Branch retinal vein occlusion associated with platelet activation

Emrullah BEYAZYILDIZ1, Mehmet ÇITIRIK2*, Mert ŞİMŞEK2, Özlem BEYAZYILDIZ3, İbrahim C. HAZNEDAROĞLU1

1 Samsun Training and Research Hospital, University of Health Sciences, Samsun, Turkey
2 Ankara Ulucanlar Eye Education and Research Hospital, University of Health Sciences, Ankara, Turkey
3 Department of Hematology, Faculty of Medicine, Hacettepe University, Ankara, Turkey

Background/aim: The aim of this study is to study subclinical platelet activation by detecting three important platelet activation parameters of mean platelet volume (MPV), platelet distribution width (PDW), and plateletcrit (PCT) in patients with branch retinal vein occlusion (BRVO) in comparison to those in healthy control subjects.

Materials and methods: This prospective study included 43 patients with BRVO (Group 1) and 40 control subjects (Group 2). The levels of MPV, PDW, and PCT were measured in both of the studied groups.

Results: The mean serum level of MPV value was 7.64 ± 0.64 in Group 1 and 7.39 ± 0.42 in Group 2. Mean serum level of PDW was 15.01 ± 1.56 in Group 1 and 14.43 ± 1.03 in Group 2. Mean serum PCT value was 0.19 ± 0.05 in Group 1 and 0.16 ± 0.04 in Group 2. MPV, PDW, and PCT levels were significantly increased in BRVO patients (P < 0.05).

Conclusion: Subclinical platelet activation reflected by MPV, PDW, and PCT may have an impact on the genesis of vessel occlusion in BRVO. The results may be important for the clinical management of patients with BRVO.

Key words: Branch retinal vein occlusion, mean platelet volume, plateletcrit, platelet distribution width, platelet parameters, retina

1. Introduction

Retinal venous occlusion is the second most common retinal vascular disorder after diabetic retinopathy causing visual loss (1). Branch retinal vein occlusion (BRVO) is a frequent retinal vascular disease with a yearly incidence of 2.14/1000 in the population over 40 years of age (2). BRVO is predisposed via various systemic and local factors (2). Hypertension and end-organ damage caused by diabetes mellitus contribute to arteriosclerosis, atherosclerosis, and endothelial dysfunction, which seem to be major risk factors for BRVO (3). Ophthalmic risk factors are ocular hypertension, glaucoma, higher ocular perfusion pressure, and changes in the retinal arteries (4). Underlying mechanisms for etiopathogenesis of BRVO are degenerative changes of the vessel wall, compression of the vein by arteriole at the arteriovenous (AV) crossing, and disturbed hematological factors (5). Endothelial dysfunction and platelet activation lead to occlusion of branch retinal veins (6,7).

Platelets have a very important role in the regulation of hemostasis and vascular integrity. Platelet indices such mean platelet volume (MPV), platelet distribution width (PDW), and plateletcrit (PCT) are standard indicators of platelet function in disease pathophysiology. The alterations in platelet parameters may affect the prothrombotic status of a patient. The platelet markers have been associated with several pathological conditions, such as increased levels of MPV shown as a risk factor for myocardial infarction and stroke (8,9). In a recent study, MPV was also shown to be increased in BRVO patients (10). To the best of our knowledge, the platelet activation parameters including PDW and PCT have not been studied in BRVO patients previously. The aim of this study is to assess the possible interrelationships of BRVO with platelet indices MPV, PDW, and PCT in order to detect subclinical platelet activation.

2. Materials and methods

This study was carried out with 43 patients with BRVO between September 2013 and January 2015. All procedures adhered to the tenets of the Declaration of Helsinki, and local approval was received from the Ethical Committee of Ankara Dışkapı Training and Research Hospital. Informed consent was obtained from each patient after
explanation of the research purposes. All patients were Turkish Caucasians. This study is registered with the Australian New Zealand Clinical Trials Registry, number ACTRN12616001246471.

The patients were included within two study groups based on the findings of clinical ocular examination. The first group included 43 BRVO patients (Group 1); the second group included 40 adult subjects serving as controls with no history of ocular and systemic disease except for hypertension and diabetes mellitus (Group 2). There were no personal or familial histories of thrombotic disease in the patients and control cases. The absence of thrombotic events or a family history of thrombosis was confirmed by means of a verified questionnaire. Patients and controls using any medication including corticosteroid and immunosuppressive therapy, who had smoking or drinking habits, or who had a history of systemic inflammatory and ocular disease were excluded. All of the patients and controls had normal liver and renal function tests and electrolytes. In order to standardize the hemostatic system parameter measurements, all sampling procedures were performed in the morning hours with fasting. The blood was collected in EDTA vacuum tubes with minimum stasis and examined within 60 min.

A complete ophthalmological examination including best corrected visual acuity, intraocular pressure measurement with Goldmann applanation tonometry, biomicroscopic examination, and dilated pupil examination of the posterior segment was performed in both groups. Subclinical BRVO was not observed in the control group.

The statistical analysis was performed by t-test according to Bonferroni procedures for multiple comparisons. The significance level was determined as P < 0.05. Differences between two groups for age and sex were evaluated using Mann–Whitney U tests. All statistics in this study were analyzed using SPSS 18.0 for Windows (SPSS Inc., Chicago, IL, USA).

3. Results
Eighty-three individuals were included in this clinical research. There were 49 women (59.1%) and 34 men (40.9%). The mean age was 59.6 ± 8.1 years (mean ± SD) in Group 1 and 61.3 ± 12.8 in Group 2. There were no difference in terms of age or sex (P = 0.5) (Table 1).

Of the 43 BRVO patients, 25 (58.1%) had hypertension and 12 (27.9%) had diabetes mellitus. The coexistence of hypertension and diabetes mellitus was present in 6 patients (13.9%). The location of vein occlusion was superotemporal in 28 eyes (65.1%), inferotemporal in 14 (32.5%), and inferonasal in 1 (2.3%). The mean best-corrected visual acuity was 20/80 (range: 20/2000 to 20/25) in the BRVO group. Biochemical parameters including glucose, lipid, and homocysteine values and clotting, plasma viscosity, and inflammatory markers were normal.

Of the 40 control subjects, 20 (52.5%) had hypertension, 9 (22.5%) had diabetes, and 4 (10%) had both hypertension and diabetes mellitus.

MPV levels showed a marked elevation in our BRVO patients (P = 0.03) compared with controls. The mean serum level of MPV was 7.64 ± 0.64 in Group 1 and 7.39 ± 0.42 in Group 2 (Table 2).

Increments in the serum level of PDW were observed in BRVO cases (P = 0.01) when compared to the control group. The mean serum level of PDW was 15.01 ± 1.56 in Group 1 and 14.43 ± 1.03 in Group 2 (Table 2).

| Table 1. Demographic data of the groups. |
|-----------------------------------------|
|                                        |
| **Group 1** (n = 43)                    | **Group 2** (n = 40) | **P** |
| **Age (years)**                         | **61.37 ± 12.87**   | **59.65 ± 8.15** | **0.47** |
| **Sex (male/female)**                   | **18/25**           | **16/24**        | **0.17** |

| Table 2. The mean values of mean platelet volume, plateletcrit, and platelet distribution width and comparison of platelet parameters (mean ± standard deviation) for groups. |
|--------------------------------------------------------------------------------|
|                                                                                   |
| **Group 1** (n = 43)                                                             | **Group 2** (n = 40) | **P** |
| **MPV (fL)**                                                                      | **7.64 ± 0.64**      | **7.39 ± 0.42** | **0.03** |
| **PDW (%)**                                                                       | **15.01 ± 1.56**     | **14.43 ± 1.03** | **0.01** |
| **PCT (%)**                                                                       | **0.19 ± 0.05**      | **0.16 ± 0.04**  | **0.04** |

MPV: Mean platelet volume, PDW: platelet distribution width, PCT: plateletcrit.
The mean PCT levels were 0.19 ± 0.05 in BRVO cases, and these were significantly higher than in controls (0.16 ± 0.04; P = 0.04) (Table 2).

4. Discussion
Branch retinal vein occlusion is a common cause of retinal vascular disorder and is said to be the second most common disease after diabetic retinopathy (11). Major pathogenetic factors confined to the pathogenesis of BRVO are abnormal hematological parameters, degenerative changes of the retinal vessel wall, and compression of the retinal vein at the AV crossing (12). Although multiple hematological parameters were discussed in the pathogenesis of BRVO, the role of coagulation parameters remains unclear. Thrombosis may develop due to platelet abnormalities, endothelial injury, increased procoagulant activity, and abnormal fibrinolysis. Larger platelets are more active and contain more intense granules and they have more powerful prothrombotic activity (13).

Increased MPV may have a role in the vessel occlusion process of BRVO. Platelet activation also correlates with increased MPV levels. The high value of MPV is an indication of increased thrombocyte size. Large platelets as diameters are more active, functional, and dense than small ones. Thus, higher MPV levels may increase the possibility of vascular complications (14,15). Previous studies showed that MPV was associated with situations such as coronary and peripheral artery disease, myocardial infarction, and cerebral ischemia (16–19).

PDW is also related to platelet activation and it is a more specific marker than MPV, because blood values of PDW are not high during simple platelet swelling (20). PDW shows variations in platelet size that may be indicators of active platelet release (21). PDW levels could be altered in several conditions and increased levels were observed in sickle cell patients with vaso-occlusive crisis (22,23). PDW may be a more specific marker than MPV for showing platelet activation and increased levels of PDW might show impaired deformability of thrombocytes and be related to microvascular resistance (24–26). Vagdatli et al. (20) found that PDW is a specific and simple marker for coagulation activation. Thus, increased PDW levels could be a risk factor and play a role in the pathogenesis of BRVO. Amin et al. (23) showed elevated levels of PDW during vaso-occlusive complications in sickle cell anemia patients.

PCT is a marker of blood-circulating platelets in a unit volume. PCT reveals quantitative abnormalities of platelets and is calculated as platelets × MPV/107 (27). Akpinar et al. (13) found that PCT had a significant predictive value for saphenous vein graft disease and emphasized that it could be used as a marker for antiplatelet therapy to prevent graft atherosclerosis in patients undergoing bypass surgery. Onder et al. (10) showed decreased levels of platelets in hypertensive BRVO patients, but not significantly. In another previous study, PCT was correlated with C-reactive protein in chronic inflammatory diseases (28). An increased MPV level in patients with hypertension and diabetes mellitus without BRVO was reported previously (29). Therefore, our control subjects were selected from among individuals with a history of hypertension and diabetes mellitus.

In the present study, BRVO patients showed significantly higher levels of MPV, PDW, and PCT. The increased levels of these parameters may play a role in the pathogenesis of BRVO. Increment in these platelet parameters may increase activation and aggregation of platelets and increase vasoactive mediators’ secretion by these platelets such as thromboxane A2, resulting in vasoconstriction, endothelial dysfunction, and impaired blood flow, which results in microvascular occlusion. Some studies suggested that thrombophilic parameters are altered in groups of patients with BRVO and control groups (10,30). However, to the best of our knowledge, PDW and PCT have not been evaluated before in patients with BRVO.

MPV is calculated by dividing the PCT by the number of platelets, which is the same calculation as for the mean red cell volume, namely dividing hematocrit by the red blood cells, and therefore PCT is analogous to the red cell hematocrit (31). Red cell indices are widely used in clinics. On the other hand, platelet indices (other than the platelet count itself) are probably the most regularly ignored part of the automated complete blood count parameters (32). Positive studies and metaanalyses of automated platelet analyses focusing on coronary heart diseases highlighted long-term efforts about the validation of those ‘routine’ lab parameters in eye vessel occlusions including BRVO (33).

The pathogenesis of BRVO includes local microenvironmental factors, including degenerative changes of vascular endothelium, compression at the AV crossing, and others. Systemic hypercoagulable factors of disturbed hematological factors including platelet activity may also have impact as a triggering factor. The treatment of this challenging occlusive event may not rely solely on antiplatelet drugs. However, we must be aware of the ongoing subclinical thrombocyte activation during the clinical management of the patients with BRVO.

Squizzato et al. (34) investigated the best available evidence on the acute treatment and the secondary prevention of RVO with antithrombotic and fibrinolytic drugs. A search of the MEDLINE and EMBASE electronic databases up to January 2009 was performed. They suggested that antithrombotic therapy, in particular low-molecular-weight heparin, may be a part of the therapeutic armamentarium for patients with recent-onset
RV ORVO. Houtsmuller et al. (35) established a double-blind study to evaluate the platelet aggregation inhibiting effect of ticlopidine on the course of ocular vein occlusions. They determined that the effect of ticlopidine was most pronounced in patients with increased platelet aggregation and least obvious in cases of hyperlipidemia. Hypertension and diabetes did not apparently influence ticlopidine’s effects. The platelet aggregation inhibitor ticlopidine was found effective in the treatment of recent ocular vein occlusions.

There are several limitations in the present study. The limited number of patients is the most important limiting factor. Secondly, this study is based on a simple baseline determination that may not reflect a patient’s long-term status. Based on the results of our present study, BRVO patients have significantly higher serum levels of MPV, PDW, and PCT than control subjects. MPV, PDW, and PCT seem to play an important role in the genesis of BRVO. Serial measurements of PDW, MPV, and PCT may be helpful in the diagnosis and prevention of BRVO. The sudden peak of these markers indicates platelet activation. The results may be important for the clinical management of patients with BRVO in everyday clinical practice since those platelet activation parameters are routinely detected in complete blood count analyses. Further large-scale and comprehensive studies are needed to support these results.

Based on the current data, if an ophthalmologist does have a concern that platelet activation could be impaired in a patient with BRVO, he or she could share the case with a hematologist if there is any sign of systemic platelet activation syndromes that potentially affect any micro- and/or macrocirculatory vessel systems. Antithrombotic, antiplatelet, anticoagulant, and pro fibrinolytic management strategies should then be determined for the given patient, as well as the hemostatic follow-up.

References

1. Klein R, Klein BE, Moss SE, Meuer SM. The epidemiology of retinal vein occlusion: the Beaver Dam Eye Study. Trans Am Ophthalmol Soc 2000; 98: 133-141.
2. David R, Zangwill L, Badarna M, Yassur Y. Epidemiology of retinal vein occlusion and its association with glaucoma and increased intraocular pressure. Ophthalmologica 1988; 197: 69-74.
3. Newman-Casey PA, Stem M, Talwar N, Musch DC, Besirli CG, Stein JD. Risk factors associated with developing branch retinal vein occlusion among enrollees in a United States managed care plan. Ophthalmology 2014; 121: 1939-1948.
4. Kolar P. Risk factors for central and branch retinal vein occlusion: a meta-analysis of published clinical data. J Ophthalmol 2014; 2014: 724780.
5. Rehak J, Rehak M. Branch retinal vein occlusion: pathogenesis, visual prognosis, and treatment modalities. Curr Eye Res 2008; 33: 111-131.
6. Wong TY, Larsen EK, Klein R, Mitchell P, Couper DJ, Klein BE, Hubbard LD, Siscovick DS, Sharrett AR. Cardiovascular risk factors for retinal vein occlusion and arteriolar emboli: the Atherosclerosis Risk in Communities and Cardiovascular Health Studies. Ophthalmology 2005; 112: 540-547.
7. Cheung N, Klein R, Wang JJ, Cotch MF, Islam AF, Klein BE, Cushman M, Wong TY. Traditional and novel cardiovascular risk factors for retinal vein occlusion: the Multiethnic Study of Atherosclerosis. Invest Ophthalmol Vis Sci 2008; 49: 4297-4302.
8. Bath P, Algert C, Chapman N, Neal B; Progress Collaborative Group. Association of mean platelet volume with risk of stroke among 3134 individuals with history of cerebrovascular disease. Stroke 2004; 35: 622-626.
9. Huczek Z, Kochman J, Filipiak KJ, Horszczaruk GJ, Grabowski M, Piatkowski R, Wilczynska J, Zielinski A, Meier B, Opolski G. Mean platelet volume on admission predicts impaired reperfusion and long-term mortality in acute myocardial infarction treated with primary percutaneous coronary intervention. J Am Coll Cardiol 2005; 46: 284-290.
10. Onder H, Klicic A, Kaya M, Bulur S, Onder E, Tunc M. Relation between platelet indices and branch retinal vein occlusion in hypertensive patients. Indian J Ophthalmol 2013; 61: 160-162.
11. The Branch Vein Occlusion Study Group. Argon laser photocoagulation for macular edema in branch vein occlusion. Am J Ophthalmol 1984; 98: 271-282.
12. Senen K, Topal E, Kilinc E, ten Cate H, Tek I, Karakoc Y, Yetkin E. Plasma viscosity and mean platelet volume in patients undergoing coronary angiography. Clin Hemorheol Microcirc 2010; 44: 35-41.
13. Akpinar I, Sayin MR, Gursoy YC, Karabag T, Kucuk E, Buyukyuval MC, Aydin M, Haznedaroegluc I. A platelet marker associated with saphenous vein graft disease. Herz 2014; 39: 142-148.
14. Ates O, Kiki I, Bilcn H, Keles M, Kocer I, Kulaoglu DN, Baykal O. Association of mean platelet volume with the degree of retinopathy in patients with diabetes mellitus. Eur J Gen Med 2009; 6: 99-102.
15. Kodiatte TA, Manikyam UK, Rao SB, Jagadish TM, Reddy M, Lingaiah HK, Lakshmaiah V. Mean platelet volume in type 2 diabetes mellitus. J Lab Physicians 2012; 4: 5-9.
16. Khandekar MM, Khurana AS, Deshmukh SD, Kakrani AL, Kadtare AD, Inamdar AK. Platelet volume indices in patients with coronary artery disease and acute myocardial infarction: an Indian scenario. J Clin Pathol 2006; 59: 146-149.
17. Tavil Y, Sen N, Yazici H, Turfan M, Hizal F, Cengel A, Abaci A. Coronary heart disease is associated with mean platelet volume in type 2 diabetic patients. Platelets 2010; 21: 368-372.

18. Pikija S, Cvetko D, Hajduk M, Trikulja V. Higher mean platelet volume determined shortly after the symptom onset in acute ischemic stroke patients is associated with a larger infarct volume on CT brain scans and with worse clinical outcome. Clin Neurol Neurosurg 2009; 111: 568-573.

19. Berger JS, Eraso LH, Xie D, Sha D, Mohler ER. Mean platelet volume and prevalence of peripheral artery disease, the National Health and Nutrition Examination Survey, 1999-2004. Atherosclerosis 2010; 213: 586-591.

20. Vagdatli E, Gounari E, Lazaridou E, Katsibourlia E, Tsikopoulou F, Labrianou I. Platelet distribution width: a simple, practical and specific marker of activation of coagulation. Hippokratia 2010; 14: 28-32.

21. Karagöz B, Alacacıoğlu A, Bilgi O, Demirci H, Özgün A, Eriğiç AA, Sayan O, Yılmaz B, Kandemir EG. Platelet count and platelet distribution width increase in lung cancer patients. Anatol J Clin Investig 2009; 3: 32-34.

22. Osselaer JC, Jamart J, Scheiff JM. Platelet distribution width for differential diagnosis of thrombocytosis. Clin Chem 1997; 43: 1072-1076.

23. Amin MA, Amin AP, Kulkarni HR. Platelet distribution width (PDW) is increased in vaso-occlusive crisis in sickle cell disease. Ann Hematol 2004; 83: 331-335.

24. Yaylali YT, Susam I, Demir E, Bor-Kucukatay M, Uludag B, Kılıç-Toprak E, Erken G, Dursunoglu D. Increased red blood cell deformability and decreased aggregation as potential adaptive mechanisms in the slow coronary flow phenomenon. Coron Artery Dis 2013; 24: 11-15.

25. Leone MC, Gori T, Fineschi M. The coronary slow flow phenomenon: a new cardiac “Y” syndrome? Clin Hemorheol Microcirc 2008; 39: 185-190.

26. Muxel S, Pasola F, Radmacher MC, Jabs A, Münzel T, Gori T. Endothelial functions: translating theory into clinical application. Clin Hemorheol Microcirc 2010; 45: 109-115.

27. Bain BJ, Bates I. Basic haematological techniques. In: Lewis SM, Bain BJ, Bates I, editors. Dacie and Lewis Practical Haematology. 9th ed. Edinburgh, UK: Churchill Livingstone; 2001. pp. 19-46.

28. Sahin F, Yazar E, Yildiz P. Prominent features of platelet count, plateletcrit, mean platelet volume and platelet distribution width in pulmonary tuberculosis. Multidiscip Respir Med 2012; 7: 38.

29. Citirik M, Beyazyildiz E, Simsek M, Beyazyildiz O, Haznedaroğlu IC. MPV may reflect subclinical platelet activation in diabetic patients with and without diabetic retinopathy. Eye (Lond) 2015; 29: 376-379.

30. Coban E, Adanir H, Bilgin D. The association of mean platelet volume levels with hypertensive retinopathy. Platelets 2008; 19: 115-118.

31. Briggs CM, Machin SJ. Automated platelet analysis. In: Kottke-Marchant K, editor. Laboratory Hematology Practice. Oxford, UK: Wiley-Blackwell; 2012. pp. 48-58.

32. Patterson K. Platelet parameters generated by automated blood counters. CME Bull Haematol 1997; 1: 13-16.

33. Chu SG, Becker RC, Berger PB, Bhatt DL, Eikelboom JW, Konkle B, Mohler ER, Reilly MP, Berger JS. Mean platelet volume as a predictor of cardiovascular risk: a systematic review and meta-analysis. J Thromb Haemost 2010; 8: 148-156.

34. Squizzato A, Manfredi E, Bozzato S, Dentali F, Ageno W. Antithrombotic and fibrinolytic drugs for retinal vein occlusion: a systematic review and a call for action. Thromb Haemost 2010; 103: 271-276.

35. Houtsmuller AJ, Vermeulen JA, Klompe M, Zahn KJ, Henkes HE, Baarsma GS, Tijssen J. The influence of ticlopidine on the natural course of retinal vein occlusion. Agents Actions Suppl 1984; 15: 219-229.