Management of macular edema secondary to branch retinal vein occlusion-combined treatment with intravitreal bevacizumab and ozurdex implant

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A B S T R A C T

Purpose: To evaluate the efficacy and safety of a dexamethasone implant in combination with intravitreal bevacizumab injection in the management of macular edema secondary to branch retinal venous occlusion.

Materials and Methods: 10 eyes were prospectively investigated. Each eye was treated with intravitreal bevacizumab followed by intravitreal ozurdex at a two-week interval. Recurrence of macular edema was treated with ozurdex only. Patient were evaluated preoperatively with BCV A, IOP, OCT and fundus evaluation and followed up at 15 days, 2 months and 4 months of ozurdex injection. Follow up all patient for 1 year duration.

Results: Mean BCVA at presentation was 0.81 log mar, and that improved to after 15 days of intravitreal bevacizumab was 0.55 log mar. At 15 days, 2 months and 4 months of intravitreal ozurdex implantation mean BCVA was 0.54 log mar, 0.54 log mar and 0.6 log mar respectively. Mean central foveal retinal thickness at presentation was 538 μm and reduced to 235 μm, after intravitreal bevacizumab. At 15 days, 2 months and 4 months after intravitreal ozurdex implantation mean central foveal retinal thickness was 182.14 μm, 189.28 μm and 352.14 μm respectively. Mean Intraocular pressure (IOP) was elevated about 1 mm of Hg after bevacizumab, about 3 mm of Hg after 4 months of ozurdex with using single anti glaucoma medication and one patient had progression of cataract after 2nd injection of ozurdex.

Conclusion: Combined treatment of intravitreal bevacizumab and ozurdex shows better and a sustained functional outcome. Increased intraocular pressure and cataract formation can be potential concerns and should be monitored.

1. Introduction

Retinal vein occlusion (RVO) is a common vascular disorder of the retina second to diabetic retinopathy. Branched retinal vein occlusion (BRVO) is a most common form of retinal vein occlusion. Macular edema is a main cause for visual loss in BRVO. Other causes of visual decrease in BRVO is retinal or optic disc neovascularization and neovascular glaucoma.

The pathogenesis of macular edema in RVO have been identified to be due to factors such as the hydrostatic effects from increased venous pressure, the presence of inflammatory cytokines (e.g., prostaglandins and interleukin-6), and the up regulation of endothelial tight junction proteins,1 and /or increased vascular endothelial growth factor (VEGF) expression.2 Common risk factors for RVO include arterial hypertension, hypercholesterolemia, diabetes mellitus, and glaucoma.3

Optical coherence tomography (OCT) is an important tool in the diagnosis of macular edema (ME).

Grid laser photocoagulation was shown to be benefit for treating ME in the branch vein occlusion study (BVOS) with limited in visual improvement.4

Anti-vascular endothelial growth factor (anti-VEGF) introduction has revolutionised the management of ME secondary to RVO provide short term visual benefit.
Recurrent ME is commonly encountered problem with intravitreal monotherapy, leading to the need of multiple injection. Few patients seems to develop resistant ME and can even develop increased or rebound ME after anti-VEGF therapy.\textsuperscript{5-7} Since the intravitreal dexamethasone implant (Ozurdex; Allergan, Irvine, CA) was recently approved for RVO,\textsuperscript{8} we used this treatment as combination with anti-VEGF.

We aim to study the combination therapy in our study.

2. Materials and Methods

Our study is prospective, consecutive, non-randomized case series, 10 eyes of 9 patients with BRVO with ME, 6 males and 3 females (mean age 59.28 years) with maximum duration of symptoms of 4 months were included. 6 eyes presented with superior temporal quadrant BRVO and 4 eyes presented with inferior temporal quadrant BRVO. We excluded patients with a known history of glaucoma or steroid response, as well as history of vitrectomy and neovascularization in the anterior or posterior segment.

Informed consent was procured prior to recruitment in a consecutive manner. Each patient underwent detail eye examination include best corrected visual acuity (BCVA), anterior segment examination, intraocular pressure (IOP) and fundus examination. OCT (Spectral domain – primus Zeiss) scan was done each visit. Each patient received intravitreal bevacizumab at presentation and after 15 days intravitreal ozurdex was given. Central foveal thickness (CFT) and parafoveal thickness (PFT) was measured at the initiation of treatment, at 15 days, 2 months and 4 months after intravitreal Ozurdex implant. Second injection was considered for patients with recurrent edema at four months. Each patient follow up done for period of 1 year as shown in the Table 1.

3. Results

A total of 10 eyes of 9 patients (8 unilateral, 1 bilateral were recruited.

The mean IOP at presentation was 14.71 mm of Hg, at two weeks of intravitreal bevacizumab was 15 mm of Hg. At 15 days, 2 months and 4 months of intravitreal ozurdex mean IOP was 17 mm of Hg, 16.42 mm of Hg and 16.28 mm of Hg with one ant glaucoma medication (Dorzolamide eye drops).

At presentation, mean BCVA was 0.81 log mar, after 15 days of intravitreal bevacizumab was 0.55 log mar, after 15 days, 2 months and 4 months of intravitreal ozurdex was 0.54 logmar, 0.54 log mar and 0.6 logmar respectively.

Mean CFT at presentation was 538 $\mu$m and reduced to 235 $\mu$m, after intravitreal bevacizumab. At 15 days, 2 months and 4 months after intravitreal ozurdex implantation mean CFT was 182.14 $\mu$m, 189.28 $\mu$m and 352.14 $\mu$m respectively.

Mean PFT at presentation was 610 $\mu$m, after 15 days of intravitreal bevacizumab mean parafoveal retinal thickness was 353.57 $\mu$m. At 15 days, 2 months and 4 months of intravitreal Ozurdex mean PFT was 258.57 $\mu$m, 262.85 $\mu$m and 372.14 $\mu$m respectively.

4/10 (40\%) of eyes presented with recurrence of ME at four months. Each patient received second dose of intravitreal Ozurdex. Follow up done. Mean CFT reduced from 420 $\mu$m to 210 $\mu$m at 2 months, 325 $\mu$m at 4 months. Out of 4 case 1 case had recurrent ME and received 3rd dose of intravitreal ozurdex.
Table 1:

| Presentation | 15 days | 2 months | 4 months | 8 months | 12 months |
|--------------|---------|----------|----------|----------|----------|
| Clinical evaluation + Bevacizumab | Ozurdex | Recurrent | Ozurdex |

Fig. 4:

Out of 10 eyes 6 cases were pseudophakia, out of 4 phakic eyes 1 eye developed cataract after 2nd injection of ozurdex.

Number of visit to the hospital was 6 in one year duration. Out of 10 eyes, 6 eyes had received only 2 intravitreal injection (Bevacizumab + Ozurdex), 3 eyes received 3 intravitreal injection (Bevacizumab + 2 dose of Ozurdex) and 1 eye received 4 intravitreal injection (Bevacizumab + 3 dose of Ozurdex).

4. Case 1

A female patient aged 58 years presents with reduced vision in Right eye (RE), her vision in RE was 6/36, fundus picture shows multiple superficial haemorrhage in superior temporal quadrant with few cotton wool spots and hard exudates and OCT shows cystoid macular edema (CME) with IS/OS junction disruption (Figure 5 a). After 15 days of intravitreal bevacizumab her Vision improved to 6/18. After ozurdex injection her vision improved to 6/12, fundus shows resolving superficial haemorrhage with few cotton wool spots (Figure 5b). After 4 months her BCVA was 6/9, fundus shows resolved superficial haemorrhage in superior temporal quadrant and OCT shows normal foveal contour with resolved ME. (Figure 5 c).

5. Case 2

A male patient aged 62 years presents with reduced vision in RE, BCVA was 6/60, fundus shows multiple superficial haemorrhage in inferior temporal quadrant and OCT shows CME (Figure 6 a). After 15 days of intravitreal bevacizumab his BCVA improved to 6/24, OCT scan shows CME with subretinal fluid (Figure 6 b). After Ozurdex injection his Vision improved to 6/18, OCT scan shows reduced ME with few parafoveal few cystic changes (Figure 6 c). After 4 months of ozurdex vision worsened to 6/36 and OCT scan show multiple cystoid changes with epiretinal membrane (Figure 6 d). He received 2nd dose of intravitreal Ozurdex, his BCVA improved to 6/18 and OCT scan shows complete resolution of macular edema (Figure 6 e).

6. Discussion

Presently macular edema due to BRVO is treated with three pharmacological drugs including anti-VEGF such as bevacizumab (Off label use) or ranibizumab, aflibercept and a corticosteroid dexamethasone implant that is ozurdex.

In clinical trial, anti-VEGF such as ranibizumab revealed a beneficial effect on visual function and reduction in central macular thickness in eye with BRVO and diabetic macular edema. However, with respect to shorter half-life of ranibizumab, numerous injections required to achieve and maintains the therapeutic effect. This is also valid for bevacizumab as shown by Epstein et al., who performed injection every 6 weeks for 12 months with significant improvement of visual acuity and reduction of ME. Frequent intravitreal injection require frequent visits to the hospital and also risk of side effects and complications. Since the intravitreal dexamethasone implant (Ozurdex; Allergan, Irvine, CA) was recently approved for RVO, we used this treatment as combination with anti-VEGF. Which results in maintain ME with reduced number of injections and also reduced the number of patient visits. We aimed to evaluate the effect of combination of therapy in case ME due to BRVO. Initial intravitreal bevacizumab helps in reduction of ME. Ozurdex helps to maintain reduction in ME for longer duration. Combination therapy helps in reduced number of patient visit for procedures and monitoring visits and hence reduced cost of monotherapy.

In our case series BCVA improved and I decreased significantly after intravitreal bevacizumab later both parameters improved and maintained with use of intravitreal ozurdex. After 4 months 40% of cases were showed recurrence, treated with repeat intravitreal ozurdex. Out of 4 recurrent cases only one case needed 3rd dose of intravitreal ozurdex.

We hypothesized that treatment with anti-VEGF will have an early impact and following the ozurdex implantation the effect is sustained. However the time interval for recurrence could not be prolonged, which is in contract to recent study of Singer et al., Recurrence occurs after period of 3.2 and 3.8 months and were in line with the known
Fig. 5: a: Fundus picture shows multiple superficial haemorrhage in superior temporal quadrant with few cotton wool spots and hard exudates and OCT shows cystoid macular edema (CME) with IS/OS junction disruption; b: Fundus picture shows with few cotton wool spots and reduced superficial haemorrhage; c: Fundus shows resolving superficial haemorrhage resolved superficial haemorrhage in superior temporal quadrant and OCT shows normal foveal contour with resolved ME

pharmacokinetics of the ozurdex implant and the results of Geneva trial, which revealed a decrease of treatment effect at about 3-4 months after implantation.14

Mayer et al.,15 studied efficacy and safety of dexamethasone implant alone or in combination with bevacizumab. 16 patients of BRVO were treated with intravitreal ozurdex and 12 patients with 3 consecutive bevacizumab injections fallowed by intravitreal ozurdex. Results noticed BCVA improved by 7.8(±2.9) letter with ozurdex and 9.4(±2.1) letters in combination group. In our series we used single injection of Bevacizumab with significant of improvement of vision. There is a benefit of combining the two agents for greater visual outcome.

Mayer et al.,15 observed an elevation of >5mm of Hg compared with the baseline in approximately 40% of patients, irrespective of treatment regimen. Whereas in our case series we are using single dose anti glaucoma drug (Dorzolamide eye drops) and noticed IOP rise of 3 mm of Hg with baseline at the end of 4 months. Although we had no uncontrolled rise of IOP based on our experience in this case series. We suggest exclude patient with known history of glaucoma and steroid response to the treatment. Dorzolamide is a topical carbonic inhibitor used to reduce IOP and CME. Reduction of ME is by supressing production of proinflammatory cytokines, interleukin-6.16 However, response is better in retinal pigmentary epithelial disease than in RVO.17
Fig. 6: a: Fundus shows multiple superficial haemorrhage in inferior temporal quadrant and OCT shows CME; b: OCT scan shows CME with subretinal fluid; c: OCT scan shows reduced ME with few parafoveal cystic changes; d: OCT scan show multiple cystoid changes with epiretinal membrane; e: OCT scan shows complete resolution of macular edema

Michael A Singer et al., study showed mean changes in OCT measured retinal thickness decreased by 195 μm at 2 weeks after intravitreal bevacizumab, with additional 63 μm decrease seen at 4 and 6 weeks, with additional 40 μm decrease seen at 2 months and at 4 months 124 μm increase in macular thickness after Ozurdex compare to baseline. The mean BCVA increased from 9.4 to 12.9 letters at 4 weeks, 12.3 letters at 2 months and back to 10 letters at 4 months.

In our case series also central retinal thickness decreased by 303 μm at 2 weeks after intravitreal bevacizumab, with additional 53 μm at 4 weeks, 46 μm at 2 months and 117.14 μm increase at 4 months in macular thickness after ozurdex compared to baseline. We can expect a possible recurrence of edema at 4 months for most patients and review. Combination therapy showed that improvement in BCVA and macular thickness is sustained for four months and also increases the percentage of patients whose macula was essential fluid free compared with anti VEGF therapy alone.

In our case series number of visit to the hospital was 6 in one year duration and also needed a less number of injection. Whereas in monotherapy the number of visit would be about 10 and require multiple dose of intravitreal injection as in Seong Joon Ahn et al. study initial dose of
three monthly intravitreal injection versus PRN intravitreal injection of bevacizumab for macular edema secondary to BRVO showed mean number of injection over the 1 year period were 3.8±1(range, 3-7) in the 3 monthly initial dose group and 2.3±1.3(range, 1-6) in the PRN group. Study visit also more.

Combined treatment of intravitreal bevacizumab and ozurdex shows better and a sustained functional outcome. Also reduced the number of visit and reduced the cost compare monotherapy.

7. Source of Funding
None.

8. Conflict of Interest
None.

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