Panretinal photoocoagulation versus intravitreal injection retreatment pain in high-risk proliferative diabetic retinopathy

Dor em panfotocogulação retiniana versus injeção intravítrea em pacientes com retinopatia diabética proliferativa de alto risco

CELIA REGINA FARIAS DE ARAÚJO LUCENA1, JOSÉ AFONSO RAMOS FILHO1, ANDRÉ MÁRCIO VIEIRA MESSIAS1, JOSÉ APARECIDO DA SILVA2, FELIPE PACENTINI PAES DE ALMEIDA4, INGRID URSULA SCOTT3, JEFFERSON AUGUSTO SANTANA RIBEIRO1,2, RODRIGO JORGE1

ABSTRACT

Purpose: To compare pain related to intravitreal injection and panretinal photoocoagulation in the management of patients with high-risk proliferative diabetic retinopathy.

Methods: Prospective study including patients with high-risk proliferative diabetic retinopathy and no prior laser treatment randomly assigned to receive panretinal photocoagulation (PRP) group or panretinal photoocoagulation plus intravitreal ranibizumab (PRPplus) group. In all patients, panretinal photoocoagulation was administered in two sessions (weeks 0 and 2), and intravitreal ranibizumab was administered at the end of the first laser session in the PRPplus group. Retreatment was performed at weeks 16 and 32 if active new vessels were detected at fluorescein angiography. Patients in the PRPplus group received intravitreal ranibizumab and patients in the PRP group received 500-μm additional spots per quadrant of active new vessels. After the end of retreatment, a 100-degree Visual Analog Scale was used for pain score estimation. The patient was asked about the intensity of pain during the whole procedure (retinal photoocoagulation session or intravitreal ranibizumab injection). Statistics for pain score comparison were performed using a non-parametric test (Wilcoxon rank sums).

Results: Seventeen patients from PRPplus and 14 from PRP group were evaluated for pain scores. There were no significant differences between both groups regarding gender, glycated hemoglobin and disease duration. Mean intravitreal injection pain (±SEM) was 4.7 ± 2.1 and was significantly lower (p<0.0001) than mean panretinal photoocoagulation pain (60.8 ± 7.8). Twelve out of 17 patients from the PRPplus group referred intensity pain score of zero, while the minimal score found in PRP group was found in one patient with 10.5.

Conclusion: In patients with high-risk proliferative diabetic retinopathy who needed retreatment for persistent new vessels, there was more comfort for the patient when retreatment was performed with an intravitreal injection in comparison with retinal photoocoagulation. Further larger studies are necessary to confirm our preliminary findings.

Keywords: Pain; Intravitreal injections; Diabetic retinopathy; Light coagulation; Vascular endothelial growth factor A

RESUMO

Objetivo: Comparar a dor relacionada à injeção intravítrea e panfotocoagulação no tratamento de pacientes com retinopatia diabética proliferativa de alto risco.

Métodos: Estudo prospectivo incluindo pacientes com retinopatia diabética proliferativa de alto risco e nenhum tratamento a laser prévio aleatoriamente designados para receber panfotocoagulação retiniana (grupo PRP) ou panfotocoagulação e ranibizumabe intravítreo (grupo PRPplus). Em todos os pacientes, a panfotocoagulação foi administrada em duas sessões (semanas 0 e 2), e ranibizumabe intravítreo foi administrado no final da primeira sessão de laser no grupo PRPplus. Retratamento foi realizado nas semanas 16 e 32 se neovasos ativos fossem detectados na angiografia de fluoresceína, utilizando ranibizumabe intravítreo no grupo PRPplus e laser adicional grupo PRP. Após o fim do retratamento, uma Escala Analógica Visual de 100-unidades foi utilizada para a estimativa da pontuação da dor. O paciente foi questionado sobre a intensidade da dor durante todo o procedimento (sessão de fotocoagulação de retina ou injeção intravítrea de ranibizumabe). A comparação dos índices de dor foi realizada utilizando um teste não-paramétrico (Wilcoxon rank sums).

Resultados: Dezessete pacientes do grupo PRPplus e 14 do grupo PRP foram avaliados para os índices de dor. Não houve diferenças significativas entre os dois grupos quanto ao sexo, hemoglobina glicosilada e duração da doença. A média de dor da injeção intravitéreo (±SEM) foi 4,7 ± 2,1, significativamente menor (p<0,0001) do que a dor média da panfotocoagulação (60,8 ± 7,8). Doze dos 17 pacientes do grupo PRPplus referiram pontuação de intensidade da dor zero, enquanto que o índice mínimo no grupo PRP foi encontrado em um paciente com 10,5.

Conclusão: Em pacientes com retinopatia diabética proliferativa de alto risco que necessitaram de retratamento por neovasos persistentes, houve mais conforto para o paciente quando o retratamento foi realizado com uma injeção intravítrea em comparação com fotocoagulação da retina. Estudos posteriores são necessárias para confirmar nossos achados preliminares.

Descritores: Dor; Injeções intravítreas; Retinopatia diabética; Fotocoagulação; Fator A de crescimento endotélio vascular
INTRODUCTION

Retinal new vessels (NV) represent an important risk factor for severe vision loss in patients with diabetes mellitus (1,2). About 60% of patients with proliferative diabetic retinopathy (PDR) respond to panretinal photocoagulation (PRP) with regression of neovascularization within 3 months (3). However, many patients require additional laser treatment, and 4.5% ultimately require pars plana vitrectomy despite PRP (4).

Besides additional laser photocoagulation, intravitreal injection of anti-vascular endothelial growth factor (VEGF) agents have become an interesting alternative for new vessels regression (5,6). However, both procedures (PRP and injection) may induce discomfort and pain. Pain regarding PRP treatment is a well-recognized concern and may even prevent its completion (7), while pain regarding intravitreal injection may disturb patient compliance and peribulbar block may be necessary for uncooperative patients (8). Since intravitreal injection has become a trendy alternative therapeutic strategy and PRP is the standard of care for high-risk PDR, we decided to compare pain related to both procedures in the management of patients with PDR.

METHODS

The study protocol adhered to the tenets of the Declaration of Helsinki and was approved by the local Institutional Review Board, and all patients gave written informed consent before entering the study.

This is a prospective, open label, randomized study in which two groups of diabetic patients were followed (9). Between February 2009 and December 2009, all patients evaluated at the Retina and Vitreous Section of the Department of Ophthalmology, School of Medicine of Ribeirão Preto, who presented with high-risk PDR and had not received any prior retinal laser treatment were invited to participate in the study.

Patients were included if they had high-risk PDR, according to Early Treatment Diabetic Retinopathy Study (ETDRS) guidelines (10), as follows: 1) presence of NV at the disc (NVD) greater than ETDRS standard photograph 10A; or, 2) presence of NVD associated with vitreous or pre-retinal hemorrhage; or, 3) NV elsewhere (NVE) covering more than a half disc area associated with vitreous or pre-retinal hemorrhage. Exclusion criteria included: 1) history of prior laser treatment or vitrectomy in the study eye; 2) history of thromboembolic event (including myocardial infarction or cerebral vascular accident); 3) major surgery within the prior 6 months or planned within the next 28 days; 4) uncontrolled hypertension (according to guidelines of the seventh report of the Joint National Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure [JNC-7]) (11); 5) known coagulation abnormalities or current use of anticoagulation medication other than aspirin; or 6) any condition affecting documentation or follow-up.

During the study enrollment period, high-risk PDR was identified in one eye of 5 patients and in both eyes of 35 patients based on clinical examination and confirmed on fluorescein angiography. At baseline, each patient received a detailed ophthalmologic examination including measurement of the logarithm of the minimum angle of resolution (logMAR) ETDRS and best corrected visual acuity (BCVA) according to a standardized refraction protocol (using modified ETDRS charts 1, 2, and R), as well as applanation tonometry, dilated slit lamp biomicroscopic examination (including grading of lenticular opacities using the Lens Opacities Classification System III) (12), and binocular indirect fundoscopic examination. Digital red free fundus photography and fluorescein angiography were performed using two fundus camera systems (HRA-OCT, Heidelberg, Germany/TRC-501A/ IMAGEnet; Topcon, Tokyo, Japan).

All patients received PRP, which was performed in two sessions (at week 0 and week 2) according to ETDRS guidelines (13,14). Six hundred to eight hundred 500µm spots were performed per session, at the discretion of the treating investigator. If patients had clinically significant macular edema (15) macular focal/grid laser was performed during the first PRP session. Patients could be retreated with focal/grid laser at the 16 and 32-week study visits. If both eyes were eligible for the study, the eye with best visual acuity was included.

Patients were enrolled in groups of two. The technician was asked to pick up one of two identical opaque envelopes, one containing the designation for PRP whereas the other contained the designation for PRP/intravitreal ranibizumab injection (IVR) treatment. The second patient was automatically assigned with the second envelope. For the 20 eyes selected to receive the combined treatment (PRP plus intravitreal ranibizumab), one intravitreal injection of 0.5 mg (0.05 ml) of ranibizumab was performed approximately 60 minutes after the completion of the first PRP session (week 0) by a single retinal specialist. Ranibizumab was injected into the vitreous cavity via a 29 gauge needle inserted through the inferotemporal pars plana 3.0 - 3.5 mm posterior to the limbus using topical proparacaine drops under sterile conditions (eyelid speculum and povideone-iodine) (16). Patients were instructed to instill one drop of 0.3% ciprofloxacin into the injected eye four times daily for 1 week after the procedure.

Retreatment was performed at weeks 16 and 32 if active new vessels were detected at fluorescein angiography. Patients in the PRP group received IVR and patients in the PRP group received 500-µm additional spots per quadrant of active new vessels.

Fifteen minutes after the end of retreatment (PRP session or retreatment IVR injection), a masked examiner used a 100-degree Visual Analog Scale (VAS) for pain score estimation (17). The numbers of the scale were visible only on the examiner’s side, so that patients could not choose the same number to guide pain scores. Prior to rating level of pain, each patient was asked to slide the marker along the entire scale, with the aid of the examiner. At point 0, the examiner clarified to the patient that this point of the scale represented “no pain at all”, at point 100, the examiner clarified to the patient that this point of the scale represented “the most intense pain one could ever feel”. The patient was asked about the intensity of pain during the whole procedure (PRP session or IVR injection). Statistics for pain score comparison were performed using a non-parametric test (Wilcoxon rank sums).

RESULTS

Seventeen patients from the PRP plus patients and 16 patients from the PRP groups completed the 16-week visit, but 17 patients from PRP and 14 from PRP were evaluated for pain scores. Patients’ demographics are summarized in table 1. Mean age of PRP patients was significantly higher than PRP plus patients; there were no significant differences between both groups regarding gender, glycosylated hemoglobin (HbA1c) and disease duration. Mean ± SD age (years) was 63.5 ± 8.9 and 51.1 ± 11.3 (p=0.0018); mean ± SD HbA1c (%) was 9.3 ± 1.1 and 9.1 ± 0.8 (p=0.5391); and mean ± SD disease duration (years) was 12.9 ± 8.8 and 14.7 ± 6.9 (p=0.5326) for PRP and PRP plus respectively. Mean IVR pain (±SEM) was 4.7 ± 2.1 and was significantly lower (p<0.0001) than mean PRP pain (60.8 ± 7.8). Twelve out of 17 patients from the PRP plus group referred intensity pain score of zero, meaning that they had no pain at all during intravitreal injection, while the minimal score found in PRP group was found in one patient with 10.5, but no other patient in this group had score under 30. Figure 1 shows the distribution of intensity pain scores in both groups; there was a statistically significant difference between groups (P<0.0001; Wilcoxon).

DISCUSSION

Despite the difference in age between groups, that would have favored lower scores in PRP group (since older patients trend to feel less pain) (16), the present results suggest that an intravitreal injection
is less painful than a PRP session for diabetic patients with high-risk PDR. In fact, pain score associated with intravitreal injections of anti-VEGF agents in patients with diabetic retinopathy and neovascular age-related macular degeneration (ARMD) ranges from 0 to 22[7], while pain level associated with a PRP session ranges from 10 to 90[8]. This difference may be explained by the longer pain stimulus during a PRP session. Another hypothesis is that there may be direct stimulation of the long ciliary nerves during a PRP session (especially if laser applied in the retina periphery at 3 and 9 o'clock meridians) usually merely under topical anesthesia for PRP contact lens positioning, while during intravitreal injections there would be needle conjunctival and uveal nociceptors stimulation, usually partially blocked by topical anesthesia[7].

**CONCLUSION**

In addition to the beneficial effects regarding BCVA and new vessels regression of PRPplus ranibizumab in comparison to PRP alone for high-risk PDR treatment reported elsewhere, there is also more comfort for the patient when retreatment is performed with an intravitreal injection. Further larger studies are necessary to confirm our preliminary findings.

**REFERENCES**