Methotrexate therapy impacts on red cell distribution width and its predictive value for cardiovascular events in patients with rheumatoid arthritis

Julia Held, Birgit Mosheimer-Feistritzer, Johann Gruber, Erich Mur and Günter Weiss

Abstract

Background: Methotrexate (MTX) is well known to affect folic acid metabolism, so MTX treatment can result in alterations of mean corpuscular volume (MCV), which may impact on red cell distribution width (RDW), as MCV levels feed into RDW calculation. We thus questioned whether RDW levels and subsequently its diagnostic utility in RA subjects, as reported before, are influenced by ongoing MTX therapy.

We assessed the impact of disease modifying drug (DMARD) treatment, especially MTX, on RDW and evaluated their influence on the predictive value of RDW for cardiovascular (CV) events in patients with rheumatoid arthritis (RA). As far as we know, this is the first study evaluating the influence of MTX on RDW.

Methods: Medical treatment, disease activity, laboratory parameters and history of CV events were retrospectively analysed in 385 RA patients at disease onset and at last follow up at our clinic. Additionally, in patients with CV event, data were recorded at last follow up prior the CV event.

Results: Disease parameters and laboratory findings associated with a serious vascular event were older age (p < 0.001), longer disease duration (p = 0.002) and a higher RDW at diagnosis (p = 0.025). No differences in RDW levels became evident with any other treatment regimen beside MTX. MTX treated patients had significantly higher RDW compared to subjects without this drug (p < 0.001). In RA patients without MTX treatment, we found RDW level significantly different between those with versus without a CV event, whereas this difference disappeared in subjects receiving MTX.

Conclusion: MTX impacts on RDW and might therefor reduce its prognostic value for CV events in patients taking MTX, whereas an increased RDW at diagnosis remains an early risk predictor for myocardial infarction and stroke in RA patients.

Keywords: Rheumatoid arthritis, Red cell distribution width, Methotrexate, Cardiovascular events

Background

Rheumatoid arthritis (RA) is one of the most prevalent systemic inflammatory diseases which involve joints and extra-articular tissues, thereby causing organ damage. Based on the chronic inflammation and immune dysregulation, the presence of RA has been associated with cardiovascular (CV) disease and an increased CV associated mortality [1, 2]. Classical risk factors for CV disease have been investigated in RA patients, however, epidemiological studies indicate that they cannot provide a sufficient explanation for the poorer CV prognosis of RA patients as compared to non-RA subjects [3, 4]. Therefore, a combination of yet not fully elucidated factors and regulatory mechanisms may contribute to CV morbidity in RA subjects. However, it is of clinical importance to identify markers which indicate an increased CV risk or predict CV mortality in RA patients. In this regard, recent studies reported an association of elevated red cell distribution width (RDW) with CV disease in...
the general population [5] but also in patients with RA [6, 7].

RDW is an automated measure of the range of variation of red blood cell (RBC) volume and is calculated as the standard deviation (SD) in red blood cell size divided by the mean corpuscular volume (MCV) (RDW (%) = 1 SD of RBC volume/MCV × 100). RDW is part of the complete blood count and has traditionally been used in anemia diagnosis [8–10] and to predict the response to iron treatment [11]. A retrospective analysis of > 20,000 patients with RA indicated that higher RDW, as well as increased levels of markers of inflammation, like C-reactive protein (CRP) or erythrocyte sedimentation rate (ESR), were associated with an increased risk of a subsequent CV event in RA-patients [12].

The mechanism behind the association of elevated RDW and CV risk in RA is yet incompletely understood. Possible explanations include that RDW reflects endothelial damage and impaired vascular repair, but also that RDW mirrors vascular inflammation underlying atherosclerosis and thereby effects myocardial infarction and stroke [13]. The described positive association between IFN-alpha, a cytokine contributing to endothelial damage, and RDW would be in agreement with this concept [14].

Of note, RDW is influenced by multiple factors related to erythropoiesis, such as iron, vitamin B12 or folic acid availability, as well as by hemolysis [9]. Moreover, RDW is also affected by organ dysfunctions (e.g. liver or renal dysfunction), inflammatory activity and some specific medications [15–17]. The latter might also affect the diagnostic potential of RDW in RA patients, as these subjects are treated with numerous disease modifying antirheumatic drugs (csDMARDs), like Methotrexate (MTX), or biological DMARDs (bDMARDs), including several cytokine antibodies [18]. According to EULAR recommendations, treatment of RA should be initiated with csDMARDs, most notably MTX in combination with low dose glucocorticoids. Although low dose MTX therapy is regarded as an anchor therapy in RA, full details of its mechanism of action and off target effects are still incompletely understood [19].

Because, MTX is well known to affect folic acid metabolism, MTX treatment can result in alterations of MCV, which may impact on RDW, as MCV levels feed into RDW calculation [6]. We thus questioned, whether RDW levels and subsequently its diagnostic utility and potential in RA subjects, as reported before [7, 11, 19, 20], are influenced by ongoing MTX therapy.

Methods

Patients

We evaluated a total of number of 385 patients with RA. These patients were either consecutively registered in the database for evaluating iron homeostasis in RA patients (n = 261), or were evaluated retrospectively following their clinical examination at our outpatient clinics (n = 124). All patients fulfilled the 2010 ACR classification criteria for the diagnosis of RA [21]. At inclusion in the database, clinical and laboratory parameters were collected of these 261 patients. The 124 patients, who consulted the outpatients’ clinic in 2014, were retrospectively evaluated. Full blood count, disease activity parameters and medication were available from that appointment. Additionally, we retrospectively evaluated clinical and laboratory findings at last visit before a CV event occurred, this visit was defined as last follow up in these patients. In patients without a CV event, either the date the patients were included in the database or the routine follow up at our outpatient clinic was defined as last follow up. The study was approved by the Ethics committee at Medical University Innsbruck, Austria (study number AN2014-0277).

CV-Event

The medical history was examined until Nov. 2016. A severe CV-Event was defined as myocardial infarction with or without ST-wave elevation or as an ischemic stroke. According to previous studies, CV events were clustered together [7, 22].

RDW

RDW is mathematically calculated based on the results of a routine blood count as the one SD of RBC volume/MCV × 100 [23]. The laboratory analyses were all performed by the Central Institute of Medical and Chemical Laboratory Diagnostics, University Hospital, Innsbruck. RDW was evaluated at initial diagnosis, during follow up between 2009 and 2016 and prior to a CV event. ΔRDW showed the change between the RDW at diagnosis and prior to CV event or at last follow up in patients without CV event.

Statistical analysis

Statistical analysis was performed using IBM SPSS Statistics 24 software. Normal distribution of laboratory parameters was assessed and retained by Kolmogorov-Smirnov-Test and One-Sample Chi-Square Test, respectively. Correlation among parameters was determined using Spearman-Rank-analysis. For comparative analysis between groups we applied Mann-Whitney-U-test, respectively cross tables. Linear regression model was applied to evaluate effects of medical treatment and laboratory findings on RDW.

Results

We retrospectively analysed 385 RA-outpatients, of whom 77 (20%) were male and 308 (80%) female. The
mean duration of RA was 14,5 years (SD 11,9), 77,2% of patients were positive for anti-citrullinated-peptide-antibodies (ACPA) and 73,5% of patients had a positive rheumatoid factor (RF) test, both parameters linked to disease severity in RA [24]. Upon evaluation of their medical history, we found that 23/385 (6%) had a documented severe CV event during the observation period (17 patients with myocardial infarction, six patients with stroke). We could not find an association between ACPA and/or RF positivity and the risk for a CV event. Disease parameters and laboratory findings associated with a vascular event were older age ($p < 0,001$) and a longer disease duration ($p = 0,002$). Patients with a cardiovascular event during follow up had a higher RDW at the date of initial diagnosis of RA as compared to subjects without a subsequent cardiovascular event ($p = 0,025$; Table 1).

We then studied for associations between disease activity, measured by DAS-28-CRP, haematological parameters (RDW and MCV at last visit or last visit before a CV event) and csDMARDs therapy in patients with or without a CV event. While no significant difference was seen for laboratory parameters, disease activity and all other medications including biological DMARDs between patients with and without a subsequent CV event, we found that leflunomid was administrated more frequently in patients with CV events, however, the total number of patients receiving leflunomid was very low ($p = 0,05$; Table 2).

Out of 385 RA patient included in the evaluation 284/385 (73,8%) were under treatment with csDMARDs and 83/284 (29,2%) patients were under combination therapy with biological DMARDs and csDMARDs. Because certain csDMARDs may impact on erythropoiesis or modulate the availability of factors involved in erythropoiesis, such as folic acid [25], we next studied for differences in RDW according to underlying therapy. Whereas no differences in RDW levels became evident with any other DMARD treatment, patients receiving MTX therapy had significantly higher RDW as compared to subjects without this drug ($p < 0,001$), although all patients under MTX treatment had folic acid supplementation prescribed (Table 3).

We then studied whether or not MTX treatment had an effect on the predictive value of RDW for severe CV events. The last routine follow up of patients with evaluation of laboratory parameters prior to the CV events was 82 days (mean) prior to such a CV complication. No significant differences in RDW, hemoglobin levels, CRP or ΔRDW (change in RDW between initial diagnosis and follow up prior to the CV event/last patient visit)

| Table 1 Demographics and laboratory parameters at initial diagnosis of RA |
|------------------------|--------|-----------|
| CV-Event               |        | P         |
|                       | Yes (n = 23) | No (n = 362) |
| Gender Female, n (%)  | 15 (65,2) | 293 (80,9) | 0,068 |
| Age, mean (SD), years | 74,1 (7,1) | 61,8 (13,2) | < 0,001 |
| Disease duration, mean (SD), years | 23,7 (13,2) | 14,1 (11,7) | 0,002 |
| RF positive, n (%)    | 17 (73,9) | 268 (74) | 0,990 |
| ACPA positive, n (%)  | 18 (78,3) | 279 (77,1) | 0,896 |
| RDW, mean (SD)        | 15,6 (0,78) | 13,6 (1,28) | 0,025 |

Demographics and laboratory parameters at initial diagnosis of RA according to a CV event during follow up, p values for statistical significances between the two groups are given by Mann Whitney U test, respectively cross tables, significance level at p level ≤ 0,05. SD standard deviation, n sample size, RF rheumatoid factor, ACPA anti citrullinated peptide antibodies, RDW red cell distribution width.

### Table 2 Clinical and laboratory findings during follow up

| CV-Event | Yes (n = 23) | No (n = 362) | p |
|----------|--------------|--------------|---|
| DAS 28 at follow up, mean (SD) | 3,12 (0,93) | 2,82 (1,37) | 0,252 |
| MCV at follow up, mean (SD) | 86,6 (5,97) | 87,2 (5,7) | 0,533 |
| RDW at follow up, mean (SD) | 15,1 (2,2) | 14,3 (1,5) | 0,130 |
| DMARDs at follow up, n (%) | Methotrexate | 15 (65,2%) | 207 (57,2%) | 0,451 |
| | Sulfasalazine | 0 | 9 (2,5%) | 0,445 |
| | Hydroxychloroquine | 2 (8,7%) | 27 (7,5%) | 0,868 |
| | Leflunomide | 3 (13%) | 15 (4,1%) | 0,050 |
| | Azathioprine | 0 | 6 (1,7%) | 0,533 |
| | Glucocorticoids | 8 (34,8%) | 133 (37,5%) | 0,810 |
| bDMARD | 8 (34,8%) | 130 (36%) | 0,892 |

Clinical and laboratory findings during follow up associated with CV event, significance level at p level ≤ 0,05 – see legend to Table 1, DAS 28 disease activity score 28, MCV mean corpuscular volume, (b)DMARD (biological) disease modifying antirheumatic drug.

### Table 3 RDW distribution as a function of underlying DMARD therapy

| DMARD | Intake | n | RDW, mean (SD) | p |
|-------|--------|---|----------------|---|
| Methotrexate | Yes | 222 | 14,5 (1,44) | < 0,001 |
| | No | 163 | 14,0 (1,55) | |
| Sulfasalazine | Yes | 9 | 13,7 (1,56) | 0,119 |
| | No | 376 | 14,3 (1,5) | |
| Hydroxychloroquine | Yes | 30 | 13,9 (1,19) | 0,085 |
| | No | 355 | 14,4 (1,52) | |
| Leflunomide | Yes | 18 | 14,7 (2,06) | 0,429 |
| | No | 367 | 14,3 (1,47) | |
| Azathioprine | Yes | 6 | 14,7 (1,23) | 0,273 |
| | No | 379 | 14,3 (1,51) | |
| bDMARDs | Yes | 139 | 14,2 (1,45) | 0,381 |
| | No | 246 | 14,4 (1,53) | |

RDW distribution as a function of underlying DMARD therapy, significance level at p level ≤ 0,05, bDMARDs biological DMARDs, see tables above, n number of patients in respective groups.
were found in our cohort between patients incurring a CV event or not. Of note, we found a highly significant difference in RDW levels at last follow up, see definition above, in subjects without MTX treatment which was absent in subjects under MTX treatment comparing patients with and without a CV event (Fig. 1). Moreover, MTX-treated patients had significantly higher RDW levels than patients without MTX therapy (Table 4).

In patients without MTX therapy RDW was significantly higher in those with a subsequent CV event. \( p = 0.006; \) Fig. 1). This predictive value of RDW was abolished in patients taking MTX \( p = 0.448, \) Fig. 1).

Discussion
As far as we know, this is the first study evaluating the influence of csDMARDs on RDW. MTX impacts on

Multiple linear regression analysis confirmed the relationship between RDW prior to the CV event and MTX treatment as well as associations of haemoglobin levels and age with RDW (Table 5A).

When performing binary regression for the risk of a CV event, we found that MTX naïve patients had a significant correlation between RDW and a CV event (Table 5B) which was not true for patients receiving MTX.

Discussion
As far as we know, this is the first study evaluating the influence of csDMARDs on RDW. MTX impacts on
RDW hence on the predictive value of RDW for CV events. Previous studies suggest that RDW is a good prognostic marker for CV disease and survival, but none of them evaluated concomitant treatment [6, 7, 12, 17]. This was also confirmed in our study indicating that an enhanced RDW at initial diagnosis, but neither ACPA nor RF positivity, is associated with an increased risk of a severe CV event [26].

However, we found that the predictive potential of RDW during follow up largely depends on the treatment of patients. Specifically, we identified that the diagnostic value of RDW as a risk indicator for subsequent CV disease is abolished in RA patients receiving MTX therapy. Although MTX is regarded as the anchor drug in RA, the mechanism of action is incompletely understood and it has been associated with negative effects on hematopoiesis, mainly via its impact on folic acid pathways, but also via direct toxic effects on hematopoietic progenitors. Accordingly, MTX but neither other csDMARDs nor bDMARD treatment resulted in alterations of red blood cell volume (MCV) and haemoglobin content of erythrocytes [27, 28]. To avoid such negative effects, patients under MTX therapy are supplemented with folic acid which was also the case in our subjects under MTX treatment [29]. However, RA patients under MTX treatment had increased RDW and MCV levels as compared to RA patients without MTX. It remains to be clarified, whether this can be referred to effects of MTX not linked to folic acid deficiency, or a reduced compliance of patients in regard to folic acid supplementation, which we could not study, because incomplete results of folic acid determination in blood were available in our study cohort. However, while MTX treatment resulted in a loss of the predictive value of RDW for subsequent CV events, higher RDW at initial diagnosis of all patients, at follow up prior to a CV event in patients without MTX treatment were significantly associated with an increased risk for a CV event.

Our study has limitations because of the retrospective design, the low number of patients with CV events and the fact that we had only insufficient data to evaluate the association of RDW, MTX therapy and CV events with other important variables including classical CV factors, the implication of a genetic component, or iron homeostasis [11, 30, 31]. In this regard, patients with RA who carried the methylene tetrahydrofolate reductase (MTHFR) 1298 allele C frequency were previously found to have an increased frequency of CV events after 5 and 10 years of follow-up. Moreover, patients carrying the MTHFR 1298 AC and CC genotypes had a significantly decreased flow-mediated endothelium-dependent vasodilatation, a marker of endothelial dysfunction that is an early indicator of atherogenesis, when compared with those carrying the MTHFR 1298 AA genotype [32]. More recent results also indicate that MTHFR expression is significantly reduced in patients with RA compared to controls. It was found to be especially true for RA patients with ischemic heart disease [33]. Taken these considerations together, these

Table 4: Effects of MTX intake on laboratory parameters at last follow up

| MTX-use | p         | CV-Event | p         |
|---------|-----------|----------|-----------|
| Yes (n = 222) |          | No (n = 163) |          |
| RDW, mean(SD), % | 14.5 (1.4) | 14.1 (1.6) | < 0.001 | 15.05 (2.2) | 14.25 (1.4) | 0.116 |
| Hb, mean(SD), g/l | 134.8 (14.5) | 132.9 (14) | 0.101 | 130.2 (14.4) | 134.2 (14.3) | 0.058 |
| CRP, mean(SD), mg/dl | 0.65 (0.97) | 1.03 (2.1) | 0.559 | 0.99 (1.1) | 0.8 (1.6) | 0.060 |
| ΔRDW, mean(SD) | 0.53 (1.24) | 0.23 (1.0) | 0.057 | 0.45 (0.6) | 0.43 (1.2) | 0.809 |

Effects of MTX intake on laboratory parameters at last visit prior to the CV event/last follow up. ΔRDW Change in RDW between initial diagnosis and follow up prior to the CV event/last follow up in patients without CV event. Hb hemoglobin level. In the overall cohort RDW at established disease was not applicable as predictive marker for a CV event. A tendency with lower hemoglobin levels and higher CRP was shown. MTX intake significantly affected RDW

Table 5: A) multiple linear regression modelling the relationship with RDW, B) Binary regression relationship between CV events and RDW

A: Multiple linear regression B: Binary regression

| Dependent variable: RDW at follow up | CV-Event |
|-------------------------------------|----------|
| MTX | p | 95% CI | RDW follow up | p | 95% CI |
| MTX | < 0.001 | 0.290–0.818 | no | RDW follow up | 0.018 | 0.491–0.935 |
| Age, y follow up | < 0.001 | 0.011–0.031 | yes | RDW follow up | 0.511 | 0.624–1.247 |
| Hb, g/l follow up | < 0.001 | −0.057−0.039 |

A) Multiple linear regression modelling the relationship with RDW at follow up, B) Binary regression, relationship between CV events and RDW at follow up according to MTX intake, y years, Hb hemoglobin level, MTX methotrexate, CV cardiovascular, CI confidence interval
results indicate that MTHFR gene may influence the risk of subclinical atherosclerosis and CV disease in patients with RA.

Therefore, we need prospective evaluations of such risk markers and profiles to gain further insights into the functional and diagnostic role of RDW and alterations of hematological parameters as risk predictors for CV events in patients with RA. Accordingly, such confirmation may also translate into clinical practice, because patients at a higher risk, based on increased RDW may deserve an intensified clinical follow up with an improved control of classical CV risk factors, such as lipid status, hypertension, smoking status or hyperuricemia.

**Conclusion**

Our study approves RDW at initial diagnosis of RA as a risk predictor for serious CV events but also indicates that its predictive value is lost during follow up in patients receiving MTX therapy, whereas it remains valid in subjects receiving non-MTX containing treatments.

**Abbreviations**

ACR: American College of Rheumatology; CRP: C-reactive protein; CV: Cardiovascular; DAS-28: Disease activity score 28; DMARD: Disease modifying drug; ESR: Erythrocyte sedimentation rate; EULAR: European League against Rheumatism; MCV: Mean corpuscular volume; MTHFR: Methylenetetrahydrofolate reductase; MTX: Methotrexate; RA: Rheumatoid arthritis; RDW: Red cell distribution width; SD: Standard deviation

**Acknowledgements**

Support by the Christian Doppler Society, Austria is gratefully acknowledged.

**Funding**

This work was supported by the Austrian research funds, project FWF-TRP 188 to GW.

**Availability of data and materials**

The datasets generated and/or analysed during the current study are not publicly available due they are part of a database, containing further, not yet published, data, but all data analysed during this study are included in this published article.

**Authors’ contributions**

JH: acquisition of data, analysis and interpretation of data; drafting the manuscript, given final approval of the version to be published, agreed to be accountable for all aspects of the work. BM: acquisition and analysis of data, given final approval of the version to be published, agreed to be accountable for all aspects of the work. JG: acquisition and analysis of data, given final approval of the version to be published, agreed to be accountable for all aspects of the work. BM: acquisition and analysis of data, given final approval of the version to be published, agreed to be accountable for all aspects of the work. JH: acquisition of data, analysis and interpretation of data; drafting the manuscript, given final approval of the version to be published, agreed to be accountable for all aspects of the work. JG: acquisition and analysis of data, given final approval of the version to be published, agreed to be accountable for all aspects of the work. JH: acquisition of data, analysis and interpretation of data; drafting the manuscript, given final approval of the version to be published, agreed to be accountable for all aspects of the work. JG: acquisition and analysis of data, given final approval of the version to be published, agreed to be accountable for all aspects of the work.

**Ethics approval and consent to participate**

The study was approved by the Ethics committee at Medical University Innsbruck, Austria (study number AN2014-0277). Patients in the database provided written informed consent, for patients who were retrospectively analysed, no consent to participate was obtained. This was approved by the Ethics committee at Medical University Innsbruck, Austria.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

**Publisher’s Note**

Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

**Author details**

1Department of Internal Medicine II, Infectious Diseases, Immunology, Rheumatology, Pneumology, Medical University of Innsbruck, Anichstr. 35, A-6020 Innsbruck, Austria. 2Department for Physical Medicine and Rehabilitation, University of Innsbruck, Innsbruck, Austria. 3Christian Doppler Laboratory for Iron Metabolism and Anemia Research, Innsbruck, Austria.

**Received: 25 November 2017 Accepted: 9 February 2018**

**Published online:** 07 March 2018

**References**

1. Smolen JS, Aletaha D, McInnes IB. Rheumatoid arthritis. Lancet. 2016;388: 2023–38.
2. Maradit-Kremers H, Crowson CS, Nicola PJ, Ballman KV, Roger VL, Jacobsen SJ, et al. Increased unrecognized coronary heart disease and sudden deaths in rheumatoid arthritis: a population-based cohort study. Arthritis Rheum. 2005;52:402–11.
3. del Rincón ID, Williams K, Stern MP, Freeman GL, Escalante A. High incidence of cardiovascular events in a rheumatoid arthritis cohort not explained by traditional cardiac risk factors. Arthritis Rheum. 2001;44: 237–45.
4. Gonzalez A, Maradit Kremers H, Crowson CS, Ballman KV, Roger VL, Jacobsen SJ, et al. Do cardiovascular risk factors confer the same risk for cardiovascular outcomes in rheumatoid arthritis patients as in non-rheumatoid arthritis patients? Ann Rheum Dis. 2008;67:64–9.
5. Patel KV, Semba RD, Ferrucci L, Newman AB, Fried LP, Wallace RB, et al. Red cell distribution width and mortality in older adults: a meta-analysis. J Gerontol A Biol Sci Med Sci. 2010;65:258–65.
6. Felker GM, Allen LA, Pocock SJ, Shaw LK, McMurray JJV, Pfeffer MA, et al. Red cell distribution width as a novel prognostic marker in heart failure: data from the CHARM Program and the Duke Databank. J Am Coll Cardiol. 2007;50:40–7.
7. Rodríguez-Carrio J, Alperi-López M, López P, Alonso-Castro S, Ballina-García FJ, Suárez A. Red cell distribution width is associated with cardiovascular risk and disease parameters in rheumatoid arthritis. Rheumatology (Oxford). 2015;54(4): 641–6.
8. Demir A, Yaralı N, Fıșın T, Duru F, Kara A. Most reliable indices in differentiation between thalassemia trait and iron deficiency anemia. Pediatr Int. 2002;44:612–6.
9. Thomas C, Thomas L. Biochemical markers and hematologic indices in the diagnosis of functional iron deficiency. Clin Chem. 2002;48:1066–76.
10. Besim JD, Gilmer PR, Gardner FH. Improved classification of anemias by MCV and RDW. Am J Clin Pathol. 1983;80:322–6.
11. Weiss S. Monitoring iron therapy in chronic heart failure. Eur J Heart Fail. 2013;15:711–2.
12. Hassan S, Antonelli M, Ballou S. Red cell distribution width: a measure of cardiovascular risk in rheumatoid arthritis patients? Clin Rheumatol. 2015; 34(6):1053–7.
13. Iadeoca C, Anrather J. The immunology of stroke: from mechanisms to translation. Nat Med. 2011;17:796–808.
14. Rodríguez-Carrio J, Alperi-López M, López P, Alonso-Castro S, Carro-Esteban SR, Ballina-García FJ, et al. Red cell distribution width is associated with endothelial progenitor cell depletion and vascular-related mediators in rheumatoid arthritis. Atherosclerosis. 2015;240:131–6.
15. Baltas S, Demirkol S, Cakar M, Aydogan M, Akhan M. The red cell distribution width may be affected by many factors in the clinical practice. J Clin Diagn Res. 2013;7:1830.
16. Mori S, Hidaka M, Kawakita T, Hidaka T, Tsuda H, Yoshitama T, et al. Factors associated with myelosuppression related to low-dose methotrexate therapy for inflammatory rheumatic diseases. PLoS One. 2016;11:e0154744.
17. Luo R, Hu J, Jiang L, Zhang M. Prognostic value of red blood cell distribution width in non-cardiovascular critically or acutely patients: a systematic review. PLoS One. 2016;11:e0167000.

18. Smolen JS, Landewé R, Bijlsma J, Burmester G, Chatzidionysiou K, Dougados M, et al. EULAR recommendations for the management of rheumatoid arthritis with synthetic and biological disease-modifying antirheumatic drugs: 2016 update. Ann Rheum Dis. 2017;76:960–77.

19. Brown PM, Pratt AG, Isaacs JD. Mechanism of action of methotrexate in rheumatoid arthritis, and the search for biomarkers. Nat Rev Rheumatol. 2016;12:731–42.

20. Al Taii H, Yaqoob Z, Al-Kindi SG. Red cell distribution width (RDW) is associated with cardiovascular disease risk in Crohn’s disease. Clin Res Hepatol Gastroenterol. 2017;41(4):490–492.

21. Aletaha D, Neogi T, Silman AJ, Funovits J, Felson DT, Bingham CO, et al. 2010 rheumatoid arthritis classification criteria: an American College of Rheumatology/European League Against Rheumatism collaborative initiative. Arthritis Rheum. 2010;62:2569–78.

22. Gonzalez-Gay MA, Gonzalez-Juanatey C, Lopez-Diaz MJ, Piñeiro A, Garcia-Porrua C, Miranda-Filloy JA, et al. HLA-DRB1 and persistent chronic inflammation contribute to cardiovascular events and cardiovascular mortality in patients with rheumatoid arthritis. Arthritis Rheum. 2007;57:125–32.

23. Lippi G, Mattiuzzi C, Ceravolo G. Learning more and spending less with neglected laboratory parameters: the paradigmatic case of red blood cell distribution width. Acta Biomed. 2017;87:323–32.

24. Song YW, Kang EH. Autoantibodies in rheumatoid arthritis: rheumatoid factors and anticitrullinated protein antibodies. J Autoimmun. 2010;34:139–46.

25. Lobb BF, Witzmann G, Ogir E, Studnicka-Benke A, Andel I, Schweitzer H, et al. Folic acid and cyanocobalamin levels in serum and erythrocytes during low-dose methotrexate therapy of rheumatoid arthritis and psoriatic arthritis patients. Clin Exp Rheumatol. 1995;13:569–63.

26. Berendsen MLT, van Maaren MC, Arts EEA, den Broeder AA, Popa CD, Fransen J. Anticyclic citrullinated peptide antibodies and rheumatoid factor as risk factors for 10-year cardiovascular morbidity in patients with rheumatoid arthritis: a large inception cohort study. J Rheumatol. 2017; https://doi.org/10.3899/jrheum.160670.

27. Gilani STA, Khan DA, Khan FA, Ahmed M. Adverse effects of low dose methotrexate in rheumatoid arthritis patients. J Coll Physicians Surg Pak. 2012;22:101–4.

28. Dubey L, Chatterjee S, Ghosh A. Hepatic and hematological adverse effects of long-term low-dose methotrexate therapy in rheumatoid arthritis: an observational study. Indian J Pharmacol. 2016;48:591.

29. McInnes IB, Schett G. Pathogenetic insights from the treatment of rheumatoid arthritis. Lancet. 2017;389:2328–37.

30. Fernández-Gutiérrez B, Perrotti PP, Gibert JP, Domènech E, Fernández-Nebro A, Cañete JD, et al. Cardiovascular disease in immune-mediated inflammatory diseases. Medicine (Baltimore). 2017;96:e7308.

31. López-Mejías R, Castañeda S, González-Juanatey C, Corrales A, Ferraz-Amaro I, Genie F, et al. Cardiovascular risk assessment in patients with rheumatoid arthritis: the relevance of clinical, genetic and serological markers. Autoimmun Rev. 2016;15:1013–30.

32. Palomo-Morales R, Gonzalez-Juanatey C, Vazquez-Rodriguez TR, Rodriguez L, Miranda-Filloy JA, Fernandez-Gutierrez B, et al. A1298C polymorphism in the MTHFR gene predisposes to cardiovascular risk in rheumatoid arthritis. Arthritis Res Ther. 2010;12(2):R71.

33. Remuñez-Martínez S, Genie F, López-Mejias R, Ubilla B, Mijares V, Pina T, et al. Decreased expression of methylene tetrahydrofolate reductase (MTHFR) gene in patients with rheumatoid arthritis. Clin Exp Rheumatol. 2016;34:106–10.