The effect of panretinal photocoagulation on confocal laser scanning ophthalmoscopy and stereo photographic parameters of optic disk topography in patients with diabetic retinopathy

Efeito da panfotocoagulação retiniana na oftalmoscopia confocal de varredura a laser e parâmetros da estereofotografia do disco óptico em pacientes com retinopatia diabética

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ABSTRACT | Purpose: To determine the effect of panretinal photocoagulation on optic disk topographic parameters in non-glaucomatous patients with proliferative diabetic retinopathy.

Methods: This was a prospective, single-center, observational study. Thirty-eight eyes of 26 patients with diabetes underwent panretinal photocoagulation for proliferative diabetic retinopathy. Stereoscopic disk photographs and optic nerve head parameters were evaluated using the Zeiss fundus camera and the confocal scanning laser ophthalmoscope (Heidelberg Retinal Tomograph), respectively, at baseline and 12 months after the completion of panretinal photocoagulation treatment (p=0.461 and 0.839, respectively). The global values of the optic nerve head parameters analyzed with the HRT3 showed no significant change from baseline to 12 months, including the disk area, cup area, rim area, cup volume, rim volume, C/D area ratio, linear C/D ratio, mean cup depth, maximum cup depth, cup shape measure, height variation contour, mean retinal nerve fiber layer thickness, and cross-sectional area. Conclusion: Our results suggest that panretinal photocoagulation does not cause morphological optic disk changes in patients with diabetic proliferative retinopathy after 1 year of follow-up.
Keywords: Light coagulation; Optic disk; Microscopy, confocal; Diabetic retinopathy; Scanning laser polarimetry

INTRODUCTION

Diabetic retinopathy (DR) is the most common ocular complication of diabetes mellitus (DM) and is one of the leading causes of blindness in developed countries(1). It is also known to be an important risk factor for chronic open-angle glaucoma, and both diseases often coexist.

In cases of proliferative diabetic retinopathy (PDR), panretinal photocoagulation (PRP) is the first-line treatment. Although PRP reduces the risk of severe vision loss(2), it laser energy has been shown to cause destruction to all layers of the retina, including the ganglion cells and the retinal nerve fiber layer (RNFL), and therefore generate visual field defects similar to those observed in glaucomatous damage(3). In such cases, visual field testing can be less helpful to evaluate glaucomatous damage in patients with PDR treated with PRP.

Pathological cupping of the optic disk is often associated with glaucoma, and it has been described in several other optic neuropathies, but rarely in retinal diseases(4). DR has been suggested to enhance the apoptosis of retinal ganglion cells, causing neurodegenerative changes in the retina(5). With the development of clinically available imaging technology, recent studies have shown that patients with diabetes, with or without DR, have thinner RNFL thickness than the normal population(6–8). Moreover, it has been suggested that ganglion cell loss secondary to PRP and ascending RNFL atrophy could change the appearance and topography of the optic disk(9). Therefore, evaluating the optic disk cupping and possible glaucomatous damage in patients with DR can be difficult, especially after PRP treatment.

In this scenario, confocal scanning laser ophthalmoscopy (CSLO), with a three-dimensional topographic analysis, can be a valuable diagnostic tool to assess the topographic changes of the optic disk, and obtain objective measurements of the rim area and cup parameters. Previous studies have shown conflicting results in both RNFL and optic disk topographic measurements after PRP treatment in patients with diabetes(10–11). The purpose of this study was to prospectively determine the effect of PRP on CSLO optic disk parameters and stereo photographic analysis in patients with non-glaucomatous PDR.

METHODS

This prospective observational cohort study enrolled patients from the retina service of the Ophthalmology Division of the University of São Paulo Medical School. The study protocol was approved by the local ethics committee and adhered to the tenets of the Declaration of Helsinki. Informed consent was obtained from all subjects.

The inclusion criteria for the study were a diagnosis of PDR (due to type 1 or 2 DM), intraocular pressure <18 mmHg, non-glaucomatous optic disk characteristics at fundus examination, a vertical cup-to-disc (C/D) ratio <0.7, and the absence of media opacities. Subjects were excluded from this study if they had a previous diagnosis of glaucoma or a family history of glaucoma, or any coexisting neuro-ophthalmic disease, uveitis, retinal artery or vein occlusion, optic disk neovascularization, diabetic macular edema (DME), corneal opacity, or previous laser photocoagulation treatment.

All participants underwent a complete ophthalmological examination at baseline, including best-corrected visual acuity with Snellen charts, Goldmann applanation tonometry, slit-lamp biomicroscopy of anterior and fundus segment using a 78D lens (Volk, Mentor, OH, USA), and indirect binocular ophthalmoscopy. Stereoscopic disk photographs and Heidelberg Retinal Tomograph (HRT3; Heidelberg Retinal Tomography, Heidelberg Engineering, Heidelberg, Germany) images were obtained at baseline and 12 months after the completion of PRP.

PRP treatment was carried out with a single spot green laser (Purepoint® laser system, 532-nm wavelength, Alcon, Fort Worth, TX, USA). All patients received at least 1,500 peripheral laser photocoagulation burns. The laser parameters used were a spot size of 250 µm, a pulse duration of 0.2 s, and enough power to cause grayish white burns following the Early Treatment Diabetic Retinopathy Study (ETDRS) guidelines. PRP was carried out in three sessions, with each session 1 week apart. Patients were examined after 6 weeks to evaluate the effect of laser treatment and the need for further laser sessions. Figure 1 exemplifies the pattern of laser photocoagulation applied in the study and the distance between the photocoagulation burns and the optic disk margins.

All stereoscopic disk photographs were taken with a Zeiss fundus camera (Carl Zeiss, Inc., Thornwood, NY, USA) and were centered on the optic disk. The horizontal and vertical diameters of the C/D ratio were determined for each stereo pair by an experienced glaucoma specialist. In order to assure a blinded analysis, all image files received a numeric code. Any laser burn observed in the picture was cropped out to ensure that the glaucoma specialist was blinded to the patient name and also the pre- or post-PRP status. A stereo viewer was used to enhance details.

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Optic nerve head (ONH) topography was analyzed with the HRT3 using confocal scanning parameters. A 15-degree angle view was used under the same intensity of dim room light. After generating a mean topographic image, contour lines were drawn by the same experienced technician. Subjects with standard deviations less than 30 µm were included in the study. The reference plane was automatically set at the standard value of 50 µm below the contour line at the temporal sector of the disk margin between 350° and 356°. Figure 2 is an example of a printout obtained from an eye using the HRT3, before laser treatment.

The following topographic parameters calculated by the HRT3 hardware were evaluated at baseline and 12 months after PRP completion: disk area (mm²), cup area (mm²), rim area (mm²), cup volume (mm³), rim volume (mm³), C/D area ratio, linear C/D ratio, mean cup depth (mm), maximum cup depth (mm), cup shape measure, height variation contour (mm), mean RNFL thickness (mm), and RNFL cross-sectional area (mm²). These variables were used for the topographic global analysis.

Statistical analyses were performed using a commercially available computer software package (SPSS, ver. 23.0; SPSS, Chicago, IL, USA). Normality was tested using the one-sample Kolmogorov-Smirnov test. The Wilcoxon signed-rank test was used to compare the stereoscopic pre- and post-PRP C/D ratios (horizontal and vertical) and HRT parameters. The statistically significant level was accepted as p<0.05.

RESULTS

This study included a total of 42 eyes of 30 patients. Four eyes of four patients were excluded during the follow-up visits (one patient developed preretinal membranes causing tractional detachment; another eye developed vitreous hemorrhage and both patients were submitted to pars plana vitrectomy; two eyes developed macular edema and were treated with intravitreal anti-vascular endothelial growth factor [VEGF] injections). Thirty-eight eyes of 26 individuals (15 female, 11 male) completed the 1-year follow-up. The mean age was 53.7 years, ranging from 26 to 74 years. Stereoscopic disk photographs resulted in a mean (± standard deviation) pretreatment horizontal C/D ratio of 0.316 ± 0.065 and a post-treatment horizontal C/D ratio of 0.303 ± 0.065. The difference between these means (-0.013) was not statistically significant (p=0.461). Similarly, the difference between the mean vertical C/D ratio in the pretreatment (0.363) and post-treatment (0.366) visits was also not statistically significant (p=0.839). These data are summarized in table 1.

The global values of the ONH parameters obtained from the HRT3 examination are presented in table 2. There was no statistically significant difference between the baseline and the 12-month HRT parameters. The cup area showed a non-significant (p=0.510) increase from the baseline (0.366 ± 0.216 mm²) to 12 months (0.376 ± 0.239 mm²). Similarly, there was an increase in the mean values of the maximum cup depth from the baseline (0.600 ± 0.244 mm) to the 12-month visit (0.636 ± 0.232 mm), without statistical significance (p=0.135). Rim area parameters, moreover, showed smaller values in the 12-month evaluation (1.638 ± 0.349 mm²) than at the baseline (1.649 ± 0.339 mm²) measurement, although this difference was also not statistically significant (p=0.451). The ratio between the cup area and disk area also showed a non-significant increase between examinations, from 0.178 ± 0.098 to 0.183 ± 0.108, with a p-value of 0.477 (Table 2).

DISCUSSION

PRP decreases the risk of severe visual loss in patients with PDR and, despite the evidence regarding the efficacy of antiangiogenic drugs, it is still the standard of care for the management of proliferative disease, according to the American Academy of Ophthalmology’s Preferred Practice Pattern for Diabetic Retinopathy[12], which is based on the Diabetic Retinopathy Study (DRS) and the
Figure 2. Example of a printout obtained from an eye using the Heidelberg Retinal Tomograph 3, before laser treatment.
ETDRS (level 1 evidence). While most visual complications in patients with PDR are related to retinal damage, it is not uncommon for patients to have glaucoma-associated visual loss, either because of disease unrelated to diabetes or because of intraocular pressure elevation from PRP or other treatment modalities, such as corticosteroid injections. Therefore, while the causal relationship between DM and open-angle glaucoma in many cases remains unclear, the diagnosis of chronic open-angle glaucoma in patients who have PDR can be challenging, particularly when submitted to PRP, as this type of treatment can produce visual field changes that may mimic glaucomatous field loss.

Ascending optic atrophy may assume several characteristic forms that can be distinguished both histopathologically and ophthalmoscopically. It occurs when a primary lesion involves the retina or optic nerve, producing an atrophic process that proceeds toward the brain. In experimental models, the distal aspect of a damaged axon was shown to degenerate 2-4 weeks after injury. In the pattern of atrophy and excavation of the ONH associated with glaucomatous damage, the loss of ganglion cell axonal fibers anterior to the lamina cribrosa gives rise to an increased C/D ratio. In general, this characteristic pattern of glaucomatous disk cupping is seen in association with elevated intraocular pressure but it has also been reported in a number of other conditions. In patients with diabetes submitted to PRP, the laser could damage photoreceptors that would secondarily affect ganglion cells, possibly producing increased cupping of the optical disk. Therefore, laser treatment could potentially induce ascending optic atrophy and mimic glaucomatous conditions. The 1-year interval between stereo photographs in our study was chosen because ample time would have elapsed to allow a possible atrophy to occur. On the other hand, an even longer interval would potentially introduce an aging change bias that could interfere with the interpretation of the C/D ratio.

Table 1. Cup-to-disc ratios before and after panretinal photocoagulation (n=38)

<table>
<thead>
<tr>
<th>Pretreatment C/D ratio</th>
<th>Post-treatment C/D ratio</th>
<th>Difference in C/D ratio</th>
<th>p*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean ± SD</td>
<td>Mean ± SD</td>
<td>Mean ± SD</td>
</tr>
<tr>
<td>Horizontal</td>
<td>0.316 ± 0.065</td>
<td>0.303 ± 0.065</td>
<td>-0.013 ± 0.026</td>
</tr>
<tr>
<td>Vertical</td>
<td>0.363 ± 0.064</td>
<td>0.366 ± 0.066</td>
<td>0.003 ± 0.028</td>
</tr>
</tbody>
</table>

C/D= cup-to-disc; SD= standard deviation.
* Wilcoxon signed-rank test.

Table 2. Comparison of HRT parameters before and after panretinal photocoagulation (n=38)

<table>
<thead>
<tr>
<th>HRT parameters</th>
<th>Pre-treatment</th>
<th>Post-treatment</th>
<th>P*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean ± SD</td>
<td>Range</td>
<td>Mean ± SD</td>
</tr>
<tr>
<td>DA</td>
<td>2.014 ± 0.387</td>
<td>1.30-2.75</td>
<td>2.014 ± 0.387</td>
</tr>
<tr>
<td>CA</td>
<td>0.366 ± 0.216</td>
<td>0.0-0.82</td>
<td>0.376 ± 0.239</td>
</tr>
<tr>
<td>RA</td>
<td>1.649 ± 0.339</td>
<td>1.15-2.36</td>
<td>1.638 ± 0.349</td>
</tr>
<tr>
<td>CV</td>
<td>0.084 ± 0.072</td>
<td>0.01-0.30</td>
<td>0.093 ± 0.080</td>
</tr>
<tr>
<td>RV</td>
<td>0.458 ± 0.145</td>
<td>0.25-0.90</td>
<td>0.458 ± 0.142</td>
</tr>
<tr>
<td>C/D area</td>
<td>0.178 ± 0.098</td>
<td>0-0.34</td>
<td>0.183 ± 0.108</td>
</tr>
<tr>
<td>LinCD</td>
<td>0.396 ± 0.144</td>
<td>0.02-0.58</td>
<td>0.401 ± 0.149</td>
</tr>
<tr>
<td>MCD</td>
<td>0.197 ± 0.092</td>
<td>0.04-0.36</td>
<td>0.188 ± 0.111</td>
</tr>
<tr>
<td>MsCD</td>
<td>0.600 ± 0.244</td>
<td>0.15-1.18</td>
<td>0.636 ± 0.232</td>
</tr>
<tr>
<td>CSM</td>
<td>-0.227 ± 0.085</td>
<td>-0.45 to -0.09</td>
<td>-0.220 ± 0.076</td>
</tr>
<tr>
<td>HVC</td>
<td>0.413 ± 0.098</td>
<td>0.24-0.64</td>
<td>0.419 ± 0.082</td>
</tr>
<tr>
<td>MRNFLFT</td>
<td>0.251 ± 0.070</td>
<td>0.07-0.42</td>
<td>0.251 ± 0.076</td>
</tr>
<tr>
<td>RNFLcsA</td>
<td>1.258 ± 0.368</td>
<td>0.35-1.90</td>
<td>1.256 ± 0.383</td>
</tr>
</tbody>
</table>

SD= standard deviation; DA= disc area; CA= cup area; RA= rim area; CV= cup volume; RV= rim volume; C/D= cup/disc; LinCD= linear C/D ratio; MCD= mean cup depth; MsCD= maximum cup depth; CSM= cup shape measure; HVC= height variation contour; MRNFLFT= mean retinal nerve fiber layer thickness; RNFLcsA= RNFL cross-sectional area.
* Wilcoxon signed-rank test.
with non-PDR have a thinner RNFL than healthy subjects\(^6,8,15,16\). This finding might be explained by the death of retinal ganglion cells, which occurs early in diabetic eyes due to enhanced apoptosis-promoting factors\(^7,17,18\). Amano et al.\(^18\) detected the presence of abnormal looking ONH in patients with diabetes and proposed that glycation end products might directly damage the optic nerve. However, Königreuther and Jonas\(^19\) analyzed the appearance of the optic disk in patients with and without diabetes using color photographs and found no difference between both these groups. Similar results were obtained by Tekeli et al.\(^20\) and Lim et al.\(^10\) using different imaging techniques.

Cross-sectional studies have indicated that PRP may cause morphologic changes in the ONH. Lim et al.\(^10\) compared diabetic eyes submitted or not to PRP using optical coherence tomography imaging. Optic nerves in eyes treated with PRP were more likely to be graded as abnormal, but the authors did not perform longitudinal comparisons and therefore disease severity could not be ruled out as a causative factor. Cancaya et al.\(^21\) found similar results using CSLO technology.

We could not identify topographic ONH changes using either CSLO or stereoscopic analysis after a 1-year follow-up of PRP for PDR. However, other longitudinal prospective studies comparing pre- and post-PRP ONH aspects have already been performed, with conflicting results. Using stereoscopic disk photographs, Johns et al.\(^9\) observed that the C/D ratio showed no significant change 1 year after PRP, but the authors reported increased optic disk pallor in the study period. On the other hand, Singh et al.\(^11\) found significant changes in CSLO parameters 6 months after PRP treatment. The results of our investigation are in conflict with this last study, since we have used the same technology to analyze the ONH morphology changes. One hypothesis to explain these inconsistent findings is that different laser delivery systems may alter the amount of ascending optic atrophy observed. While we used a spot size of 250 \(\mu m\) in our study, Singh et al. used a spot size of 300 \(\mu m\). A larger spot size may be related to greater inner retinal damage and therefore more ascending optic atrophy. This possible explanation is confirmed by the findings of Lee et al.\(^22\), which suggest that PASCAL photocoagulation, a new form of multiphoton laser treatment for DR, may not cause changes in optic disk morphology due to its shorter pulse duration (and consequently more restricted retinal damage) than conventional single spot PRP. Additionally, Kim and Cho\(^23\) measured RNFL thickness before and 6 months after PRP and concluded that laser photocoagulation of a moderate degree does not damage the RNFL.

A limitation of our investigation is its relatively small sample size, although the number of patients recruited was similar to other prospective interventional studies\(^9,11\). Moreover, we adopted rigid inclusion criteria that excluded patients with optic disk neovascularization and/or baseline DME. The presence of disk neovascularization does not allow an adequate analysis of ONH parameters by CSLO, because fibrovascular tissue can obscure the contour of the optic cup\(^8\). Moreover, DME is frequently present when PDR is detected in patients with diabetes, and often requires additional treatment with intravitreal anti-VEGF injections, which could somehow interfere with the parameters analyzed in our study. In order to allow proper HRT documentation of the optic disc and to avoid the bias of anti-VEGF injections, only patients with PDR without disc neovascularization and without macular edema were eligible for this particular study. We know from the DRS that only 40% of PDR cases do not present with disc neovascularization\(^2,24\).

The recently published protocol S\(^24\) from the DCRC network, a multicentric and randomized clinical trial that compared PRP or anti-VEGF for PDR cases, reported that around 22% of the participants with PDR had DME at baseline. Therefore, most PDR cases presented at our service were not eligible for this study. The results of protocol S\(^24\) showed that both therapies produced similar results in preventing complications of DR and visual acuity after 5 years of follow-up, but the group that received anti-VEGF intravitreal injections had lower rates of developing vision-impairing DME, and less visual field loss when compared to patients receiving PRP. However, the costs and frequency of visits are significantly higher and low adherence to treatment is an issue because it requires more constant follow-up.

In clinical practice, it is common to see many patients who were treated with PRP in the past with severe optic disc atrophy, detected several years after treatment. However, such damage can often be attributed to the higher intensity of the lasers used in the past. In contrast, data from our study show that after performing photocoagulation with moderate intensity power, PRP did not cause significant changes in the ONH morphology of patients with PDR after 1 year of follow-up.

REFERENCES


