Case Report

Branch Retinal Vein Occlusion, Macular Ischemia, and Intravitreal Anti-VEGF Therapy

Thomas Bertelmanna, b Hans Ulrich Franka Hendrik Ansgar Fuchsa
Nicolas Feltgenb

aBelenus Eye Center Siegen, Siegen, Germany; bDepartment of Ophthalmology, University Medical Center Goettingen, Goettingen, Germany

Keywords
Branch retinal vein occlusion · Macular ischemia · Anti-VEGF · Corticosteroids · Intravitreal injection · Visual acuity · Central retinal thickness · Macular thickness · COMRADE

Abstract
Purpose: To report a case with ischemic macular edema (ME) due to an acute branch retinal vein occlusion (BRVO) which was treated with repeated intravitreal anti-VEGF injections.

Methods: Retrospective case presentation. Results: A 66-year-old female patient was treated with repeated intravitreal anti-VEGF injections due to ischemic ME following an acute BRVO. Over a period of 2.5 years best corrected visual acuity increased from 0.06 to 0.6 (decimal notation) accompanied by a reduction in central retinal thickness from 546 to 292 µm. Overall 17 anti-VEGF injections were administered to treat repeated recurrence of ME. Macular ischemia did not worsen during this profound intravitreal anti-VEGF therapy. Conclusion: Intravitreal anti-VEGF therapy can be a beneficial treatment strategy even in ischemic ME following an acute BRVO.
Introduction

Branch retinal vein occlusion (BRVO) is the second most common retinal vascular disorder following diabetic retinopathy with an overall incidence of 0.5–1.2% [1, 2]. One major complication of BRVO is the development of a cystoid macular edema (ME) in more than 90% of the eyes affected, which in turn can cause severe visual disturbances and vision loss [2]. Despite the fact that BRVO and the consecutive ME can resolve spontaneously within 1 year in about 50% of patients, a prolonged ME can often hamper visual recovery [2]. Beside perfused, non-ischemic ME that can therapeutically be addressed with intravitreal anti-VEGF substances, corticosteroids, or focal laser treatment [3, 4], there is little information available about treatment success in ischemic cases. Recently, the first report of the 24-month, prospective, open-label, randomized, active-controlled, multicenter, phase IIIb study BRIGHTER ("Individualized Stabilization Criteria-Driven Ranibizumab versus Laser in Branch Retinal Vein Occlusion: Six-Month Results of BRIGHTER") was published [5]. Herein, favorable results were described for eyes suffering from ischemic ME for the first time. To date it remains questionable if such treatment effects can also be observed in routine clinical care. The aim of this case report was to demonstrate the treatment success with intravitreally injected anti-VEGF in a patient suffering from ischemic ME due to an acute BRVO.

Case Report

A 66-year-old female patient, who complained about vision loss in her left eye, was initially seen in our department in April 2014. The conducted ophthalmologic examination revealed an acute BRVO in the inferior-temporal vein with consecutive ME development. Best corrected visual acuity (BCVA) in her left eye was 1/15 (0.06; decimal notation). Fluorescein angiography (FAG) showed an acute BRVO with ischemia in the macular area and along the inferior temporal retinal arcade (Fig. 1).

Optical coherence tomography (OCT) analysis detected a central retinal thickness (CRT) of 546 µm (Fig. 2). Except a beginning nuclear sclerosis of the lens, no further abnormalities were detected. Intraocular pressure was measured to be 17 mm Hg. The patient was sent to her family doctor for a complete medical workup and rheological therapy was conducted. Hypertension was diagnosed and subsequently treated.

Two weeks later intravitreal anti-VEGF therapy with bevacizumab (BEV) was initiated. After 3 injections, a marked increase in BCVA accompanied by a considerable CRT reduction was observed. A follow-up FAG in August 2014 and in September 2016 (Fig. 3) showed no changes in central retinal ischemia status; there was neither a decrease nor an increase in the size of the central ischemic retinal area. Retinal collaterals developed along the lower arcade as well as between the upper and lower arcade from month 4 on. A peripheral laser photocoagulation was advised in April 2014, but the patient did not give her consent for this procedure. One month later a recurrent cystoid ME was observed that was again successfully treated with 3 intravitreal BEV injections. After another ME relapse, therapy was switched to ranibizumab (RAN) and continued till October 2016. At the last visit, ME had again completely resolved and BCVA increased to 0.6 (Fig. 2).
Discussion

The recently published BRIGHTER data showed the beneficial effect of intravitreally administered RAN to treat ischemic ME due to BRVO for the first time. Overall, eyes with ischemic ME gained BCVA of 14 letters within the 6-month study period. Furthermore, the comparison of treatment effects between ischemic and non-ischemic ME did not significantly differ [5]. This is surprising, because for ischemic ME the visual prognosis, irrespective of applied treatment modalities, was thought to be unfavorable [6]. In the presented case taken from routine clinical care, the functional improvement was comparable to the treatment success in the BRIGHTER study as shown by an increase of BCVA from 0.06 at the first visit to 0.6 almost 2.5 years later.

The favorable treatment success in our case might partially be attributed to the fact that the onset of BRVO was only 2 weeks before starting intravitreal anti-VEGF injections, and rheological therapy was immediately conducted. It is well known that recent occlusions respond better than older ones [7]. Older BRVOs can lead to profound alterations of the retinal structure which can be detected with OCT (e.g., discontinuation of the external limiting membrane, myoid and ellipsoid zone). This aspect had so far not been considered in the BRIGHTER analysis even though older BRVOs were included as well. However, the second OCT scan in our case taken in August 2014 showed no alterations of the outer retinal structures. Further research is indicated to clarify this as yet unanswered question.

Another important aspect refers to the treatment regimen. In our case 3 intravitreal anti-VEGF injections were performed until ME resolved, followed by observation and reinjections as soon as the recurrence of ME was detected. This might be the cause of a total of 5 episodes of a reappearance of ME. Applying a treat-and-extend scheme [7, 8] or switching to a dexamethasone implant [9] might be more favorable to avoid this zigzag pattern of CRT. At least with intravitreally injected steroids common complications like cataract worsening or an increase of intraocular pressure can occur, all of which we aimed to avoid [9].

Switching from BEV to RAN might have been too early in the presented case although previous research indicated that switching from one anti-VEGF substance to another might intensify the anti-VEGF effect. However, when reviewing the literature, there are to date no definite guidelines on when and how to switch drugs for eyes suffering from BRVO. Both, RAN compared to BEV, could resolve ME, and even after a total of 17 anti-VEGF injections and 2.5 years of therapy, no ME was evident in October 2016, and thus, anti-VEGF therapy was quite effective in this case.

Whereas focal laser treatments in cases of macular ischemia should be avoided [4], we recommended that a peripheral laser photocoagulation should be performed within the area of retinal ischemia. Unfortunately, this was refused by the patient. Recent reports indicate that a combination therapy including anti-VEGF and peripheral targeted laser treatment significantly reduces the injection frequency and stabilizes a dry macula, which was not achieved by anti-VEGF injections alone [10, 11]. It might be hypothesized that with this combined treatment the frequently occurring ME relapses might have been avoided. So far, the literature does not provide a definite answer.

The most important aspect of this case is the observation of macular ischemia development during the profound anti-VEGF therapy. Herein, anti-VEGF blockade had no effect on the central area of non-perfused retinal tissue. These results are in accordance with a report by Rishi et al. [12] demonstrating that anti-VEGF treatment had no negative impact on ischemia development and a recently published review article about anti-VEGF treatment and macular ischemia [6]. Whereas Rishi et al. reported on their results after 1 single BEV injec-
tion, our case clearly demonstrates the safety of anti-VEGF with regard to retinal ischemia development in eyes with ischemic BRVO, because overall 17 injections in a timeframe of 2.5 years were administered.

There are some reports published recently which show a prognostic value of OCT angiography and ischemic areas [13, 14]. This new technique is still under debate and was not available in the present case.

Conclusions

As demonstrated with this case report and the data of the BRIGHTER study, intravitreally injected anti-VEGF drugs might be an excellent treatment option for ischemic macular edema due to an acute BRVO. Considerable BCVA gains and CRT reductions could be expected without increasing size and severity of central retinal ischemia in the short and long term. This seems to be plausible not only in phase III, randomized clinical trials, but also in routine clinical care. Further research in a routine clinical setting is indicated to confirm the results as described herein. Furthermore, the ischemia data taken from the COMRADE studies [15] are awaited to answer the question whether anti-VEGF or corticosteroids act similarly in such scenarios.

Statement of Ethics

The authors have no ethical conflict to disclose.

Disclosure Statement

Thomas Bertelmann has received funding for research and clinical trials from Alcon (USA), Alimera Sciences (USA), Allergan (Ireland), Bayer HealthCare (Germany), and Novartis (Switzerland), as well as consulting fees, honoraria, and travel reimbursement from Alcon (USA), Alimera Sciences (USA), Allergan (Ireland), Bayer HealthCare (Germany), and Novartis (Switzerland). He is a scientific staff member of the Department of Ophthalmology, University Medical Center Goettingen, Germany and was a Medical Advisor for Novartis Pharma GmbH (Nuremberg, Germany) from April 2015 till August 2016.

Nicolas Feltgen has received funds from Novartis, Allergan, Bayer, and Heidelberg Engineering.

Funding Sources

No funding was received for this research.

References


Fig. 1. Fluorescein angiography showed profound macular and peripheral ischemia in the early phase (left), in the middle phase (middle), and in the late phase (right, composite view).
### Fig. 2.
Intravitreal injections, development of BCVA and CRT, and corresponding OCT images between April 2014 and October 2016.
Fig. 3. Follow-up fluorescein angiography showed profound macular and peripheral ischemia in the early phase (left), in the middle phase (middle), and in the late phase (right, composite view).