Correlation between peripapillary retinal nerve fiber layer thickness and fundus autofluorescence in primary open-angle glaucoma

Purpose: To investigate the relationship between retinal nerve fiber layer (RNFL) thickness and retinal pigment epithelium alterations in patients with advanced glaucomatous visual field defects.

Methods: A consecutive, prospective series of 82 study eyes with primary open-angle glaucoma and advanced glaucomatous visual field defects were included in this study. All study participants underwent a full ophthalmic examination followed by visual field testing with standard automated perimetry as well as spectral-domain optical coherence tomography (SD-OCT) for peripapillary RNFL thickness and Optos wide-field fundus autofluorescence (FAF) images. A pattern grid with corresponding locations between functional visual field sectors and structural peripapillary RNFL thickness was aligned to the FAF images at corresponding location. Mean FAF intensity (range: 0 = black and 255 = white) of each evaluated sector (superotemporal, temporal, inferotemporal, inferonasal, nasal, superonasal) was correlated with the corresponding peripapillary RNFL thickness obtained with SD-OCT.

Results: Correlation analyses between sectoral RNFL thickness and standardized FAF intensity in the corresponding topographic retina segments revealed partly significant correlations with correlation coefficients ranging between 0.004 and 0.376 and were statistically significant in the temporal inferior central field ($r = 0.324$, $P = 0.036$) and the nasal field ($r = 0.376$, $P = 0.014$).

Conclusion: Retinal pigment epithelium abnormalities correlate with corresponding peripapillary RNFL damage, especially in the temporal inferior sector of patients with advanced glaucomatous visual field defects. A further evaluation of FAF as a potential predictive parameter for glaucomatous damage is necessary.

Keywords: glaucoma, fundus autofluorescence, FAF, retinal nerve fiber layer, RNFL, optical coherence tomography, OCT, imaging

Introduction

Structural changes in primary open-angle glaucoma (POAG) occur in the optic nerve head as well as peripapillary retinal nerve fiber layer (RNFL).\(^1\)\(^-\)\(^4\) While it has been assumed that structural loss of the RNFL anticipates functional visual field loss, the relationship between structure and function in glaucoma is not yet fully understood and remains the subject of intense research.\(^2\)\(^-\)\(^9\)

While a close relationship between RNFL and the retinal pigment epithelium (RPE) is a prerequisite for normal vision,\(^10\)\(^-\)\(^11\) little is known about RPE involvement in patients suffering from glaucoma. A previously published study on dogs has shown pathologic abnormalities and RPE damage in glaucomatous eyes.\(^12\) Few published studies on human glaucomatous eyes have shown correlations between peripapillary
RPE abnormalities and optic nerve degeneration in glaucoma patients. A recent study has shown that RPE atrophies seem to progress over the course of the disease.

Fundus autofluorescence (FAF) has been used for the evaluation of degenerative, inflammatory, and neoplastic diseases of the posterior fundus. The FAF signal is supposedly derived primarily from fluorophores found in lipofuscin within the RPE layer – mainly fluorophore N-retinylidene-N-retinyl ethanolamine (A2E) – and indicates altered structure as a result of atrophic changes due to inflammation resulting in increased or decreased FAF in the affected areas. Open-angle glaucoma (OAG) is generally understood as a degenerative disease and remains one of the leading causes of blindness in the elderly population. POAG is the most common subtype of OAG, with glaucoma-typical structural as well as functional glaucomatous alterations. To date, there is only very limited evidence on altered RPE signal intensity in glaucoma patients.

The recently developed ultra-wide-field scanning laser ophthalmoscope Optomap Panoramic 200Tx (Optos, Dunfermline, Fife, Scotland) allows non-mydriatic wide-field imaging and provides an additional wide-field FAF detection modality.

The goal of this study was to investigate the relationship between RNFL thickness and RPE alterations as detected by wide-field FAF in patients suffering from manifest POAG.

**Patients and methods**

**Patients**

A consecutive, prospective series of 82 study eyes, mean age 64 years (range: 35–86 years), with POAG and advanced glaucomatous visual field defects was included in this study (47 female and 35 male, 42 right and 40 left eyes).

According to the definition of the European Glaucoma Society and a previous work by Katz et al., a visual field defect suspicious for glaucoma is defined as a cluster of at least three or more contiguous non-edge points with significantly reduced sensitivity ($P < 0.05$), one of which with a significance of at least $P < 0.01$ on the same side of the horizontal meridian in the pattern deviation plot.

All study participants underwent a full ophthalmic examination including best-corrected visual acuity (BCVA) in decimals obtained with a Snellen projection chart; objective and subjective refraction; slit-lamp biomicroscopy; intraocular pressure measurement with Goldmann applanation tonometry; gonioscopy; and fundus examination by indirect ophthalmoscopy.

Each included patient underwent visual field testing followed by spectral-domain optical coherence tomography (SD-OCT) for peripapillary RNFL thickness and Optos wide-field FAF images.

Only eyes suffering from POAG were included, defined by a documented history of elevated intraocular pressure, characteristic optic nerve head damage, and/or characteristic visual field defects. Exclusion criteria were any evidence for optic nerve head degeneration derived from any other subtype of glaucoma than POAG; opticopathy of any reason other than glaucoma; diabetic retinopathy; macular disease; posterior uveitis; or retinal surgery.

All research was conducted in accordance with institutional guidelines and board approval and conformed to the tenets of the World Medical Association Declaration of Helsinki.

**SD-OCT measurements**

Conventional predefined circular peripapillary optic nerve head (ONH) cross-sectional scans were obtained from each included study eye with a Spectralis optical coherence tomography device (Heidelberg Engineering, Heidelberg, Germany) with an excitation wavelength of 870 nm running at a speed of 40,000 A-scans per second. Global and sectoral (superotemporal, temporal, inferotemporal, inferonasal, nasal, and superonasal) RNFL thickness values in µm were calculated with the inbuilt Heidelberg software.

**Optomap imaging**

Optomap imaging was performed without pupil dilation. Optomap imaging consisted of taking several images and saving the best image of each included eye for grading. The device takes one image in approximately 0.25 seconds, thus avoiding motion artifacts. Total scanning time was about 3–5 minutes, which included patient positioning, and was performed by an experienced technician (TM). Basic operation of the Optomap for ultra-wide-field composite color fundus images involves a scanning laser ophthalmoscope with two laser wavelengths scanning at 532 nm (“green laser separation”) and 633 nm (“red laser separation”). The optical resolution was 3,900–3,072 pixels, resulting in approximately 17–20 pixels per degree. Wide-field FAF images were obtained with an excitation wave length of 532 nm, and autofluorescence was detected by a broadband detector at 540 to 800 nm.

**Quantification of FAF signal intensity and correlation with peripapillary RNFL thickness**

For further analysis, the pattern grid with corresponding locations between functional visual field sectors and
structural peripapillary RNFL thickness originally described by Garway-Heath et al. was aligned to the FAF images at corresponding locations (Figure 1). FAF brightness with values ranging between 0 (black) and 255 (white) was obtained for each FAF sector (superotemporal peripheral [1], superotemporal central [2], temporal central [3], inferotemporal central [4], inferotemporal peripheral [5], and nasal [6]; Figure 1C and D) using standalone software based on Matlab (7.0 R14; Mathworks Inc, Natick, MA, USA), as described in our previous works. Mean FAF intensity of each sector was standardized for each included study eye to eliminate individual FAF intensity differences due to aging and correlated with the corresponding peripapillary RNFL thickness obtained with SD-OCT.

**Statistical analysis**

Data were collected and analyzed using SPSS software (version 19.0; IBM Corporation, Armonk, NY, USA). A P-value of < 0.05 was considered statistically significant. Univariate parametric analyses were applied.

**Results**

Eighty-two glaucomatous eyes were included in our prospective study. Patient characteristics are displayed in Tables 1 and 2. Mean FAF intensities and mean sectoral peripapillary RNFL thickness values for analyzed subfields 1–6 of each included study eye can be seen in Table 3. The highest mean FAF intensity was observed in the peripheral inferotemporal sector, while the central superotemporal sector showed the lowest mean FAF intensity of all investigated subfields (Table 3). Mean RNFL thickness was highest in the peripheral inferotemporal sector and lowest in the nasal sector (Table 3). Correlation analyses between sectoral RNFL thickness and standardized FAF intensity in the corresponding topographic retina segments revealed partly significant correlations (Table 4). Correlation coefficients ranged between 0.004 and

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**Figure 1** Structure–function relationship between visual field (A) and peripapillary nerve fiber layer obtained with spectral-domain optical coherence tomography (C and D), adapted for corresponding locations of the posterior fundus (D). **Note:** The principle of the evaluation is based on the works of Strouthis et al.
and 0.376 and were statistically significant in the temporal inferior central field ($r = 0.324, P = 0.036$) and the nasal field ($r = 0.376, P = 0.014$).

**Discussion**

In retinal metabolism, a close symbiotic interaction between neurosensory retina and RPE exists.$^{26}$ RPE has various support functions, including neuroprotection, and its damage results in functional alteration of the neurosensory retina, including the RNFL.$^{28-32}$ Therefore, FAF has become a valuable tool in monitoring the development and progression of atrophic RPE dysfunction and consecutive neurosensory degeneration, such as in age-related macula degeneration (AMD).$^{18,19}$ A close correlation between RPE damage measured with FAF intensity and visual deterioration has been described.$^{33-36}$ In reverse, RNFL damage may be associated with RPE dysfunction. However, little is known as to whether RNFL damage, which has been demonstrated to correlate with visual field function,$^{7,26,37}$ is connected with RPE malfunction or degeneration potentially detectable by FAF. Our observations of a certain correlation between sectoral RNFL thickness and FAF intensity at corresponding locations in glaucomatous eyes are in accordance with published studies on dogs and reveal a close interaction between RPE and the neurosensory retina.$^{12}$ The results of this study provide early evidence that peripapillary RNFL degeneration in glaucoma patients may be accompanied by corresponding RPE alterations. These findings apply especially for the temporal inferior and nasal sectors and thus are, at least in part, in agreement with results and observations of previously published data on structure–function relationships in glaucomatous eyes, which suggest the temporal inferior peripapillary region in particular to be a predilection site for structural damage in glaucoma.$^{38,39}$

Those studies on evaluated functional visual field abnormalities in glaucoma patients correlated the observed functional abnormalities to structural RNFL changes and showed the temporal inferior sector, in particular, to be subject to structural changes in patients with visual field defects due to glaucoma.

Further studies are needed to give a better understanding of peripapillary sectors more susceptible to degenerative changes than others and to possibly address the question as to whether this observation is due to biomechanic or nutritive predisposing anatomic circumstances.

Significant correlations between nasal RNFL thickness and corresponding FAF intensity in human POAG eyes have

| Parameter | n = 84 |
|-----------|--------|
| Age (years) | $64 \pm 13$ |
| BCVA (Snellen) | $0.78 \pm 0.25$ |
| CDR | $0.82 \pm 0.15$ |
| US-CCT (µm) | $541 \pm 35$ |

**Note:** Values are given as mean ± standard deviation.

**Abbreviations:** BCVA, best-corrected visual acuity; CDR, vertical cup/disc ratio; US-CCT, ultrasound-based measurement of central corneal thickness.

| Table 1 Patient characteristics: age, BCVA, vertical CDR, and US-CCT |
|----------------|-------------|
| Parameter | n = 84 |
| Age (years) | $64 \pm 13$ |
| BCVA (Snellen) | $0.78 \pm 0.25$ |
| CDR | $0.82 \pm 0.15$ |
| US-CCT (µm) | $541 \pm 35$ |

**Note:** Values are given as mean ± standard deviation.

**Abbreviations:** BCVA, best-corrected visual acuity; CDR, vertical cup/disc ratio; US-CCT, ultrasound-based measurement of central corneal thickness.

**Table 2** Age distribution of all included patients

| Age subgroup (years) | n |
|---------------------|---|
| 31–40 | 4 |
| 41–50 | 10 |
| 51–60 | 14 |
| 61–70 | 31 |
| 71–80 | 19 |
| 81–90 | 6 |

**Discussion**

In retinal metabolism, a close symbiotic interaction between neurosensory retina and RPE exists.$^{26}$ RPE has various support functions, including neuroprotection, and its damage results in functional alteration of the neurosensory retina, including the RNFL.$^{28-32}$ Therefore, FAF has become a valuable tool in monitoring the development and progression of atrophic RPE dysfunction and consecutive neurosensory degeneration, such as in age-related macula degeneration (AMD).$^{18,19}$ A close correlation between RPE damage measured with FAF intensity and visual deterioration has been described.$^{33-36}$ In reverse, RNFL damage may be associated with RPE dysfunction. However, little is known as to whether RNFL damage, which has been demonstrated to correlate with visual field function,$^{7,26,37}$ is connected with RPE malfunction or degeneration potentially detectable by FAF. Our observations of a certain correlation between sectoral RNFL thickness and FAF intensity at corresponding locations in glaucomatous eyes are in accordance with published studies on dogs and reveal a close interaction between RPE and the neurosensory retina.$^{12}$ The results of this study provide early evidence that peripapillary RNFL degeneration in glaucoma patients may be accompanied by corresponding RPE alterations. These findings apply especially for the temporal inferior and nasal sectors and thus are, at least in part, in agreement with results and observations of previously published data on structure–function relationships in glaucomatous eyes, which suggest the temporal inferior peripapillary region in particular to be a predilection site for structural damage in glaucoma.$^{38,39}$

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Further studies are needed to give a better understanding of peripapillary sectors more susceptible to degenerative changes than others and to possibly address the question as to whether this observation is due to biomechanic or nutritive predisposing anatomic circumstances.

Significant correlations between nasal RNFL thickness and corresponding FAF intensity in human POAG eyes have

| Table 3 Mean FAF intensities and peripapillary RNFL thickness values for each analyzed sector (1–6) |
| Sector | FAF intensities (0–255) | RNFL thickness (µm) |
|--------|----------------|------------------|
| Superotemporal peripheral (1) | 160 ± 27 | 55.9 ± 15.8 |
| Superotemporal central (2) | 146 ± 24 | 74.2 ± 23.7 |
| Temporal central (3) | 163 ± 28 | 78.6 ± 24.5 |
| Inferotemporal central (4) | 165 ± 26 | 90.3 ± 28.7 |
| Inferotemporal peripheral (5) | 175 ± 29 | 94.0 ± 33.8 |
| Nasal (6) | 171 ± 28 | 54.1 ± 15.2 |

**Note:** Values are given as mean ± standard deviation.

**Abbreviations:** FAF, fundus autofluorescence; RNFL, retinal nerve fiber layer.

**Table 4 Correlations (Pearson correlation coefficient) between FAF intensity and peripapillary RNFL thickness at corresponding sectors (1–6)**

| Correlation between sectoral mean FAF intensity and RNFL thickness |
|----------------|----------------|
| Sector | Correlation |
| Superotemporal peripheral (1) | 0.273; $P = 0.080$ |
| Superotemporal central (2) | 0.114; $P = 0.474$ |
| Temporal central (3) | 0.004; $P = 0.979$ |
| Inferotemporal central (4) | 0.324; $P = 0.036^*$ |
| Inferotemporal peripheral (5) | 0.280; $P = 0.073$ |
| Nasal (6) | 0.376; $P = 0.014^*$ |

**Note:** *Values that were statistically significant.

**Abbreviations:** FAF, fundus autofluorescence; RNFL, retinal nerve fiber layer.
not, to our knowledge, been published so far and remain our focus of continuing research.

A limitation of our study is the fact that, due to its cross-sectional character, we do not have data from over the course of glaucoma disease and are not able to predict any of our observed developments. Additionally, FAF intensity changes during aging, so that interindividual comparison of measured FAF intensity values should be done with caution.

Conclusion
The presented early data from POAG patients shows that, at first – and from the structural point of view – a certain correlation between FAF, reflecting RPE function, and peripapillary RNFL thickness exists. As a next step, it will have to be explored whether these findings are in correlation to the actual function deficit measured by visual fields and whether our observation can be interpreted with regards to a potential predictive parameter for glaucomatous damage.

In summary, we were able to correlate RPE abnormalities to corresponding peripapillary RNFL damage especially in the temporal inferior sector of patients with manifest glaucomatous visual field defects. A further evaluation of FAF as a potential predictive parameter for glaucomatous damage is necessary.

Disclosure
The authors report no conflicts of interest in this work.

References
1. Douglas GR, Drance SM, Schulzer M. A correlation of fields and discs in open angle glaucoma. Can J Ophthalmol. 1974;9(4):391–398.
2. Zeyen TG, Caprioli J. Progression of disc and field damage in early glaucoma. Arch Ophthalmol. 1993;111(1):62–65.
3. Quigley HA, Broman AT. The number of people with glaucoma worldwide in 2010 and 2020. Br J Ophthalmol. 2006;90(3):262–267.
4. Quigley HA, Katz J, Derick RJ, Gilbert D, Sommer A. An evaluation of optic disc and nerve fiber layer examinations in monitoring progression of early glaucoma damage. Ophthalmology. 1992;99(1):19–28.
5. Kerrigan-Baumrind LA, Quigley HA, Pease ME, Kerrigan DF, Mitchell RS. Number of ganglion cells in glaucoma eyes compared with threshold visual field tests in the same persons. Invest Ophthalmol Vis Sci. 2000;41(3):741–748.
6. Airaksinen PJ, Tuulonen A, Alanko HI. Rate and pattern of neuroretinal rim area decrease in ocular hypertension and glaucoma. Arch Ophthalmol. 1992;110(2):206–210.
7. Swanson WH, Felius J, Pan F. Perimetric defects and ganglion cell damage: interpreting linear relations using a two-stage neural model. Invest Ophthalmol Vis Sci. 2004;45(2):466–472.
8. Keltner JL, Johnson CA, Anderson DR, et al. Ocular Hypertension Treatment Study Group. The association between glaucomatous visual fields and optic nerve head features in the Ocular Hypertension Treatment Study. Ophthalmology. 2006;113(9):1603–1612.
9. Hood DC, Kardon RH. A framework for comparing structural and functional measures of glaucomatous damage. Prog Retin Eye Res. 2007;26(6):688–710.
10. Hosoya K, Tachikawa M. The inner blood-retinal barrier: molecular structure and transport biology. Adv Exp Med Biol. 2012;763:85–104.
11. Sparrow JR, Hicks D, Hamel CP. The retinal pigment epithelium in health and disease. Curr Mol Med. 2010;10(9):802–823.
12. Mangan BG, Al-Yahya K, Chen CT, et al. Retinal pigment epithelial damage, breakdown of the blood-retinal barrier, and retinal inflammation in dogs with primary glaucoma. Vet Ophthalmol. 2007;10 Suppl 1:117–124.
13. Jonas JB, Naumann GO. Parapapillary retinal vessel diameter in normal and glaucoma eyes. II. Correlations. Invest Ophthalmol Vis Sci. 1989;30(7):1604–1611.
14. Jonas JB, Nguyen NX, Gusek GC, Naumann GO. Parapapillary choroidal atrophy in normal and glaucoma eyes. I. Morphometric data. Invest Ophthalmol Vis Sci. 1989;30(5):908–918.
15. Wilensky JT, Koller AE. Peripapillary changes in glaucoma. Am J Ophthalmol. 1976;81(3):341–345.
16. Guo Y, Wang YX, Xu L, Jonas JB. Five-year follow-up of parapapillary atrophy: the Beijing Eye Study. PLoS One. 2012;7(5):e32005.
17. Yannuzzi LA, Ober MD, Slakter JS, et al. Ophthalmic fundus imaging: today and beyond. Am J Ophthalmol. 2004;137(3):511–524.
18. Schmitz-Valckenberg S, Holz FG, Bird AC, Spade RF. Fundus autofluorescence imaging: review and perspectives. Retina. 2008;28(3):385–409.
19. Reznicek L, Wasfy T, Stumpf C, et al. Peripheral fundus autofluorescence is increased in age-related macular degeneration. Invest Ophthalmol Vis Sci. 2012;53(4):2193–2198.
20. Reznicek L, Kernt M, Seidensticker F, et al. Autofluorescence in posterior uveitis. Ophthalmology. 2012;119(8):1712–1713. e3.
21. Delori FC, Dorey CK, Starenghi G, Arend O, Goger DG, Weiter J. In vivo fluorescence of the ocular fundus exhibits retinal pigment epithelium lipofuscin characteristics. Invest Ophthalmol Vis Sci. 1995;36(3):718–729.
22. Delori FC, Goger DG, Dorey CK. Age-related accumulation and spatial distribution of lipofuscin in RPE of normal subjects. Invest Ophthalmol Vis Sci. 2001;42(8):1855–1866.
23. Kitagawa K, Nishida S, Ogura Y. In vivo quantitation of autofluorescence in human retinal pigment epithelium. Ophthalmologica. 1989;199(2–3):116–121.
24. Katz J, Sommer A, Gaasterland DE, Anderson DR. Comparison of analytic algorithms for detecting glaucomatous visual field loss. Arch Ophthalmol. 1991;109(12):1684–1689.
25. Garway-Heath DF, Poinoosawmy D, Fitzke FW, Hitchings RA. Mapping the visual field to the optic disc in normal tension glaucoma eyes. Ophthalmology. 2000;107(10):8109–8115.
26. Nilforoushan N, Nassiri N, Moghimi S, et al. Structure–function relationships between spectral-domain OCT and standard achrromatic perimetry. Invest Ophthalmol Vis Sci. 2012;53(6):2740–2748.
27. Reznicek L, Dabov S, Hartigolou C, Kampik A, Kernt M, Neubauer AS. Green-light fundus autofluorescence in diabetic macular edema. Int J Ophthalmol. 2013;6(1):75–80.
28. Hadziahmetovic M, Kumar U, Song Y, et al. Microarray analysis of murine retinal light damage reveals changes in iron regulatory, complement, and antioxidant genes in the neurosensory retina and isolated RPE. Invest Ophthalmol Vis Sci. 2012;53(9):5231–5241.
29. Strauss O. The retinal pigment epithelium in visual function. Physiol Rev. 2005;85(3):845–881.
30. Boulton M, Dayah-Barker P. The role of the retinal pigment epithelium: topographical variation and ageing changes. Eye (Lond). 2001;15(3 Pt 3):384–389.
31. Boulton M, Rózanowska M, Rózanowski B. Retinal photodamage. J Photochem Photobiol B. 2001;64(2–3):144–161.
32. Araki T, Taniwaki T, Becerra SP, Chader GJ, Schwartz JP. Pigment epithelium-derived factor (PEDF) differentially protects immature but not mature cerebellar granule cells against apoptotic cell death. J Neurosci Res. 1998;53(1):7–15.
33. Acton JH, Gibson JM, Cubbidge RP. Quantification of visual field loss in age-related macular degeneration. PloS One. 2012;7(6):e39944.
34. Acton JH, Smith RT, Hood DC, Greenstein VC. Relationship between retinal layer thickness and the visual field in early age-related macular degeneration. Invest Ophthalmol Vis Sci. 2012;53(12):7618–7624.
35. Seidensticker F, Neubauer AS, Wasfy T, et al. Wide-field fundus autofluorescence corresponds to visual fields in chorioretinitis patients. Clin Ophthalmol. 2011;5:1667–1671.
36. Coleman HR, Chan CC, Ferris FL 3rd, Chew EY. Age-related macular degeneration. Lancet. 2008;372(9652):1835–1845.
37. Rao HL, Zangwill LM, Weinreb RN, Leite MT, Sample P A, Medeiros FA. Structure-function relationship in glaucoma using spectral-domain optical coherence tomography. Arch Ophthalmol. 2011;129(7):864–871.
38. Vermeer KA, van der Schoot J, Lemij HG, de Boer JF. RPE-normalized RNFL attenuation coefficient maps derived from volumetric OCT imaging for glaucoma assessment. Invest Ophthalmol Vis Sci. 2012;53(10):6102–6108.
39. Leaney J, Healey PR, Lee M, Graham SL. Correlation of structural RNFL parameters and functional measures using HRT3 and Spectralis SD-OCT at different levels of glaucoma severity. Clin Experiment Ophthalmol. 2012;40(8):802–812.
40. Strouthidis NG, Vincotti V, Tucker AJ, Gardiner SK, Crabb DP, Garway-Heath DF. Structure and function in glaucoma: the relationship between a functional visual field map and an anatomic retinal map. Invest Ophthalmol Vis Sci. 2006;47(12):5356–5362.