First-Visit Diagnosis of Preperimetric Glaucoma

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Abstract: Purpose: To present a revised interpretation of the work-up data that enabled diagnosis of preperimetric glaucoma (PPG) at the first examination.

Methods: a) Literature analysis on PPG; b) 6-year follow-up of a glaucoma-suspect patient.

Results: Two new concepts may be adapted: (a) the objective finding of retinal nerve fiber layer (RNFL) thinning below the normal border in the opposing typical glaucoma locations, the inferior and superior quadrants, and in a non-diffuse pattern, appears asymptotically and simultaneously only in glaucoma; and (b) the imaging-related RNFL thickness may be considered the reference glaucoma standard, whereas the suspicious early glaucomatous optic neuropathy, having a potential diagnostic inaccuracy, would serve as a complementary revealing finding. That approach enabled, in retrospect, a first-visit diagnosis of low-tension PPG in the patient. Diagnosis was confirmed after 6 years, when cecocentral scotoma and further RNFL thinning emerged despite treatment.

Conclusions: A revised approach enabled PPG diagnosis during the first visit.

Keywords: Preperimetric glaucoma, early glaucoma, low-tension glaucoma, optical coherence tomography, retinal nerve fiber layer.

INTRODUCTION

In glaucoma, structural injury has been documented to precede functional injury in most, but not all, eyes [1]. Recently, a change in the diagnostic criteria of glaucoma has been promoted such that glaucoma diagnosis may be made before the old prerequisites’ functional criteria of standard automatic perimetry visual field (SAP-VF) loss are apparent, namely ‘preperimetric glaucoma’ (PPG). In PPG, structural injury, and especially glaucomatous optic neuropathy (GON), ought to be clinically proven. However, because of the inter-individual variability of the normal optic nerve head (ONH), the subjective diagnosis of GON at its early stages, either clinically or by means of 3-D photos, is often challenging [2, 3]. Detection of early loss of retinal nerve fibers, both clinically or by means of red-free photos, is also subjective and may be indecisive; more so in diffuse than in localized retinal nerve fibers loss [4]. In contrast, numerous studies have found the objective, quantitative assessment of retinal nerve fiber layer (RNFL) loss in early and advanced glaucoma by either optical coherence tomography (OCT, -1, -2 or -3) or scanning laser polarimetry (GDx-VCC) to be both accurate and reproducible [3-14]. In spite of these findings, the ONH evaluation is still considered the reference standard in PPG diagnosis, but proof of progressive worsening at the ONH, the retinal nerve fibers or SAP-VF is required to verify diagnosis of early GON [2, 15]. However, progression would naturally be coupled with further irreversible loss of RNFL and may occasionally be associated with VF injury.

The OCT and GDx-VCC provide micron-resolution assessment of the peripapillary (3.4-mm in diameter) RNFL thickness. The RNFL loss in glaucoma, as depicted by these imaging devices, has been found to be directly associated with the severity of the disease [6-11]: In advanced glaucoma, RNFL loss is generally found to be significantly thinner than normal in all or most (except horizontal) sites. In moderate and early glaucoma associated with typical VF loss, peripapillary RNFL loss regularly occurs in typical locations, i.e., at the inferior and/or superior quadrants and/or globally. In eyes with PPG, in which SAP-VF is normal and ONH is determined as GON by other diagnostic modalities, such as the use of frequency-doubling technology, RNFL loss regularly occurs at the same typical locations as in perimetric glaucoma [2, 3, 9, 16].

Retinal nerve fiber layer thinning has been found to be a predictor of the development of glaucomatous visual field damage in glaucoma-suspect eyes, i.e., when the ONH looked clinically suspicious of being glaucomatous [17], but this group of authors stated that proof of progressive worsening of the disease is a prerequisite for diagnosing PPG, as mentioned above [2, 15]. To avoid an irreversible disease progression in such eyes, in association with normal VF, it should be our goal to prompt or rule out a diagnosis of PPG. An interpretation of the cumulative data found during the evaluation of a 49-year-old patient, which is based on a revised literature analysis, enabled a diagnosis of PPG at his first visit. To the best of my knowledge (search via Entrez Pubmed), this revised diagnostic approach is presented here for the first time.

METHODS

The study is based on analysis of the literature review on RNFL loss in association with perimetric glaucoma and PPG...
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diagnosis. Conclusions refer to a revised interpretation of these cumulative data. That analysis was retrospectively carried out and a PPG diagnosis confirmed after 6 years of follow-up of a patient who presented initially with suspicious GON, low intraocular pressure, normal SAP-VF and abnormally thinned RNFL in each eye.

RESULTS

A) Literature Analysis

The cumulative data on the micron-resolution RNFL imaging studies, in association with the various stages of glaucoma, point out the following:

1. There is potential for inaccuracy in the subjective clinical determination of suspicious ONH at its early glaucoma stage as GON [2, 3, 8, 15];

2. Experts of the Optic Disk Reading Center found that even detection of GON progression is challenging if changes are subtle [17];

3. Looking for a proof of disease progression in order to define a glaucoma-suspect as having PPG might be risky. For example, complicated progression ensued in the presented patient when the foremost detectable progression was associated with cecocentral scotoma and reduction in visual acuity (see below);

4. On the other hand, the RNFL thickness assessment by high-resolution imaging devices is most often objective and accurate [5-16]. Several imaging-related RNFL studies on PPG-suspect eyes overcame the obstacle of doubtful GON diagnosis by either (a) assessing eyes with abnormal doubling technology perimetry as glaucomatous (PPG) [10]; (b) studying the fellow eyes of glaucoma patients with unilateral VF loss [3]; or (c) determining GON only in eyes with documented evidence of progressive ONH changes [2]. In these PPG studies, significant RNFL loss emerged at the same locations as typically occur in the more advanced perimetric glaucoma, i.e., at the inferior quadrant, superior quadrant and/or globally [2, 3, 9, 16];

5. A comparative study has shown that an objective structural assessment of the RNFL thickness by the imaging devices is more accurate than clinical evaluation of disc photographs in PPG [3].

Based on these five points, a revised diagnostic approach of glaucoma at the preperimetric stage is considered such that the subjective clinical diagnosis of GON as the reference standard in diagnosing PPG would probably be changed by adopting the available and more accurate alternative: objective and quantitative assessment of the peripapillary RNFL thickness loss. This change may be undertaken by recognizing that three findings appear concurrently only in glaucoma: (a) the presence of suspicious GON; (b) in an asymptomatic patient; together with (c) simultaneous RNFL loss in a non-diffuse pattern at the two opposing locations, the superior and inferior quadrants (Fig. 1), without RNFL nasal quadrant attenuation, to a level below the lower border of the normative database (in OCT-2 and to the red area in OCT-3). These observations have not been reported (search via Entrez Pub-Med) in any RNFL study on other optic neuropathies, including multiple sclerosis, Parkinson’s disease, ischemic optic neuropathy, retrobulbar neuritis, branch retinal artery occlusion, traumatic optic neuropathy, dominant optic atrophy, Leber’s hereditary optic neuropathy (LHON) and AIDS optic neuropathy [18-28]. All of these are, by and large, symptomatic at presentation and/or present with diffuse RNFL thinning, rather than presenting as a loss at the two opposing, typical glaucomatous sites.

The revised interpretation presented here enabled, in retrospect, a first-visit diagnosis of low-tension PPG. The OCT initially disclosed bilateral typical RNFL thinning, and diagnosis was confirmed after 6 years, when cecocentral scotoma and further RNFL thinning emerged despite topical treatment.

B) Case Report

A 49-year-old healthy man was referred to the outpatient clinic of our institution because of bilateral suspicious GON (“glaucoma suspect”) and normal standard VFs. The patient reported no history of trauma or family history of visual loss, and he had not received any medication. On examination in February 2002, his best-corrected visual acuity (BCVA) was 6/12 in each eye with myopic astigmatism; IOP was 12 mmHg in the RE and 14 in the LE, respectively, without hypotensive treatment. The anterior chamber angle was open +4 bilaterally. The ONHs had vertical cup/disc ratios of 0.7 and thin rims, and the optic cup in the right eye (RE) looked deeper than that in the other disk. An OCT examination (OCT-2000, Carl Zeiss Meditec, Dublin, CA, USA) was carried out. The Circle group program, 3.4-mm in diameter, was centered on the ONH for the assessment of the peripapillary RNFL. Results were based on the mean of five subsequent good scans per eye. Comparison was made in the graphic presentation with the age-matched OCT normative database (Fig. 1). The 24-2 Swedish Interactive Testing Algorithm of the SAP (Humphrey Zeiss Meditec, Dublin, CA, USA) was used for VF examination.

Quantitative, numerical analysis of the first OCT findings can be made at this time only with our own RNFL age-matched normative database (n = 30; Fig. 1) since numerical data of the OCT manufacturer normative database are not available. Furthermore, two instruments with the same software can have different RNFL-t outputs [3]. The mean global normal RNFL-t in our instrument is 138 ± 30 μm, and its mean lower border is 113 ± 31 μm. The range between the upper and lower borders in each site (i.e., clock-hour) in our normative database is thinner than that of the OCT manufacturer normative database.

In February 2002, the RNFL-t was bilaterally thinner than the lower border of the OCT- manufacturer normative database, mainly in the inferior quadrants, the superior-temporal site, the temporal quadrant and globally in each eye (Figs. 1, 2A). Quantitative comparison at that initial stage with our own normative database revealed thinning of the RNFL below its lower border in the two typical RNFL-loss glaucomatous locations by 37% (from 150 to 94 μm; Fig. 1) in the inferior quadrant and 43% in the superior-temporal site in the RE, and by 32% and 42% in the LE, respectively (Fig. 1). Mean global (4 quadrants) RNFL-t (Table at Fig. 2A) was 95 μm in the RE and 104 μm in the LE, which was lower by 16% and 8%, respectively, than the mean RNFL-t
Lower border, 113 μm (Fig. 1). The mean RNFL of only the temporal quadrant was markedly thinner than the mean of the lower border self-normative database, by 46% and 53%, respectively. The VFs were full (Fig. 2).

A 24-hour diurnal-curve IOP took place thereafter, and IOPs ranged between 12-14 mmHg in each eye. Based on the clinical suspicion of early GON associated with RNFL thinning in two opposing typical glaucoma locations, i.e., in the inferior quadrant and the superior-temporal site, as well as globally, diagnosis of low-tension PPG was made. Col. betaxolol 0.5%, a hypotensive and possibly neuroprotective agent, was prescribed b.i.d. following the diurnal curve examination for safety reasons.

According to the patient, during 6 years of follow-up, examinations were usually carried out in a community clinic every 3 to 4 months, while he was periodically referred for clinical and VF examinations. During follow-up, obstructive sleep apnea was ruled out per anamnesis. Central corneal thicknesses were 512 and 520 μm, for the LE and RE, respectively. The IOP ranged between 10-14 mmHg in each eye. The VF data was full (Fig. 2).

The subsequent work-up in our clinic in September 2007, the cup/disc ratio was found to have worsened to 0.8 in each eye with thin rims, and cecocentral scotoma was found, mainly in the lower border self-normative database, by 46% and 53%, respectively. The VF data was full (Fig. 2).

Topical brimonidine tartrate 0.2% was bilaterally added. After 8 months, the scotoma widened in the RE (Fig. 2C) and BCVA was 6/18 in that eye; the scotoma in the LE became non-specific and more diffuse, while BCVA remained 6/12.

An OCT-RNFL evaluation was carried out for the second time in May 2008. Further RNFL thinning was found in the inferior (26%) and temporal (67%) quadrants and globally (29%) of the RE, while it occurred in the superior quadrant (14%) in the LE. In addition, further RNFL thinning was detected bilaterally at various other sites (Fig. 2B). A third OCT examination was done four months later, revealing similar data. The IOP in a repeat diurnal curve ranged between 9-14 mmHg in each eye and was followed later by further worsening of the cecocentral scotoma thinning in the RE (not presented). Approval of the local Ethics Committee was not required for this report.

DISCUSSION

Based on a revised deduction from the literature review, the objective finding of a simultaneous RNFL thinning in two typical glaucoma locations that appeared asymptptomatically and in the presence of suspicious early GON in the presented patient raised the possibility of PPG diagnosis during the first work-up visit. After 6 years, diagnosis was confirmed when progression of the disease had been found, manifesting in the emergence of cecocentral scotoma, bilateral worsening of GON and further RNFL thinning, despite topical treatment. This revised deduction may be supported by the following:

Studies have suggested that RNFL assessment may become a stand-alone criterion for diagnosing PPG. Lu et al. stated that the best stand-alone diagnostic strategy for OCT RNFL data is to classify an eye as glaucomatous if the combination of the overall, inferior quadrant and superior quadrant RNFL thickness average is below normal [8]. In regard to RNFL loss vs. function relationship, Horn et al. found a clear association of the inferior and superior RNFL loss with the corresponding area of VF loss [10]. Hood et al. suggested that hemifield may be considered glaucomatous if one of the 5 clock-hour RNFL sectors (3 and 9 excluded) is below the normative database, i.e., at <1% area (red area of the OCT-3 output graph), or two are at <5% (yellow) [11]. In that regard, Caprioli et al. found that OCT is superior to both GDx-VCC and HRT II in its ability to detect RNFL loss in glaucoma [3]. The presented diagnostic interpretation is in accordance with these studies. However, several studies have shown that none of the new imaging instruments has a perfect diagnostic accuracy or is flawless. It is not unusual to find false positives or cases detected as abnormal by the imaging instruments and not confirmed by more conventional methods.
Fig. (2). (A) Top: The mean initial OCT RNFL examination of 5 scans is presented in clock-hours, quadrants, and graphically, in comparison with the OCT age-matched normative database software provided by the manufacturer (bound in yellow). The RNFL is thinner than the lower normal border in both the right (OD) and left eye (OS) superiorly, inferiorly and temporally. Bottom: Standard visual fields are normal. (B) Visual fields in September 2007. Cecocentral scotoma is evident bilaterally, more in the left eye. (C). Top: In May 2008, further thinning of the RNFL is detected in the inferior, superior and temporal quadrants of the right eye and in the superior quadrant of the left eye. Bottom: The cecocentral scotoma widened in the right eye and became non-specific and less distinct in the left eye.
We used the OCT-2 instrument in our study. Its reproducibility between examinations in perimetric glaucoma was found to be 11.8% [5]. The aforementioned PPG studies used OCT-3 or GDx-VCC [2, 3, 9, 16]. The output of RNFL thickness in OCT-3 is parallel to that of OCT-2, although RNFL thickness data in the OCT-2 are generally thicker [14]. We assume that further studies would quantitatively clarify (a) at which level a localized RNFL loss should be considered pathological and (b) the minimum required area, quadrants or sites of substantial RNFL thinning, including mean global loss, for designating the eye as glaucomatous.

In addition to disease progression, a clinical aid in diagnosing PPG may be the frequency-doubling technology or short-wavelength automated perimetry, although RNFL loss commonly precedes VF loss as detected by these perimetric methods [3, 16]. In the case of diffuse RNFL thinning, loss should be considered pathological and (b) the minimum quantitatively clarify (a) at which level a localized RNFL loss is clinically significant. We assume that further studies would clarify (a) at which level a localized RNFL loss should be considered pathological and (b) the minimum required area, quadrants or sites of substantial RNFL thinning, including mean global loss, for designating the eye as glaucomatous.

In summary, a revised interpretation of the data in the diagnosis of PPG is presented. That approach enabled, in retrospect, a diagnosis of PPG in an adult during the first visit, which was verified 6 years later. It was based on the literature findings that the pattern of RNFL-t loss at the typical glaucomatous sites was repeatedly found to reflect glaucoma, including its preperimetric stage. More importantly, the OCT affords an objective and accurate real-time structural state of the RNFL-t that indicates whether RNFL loss at a specific glaucomatous site has already occurred in the glaucoma suspect. Further studies are required to confirm the presented interpretation of RNFL data and to provide quantitative criteria for early diagnosis of PPG.

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