

Retinal Sensitivity and Ganglion Cell-Related Retinal Layer Thickness in the Normal Aging Process

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Research Article

Keywords: visual field (VF), circumpapillary retinal nerve fiber layer thickness (cpRNFLT), spectral-domain optical coherence tomography (SD-OCT)

Posted Date: January 21st, 2021

DOI: <https://doi.org/10.21203/rs.3.rs-149830/v1>

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1 **Retinal sensitivity and ganglion cell-related retinal layer thickness in the normal aging**
2 **process**

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22 **Abstract**

23 Aging-associated changes in visual field (VF) sensitivity were compared prospectively
24 and longitudinally with the circumpapillary retinal nerve fiber layer thickness (cpRNFLT)
25 and macular ganglion cell-inner plexiform layer thickness (GCIPLT) changes in the
26 corresponding retinal areas of the same eyes (72 eyes of 37 normal Japanese subjects;
27 mean age, 51.3 years). The Humphrey Field Analyzer 24-2 test (HFA 24-2) and
28 spectral-domain optical coherence tomography (SD-OCT) measurements of the
29 cpRNFLT and GCIPLT in a 0.6-mm-diameter circle corresponding to the four central
30 points of HFA 24-2 adjusted for retinal ganglion cell displacement (GCIPLT_{4TestPoints}) were
31 performed every 3 months for 3 years. The time changes of the mean sensitivity over
32 the entire field (VF_{mean}) and the four central points (VF_{4TestPoints}), cpRNFLT, and
33 GCIPLT_{4TestPoints} were analyzed using a linear mixed model. The aging-associated decline
34 rates of VF_{mean} and VF_{4TestPoints} were 0.12 and 0.19 decibels/year ($p < 0.001$), which
35 significantly accelerated with increased subjects' age (0.009 and 0.010 decibels/year,
36 $p < 0.001$, respectively) without changes in the ocular media. Those of the CpRNFLT and
37 GCIPLT_{4TestPoints} were not significant in both ($p > 0.114$), but significantly accelerated with
38 increased subjects' age (0.021 and 0.010 $\mu\text{m}/\text{year}$, $p = 0.001$ and 0.004, respectively).
39 These results have implications in studying physiological aging- or disease-related
40 changes in these parameters.

41

42 Histologic studies in human eyes have estimated that about 7,000 retinal ganglion cells
43 (RGCs) are lost during normal aging.[1-5] In accordance with these histologic findings,
44 previous cross-sectional or longitudinal studies in normal human eyes have shown that
45 visual field (VF) sensitivity measured using standard automated perimetry (SAP) [6-9]
46 and the thickness of the RGC-related retinal layers determined by spectral-domain
47 optical coherence tomography (SD-OCT) showed aging-associated declines.[10-19]
48 Previous studies of glaucomatous eyes with early or pre-perimetric damage have
49 reported that changes in the circumpapillary retinal nerve fiber layer thickness
50 (cpRNFLT) measured by SD-OCT generally preceded glaucomatous changes in the VF
51 sensitivity detected using SAP standard test programs such as the Humphrey Field
52 Analyze 24-2 Swedish Interactive Thresholding Algorithm (HFA-24, Carl Zeiss Meditec,
53 Dublin, CA).[20-23] Both glaucoma and the normal aging process reduce the thickness
54 of the RGC-related retinal layers measured using SD-OCT and VF sensitivity measured
55 by SAP.[6-19] However, the comparisons of the aging-associated declines measured by
56 the two tests have not been reported in the same normal subjects' eyes, while
57 comparisons of declines in the two tests in the same glaucomatous eyes have been
58 studied.[20-23] In the current study, we performed prospective measurements of the
59 longitudinal time changes in the cpRNFLT and macular ganglion cell-inner plexiform
60 layer thickness (GCIPLT) and the SAP sensitivity in the corresponding retinal areas in
61 the same normal eyes.

62

63 Results

64 A total of 76 eyes of 38 normal subjects were enrolled; two eyes of one subject and
65 one eye of one subject were excluded because of development of small retinal
66 hemorrhages of unknown origin and epiretinal membrane or vitreoretinal traction,
67 respectively, during follow-up, and one eye of one subject was excluded because
68 reliable SD-OCT measurements could not always be obtained mainly due to saccades
69 and blinking. The demographics of the remaining 72 eyes of 37 subjects are shown in
70 Table 1. Inter-visit test-retest variability calculated as the minimal detectable change
71 (MDC) ($MDC = 1.96 \times \sqrt{2} \times \text{standard error of measurements}$) [24] was 3.3 μm for the
72 cpRNFLT and 1.6 μm for the mean of the GCIPLT in a circular retinal area with a
73 diameter of 0.6 mm (approximately 2 degrees of visual angle) corresponding to the
74 four central test points of the HFA 24-2, adjusted for RGC displacement [25]
75 ($GCIPLT_{4\text{TestPoints}}$), respectively.

76 During the 3-year prospective follow-up, ocular transparent media including lens
77 showed no changes on biomicroscopic examination. The mean sensitivity over the
78 entire field (VF_{mean}) and that of the four central test points of the HFA 24-2 ($VF_{4\text{TestPoints}}$)
79 of an eye with average parametric values of this cohort (Table 1) showed significantly
80 negative longitudinal time changes (age-associated declines) of -0.12 decibels
81 [dB]/year ($p < 0.001$) and -0.19 dB/year ($p < 0.001$), respectively, which were significantly
82 more negative by -0.009 and -0.010 dB/year with older baseline ages ($P < 0.001$), and by
83 -0.11 and -0.14 dB/dB with higher baseline sensitivities ($P < 0.001$), respectively. Further,

84 the VF_{mean} and $VF_{4\text{TestPoints}}$ measurement results themselves were also significantly
85 lower by 0.021 and 0.028 dB/year with older baseline age ($p < 0.001$), and higher by
86 0.60 and 0.43 dB/dB with higher baseline sensitivity, respectively (Tables 2 and 3).

87 On the other hand cpRNFLT and the mean of the GCIPLT in a circular retinal area
88 with a diameter of 0.6 mm (approximately 2 degrees of visual angle) corresponding to
89 the four central test points of the HFA 24-2, adjusted for RGC displacement [25],
90 $GCIPLT_{4\text{TestPoints}}$, of an eye with average parametric values of this cohort (Table 1)
91 showed no significant longitudinal time changes (aging-associated declines) ($p = 0.352$
92 and 0.114, respectively); however, the aging-related declines of cpRNFLT and
93 $GCIPLT_{4\text{TestPoints}}$ were significantly more greater by 0.021 and 0.010 $\mu\text{m}/\text{year}$ with older
94 baseline age ($p = 0.001$ and 0.004, respectively). The cpRNFLT and $GCIPLT_{4\text{TestPoints}}$
95 measurement results themselves were also significantly thinner by 0.025 μm with
96 older baseline age ($p < 0.001$) in both, by 0.15 and 0.09 μm with longer axial length (AL)
97 ($p = 0.030$ and 0.019, respectively), and by 0.62 and 0.19 $\mu\text{m}/\text{mm}^2$ with greater disc area
98 ($p < 0.001$ and $P = 0.048$, respectively) (Tables 4 and 5).

99

100 In the current study, the longitudinal aging-associated declines in the VF sensitivity and
101 SD-OCT-measured thicknesses of the RGC-related retinal layers in the corresponding
102 area were measured in the same eyes of normal Japanese subjects (mean age, 51
103 years). The longitudinal aging-related decline rates of the VF_{mean} , the mean sensitivity
104 of the all test points of HFA 24-2, and $VF_{4\text{TestPoints}}$, the mean sensitivity of the four

105 central test points of HFA 24-2, were -0.12 and -0.19 dB/year, respectively, which were
106 significantly more negative along with older baseline age and higher VF sensitivity. This
107 aging-associated declines in the VF sensitivity currently estimated from longitudinal VF
108 measurement results from normal subjects were comparable to those estimated by
109 previous cross-sectional studies (-0.1 ~-0.2 dB/year). [6-9] The current study further
110 confirmed significant acceleration of the aging-associated decline rates in the VF
111 sensitivity previously observed only in cross-sectional datasets [7-9] by longitudinal
112 observation. Since no significant changes were seen in the ocular media including lens
113 in the current subjects during the prospective follow-up period, acceleration of
114 aging-associated declines in the VF sensitivity currently observed would be mainly
115 attributed to post-retinal neuronal factors rather than pre-retinal ones.

116 The aging-associated changes in the thickness of the RGC-related retinal layers in
117 the retinal area corresponding to the VF_{mean} and $VF_{4\text{TestPoints}}$ measurement sites
118 adjusted for RGC displacement [25], i.e., the $cpRNFLT$ and $GCIPLT_{4\text{TestPoints}}$, respectively,
119 showed the same trends as each other; the longitudinal aging-associated decline rates
120 during follow-up (coefficients for duration) were not significant, but they were
121 significantly more negative by $-0.01 \sim 0.02 \mu\text{m}/\text{year}$ along with older subjects' age
122 (coefficients for interaction between the baseline age and duration were significantly
123 negative), being compatible with increases aging-associated decline in the VF
124 sensitivity along with older subjects' age observed in the same eyes. The baseline age,
125 AL, and disc area had significantly negative effects on the $cpRNFLT$ and $GCIPLT_{4\text{TestPoints}}$

126 measurement results themselves, which agreed with previous cross-sectional SD-OCT
127 study results for the cpRNFLT and age [10,13,14], AL [14], and disc area [10] and for
128 GCIPLT and age [11,12,15] and AL,[11,15] suggesting little differences in the SD-OCT
129 characteristics between the current normal subjects and those in previous studies.

130 Leung et al. followed 70 eyes of 35 normal subjects every 4 months for 30 months
131 using Cirrus HD-OCT (Carl Zeiss Meditec), and reported a significant longitudinal
132 aging-associated decline in the cpRNFLT of $-0.52 \mu\text{m}/\text{year}$, [16]; they also followed 72
133 eyes of 40 normal subjects every 4 months for a mean of 40 months and reported a
134 significant aging-associated decline of the macular GCIPLT of $-0.32 \mu\text{m}/\text{year}$. [17]

135 Vianna et al. followed 37 eyes of 37 normal subject every 6 months for a median of 4.5
136 years using Spectralis OCT (Heidelberg Engineering, Heidelberg, Germany) and
137 reported significant longitudinal aging-associated decline in the cpRNFLT

138 $-0.44 \mu\text{m}/\text{year}$. [18] Hammel et al. measured the aging-related decreases in the
139 cpRNFLT and macular GCIPLT in 28 normal subjects using Cirrus HD-OCT for a median
140 of 1.7 years with 6 visits and also reported a significant longitudinal aging-associated
141 decline in the cpRNFLT of $-0.48 \mu\text{m}/\text{year}$ but not in the macular GCIPLT. [19] In the

142 current study, we used the 3D-OCT 2000 (Topcon) and measured the cpRNFLT and
143 GCIPLT corresponding to the central four test points of the HFA 24-2 (GCIPLT_{4TestPoints})
144 every 3 months for 3 years in 72 eyes of 37 normal Japanese subjects and could not
145 detect significant longitudinal aging-associated changes in these parameters. This
146 finding was rather unexpected, since both the cpRNFLT and GCIPLT_{4TestPoints}

147 measurement results themselves were significantly and negatively correlated with the
148 subjects' ages in the current cohort (coefficients for age at baseline were significantly
149 negative). To re-confirm that time changes in the cpRNFLT and GCIPLT_{4TestPoints} during
150 the follow-up period were not significant, we conducted simple linear regression
151 analyses using only duration (time lapse after entry) as an explanatory variable, i.e.,
152 time changes in the cpRNFLT or GCIPLT_{4TestPoints} = A × duration + B. The coefficient for
153 duration, A, was calculated to be 0.0042 μm/year ($p=0.470$) and $p=0.0024$ μm/year
154 ($p=0.538$) for the cpRNFLT and GCIPLT_{4TestPoints}, respectively.

155 Larger test-retest variability, fewer measurements, and a shorter follow-up period
156 should reduce the power to detect a significant trend. The inter-session test-retest
157 variability values of the measurement results with the current SD-OCT instrument were
158 3.3 and 1.6 μm, respectively, for the minimum detectable change in the cpRNFLT and
159 GCIPLT_{4TestPoints}, which were favorably compared with those reported in the literature
160 for measurement results with other SD-OCT instruments.[24,30] The follow-up period
161 and interval in the current study were similar to (3 years vs. 1.7 ~4.5 years,
162 respectively) and shorter than (3 months vs. 3.5 ~ 6 months, respectively) those
163 adopted by previous studies.[16-19] Thus, it seems rather unlikely that a shorter
164 follow-up period and fewer measurements mainly accounted for an undetected
165 significant trend in longitudinal decline in the cpRNFLT and GCIPLT_{4TestPoints} in the
166 current subjects.

167 The current study indicated that in older subjects, the longitudinal aging-associated

168 declines in the cpRNFLT and GCIPLT_{4TestPoints} were significantly more negative
169 (coefficients for interaction between the baseline age and duration were significantly
170 negative) by -0.02 and -0.01 $\mu\text{m}/\text{year}$, respectively, suggesting that in subjects
171 somewhat older than the current ones with mean age of 51 years, the longitudinal
172 aging-associated change rates of the cpRNFLT and GCIPLT_{4TestPoints} would be sufficiently
173 negative to be detected. For example, at age 65 years (14 years older than the mean
174 age of the current subjects), a longitudinal aging-associated decline rate of the
175 cpRNFLT (GCIPLT_{4TestPoints}) would be $0.067 - 0.02 \times 14 = -0.21 \mu\text{m}/\text{year}$ ($-0.064 -$
176 $0.01 \times 14 = -0.20 \mu\text{m}/\text{year}$). Assuming similar variations in the cpRNFLT and
177 GCIPLT_{4TestPoints} measurement results, i.e., similar standard error values for estimates,
178 and the same values for other explanatory variables, a coefficient for duration of -0.15
179 $\mu\text{m}/\text{year}$ would be significantly different from zero, i.e., a significant decline rate of
180 -0.15 $\mu\text{m}/\text{year}$ would be obtained for the cpRNFLT or GCIPLT_{4TestPoints}. Thus, it is likely
181 that subjects older than the current ones could show significant longitudinal
182 aging-related declines in the cpRNFLT and GCIPLT_{4TestPoints}. The somewhat younger ages
183 of subjects in the current study than in the previous studies by Leung et al.[16,17] and
184 Vianna et al.[18] (51 vs. 56, 58 and 65 years, respectively) might be partly responsible
185 for the discrepancy between the current and their results. Relative rates of
186 aging-associated declines in the cpRNFLT were reportedly less than predicted by
187 aging-associated decline in the number of RGCs, which was attributed to the presence
188 of differential aging-associated declines of the non-neuronal components in the

189 RNFL.[31,32] A racial difference in the cpRNFLT has been reported [13,14] and a
190 comparison of the BMO-MRW and cpRNFLT between normal Japanese and Caucasians
191 subjects suggested a difference in the ratio of the amount of RGC axons to that of the
192 non-neuronal components in the RNFL between them.[33] If there are racial
193 differences in the amount of aging-associated changes in the non-neuronal component
194 between Japanese subjects and those of other races, this might partly explain the
195 discrepancy between the current result and those of Leung et al, Vianna et al., and
196 Hammel et al.[16-19]. Not only aging-associated changes in the retinal neurons but
197 also those in the neurons in the central visual pathway should be related to a
198 physiologic aging-associated decline in the perceived VF sensitivity in a specific retinal
199 area. Since the GCIPLT or cpRNFLT in a specific retinal area is related to the number of
200 RGCs or RGC axons but not neurons in the central visual pathway, the physiologic aging
201 processes at age 50 years might be reflected with more sensitivity in the perceived VF
202 sensitivity decline than in the thickness of the RGC-related retinal layers in the
203 corresponding retinal area. A significant aging-associated decline in the perceived VF
204 sensitivity not associated with a significant aging-associated decline in the cpRNFLT or
205 macular GCIPLT in the corresponding area currently observed in normal Japanese eyes
206 might be at least partly explained as above. On the other hand, glaucomatous damage
207 primarily affects the RGCs and their axons and secondarily the neurons in the central
208 visual pathway. In this case, the perceived SAP sensitivity decline may be dampened by
209 the plasticity of the visual cortex and normal cerebral adaptation to chronic

210 deterioration in the visual input caused by slowly progressing glaucomatous RGC
211 damage.[34]

212 Whatever the causes of the discrepancy in the aging-associated decline rates in the
213 cpRNFLT or macular GCIPLT between the current and previous longitudinal studies, the
214 current finding that a significant acceleration of aging-associated decline rate was
215 detected not only in the VF sensitivity, but also in the cpRNFLT and macular GCIPLT is
216 interesting. Acceleration of aging-associated decline rates of cpRNFLT and macular
217 GCIPLT along with increase in subjects' age is compatible with, and may be one of the
218 post-retinal neural factors for acceleration of ageing-associated decline rates of VF
219 sensitivity currently found and previously reported in cross-sectional datasets [7-9]
220 Further, it underlines again that an age-matched normal control group must be always
221 provided in studying diseases-caused thinning of cpRNFLT or macular GCIPLT, since
222 physiological aging-associated thinning rates of these RGC-related retinal layers
223 thicknesses were dependent on subjects' age.

224 The current study had several limitations. First, the number of normal subjects was
225 small and the follow-up period short, which would result in a lower statistical power to
226 detect significant time changes in the RGC-related retinal layer thickness measured
227 using a current SD-OCT instrument. As discussed previously, however, the cpRNFLT or
228 GCIPLT time change rates of about 0.3~0.5 $\mu\text{m}/\text{year}$ have been reported in previous
229 studies with a similar number of normal subjects and follow-up periods,[16-19] and the
230 test-retest variation of the current SD-OCT measurement results was thought to be

231 reasonably satisfactory based on modern technology. These results suggested that the
232 statistical power to detect the time change rates of the cpRNFLT or GCIPLT in the
233 current normal subjects should be comparable to those of previous studies that
234 detected significant aging-related declines of the cpRNFLT or macular GCIPLT.[16-19]
235 The mean age of the current normal Japanese subjects was about 50 years and as
236 discussed previously, the physiologic aging-related decline of the cpRNFLT and
237 GCIPLT_{4Testpoints} were calculated to be about 0.2 $\mu\text{m}/\text{year}$ at age 65 years, which could
238 be detected under the same study conditions. Thus, it should be noted that the current
239 results were applicable in Japanese subjects aged around 50 years.

240 In summary, acceleration of aging-associated decline rates of VF sensitivity
241 associated with acceleration of aging-associated decline rate or cpRNFLT or macular
242 GCIPLT in the corresponding retinal area along with increased subjects' age may have
243 implications for studying not only the physiologic aging-related changes in visual
244 function, but also the diseases-caused rates of thinning in the cpRNFLT or macular
245 GCIPLT. Further, the physiologic aging processes was more evidently found in the
246 declines in the HFA-measured VF sensitivity than in the cpRNFLT or macular GCIPLT in
247 corresponding retinal areas measured using a current SD-OCT instrument in norma
248 Japanese aged around 50 years.

249

250 **Methods**

251 **Subjects.** Self-reported healthy Japanese individuals were recruited at the Tajimi Eye

252 Clinic (Tajimi, Gifu, Japan). After subjects were screened verbally and the medical
253 histories recorded, an ocular examination was performed that included measurements
254 of the uncorrected and autorefraction-corrected visual acuity (VA) with the Landolt
255 chart at 5 meters and the corneal curvature using an autorefractometer (KR-800A,
256 Topcon, Tokyo, Japan). In addition, the central corneal thickness and AL were measured,
257 respectively, using a specular microscope (SP-3000P, Topcon) and the IOLMaster (Carl
258 Zeiss Meditec). The VFs were examined using SAP (HFA 24-2 SITA program, Carl Zeiss
259 Meditec). The VF examination was repeated if it was considered unreliable or outside
260 the normal limits. The SD-OCT examination was followed by dilated optic disc stereo
261 photography and fundus photography, dilated funduscopy, slit-lamp biomicroscopy,
262 and intraocular pressure (IOP) measurements by Goldmann applanation tonometry. A
263 pair of sequential stereoscopic optic nerve head photographs at a parallax of about 8
264 degrees (30-degree angle of view) and non-stereoscopic fundus photographs
265 (45-degree angle of view) was obtained using a digital fundus camera (TRC-NW7,
266 Topcon) after pupillary dilation with 1.0% tropicamide. All ocular examinations were
267 performed bilaterally.

268 The inclusion criteria were age between 20 and 80 years; normal eye examinations
269 without any clinically significant cataract, ocular media, vitreoretinal, or choroidal
270 abnormalities; IOP of 21 mmHg or lower; best-corrected decimal VA of 1.0 or higher;
271 spherical refraction of ± 6 diopters (D) or less; astigmatism of 2 D or less; AL of 26 mm
272 or less; no previous ocular surgeries; normal VF test results with the glaucoma

273 hemifield test, and mean deviation and pattern standard deviation within normal limits.
274 Subjects were excluded if the VF results were unreliable based on the perimetrist's
275 notes and reliability indices with fixation loss and false positive rates of over 20% and
276 over 15%, respectively; the optic disc stereo photographs were of insufficient quality;
277 or the OCT images were of insufficient quality (typically truncated B-scans and scans
278 with a manufacturer-authorized image quality score of 30 or lower). After enrollment,
279 routine ophthalmic examinations, SD-OCT and VF measurements were performed
280 every 3 months for 3 years.

281 The Review Board and Ethics Committee of Gifu Prefecture Medical Association
282 approved the study (reference number, 25-1-001), which adhered to the tenets of the
283 Declaration of Helsinki. The study was registered in the University Hospital Medical
284 Information Network Clinical Trial Registry (UMIN-000012412). Each subjects provided
285 written informed consent after they received a full explanation of the study protocol.

286

287 **SD-OCT.** SD-OCT data sets were obtained using a 3D-OCT 2000 (Topcon) with the
288 horizontal 3-dimensional (3D) scan protocol in which data were obtained from 6.0 ×
289 6.0-mm-square areas (512 A-scans × 128 frames) centered on the disc and with the
290 vertical 3D scan protocol in which data were obtained from a 7.0 × 7.0-mm-square area
291 (512 A-scans × 128 frames) centered on the fovea over a period of about 1.5 seconds
292 for each scan. The magnification was corrected according to the
293 manufacturer-provided formula that was based on refractive error, corneal radius, and

294 AL.[26] The data obtained in the presence of eye movements were discarded and the
295 examination was repeated. Images also were excluded if they were affected by
296 involuntary blinking or saccades, indicated by breaks, shifting of the vessels, or the
297 presence of a straight line across the fundus OCT image, or had an image quality score
298 of 30 or less. OCT measurements were repeated 3 times with several-second intervals,
299 and the image with the best quality was used. The disc barycenter was determined
300 using the raster-scan data, and the fovea identified in the OCT image as the pixel that
301 was thinnest between the inner limiting membrane and the photoreceptor inner/outer
302 segment junction adjacent to the fixation point. The RNFL and GCIPL were segmented
303 automatically in all B-scan images.[27] To minimize variability due to misplacement of
304 the measurement location and/or segmentation error, an experienced researcher (T.K.)
305 checked the locations of the disc barycenter and fovea and all layer segmentation in all
306 images. The cpRNFLT was measured along the 3.4-mm-diameter circle centered on the
307 disc center. The macular GCIPLT in a 0.6-mm-diameter circular retinal area
308 (corresponding to about 2 degrees of visual angle) corresponding to each of the four
309 central test points of the HFA 24-2 and adjusted for the RGC displacement [25] was
310 sequentially obtained and the mean of these 4 measurement results were obtained
311 ($GCIPLT_{4TestPoints}$). The diameter of the retinal area (2 degrees) was similar to the grid
312 size of the HFA 10-2 test program and roughly twice as large as the size-III stimulus
313 point movements during fixation, such as drift.

314 Inter-visit test-retest variabilities of the cpRNFLT and $GCIPLT_{4TestPoints}$ measurements

315 with the current instrument were calculated using the two measurements obtained at
316 enrollment during two separate sessions.

317

318 **Data analysis.** The results are expressed as the mean (standard deviation). The effects
319 of aging (duration or time lapse from the baseline measurement) on the cpRNFLT,
320 mean VF sensitivity (dB) on the HFA 24-2 VF (VF_{mean}), GCIPLT_{4TestPoints}, and the mean VF
321 sensitivity of the central 4 test points of the HFA 24-2 ($VF_{4\text{TestPoints}}$) were analyzed, using
322 the time changes of the cpRNFLT, VF_{mean} , GCIPLT_{4TestPoints}, or $VF_{4\text{TestPoints}}$ values during
323 follow-up as dependent variables and using the linear mixed model, which considers
324 correlations between the paired eyes and measurement values from the same eyes.
325 The explanatory variables were duration (time lapse from the baseline measurement),
326 baseline age, baseline VF_{mean} or $VF_{4\text{TestPoints}}$, and AL [28,29] for the time changes of the
327 VF_{mean} and $VF_{4\text{TestPoints}}$, respectively. For the time changes of the cpRNFLT or
328 GCIPLT_{4TestPoints}, the variables were duration (time lapse from the baseline
329 measurement), baseline age, baseline cpRNFLT or GCIPLT_{4TestPoints}, AL [11,15], and disc
330 area [10,16,17], respectively. The JMP[®] Pro 13 software (SAS Institute Inc., Cary, NC,
331 USA) was used for analysis and $p < 0.05$ was considered statistically significant.

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432 **Data Availability Statement**

433 Datasets generated and/or analyzed during the current study are available from the
434 the orresponding author on reasonable request.

435 **Author Contributions**

436 M.A., A.I. and Y.T. designed the current study, A.I. collected the data, M.A., A.I. and
437 Y.T. analyzed the data, M.F. and Y.O. carried out statistical calculation, and M.A. and
438 A. I. drafted the manuscript. All authors reviewed the manuscript.

439 **Additional Information**

440 Competing financial interests: Makoto Araie holds a patent licensed to Topcon
441 without any royalties, is a consultant of Santen and Topcon. Makoto Fujii and Yuko
442 Ohno declare no competing financial interests. Yuki Tanala is an employee of
443 Santen, and Tsutomu Kikawa is an employee of Topcon. Aiko Iwase holds a patent
444 licensed to Topcon without any royalties, is a consultant of Santen and has received
445 speaker honoraria from Carl Zeiss Meditec and Santen. All authors attest that they
446 meet the current ICMJE requirements to qualify as authors.

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