The Association Between Red Cell Distribution Width and Bone Marrow Fibrosis in Patients with Philadelphia-Negative Myeloproliferative Neoplasms

Philadelphia Kromozomu Negatif Myeloproliferatif Neoplazili Hastalarda Eriosit Dağılım Genişliği İndeksi ile Kemik İliği Fibrozisi Arasındaki İlişkinin Değerlendirilmesi

Objective: Red cell distribution width (RDW) was shown to be increased in primary myelofibrosis (PMF) patients and it is intriguing whether RDW could be used instead of biopsy in predicting presence of bone marrow fibrosis (BMF) to some extend in Philadelphia-negative myeloproliferative neoplasms (MPNs) comprising polycythemia vera (PV), essential thrombocytosis (ET) and PMF. Our aim is to evaluate the relationship between BMF degree and RDW values in patients with MPNs.

Method: We retrospectively reviewed the data of 118 patients, who were followed with the diagnosis of MPNs at our Hematology Clinic between 2010 and 2017.

Results: 52 patients had PV, 60 had ET, 4 had PMF and 2 had unclassifiable MPN. Twentynine (24.6%) patients were with grade 0 and grade 1 reticulin fibrosis were considered to be free of BMF, and the remaining 89 (75.4%) patients with ≥ grade 2 reticulin fibrosis were considered to have BMF. The median RDW value was 14.6% (range 12.4-23.1%). The median RDW value revealed with 14.1% (range, 12.4-17.8) in patients without BMF and 15% (range, 12.4-23.1) in patients with BMF (p=0.054). In subgroup analysis of 8 patients with advanced BMF of grade 3, the median RDW value was 18.45% (range, 16.4-23.1) and it was 14.45% (range, 12.4-23) in the remaining 110 patients (p=0.008).

Conclusion: Although the present study does not provide a precise conclusion about the association between RDW and BMF, it seems that increased RDW can point out the presence of advanced BMF in patients with MPNs.

Keywords: red cell distribution width, chronic myeloproliferative neoplasms, bone marrow fibrosis

Giriş: Eriosit dağılım genişliğinin (RDW) primer miyelofibroz hastalarında artış gösterilmiştir ve Philadelphia-negatif miyeloproliferatif neoplazilerde (MPN) kemik iliği fibrozisi varlığında etmede biyopsis yerine RDW’ın kullanılabilmesi ilgi çekicidir. Çalışmanın amacı: Polisitemi vera (PV), esansiyel trombositositoz (ET) ve PMF alt tiplerini içeren MPN’li hastalarda myelofibroz derecesi ile RDW değerleri arasındaki iliği değerlendirilmektir.

Yöntem: Hastanemizin Hematoloji Kliniği’nde 2010-2017 tarihleri arasında MPN tanısı taşıyan 118 hastanın verilerini retrospektif olarak inceledik

Bulgular: Elli iki hastada PV, 60 hastada ET, 4 hastada PMF ve 2 hastada sınıflandırılmayan MPN saptandı. Derece 0 ve derece 1 retikülin fibrozisi bulunan 29 hasta (% 24,6) myelofibrozisi olmayan olarak, kalan 2 ≥ derece retikülin fibrozisi olan 89 (% 75,4) hasta ise myelofibrozisi bulunan olarak kabul edildi. Medyan RDW değeri %14,6 (%12,4-23,1) idi. Ortanca RDW değeri myelofibrozisi olmayan hastalarda %14.1 (%12,4-17,8) ve myelofibrozisi hastalarda %15 (12,4-23,1) olarak odaya sonuçlandı (p = 0.054). Derece 3 fibrozisi olup ileri myelofibrozisi olan 8 hastanın alt grub analizinde medyan RDW değeri % 18.45 (16,4-23,1) ve kalan 110 hastada % 14.45 (%12,4-23,1) saptandı (p = 0.008).

Sonuç: Bu çalışma, RDW ve myelofibroz arasındaki ilişki hakkında kesin bir sonuç sağlamada da,arms RDW’ın MPN’li hastalarda ileri kemik iliği fibrozisi varlığında işaret edebileceğini göründüktedir.

Anahtar Kelimeler: eriosit dağılım genişliği, kronik myeloproliferatif neoplaziler, kemik iliği fibrozu

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INTRODUCTION

Polycythemia vera (PV), essential thrombocythemia (ET) and primary myelofibrosis (PMF) which are among the Philadelphia chromosome negative myeloproliferative neoplasms (MPNs) occur as the result of uncontrolled clonal stem cell-derived myeloid serial proliferation (1). Bone marrow fibrosis (BMF) is characterized by reticulin fibrosis and/or collagen fiber accumulation and can be seen in MPNs in varying grades in onset also can develop later (2). And the presence of ≥ grade 2 reticulin/collagen fibrosis is the major criteria for the diagnosis of PMF according to the World Health Organization (WHO) 2016 criteria, so performance of bone marrow biopsy is indispensable for the diagnosis of both PMF, post-PV and post-ET myelofibrosis(3). However, instead of biopsy a blood test ensuring information about the presence of BMF would be more comfortable for PV and ET patients especially on follow up.

Anisocytosis is a feature of PMF and can be monitored by red cell distribution width (RDW) which was shown to be increased in PMF patients, previously (4,5). Red cell distribution width is usually reported within the routine complete blood cell count (CBC) panel and makes it easily available. It is intriguing whether RDW could be used in predicting the presence of BMF to some extent. However, there is lack of data about the relationship between RDW and BMF in patients with MPNs.

The aim of this study is to evaluate the relationship between BMF and RDW value in patients with MPNs.

METHODS

We retrospectively reviewed the data of 146 patients, who were followed with the diagnosis of PV, ET and PMF and had bone marrow biopsy at the time of diagnosis, at University of Health Sciences Istanbul Training and Research Hospital Hematology Clinic between August 2010 and January 2018. Total of twenty-eight patients who had concomitant iron/vitamin B12 deficiency or not fully meeting the diagnostic criterions were excluded from the study. The diagnosis of MPNs were made according to the WHO criteria (3). Reticulin and trichromestaining were applied to the specimens and grading was done by an expert pathologist as follows: Grade 0: Scattered linear reticulin with no intersections corresponding to normal bone marrow; grade 1: Loose network of reticulin with many intersections, especially in perivascular areas; grade 2: Diffuse and dense increase in reticulin with extensive intersections, occasionally with only focal bundles of collagen and/or focal osteosclerosis; grade 3: Diffuse and dense increase in reticulin with extensive intersections with coarse bundles of collagen, often associated with significant osteosclerosis. The data comprising age, gender, RDW values, hemoglobin levels, white blood cell count (WBC), platelet count, lactate dehydrogenase (LDH) levels, JAK2V617F mutation status, presence of splenomegaly and thromboembolic event history at the time of bone marrow biopsy before any treatment or phlebotomy procedure, were obtained from the hospital documentation system. During the eight-years of data collection period, the RDW values were reported as RDW-CV (%) rather than RDW-SD (FL) parameter in the complete blood count of most of our patients and differences in laboratory reference intervals for RDW-CV were observed. Therefore, we determined the median RDW-CV value of the entire patient group as the RDW cut-off value.

Statistical evaluation was made by SPSS 24 program. Data were described as numbers and percentage or median and range, when appropriate. x^2 Fisher's exact test was used for evaluating categorical values and Mann Whitney U test for continuous values in patient groups. All p-values were 2-sided with statistical significance at 0.05 alpha levels.

RESULTS

A hundred and eighteen patients were included into the study. Among them 52 had PV, 60 had ET, 4 had PMF and 2 had unclassifiable CMPN at the time of diagnosis. The patient characteristics are shown in table 1.
Twenty-nine (24.6%) patients whose bone marrow biopsy revealed with grade 0 and grade 1 reticulin fibrosis were considered to be free of BMF, and the remaining 89 (75.4%) patients with ≥ grade 2 reticulin fibrosis were considered to have BMF. The median RDW value was 14.6% (range 12.4-23.1%). When RDW values were examined according to the presence of BMF, the median RDW value was 14.1% (range, 12.4-17.8) in patients without BMF and 15% (range, 12.4-23.1) in patients with BMF (p=0.054).

In subgroup analysis of 8 advanced BMF patients with grade 3 fibrosis, the median RDW value was 18.45% (range, 16.4-23.1) and the median RDW value in the remaining 110 patients was 14.45% (range, 12.4-23) (p=0.008).

| Table 1 : The Patient Characteristics |
|--------------------------------------|
| Characteristis                        | N=118 |
|                                       | n    | %   |
| Gender                                |      |     |
| Female                                | 51   | 43.2|
| Male                                  | 67   | 56.8|
| Age (years)                           |      |     |
| median                                | 53   |     |
| range                                 | 18-79|     |
| CMPN                                  |      |     |
| PV                                    | 52   | 44.1|
| ET                                    | 60   | 50.8|
| PMF                                   | 4    | 3.4 |
| UC                                    | 2    | 1.7 |
| Hemoglobin level, g/dl                |      |     |
| median                                | 14.45|     |
| range                                 | 8.6-21.3|    |
| WBC, 10⁹/L                            |      |     |
| median                                | 10   |     |
| range                                 | 2.82-19.94|  |
| Platelet, 10⁹/L                       |      |     |
| median                                | 625  |     |
| range                                 | 33-1408|    |
| LDH                                   |      |     |
| High                                  | 38   | 32.2|
| Normal                                | 68   | 57.6|
| Unknown                               | 12   | 10.2|
| Splenomegaly                          |      |     |
| Present                               | 23   | 19.5|
| Absent                                | 61   | 51.7|
| Unknown                               | 34   | 28.8|
| JAK 2 V617F                           |      |     |
| Positive                              | 68   | 57.6|
| Negative                              | 49   | 41.5|
| Unknown                               | 1    | 0.9 |
| History of thrombosis                 |      |     |
| Positive                              | 5    | 4.2 |
| Negative                              | 112  | 94.9|
| Unknown                               | 1    | 0.9 |
| RDW-CV %                              |      |     |
| median                                | 14.6 |     |
| range                                 | 12.4-23.1|   |
| RDW-CV                                |      |     |
| RDW-CV > 14.6                         | 37   | 31.4|
| RDW-CV ≤ 14.6                         | 81   | 68.6|
| BMF                                   |      |     |
| Present                               | 89   | 75.4|
| Absent                                | 29   | 24.6|
The cut-off point for RDW was determined as 14.6% according to the median level of RDW. Elevated RDW (>14.6) values were present in 37 (31.4%) of patients and RDW values were normal in 81 (68.6%) patients. The relationship between BMF and RDW was investigated. While BMF was found in 32 (86.5%) patients with high RDW level; 57 (70.4%) patients had BMF in the patient group with normal RDW values (p=0.068) (table 2).

|                      | High RDW-CV >14.6 | Normal RDW-CV ≤ 14.6 | P value |
|----------------------|-------------------|----------------------|---------|
| BMF Absent (grade 0-1) | n=5, %13.5        | n=24, %29.6          |         |
| BMF Present (grade ≥ 2) | n=32, %86.5     | n=57, %70.4          | 0.067   |

DISCUSSION

Red cell distribution width is an index measuring variability of peripheral blood erythrocyte volumes and represents anisocytosis (4). Besides its function in CBC, RDW has been recognized as a marker of subclinical inflammation by reflecting an increase in level of C-reactive protein, sedimentation rate (6), and cytokines such as hepcidin and interleukin 6 in recent years (7,8). In addition, there has been growing evidence about the negative impact of elevated RDW on inflammatory diseases, cardiovascular diseases, solid organ malignancies and some hematological disorders such as chronic lymphocytic leukemia, multiple myeloma, chronic myeloid leukemia, diffuse large b cell lymphoma and PMF (5,9-15). Unlike previous mentioned functions of RDW, we investigated whether it could give information about the BMF status of the patients with the diagnosis of MPNs. We found that, though not significant statistically, the number of patients having BMF was higher in the group with increased RDW values and the median RDW value increased in patients with BMF. In subgroup analysis, the increase in RDW was more prominent in patients with grade 3 fibrosis.

The presence of BMF, contributing to morbidity and mortality with accompanying risk factors, has a substantial role in patients with MPNs(16,17).

While BMF is an essential prominent feature of PMF, it occurs in 5-14% PV patients and 15-20% ET patients at diagnosis (3,18,19). Also, rapid progression to overt myelofibrosis can develop during PV and ET course and requires bone marrow biopsy for the diagnosis of post-PV and Post-ET myelofibrosis (20). Anisocytosis, which can be identified by means of increased RDW, is attributed to the myelofibrosis (4) and has a potential to substitute the bone marrow biopsy in estimating BMF, at least in selected patient groups. Lucijanic et al. demonstrated that increased median RDW, with a median level of 19% in PMF patients was associated with decreased overall survival(5). Similarly, in our study we found that RDW increased significantly in patients with grade 3 myelofibrosis which is consistent with PMF or Post PV, ET myelofibrosis. However, the number of patients with PMF was quite low compared to the patients with other MPNs.

Limitations of the Study

Due to the retrospective nature of our study comparatively low number of patients with primary myelofibrosis might have hindered the influence of RDW on estimating the grade of BMF.

Conclusion

Consequently, although the present study does not provide a precise conclusion about the association between RDW and BMF, it seems that increased RDW can point out the presence of advanced BMF in patients with MPNs.

However, the contribution of RDW value to the estimation of myelofibrosis grade in MPNs should be elucidated with studies including large number of patients with PMF and post-PV/post-ET myelofibrosis.
Competing Interests
The authors declare that there is no conflict of interest regarding the publication of this article.

Authors Contributions
All the authors declare that they have participate in the preparation of this study.

Ethics Committee Approval: This study was approved by local ethic committee (2011-KAEK-50)

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