Comparison of one and three initial monthly intravitreal ranibizumab injection in patients with macular edema secondary to branch retinal vein occlusion

Alper Halil Bayat, Akin Çakır, Seyma Gülçenur Özturan, Selim Bölükbaşı, Burak Erden, Mustafa Nuri Elçioğlu

Department of Ophthalmology, Okmeydani Research & Training Hospital, Istanbul 34384, Turkey

Correspondence to: Alper Halil Bayat. Department of Ophthalmology, Okmeydani Research & Training Hospital, Darulaceze Street No:25 Sisli, Istanbul 34384, Turkey. alperhalil76@hotmail.com

Received: 2017-10-11        Accepted: 2018-04-23

Abstract
● AIM: To compare three initial monthly intravitreal ranibizumab (IVR) injections followed by pro re nata (PRN) dosing with one initial monthly IVR injections followed by PRN dosing for macular edema (ME) secondary to branch retinal vein occlusion (BRVO).
● METHODS: Forty-two eyes of 42 patients who had IVR injections for BRVO were retrospectively studied. Eighteen eyes received 1 initial IVR injection (1+PRN group) and 24 eyes received 3 monthly IVR injections (3+PRN). At 1, 3, 6 and 12mo; spectral-domain optical coherence tomography (SD-OCT) was performed. Central macular thickness (CMT), the integrity of the external limiting membrane (ELM), the presence of subretinal fluid, cyst size, the presence of inner segment/outer segment (IS/OS) defect were determined.
● RESULTS: At baseline the mean CMT was 521.3±153.2 µm in the 3+PRN group while it was 438.1±162.4 µm in 1+PRN group. At the final visit, mean CMT was 278.3±87.8 µm in the 3+PRN group and 285.2±74.2 µm in the 1+PRN group (P=0.079). The changes in CMT over the entire study period were also comparable in both groups (243±160 µm in the 3+PRN group, and 152.9±175.3 µm in the 1+PRN group; P=0.090). At baseline, best-corrected visual acuity (BCVA) was 0.92±0.60 logarithm of the minimal angle of resolution (logMAR) in the 3+PRN group, while it was 0.72±0.46 logMAR in the 1+PRN group. Final BCVA was 0.42±0.55 logMAR in the 3+PRN group and 0.32±0.50 logMAR in the 1+PRN group (P=0.979). Additionally, the BCVA changes from baseline to final visit were not significantly different (-0.50±0.45 logMAR in the 3+PRN group, and -0.33±0.39 logMAR in the 1+PRN group; P=0.255).
● CONCLUSION: No significant differences in the anatomical or functional results are found between 3+PRN and 1+PRN regimens in the patients receiving ranibizumab for ME secondary to BRVO. Intact IS/OS and baseline BCVA are good predictor of the visual gain, while baseline CMT is a good predictor of the anatomical gain.
● KEYWORDS: branch retinal vein occlusion; ranibizumab; macular edema; therapy; predictive factors

DOI:10.18240/ijo.2018.09.17

INTRODUCTION

Retinal vein occlusion (RVO) is the second most frequent retinal vascular disorder after diabetic retinopathy. RVO is divided into central retinal vein occlusion (CRVO) and branch retinal vein occlusion (BRVO). Vascular compression during arteriovenous passages, degenerative changes in venous walls and hypercoagulability are the underlying pathophysiology of BRVO. Retinal ischemia after vascular occlusion can cause an increase the amount of vascular endothelial growth factor (VEGF) which can increase vascular permeability and cause macular edema (ME)\cite{1,2}. ME is the main cause of vision loss\cite{3,4}. There are some treatment modalities for ME such as intravitreal dexamethasone implants, laser treatment, and intravitreal injections of anti-VEGF agents\cite{9-11}.

Ranibizumab 0.5 mg which is an anti-VEGF agent was approved in June 2010 for the treatment of ME due to BRVO and CRVO in the United States, based on the 6mo-results of two phase III, randomized, double-masked, 12mo, controlled study-BRAVO. In this study, six sequential monthly intravitreal ranibizumab (IVR) injections followed by pro re nata (PRN) regimen improved best-corrected visual acuity (BCVA) compared with imitation\cite{12,13}. Yet the requirement of multiple intravitreal anti-VEGF agent injections for initial treatment of ME after BRVO is not well figured out and they may increase the risk of systemic or ocular complications\cite{14-15}.
In the real-life clinical practice, decreasing the number of anti-VEGF injection of the initial phase of treatment might be effective. In our study, we treated ME due to BRVO with two groups: 1 and 3 monthly initial IVR injections followed by PRN regimen. We compared them with central macular thickness (CMT), BCVA changes from pre-injection to final visit. There are not enough real-life studies about anti-VEGF treatment in ME due to BRVO. So we aimed to compare PRN treatment outcomes following three consecutive doses and single dose treatment in our clinic.

**SUBJECTS AND METHODS**

**Patients** Forty-two eyes of 42 patients (27 men and 15 women) with treatment-naive acute ME due to BRVO were studied. Symptom durations of the patients were less than 2mo before the examination. All of the patients who had intravitreal injections of ranibizumab (0.5 mg/0.05 mL) at Okmeydani Research & Training Hospital between June 2014 and December 2016 were retrospectively studied. Patients with CMT>300 μm in optical coherence tomography (OCT) were treated. Twenty four eyes received 3 monthly IVR injections (3+PRN group) and 18 eyes received one initial IVR injections (1+PRN group). The patients who have minimum 12mo follow up period were included in this study. None of the patients had macular grid laser photocoagulation. It is known that RVO leads retinal non-perfusion. Peripheral retinal non-perfusion (PRNP) was found laser photocoagulation. It is known that RVO leads retinal non-perfusion. Peripheral retinal non-perfusion (PRNP) was found to be effective. In our study, we treated ME due to BRVO with two groups: 1 and 3 monthly initial IVR injections followed by PRN regimen. We compared them with central macular thickness (CMT), BCVA changes from pre-injection to final visit. There are not enough real-life studies about anti-VEGF treatment in ME due to BRVO. So we aimed to compare PRN treatment outcomes following three consecutive doses and single dose treatment in our clinic.

**Study Design**

We retrospectively studied. Patients with CMT>300 μm in optical coherence tomography (OCT) were treated. Twenty four eyes received 3 monthly IVR injections (3+PRN group) and 18 eyes received one initial IVR injections (1+PRN group). The patients who have minimum 12mo follow up period were included in this study. None of the patients had macular grid laser photocoagulation. It is known that RVO leads retinal non-perfusion. Peripheral retinal non-perfusion (PRNP) was found to be effective. In our study, we treated ME due to BRVO with two groups: 1 and 3 monthly initial IVR injections followed by PRN regimen. We compared them with central macular thickness (CMT), BCVA changes from pre-injection to final visit. There are not enough real-life studies about anti-VEGF treatment in ME due to BRVO. So we aimed to compare PRN treatment outcomes following three consecutive doses and single dose treatment in our clinic.

**Statistical Analysis**

Statistical analyses were performed using the SPSS software version 15. Descriptive analyses were presented using means and standard deviations for normally distributed variables. Student’s t-test and Mann-Whitney U test were used to compare the parameters between the groups. When investigating the effect of the two different treatment regimens on the change in BCVA and CMT over time, repeated measures analysis of variance (ANOVA) was used. When investigating the visual and anatomical gain by different regimens (1+PRN and 3+PRN) the effect of all predictors was assessed and analyzed with SD-OCT by two experienced ophthalmologists. The average value of each parameter was considered for statistical analyzes.

**RESULTS**

Forty-two eyes of 42 patients with ME due to BRVO were examined in this retrospective cohort study. All of the patients have a minimum 12mo follow-up time. Twenty-four eyes received 3 monthly IVR injections (3+PRN group), 18 eyes received 1 initial IVR injection (1+PRN group). Both two groups were similar in terms of age, gender and the duration of the interval between the onset of the RVO and initiation of the treatment (P=0.601, P=0.783, P=0.169 respectively). Only one (4.2%) patient in the 3+PRN group showed a statistically significant improvement in BCVA.

**Table 1 Baseline characteristic of the patients**

<table>
<thead>
<tr>
<th>Parameters</th>
<th>3+PRN group (n=24)</th>
<th>1+PRN group (n=18)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (y)</td>
<td>60.2±10.2</td>
<td>58.6±8.9</td>
<td>0.601</td>
</tr>
<tr>
<td>Gender, male</td>
<td>15 (62.5)</td>
<td>12 (66.7)</td>
<td>0.783</td>
</tr>
<tr>
<td>Follow-up period (mo)</td>
<td>14.5±2.5</td>
<td>14.8±2.6</td>
<td>0.980</td>
</tr>
<tr>
<td>No. of injections</td>
<td>4.2±1.3</td>
<td>2.8±1.6</td>
<td>0.004*</td>
</tr>
<tr>
<td>Cataract development</td>
<td>1 (4.2)</td>
<td>0 (0.0)</td>
<td>1.00</td>
</tr>
<tr>
<td>Anti-glucocorticron therapy</td>
<td>1 (4.2)</td>
<td>0 (0.0)</td>
<td>1.00</td>
</tr>
<tr>
<td>Argon laser scatter photocoagulation</td>
<td>7 (29.2)</td>
<td>5 (27.8)</td>
<td>0.921</td>
</tr>
<tr>
<td>Baseline CMT (µm)</td>
<td>521.3±153.2</td>
<td>438.1±162.4</td>
<td>0.098</td>
</tr>
<tr>
<td>Final CMT (µm)</td>
<td>278.3±87.8</td>
<td>285.2±74.2</td>
<td>0.079</td>
</tr>
<tr>
<td>Baseline BCVA (logMAR)</td>
<td>0.92±0.60</td>
<td>0.72±0.46</td>
<td>0.354</td>
</tr>
<tr>
<td>Final BCVA (logMAR)</td>
<td>0.42±0.55</td>
<td>0.38±0.50</td>
<td>0.979</td>
</tr>
<tr>
<td>Duration of symptoms (d)</td>
<td>23.79±7.15</td>
<td>20.78±6.54</td>
<td>0.169</td>
</tr>
<tr>
<td>Subretinal fluid</td>
<td>9 (37.5)</td>
<td>5 (27.7)</td>
<td>0.508</td>
</tr>
<tr>
<td>Integrity of ELM (yes/no)</td>
<td>13/11</td>
<td>11/7</td>
<td>0.327</td>
</tr>
<tr>
<td>Integrity of IS/OS (yes/no)</td>
<td>15/9</td>
<td>9/9</td>
<td>0.418</td>
</tr>
</tbody>
</table>

CMT: Central macular thickness; BCVA: Best-corrected visual acuity; ELM: External limiting membrane; IS/OS: Inner segment/outer segment. *Statistically significant.
Final CMT was 278.3±87.8 µm in the 3+PRN group and 285.2±74.2 µm in the 1+PRN group (P=0.079). Additionally, the CMT changes from baseline to final visit were not significantly different (243±160 µm in the 3+PRN group, and 152.9±175.3 µm in the 1+PRN group; P=0.090). The trend in CMT changes over time was also similar in both groups (Figure 1). When the change in CMT was investigated, initial CMT was adjusted in one-way ANCOVA for the confounding effect and there was not a statically significant difference between groups (P=0.585). Final BCVA was 0.42±0.55 logMAR in the 3+PRN group and 0.38±0.50 logMAR in the 1+PRN group (P=0.979). Additionally, the BCVA changes from baseline to final visit were not significantly different (-0.50±0.45 logMAR in the 3+PRN group, and -0.33±0.39 logMAR in the 1+PRN group; P=0.255). The trend in BCVA changes over time was also similar in both groups (Figure 2). When the change in BCVA was investigated, initial CMT was adjusted in one-way ANOVA for the confounding effect and there was not a statically significant difference between groups (P=0.693).

We investigated by using simple regression analysis whether argon laser scatter photocoagulation to PRNP had an effect at final functional and anatomical outcomes. We found that it did not have any effect (P=0.552 and P=0.685). Argon laser scatter photocoagulation did not effect also number of injections (P=0.193).

The predictor factors of the visual and anatomical gain were also investigated in the entire study group. When investigating the changes in BCVA and CMT by different regimens (1+PRN and 3+PRN) the effect of all predictors were investigated by using both simple and multiple regression analysis (with enter method). The results of regression analysis were shown in Tables 2, 3. Multiple linear regression analysis with enter method revealed that baseline CMT was the most valuable predictive factor for the change in CMT (Beta coefficient=0.938, P<0.001). In addition, baseline BCVA was found to be the most important predictive factor for the change in BCVA (Beta coefficient=-0.351, P=0.003). Integrity of IS/OS was found to correlate with visual and anatomical gain with simple regression analysis (Beta coefficient=-0.286, P=0.034 and Beta coefficient=176.681, P<0.001 respectively).

DISCUSSION

In this study, the patients initially received one or three monthly IVR for ME. At 6mo and final visit, there was no significant difference in CMT or BCVA change from baseline between the 2 regimen groups. We believe that these two treatment regimens may achieve similar results. We attributed this finding to the difference in intravitreal VEGF levels of the patients. Considering the natural pathophysiology of BRVO, ischemia induces VEGF secretion. Since the amount of ischemia is not standard in all the patients, the VEGF levels will be variable. In current literature, there is only one study comparing the single injections with 3 monthly injections of IVR in the treatment of ME due to BRVO. Miwa et al.[17] confirmed that there is no significant difference between one and three monthly IVR injections as in our study.
There is one study which compares one or three monthly injections with bevacizumab. Ito et al [18] reported that the single injection of bevacizumab’s and three monthly injections results were similar. There were not any significant differences in visual outcomes and CMT findings.

In the BRAVO trial, patients received six monthly IVR in the first 6mo. According to their retreatment criteria, additional injections were performed in the follow-up (6+PRN regimen) [12-13,19].

Surely, it is difficult to compare the BRAVO trial and the current study due to different inclusion criteria. But the real-life studies are overgrowing every single day in the literature which reveal different results compared to randomized clinical trials.

In a recent study, Osaka et al [20] studied patients with ME secondary to CRVO. Twenty nine eyes of the patients were treated with 3+PRN regimen, while 20 eyes were treated 1+PRN regimen. They followed the patients for twelve month. At final visit, they reported that 1+PRN regimen achieved visual outcomes similar to those of 3+PRN regimen with fewer injections.

In brief, we believe these three treatment regimens (1+PRN, 3+PRN, 6+PRN) can achieve similar results, especially in good responding patients to the first anti-VEGF injection. We hypothesis that the response to first anti-VEGF injection helps on forecasting the VEGF concentrations as well the prognosis of the patients.

We also investigated the predictor factors of the change in BCVA and CMT in the entire group. We found that baseline CMT was significantly associated with the anatomical gain. Similar to our results, Ach et al [21] reported that the initial CMT was a predictive factor for short and long-term responses to anti-VEGF treatment.

The multiple regression analyses showed that the pretreatment BCVA had the highest correlation with visual gain. Additional, the statistical analysis showed that the BCVA improvement at 12-month follow-up was better in the eyes with intact photoreceptor IS/OS layer. Similar to our conclusion, Shin et al [22] also found that integrity of IS/OS is significantly correlated with visual gain.

In the WAVE [16] and RELATE [23] studies, the researcher wanted to investigate the effect of peripheral laser application on treatment burden and visual outcomes. But these two studies were different in some aspects. First, in WAVE study peripheral laser application was performed after first injections, while in RELATE study it was done after two consecutive injections. Secondly, inclusion criteria were different between those the trials. In WAVE study, patients had ischemic RVO and ME poorly responsive to anti-VEGF and they investigated whether those patients could benefit after argon laser scatter photocoagulation. But in RELATE study, at baseline 41% of patients were treatment naive and perfusion status was not considered as an inclusion criteria. Despite these differences both studies concluded that peripheral laser photoocoagulation did not effect number of injections and visual outcomes in RVO patients under ongoing anti-VEGF treatment. In our study we performed argon laser scatter photocoagulation after first injections similar to WAVE study. But our patients were treatment naive and perfusion status was not an inclusion criteria similar to RELATE study. Although our study was different from these studies and had fewer patients we found that argon laser scatter photoocoagulation did not affect treatment burden and visual outcomes in BRVO patients with ongoing anti-VEGF treatment.

In conclusion, our results suggest that both 3+PRN IVR and 1+PRN IVR regimens may achieve similar results in one-year follow-up. Another result is that baseline SD-OCT characteristics can be helpful in predicting the final visual outcome after IVR injection in patients with ME secondary to BRVO. Especially the integrity of IS/OS and initial BCVA are significantly correlated with visual gain. And finally, peripheral photocoagulation did not impact treatment burden, visual and anatomical outcomes in patients with ME due to BRVO.

ACKNOWLEDGEMENTS

Conflicts of Interest: Bayat AH, None; Çakır A, None; Özturan ŞG, None; Bölükbaşı S, None; Erden B, None; Elçioğlu MN, None.

REFERENCES


5. Shin et al [24].
One versus three initial ranibizumab injections in BRVO


