL < 1% (red)       | 90.3%       | 0.453   | 70.1%       | <0.001  |
Discussion
The present study compared diagnostic accuracy for the detection of glaucomatous defects in HM eyes between wide-field OCTA density map (OCTA-PanoMap) and conventional red-free photography or wide-field maps from SS-OCT (Topcon DRI OCT). Data demonstrated the clinical utility of OCTA-PanoMap for the diagnosis of HM-OAG.

OCTA has been widely studied and used in recent years in the diagnosis of glaucoma over the peripapillary and macular areas and imaging of the superficial and deep layers.

OCTA demonstrated reproducible focal loss of the radial peripapillary capillary (RPC) in patients with early OAG compared with that in normal controls. Using OCTA, reduced peripapillary retinal perfusion in glaucomatous eyes can be visualized as focal defects and quantified as peripapillary flow index and vessel density. Significant reductions in the RPC density were correlated with the site of RNFL decrease and visual field loss in glaucoma. There was also an attempt to build a normative database based on the change in the RPC density with age and to create a deviation map using this database.

This decrease in the superficial blood vessel density in glaucoma has also been reported in the macular area. Most studies have shown decreased perfusion in the superficial vascular complex of the macular area in patients with glaucoma using OCTA and that these decreases were correlated with visual field loss.

Recently, many studies have attempted to integrate the information of peripapillary and macular areas using OCT in the field of glaucoma. With the development of OCT, it is becoming more common to analyze these two areas together rather than individually. Wide-field OCT scan and map from SS-OCT and the integrated map (PanoMap) of SD-OCT are representative examples of areas currently being commercialized. These may help to well visualize continuous glaucomatous structural damage topographically and well facilitate the understanding.
of the spatial relationship between two areas in glaucoma. Research on applying this concept to OCTA is limited, and a few study with similar attempts are introduced below.

Wide-field montage OCTA can visualize the expansion of the RPC network. The RPC density was also correlated well with the RNFL thickness. The wide-field OCTA (8 × 8 mm) scan can visualize RPC information in a large area using software montage, but it can only cover the peripapillary area. These studies attempted to reveal wider application areas using OCTA, but all were confined to the peripapillary area and did not address glaucoma.

One study showed that significant microvascular damage was present in both the macular and peripapillary areas in early OAG. This study used vessel density parameters of OCTA in both the areas but evaluated only quantitative parameters, not those in HM glaucoma. OCTA measurements detect changes in retinal microvasculature (in the density of both the peripapillary and macular vessels) before visual field damage is detectable in patients with OAG. Another study also compared the vessel density in each of the two areas among the control, OAG, and angle closure glaucoma groups. Those studies evaluated the vessel density in both the areas, but all analyses were focused on parameters and not on topographic analysis with the map.

There have also been many OCTA studies related to glaucoma in myopia and HM. OCTA measurements can be useful for diagnosing glaucoma in HM eyes, especially when using calculated indices such as macular vessel density ratio (outer macular vessel density/inner macular vessel density). Some studies compared the vascular–function with structure–function correlations in HM glaucomatous eyes. Peripapillary vessel density, as measured using OCTA, may be useful in the evaluation of glaucomatous visual field damage in HM eyes. The performance of the vessel density was remarkable even when the RNFL measured using OCT was not reliable due to segmentation errors.

The present study has several differences from existing OCTA studies. First, a topographic map was used, not the parameters of OCTA. Second, two topographic maps of the optic nerve and macula were combined and analyzed. Finally, this method was applied to HM to compare the diagnostic ability of detecting glaucoma using conventional imaging methods.

Because this study was only performed on HM eyes, the inter-grader agreement between glaucoma specialists in this study was not higher than that reported in other studies performed with the general population. This is because it is difficult to diagnose glaucoma itself in HM, and in the process of determining any factor that could lead to its diagnosis, the method used in this study (OCTA-PanoMap) was devised.

The OCTA-PanoMap proposed in this study has two main strengths. First, the OCTA-PanoMap can visualize the structural glaucomatous damage more objectively in HM. The sensitivity of the OCTA-PanoMap was significantly superior to that of red-free photography, and the specificity of the OCTA-PanoMap was significantly superior to that of other OCT categorical parameters. As mentioned earlier, the presence of the tigroid fundus poses a significant challenge in the evaluation of glaucomatous damage in HM. Although OCT affords reproducible measurement of the RNFL thickness (swpt-source technology was used in this study), the detection of RNFL abnormalities in HM eyes is complicated because of the high rate of false-positive errors (especially in OCT deviation map), which is likely related to the lack of inclusion of individuals with HM in the normative
were confirmed using two consecutive reliable tests outside 99% of the age-specific normal limits or a PSD outside 95% of the normal limits. The visual field defects of one point with a probability of < 1%; or a cluster of two points with a probability of < 1%, and a GHT result within normal limits. Eyes with glaucomatous visual field defects were defined as those with a cluster of three points with probabilities of < 5% on the pattern deviation map in at least one hemifield, including at least one point with a probability of < 1%; or a cluster of two points with a probability of < 1%, and a GHT result outside 99% of the age-specific normal limits or a PSD outside 95% of the normal limits. The visual field defects were confirmed using two consecutive reliable tests.

In conclusion, the wide-field OCTA density map (OCTA-PanoMap) showed good diagnostic ability for the discrimination of HM glaucoma from healthy HM eyes. As a complementary method of an alternative imaging modality, the OCTA-PanoMap of SS-OCT can serve as a useful tool for the detection of HM glaucoma.

Methods
The study protocol was approved by the Institutional Review Board (IRB) of Hanyang University Hospital. Informed consent was waived due to its retrospective nature and it was confirmed by IRB of our institution. The study design adhered to the tenets of the Declaration of Helsinki for biomedical research.

Participants. This retrospective cross-sectional study enrolled patients with healthy HM eyes and HM eyes with OAG who visited the Glaucoma Clinic of Hanyang University Hospital between August 2018 and December 2020.

All participants underwent complete ophthalmologic examination, including visual acuity testing, manifest refraction assessment, slit-lamp examination, intraocular pressure (IOP) measurements using Goldmann applanation tonometry, gonioscopy, dilated fundus examination, axial length measurement (IOLMaster; Carl Zeiss Meditec, Dublin, CA, USA), stereo disc photography, red-free photography (EIDON confocal scanner; CenterVue, Padua, Italy), 24–2 perimetry using the Swedish interactive thresholding algorithm (Humphrey Field Analyzer II; Carl Zeiss Meditec, Jena, Germany), and SS-OCT (DRI-OCT Triton; Topcon, Tokyo, Japan).

A visual field was considered to be reliable if the fixation losses were < 20%, the false positive rate was < 15%, and the false-negative rate was < 15%. A normal visual field was defined as that with a mean deviation (MD) and pattern standard deviation (PSD) within 95% confidence limits, and results of the glaucoma hemifield test within normal limits. Eyes with glaucomatous visual field defects were defined as those with a cluster of three points with probabilities of < 5% on the pattern deviation map in at least one hemifield, including at least one point with a probability of < 1%; or a cluster of two points with a probability of < 1%, and a GHT result outside 99% of the age-specific normal limits or a PSD outside 95% of the normal limits. The visual field defects were confirmed using two consecutive reliable tests.

The inclusion criteria were as follows: best-corrected visual acuity of 20/40 or higher, HM, and an open anterior chamber angle. HM was defined as refraction less than −6.0D or Axial length (AXL) > 26 mm. The exclusion criteria were as follows: history of ophthalmic surgery (e.g., glaucoma filtration surgery), severe glaucoma with an MD worse than −20 dB, any other ocular disease that could interfere with the visual function, any media opacity that could significantly interfere with OCT acquisition, and an inability to obtain a high-quality OCT image. All of the OCT scan images had to have an image quality score ≥ 30, according to the manufacturer's recommendation. Eyes with poor OCTA images were excluded on the basis of the following criteria: image quality < 30, poor-clarity images, localized weak signal caused by artifacts such as floaters, residual motion artifacts visible as irregular vessel patterns or disc boundary on the en face angiogram or segmentation failure. One eye was randomly chosen for the study in cases where both the eyes met all the eligibility criteria.

We identified patients with OAG using several features in addition to an open angle confirmed using gonioscopy. The first feature was the presence of a characteristic optic disc with localized or diffuse neuroretinal rim thinning, increased cupping, or a cup-to-disc ratio difference > 0.2 between the eyes observed in the stereo disc photograph. The presence of an RNFL defect on red-free fundus imaging was an alternative feature. The presence of glaucomatous visual field defects was a prerequisite for glaucoma diagnosis, and patients with preperimetric glaucoma (normal visual field) were excluded from this study. A patient diagnosed with OAG was treated with topical medication or not according to the decision of the glaucoma specialist. Healthy eyes were defined as those without any history or evidence of intraocular surgery, intraocular pressure of ≤ 21 mmHg with no history of increased intraocular pressure, absence of glaucomatous appearance of the optic disc, and normal ophthalmologic findings. The wide-field OCT scan analysis was performed according to the right-eye orientation. A senior glaucoma specialist (MS) determined whether the participants were HM-OAG or HM healthy controls. All previous medical records, including images and visual field results, were reviewed to examine disease progression and confirm the diagnosis. Detailed objective method used to determine progression is provided in the Supplementary Note.

Swept-source optical coherence tomography. We employed a wide-field scan protocol (12 × 9 mm²) using the DRI-OCT device. The DRI-OCT is an SS-OCT device that uses a wavelength-sweeping laser with a central wavelength of 1050 nm and tuning range of approximately 100 nm. We acquired 100,000 A-scans per second with an 8-µm axial tissue resolution. A 12 × 9 mm² scan contained 256 B-scans, each containing 512...
A-scans for a total of 131,072 axial scans/volume. The wide-field RNFL thickness map and wide-field deviation map were obtained and the method has been described in detail in a previous study62.

**OCTA imaging.** We imaged the optic disc and macular OCTA using a commercially available SS-OCTA system (DRI OCT Triton; Topcon), separately. Scans were obtained from 4.5 x 4.5 mm² cubes, centered on the optic disc and macula, respectively. We used the automated layer segmentation of the OCT instrument software to generate en face images of the retinal vasculature from the selected en face slabs. The density map of the selected en face slabs were presented as a color-coded map.

**Construction of the wide-field OCTA density map.** With the built-in analysis software, superficial retinal layer [from the internal limiting membrane (ILM) to the inner border of the inner nuclear layer] of the macular and the density map of radial peripapillary capillary layer (from top to 70.2 µm below ILM) were used for the present study.

We used a combined wide-field OCTA density map for single-image evaluation of glaucoma diagnosis. The map was created by the same-day superimposition of optic disc and macular OCTA density maps onto RNFL photography as aligned using the Photoshop software (Version 11.0; Adobe, San Jose, CA, USA) based on vascular landmarks. Each combined map was labeled by date, and all, as collectively integrated, were defined as the OCTA-PanoMap.

**Definition of the RNFL defect.** RNFL defect on red-free photography, wide-field RNFL thickness map, and wide-field deviation map has been described in detail in a previous study32.

RNFL defect on red-free photography was defined as having a width at a 1-disc-diameter distance from the edge of the disc larger than that of a major retinal vessel, diverging in an arcuate or wedge shape60,61. RNFL defect on the wide-field RNFL thickness map was defined as an arcuate or wedge-shaped diverging dark-blue area surrounding an abrupt color scale change that appeared less thick than neighboring areas on color-coded maps of wide RNFL thickness. The minimum defect size was larger than the diameter of a major retinal vessel33,32,62,63. RNFL defect on the SuperPixel map was arbitrarily defined as the presence of a wedge-shaped area of at least 20 contiguous yellow/red pixels along with RNFL thinning on the deviation map31,32,60,64. Since there were many yellow/red pixels on the deviation map of HM patients, the judgment was made by dividing the ones judged by the glaucoma specialist (manually) and those judged by criteria (20 contiguous pixels). RNFL defect on the OCTA-PanoMap was defined similarly to that on wide-field RNFL thickness map, as an arcuate or wedge-shaped diverging dark area surrounding an abrupt color scale change that appeared less crowded than neighboring areas did on color-coded density maps.

RNFL defects on above-mentioned images were evaluated independently by two glaucoma specialists (WJL, KIN) in a masked fashion. For prevention of bias, the graders who evaluated the wide-field maps were masked to the red-free photography. When the two experts disagreed, a third ophthalmologist (MS) evaluated the photographs and the disagreement was resolved after discussion.

**Statistical analyses.** Statistical tests were performed using IBM SPSS Statistics 24 (IBM Corp, Armonk, NY) and MedCalc (MedCalc Software, Ostend, Belgium). The independent t-test was used to compare the characteristics of the continuous variables and the chi-squared test was used for the categorical variables. Inter-rater agreement between two glaucoma specialists (WJL, KIN) was evaluated using the Kappa test. Cohen suggested the Kappa result be interpreted as follows: values ≤ 0 as indicating no agreement, 0.01–0.20 as none to slight, 0.21–0.40 as fair, 0.41–0.60 as moderate, 0.61–0.80 as substantial, and 0.81–1.00 as almost perfect agreement65. To assess the abilities of each categorical variable, such as wide-field RNFL, thickness map, or the criteria on RNFL defect on above-mentioned images were calculated for the continuous variables and the comparison of AUC values of the parameters was assessed using the method described by DeLong et al.66. P values < 0.05 were considered statistically significant. The values are presented as the mean ± standard deviation.

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**References**

1. Mitchell, P., Hourihan, F., Sandbach, J. & Wang, J. J. The relationship between glaucoma and myopia—The blue mountains eye study. *Ophthalmology* **106**(10), 2010–2015 (1999).
2. Xu, L. et al. High myopia and glaucoma susceptibility the Beijing Eye Study. *Ophthalmology* **114**(2), 216–220 (2007).
3. Marcus, M. W., de Vries, M. M., Montolio, F. G. J. & Jansonius, N. M. Myopia as a risk factor for open-angle glaucoma: A systematic review and meta-analysis. *Ophthalmology* **118**(10), 1989–U146 (2011).
4. Morgan, I. G. et al. The epidemics of myopia: Aetiology and prevention. *Prog. Retin. Eye Res.* **62**, 134–49 (2018).
5. Holden, B. A. et al. Global prevalence of myopia and high myopia and temporal trends from 2000 through 2050. *Ophthalmology* **123**(5), 1036–42 (2016).
6. Tan, N. Y. Q. et al. Glaucoma in myopia: Diagnostic dilemmas. *Br. J. Ophthalmol.* **103**(10), 1347–55 (2019).
7. Kim, T. W. et al. Optic disc change with incipient myopia of childhood. *Ophthalmology* **119**(1), 21-6.e1–3 (2012).
8. Samarawickrama, C. et al. Myopia-related optic disc and retinal changes in adolescent children from Singapore. *Ophthalmology* 118(10), 2050–7 (2011).
9. Wang, Y. X., Panda-Jonas, S. & Jonas, J. B. Optic nerve head anatomy in myopia and glaucoma, including parapapillary zones alpha, beta, gamma and delta: Histology and clinical features. *Prog. Retin. Eye Res.* 83, 100933 (2021).
10. Zhang, Q. et al. Size and shape of Bruch’s membrane opening in relationship to axial length, gamma zone, and macular Bruch’s membrane defects. *Invest. Ophthalmol. Vis. Sci.* 60(7), 2591–8 (2019).
11. Chang, J. K. & Yoo, Y. C. Correct calculation circle location of optical coherence tomography in measuring retinal nerve fiber layer thickness in eyes with myopic tilted discs. *Invest. Ophthalmol. Vis. Sci.* 52(11), 7894–900 (2011).
12. Leung, C. K. et al. Retinal nerve fiber layer imaging with spectral-domain optical coherence tomography: Interpreting the RNFL maps in healthy myopic eyes. *Invest. Ophthalmol. Vis. Sci.* 53(11), 7194–200 (2012).
13. Hwang, Y. H., Yoo, C. & Kim, Y. Y. Myopic optic disc tilt and the characteristics of peripapillary retinal nerve fiber layer thickness measured by spectral-domain optical coherence tomography. *J. Glaucoma* 21, 260–5 (2012).
14. Shoji, T. et al. Assessment of glaucomatous changes in subjects with high myopia using spectral domain optical coherence tomography. *Invest. Ophthalmol. Vis. Sci.* 52(2), 1098–102 (2011).
15. Lin, P. W., Chang, H. W. & Poon, Y. C. Retinal thickness asymmetry in highly myopic eyes with early stage of normal-tension glaucoma. *J. Ophthalmol.* 2021, 6660631 (2021).
16. Kim, Y. K. et al. Glaucoma-diagnostic ability of ganglion cell-inner plexiform layer thickness difference across temporal raphe in highly myopic eyes. *Invest. Ophthalmol. Vis. Sci.* 57(14), 5856–63 (2016).
17. Bswas, S., Lin, C. & Leung, C. K. Evaluation of a myopic normative database for detection of retinal nerve fiber layer thickness. *JAMA Ophthalmol.* 134(9), 1032–9 (2016).
18. Seol, B. R., Kim, D. M., Park, K. H. & Jeoung, J. W. Assessment of optical coherence tomography color probability codes in myopic glaucoma eyes after applying a myopic normative database. *Am. J. Ophthalmol.* 183, 147–53 (2017).
19. Nakashiri, H. et al. Sensitivity and specificity for detecting early glaucoma in eyes with high myopia from normative database of macular ganglion cell complex thickness obtained from normal non-myopic or highly myopic Asian eyes. *Graefes Arch. Clin. Exp. Ophthalmol.* 253(7), 1143–52 (2015).
20. Kim, Y. W., Lee, J., Kim, J. S., & Park, K. H. Diagnostic accuracy of wide-field map from swept-source optical coherence tomography for primary open-angle glaucoma in myopic eyes. *Am. J. Ophthalmol.* (2020).
21. Lee, E. I., Kim, M. H., Lee, S. H. & Kim, T. W. OCT-angiography of the peripapillary retina in primary open-angle glaucoma. *Invest. Ophthalmol. Vis. Sci.* 57(14), 6265–70 (2016).
22. Chen, A. et al. Measuring glaucomatous focal perfusion loss in the peripapillary retina using OCT angiography. *Ophthalmology* 127(4), 484–91 (2020).
23. Kim, J. A., Lee, E. J. & Kim, T. W. Evaluation of parapapillary choroidal microvasculature dropout and progressive retinal nerve fiber layer thinning in patients with glaucoma. *JAMA Ophthalmol.* 137(7), 810–6 (2019).
24. Lee, E. I., Kim, T. W., Kim, J. A. & Kim, J. A. Central visual field damage and parapapillary choroidal microvasculature dropout in primary open-angle glaucoma. *Ophthalmology* 125(4), 588–96 (2018).
25. Lee, E. J., Lee, K. M., Lee, S. H. & Kim, T. W. Parapapillary choroidal microvasculature dropout in glaucoma: A comparison between optical coherence tomography angiography and indocyanine green angiography. *Ophthalmology* 124(8), 1209–17 (2017).
26. Park, H. L., Kim, J. W. & Park, C. K. Choroidal microvasculature dropout is associated with progressive retinal nerve fiber layer thinning in glaucoma with disc hemorrhage. *Ophthalmology* 125(7), 1003–13 (2018).
27. Suh, M. H. et al. Deep retinal layer microvasculature dropout detected by the optical coherence tomography angiography in glaucoma. *Ophthalmology* 123(12), 2509–18 (2016).
28. Kim, G. N., Lee, E. J. & Kim, T. W. Microstructure of nonjuxtapapillary microvasculature dropout in healthy myopic eyes. *Invest. Ophthalmol. Vis. Sci.* 61(2), 36 (2020).
29. Na, H. M., Lee, E. I., Lee, J. E. & Kim, S. H. & Kim, T. W. Evaluation of parapapillary choroidal microvasculature to detect glaucomatous damage in eyes with high myopia. *J. Glaucoma* 29(1), 39–45 (2020).
30. Lee, K. et al. Diagnostic ability of vessel density measured by spectral-domain optical coherence tomography angiography for glaucoma in patients with high myopia. *Sci. Rep.* 10(1), 3027 (2020).
31. Lee, W. J. et al. Comparison of glaucoma-diagnostic ability between wide-field swept-source OCT retinal nerve fiber layer maps and spectral-domain OCT. *Eye (Lond)* 32(9), 1483–92 (2018).
32. Lee, W. J. et al. Diagnostic ability of wide-field retinal nerve fiber layer maps using swept-source optical coherence tomography for detection of preperimetric and early perimetric glaucoma. *J. Glaucoma* 26(6), 577–85 (2017).
33. Hood, D. C. Improving our understanding, and detection, of glaucomatous damage: An approach based upon optical coherence tomography (OCT). *Prog. Retin. Eye Res.* 57, 46–75 (2017).
34. Hood, D. C. et al. Glaucomatous damage of the macula. *Prog. Retin. Eye Res.* 32, 1–21 (2013).
35. Lee, W. J. et al. Serial combined wide-field optical coherence tomography maps for detection of early glaucomatous structural progression. *JAMA Ophthalmol.* 136(10), 1121–7 (2018).
36. Lee, W. J. et al. Combined use of retinal nerve fiber layer and ganglion cell-inner plexiform layer event-based progression analysis. *Am. J. Ophthalmol.* 196, 65–71 (2018).
37. Lee, W. J., Park, K. H. & Seong, M. Vulnerability zone of glaucoma progression in combined wide-field optical coherence tomography event-based progression analysis. *Invest. Ophthalmol. Vis. Sci.* 61(5), 56 (2020).
38. Mansoori, T., Sivaswamy, J., Gamalapati, J. S. & Balakrishna, N. Radial peripapillary capillary density measurement using optical coherence tomography angiography in early glaucoma. *J. Glaucoma* 26(5), 438–43 (2017).
39. Mansoori, T. et al. Measurement of radial peripapillary capillary density in the normal human retina using optical coherence tomography angiography: *J. Glaucoma* 26(3), 241–6 (2017).
40. Liu, L. et al. Optical coherence tomography angiography of the peripapillary retina in glaucoma. *JAMA Ophthalmol.* 133(9), 1045–52 (2015).
41. Mammoto, Z. et al. Quantitative optical coherence tomography angiography of radial peripapillary capillaries in glaucoma, glaucoma suspect, and normal eyes. *Am. J. Ophthalmol.* 170, 41–9 (2016).
42. Pinhas, A. et al. A method for age-matched OCT angiography deviation mapping in the assessment of disease- related changes to the radial peripapillary capillaries. *PloS ONE* 13(5), e0197062 (2018).
43. Takusagawa, H. L. et al. Projection-resolved optical coherence tomography angiography of macular retinal circulation in glaucoma. *Ophthalmology* 124(11), 1589–99 (2017).
44. Onishi, A. et al. Parafoveal vessel changes in primary open-angle glaucoma and normal-tension glaucoma using optical coherence tomography angiography. *Clin. Ophthalmol.* 13, 1935–45 (2019).
45. Hou, H. et al. Macula vessel density and thickness in early primary open-angle glaucoma. *Am. J. Ophthalmol.* 199, 120–32 (2019).
46. Wang, K. H., Lam, A. K. N. & Leung, C. K. Optical coherence tomography angiography compared with optical coherence tomography macular measurements for detection of glaucoma. *JAMA Ophthalmol.* 136(6), 866–74 (2018).
47. Wu Dunn, D. et al. OCT angiography for the diagnosis of glaucoma: A report by the American Academy of Ophthalmology. *Ophthalmology* 128(8), 1222–35 (2021).
48. Lee, W. J. et al. Clinical use of PanoMap for glaucoma: Frequently damaged areas in early glaucoma. *J. Glaucoma* 30(1), 10–6 (2021).
52. Lu, P. et al. Quantitative analysis of microvasculature in macular and peripapillary regions in early primary open-angle glaucoma. Curr. Eye Res. 45(5), 629–35 (2020).
53. Yarmohammadi, A.
54. Hou, T. Y. et al. Comparison of vascular-function and structure-function correlations in glaucomatous eyes with visual field loss. Ophthalmology 125(4), 578–87 (2018).
55. Lee, S. H., Lee, E. J. & Kim, T. W. Increase in the OCT angiographic peripapillary vessel density by ROCK inhibitor ripasudil instillation: A comparison with brimonidine. Graefes Arch. Clin. Exp. Ophthalmol. 256(7), 1257–64 (2018).
56. Shin, J. W. et al. Optical coherence tomography angiography vessel density changes in normal-tension glaucoma treated with carteolol, brimonidine, or dorzolamide. J. Glaucoma 30(6), 690–6 (2021).
57. Chihara, E., Dimitrova, G. & Chihara, T. Increase in the OCT angiographic peripapillary vessel density by ROCK inhibitor ripasudil instillation: A comparison with brimonidine. Graefes Arch. Clin. Exp. Ophthalmol. 256(7), 1257–64 (2018).
58. Lin, Y. H. et al. Optical coherence tomography angiography vessel density changes in normal-tension glaucoma treated with carteolol, brimonidine, or dorzolamide. J. Glaucoma 30(6), 690–6 (2021).
59. Anderson, D. & Patella, V. Automated Static Perimetry 2nd edn. (Mosby, 1999).
60. Kim, M. J. et al. Comparison of macular GCIPL and peripapillary RNFL deviation maps for detection of glaucomatous eye with localized RNFL defect. Acta Ophthalmol. 93(1), c22-c8 (2015).
61. Kim, T. J., Kim, Y. K. & Kim, D. M. Patterns of subsequent progression of localized retinal nerve fiber layer defects on red-free fundus photographs in normal-tension glaucoma. Korean J. Ophthalmol. 28(4), 330–6 (2014).
62. Sakamoto, A. et al. Three-dimensional imaging of the macular retinal nerve fiber layer in glaucoma with spectral-domain optical coherence tomography. Invest. Ophthalmol. Vis. Sci. 51(10), 5062–70 (2010).
63. Shin, J. W., Uhm, K. B., Lee, W. J. & Kim, Y. J. Diagnostic ability of retinal nerve fiber layer maps to detect localized retinal nerve fiber layer defects. Eye (Lond) 27(9), 1022–31 (2013).
64. Hwang, Y. H., Jeong, Y. C., Kim, H. K. & Sohn, Y. H. Macular ganglion cell analysis for early detection of glaucoma. Ophthalmology 121(8), 1508–15 (2014).
65. McHugh, M. L. Interrater reliability: The kappa statistic. Biochem. Med. (Zagreb) 22(2), 376–82 (2012).
66. Delong, E. R., DeLong, D. M. & Clarke-Pearson, D. L. Comparing the areas under two or more correlated receiver operating characteristic curves: A nonparametric approach. Biometrics 44(3), 837–48 (1988).

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The authors declare no competing interests.

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