



Neutrophil-to-Lymphocyte Ratio as an Inflammatory Marker in Familial Mediterranean Fever: A Systematic Review and Meta-analysis

Alireza Omranzadeh¹ Ashkan Baradaran¹ Alireza Ghodsi¹ Soheil Arekhi¹ Malihe Dadgarmoghaddam² Amin Mirshekaran¹ Amirreza Dehghan Tarazjani¹ Bahare Fazeli³

¹ Student Research Committee, Mashhad University of Medical Sciences, Mashhad, Iran

²Community Medicine Department, Medical School, Mashhad University of Medical Sciences, Mashhad, Iran

³Inflammation and Inflammatory Diseases Research Center, Faculty of Medicine, Department of Immunology, Inflammation and Inflammatory Diseases Research Center, Mashhad University of Medical Sciences, Mashhad, Iran

| Child Sci 2021;11:e100-e109.

Abstract

Keywords familial

Address for correspondence Bahare Fazeli, Faculty of Medicine, Department of Immunology, Inflammation and Inflammatory Diseases Research Center, Mashhad University of Medical Sciences, Mashhad, Iran (e-mail: Fazelib@mums.ac.ir).

Familial Mediterranean fever (FMF) is a genetic disease with inflammatory basis. Several studies have assessed the role of neutrophil-to-lymphocyte ratio (NLR) in detecting this inflammation; however, no systematic review or meta-analysis has assessed these studies together. The aim of this study is to systematically review and meta-analyze the NLR value between FMF patients and normal controls. Scopus, PubMed, Embase, and ISI Web of Science were searched using predesigned search strategy to find the studies that assessed NLR in FMF patients and compared the value with normal controls. There was no time limitation. Finally, two researchers extracted data including first author name, publication year, the country, study design, number of patients and controls, time of disease diagnosis, FMF diagnostic criteria, mean age of the patients, and the NLR value. The data were systematically reviewed and meta-analyzed. In total, 464 articles were found on search; however, only 12 studies qualified for enrollment in the systematic review and 10 studies, with appropriate effect size, in the meta-analysis. These studies were conducted between 2013 and 2019. Eleven studies were conducted in Turkey and one in Egypt. Out of 12 studies, 9 had enrollment criteria for FMF patients: 8 studies used Tel Hashomer criteria and 1 study used Yalçinkaya-Özen criteria. All studies, except for two, had genetic confirmation for FMF. The mean NLR values in Mediterranean fever attack-free (standard difference in means = 0.482; p < 0.0001) and attack groups neutrophil-to-(standard difference in means = 0.853; p = 0.001) were significantly higher than lymphocyte ratio control group. The mean NLR value may be related to the underlying inflammation ► inflammation in FMF.

received November 19, 2020 accepted after revision March 8, 2021

DOI https://doi.org/ 10.1055/s-0041-1728730. ISSN 2474-5871.

© 2021. The Author(s).

This is an open access article published by Thieme under the terms of the Creative Commons Attribution License, permitting unrestricted use, distribution, and reproduction so long as the original work is properly cited. (https://creativecommons.org/licenses/by/4.0/) Georg Thieme Verlag KG, Rüdigerstraße 14, 70469 Stuttgart, Germany

Familial Mediterranean fever (FMF) is a genetic autoinflammatory disorder with autosomal recessive inheritance that affects 1 out of every 500 childbirths, in Mediterranean endemic region.¹ FMF is mainly the disease in Turks, Armenians, Arabs, and Sephardic Jews that is caused by various mutations of the Mediterranean Fever (*MEFV*) gene, which is located on the short (p) arm of the 16th chromosome.² However, this disease is not only limited to the Mediterranean region, and reports from all around the world are available.^{3,4} The disease also exists in Iranian Azeri Turks, as a varying range of 21 to 42.4% of *MEFV* gene mutations have been reported in this ethnicity.⁵

t is believed that mutation in the *MEFV* gene results in autoinflammatory processes that cause fever and inflammation of serous membranes. This serous inflammation demonstrates itself as abdominal and chest pains, which are another hallmarks of the disease.² Most of the cases present their first attack during the childhood; 65 and 95% of FMF patients experience their first attack before 10 and 20 years of age, respectively.^{6,7} The duration of FMF attacks usually ranges from 1 to 3 days, and in this period, several laboratory markers of inflammation including C-reactive protein (CRP), erythrocyte sedimentation rate (ESR), serum amyloid A (SAA), and fibrinogen arise.^{8,9} All these inflammation markers usually return to normal levels in attack-free period; however, it is reported that subclinical inflammation still remains.^{8,10}

Neutrophil-to-lymphocyte ratio (NLR) is a newly developed marker of inflammation that has been assessed in several autoimmune and autoinflammatory diseases, such as ankylosing spondylitis,¹¹ rheumatoid arthritis,¹² lupus erythematous,13 Kawasaki's disease,14 and Behçet's disease.¹⁵ Recently, this marker has been used to assess baseline inflammation in FMF patients, during both attack and attackfree periods.^{16,17} However, controversies regarding levels of NLR during clinical and subclinical periods of FMF remain. Some studies believe that the NLR level in FMF patients during attack-free periods is significantly higher than in normal controls^{9,18,19}; however, other studies believe that the amount of NLR is significantly higher only during attack periods.^{16,17,20} Therefore, we aimed to systematically review and meta-analyze the available evidence regarding the role of NLR in demonstrating clinical and subclinical inflammation in FMF patients.

Materials and Methods

Study Protocol

This study was conducted according to the Preferred Reporting Items for Systematic reviews and Meta-Analyses (PRISMA) guideline.²¹ The study protocol was previously registered in International Prospective Register of Systematic Reviews (PROSPERO).²²

Search Strategy

A systematic electronic literature search was conducted in ISI Web of Science, Scopus, PubMed, and Embase databases using medical subject heading (MeSH). The following search terms were employed to include all relevant studies: ("Familial Mediterranean fever" OR "FMF" OR "familial paroxysmal polyserositis" OR "recurrent polyserositis" OR "benign paroxysmal polyserositis" OR "periodic disease" OR "periodic peritonitis") AND ((neutrophil* AND lymphocyte*) OR NLR). All databases were reviewed from inception to November 2, 2020, using the above-mentioned search strategy. There was no time limit for our search. Also, the references of the related studies were hand checked to find any remaining relevant publication.

Selection of the Studies

Two authors (A. O. and S. A.) screened related studies separately. In the first step, the studies were reviewed based on their titles and abstracts, and then their full texts were used to achieve the final screening for the studies. The results of the study selection by each reviewer were compared and disagreements and inconsistencies were resolved by an immunologist (B. F.), who was expert in this topic. All the studies that compared NLR level between patients with FMF and normal controls were included. Animal studies, case reports, review articles, and letter to editor were excluded from the review.

Data Extraction

Data extraction was performed by two authors (A. O. and A. G.) using a predesigned data extraction Excel form. The extracted information includes first author's name, year of publication, country where study was conducted, study design, inclusion criteria for FMF, number of subjects in each group, duration from the disease diagnosis, FMF diagnostic criteria, genetic confirmation, gender, mean age of subjects, age at FMF diagnosis, presence and dose of colchicine treatment, and NLR level for each of the study groups. In case of disagreement regarding the extracted data, a third reviewer (B. A.) resolved the disputes.

Quality Assessment of the Included Studies

The quality assessment of the studies was also done by two reviewers (A. O. and A. B.), and in case of any disagreements, these discrepancies were resolved by a social medicine specialist (M. D.) as an expert. The Joanna Briggs checklist²³ was used to assess the quality of the studies, all of which were case–control. This checklist consists of 10 questions that examine various areas of the methodology of the case– control studies and report the final quality. The answer to each of the questions is divided into four options, namely, yes (Y), no (N), unclear (U), and not applicable (NA). All studies with appropriate statistical analysis were included and those that lacked this item were excluded.

Heterogeneity, Risk of Bias, and Meta-analysis

Statistical analyses were performed using Comprehensive Meta-Analysis (CMA) software version 2 (Biostat, Englewood, NJ, USA). The mean NLR values were pooled in the software as effect size and a random effect model was used to compare data between FMF attack or attack-free cases and normal controls. Results were presented as forest plots. Heterogeneity was tested by using the l^2 index and Cochran's Q statistic. $l^2 > 75\%$ and significant Cochran's Q test indicate heterogeneity

between studies. Moreover, funnel plot of the included studies was designed to assess publication bias.

Results

Study Selection Process

An electronic search through the PubMed, Scopus, Embase, and ISI Web of Science databases yielded 116, 106, 127, and 115 studies, respectively. Out of a total of 464 studies, 199 were removed as duplicates and 265 studies remained and underwent title and abstract reviewing. Totally, 244 articles were excluded after this procedure. As a result, the full texts of 21 studies were reviewed, of which 9 studies were excluded from the systematic review for various reasons, including lack of relevance to our study and irrelevant outcomes with those intended in our review. Finally, 12 studies were included for systematic review. To perform meta-analysis, 2 studies were excluded and finally only 10 studies were analyzed. One study was excluded because it was conducted only on pregnant women and the population was not similar to other studies. Another study did not have our intended effect size, and due to the abnormal distribution, it was not possible to convert the effect size. **-Fig. 1** shows the PRISMA flowchart, which summarizes the study selection process.

Study Characteristics

► **Table 1** shows the characteristics of the included studies. As illustrated, all the studies were published between 2013 and 2019. Also, all studies, except for one in Egypt,²⁴ were conducted in Turkey. In 9 out of the 12 studies, the inclusion criteria were clear; 8 studies used the Tel Hashomer criteria²⁵ and one study¹⁷ employed the Yalçinkaya–Özen criteria.²⁶ Furthermore, genetic validation was used in all publications, except for two studies.^{27,28} All studies were case–controlled. ► **Table 1** shows details of the included studies.

Patient Characteristics

As demonstrated in **►Table 2**, our study population consisted of 1,980 FMF patients (970 males and 1,010 females) and 880 controls (382 males and 498 females). Out of 1,980

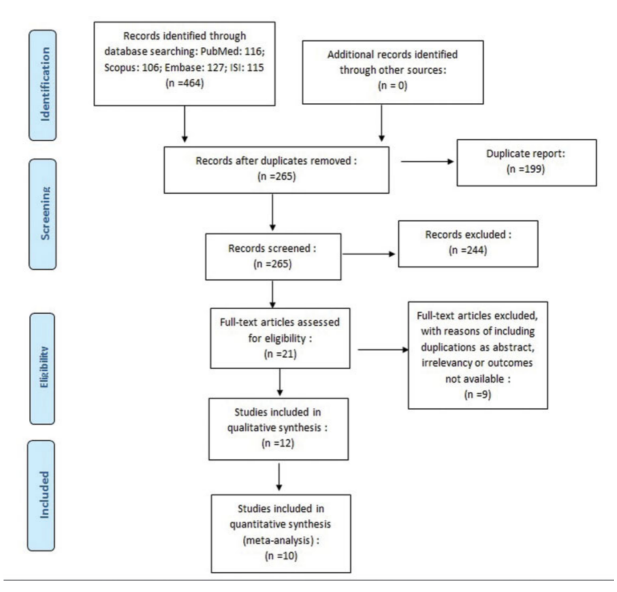


Fig. 1 PRISMA flowchart of paper selection.

Table 1 Characteristics of the enrolled studies including first author's name, publication year, country and area, type of study, inclusion criteria of the patients, FMF diagnosis criteria, and genetic confirmation

First author	Publication year	Country/area	Type of study	Inclusion and exclusion criteria	Genetic confirmation	FMF diagnosis criteria
Yorulmaz ¹⁶	2019	Turkey/Konya	Case-control	Patients older than 1 mo and younger than 18 y with a diagnosis of FMF Patients with splenomegaly, diabetes, asthma, blood disorders, kidney and liver failure, uncontrolled hypertension, proteinuria, and those receiving nonsteroidal anti-inflammatory drugs and anticoagulants were excluded	Yes	Tel Hashomer ²⁵
Kholoussi ²⁴	2018	Egypt/Cairo	Case-control	Patients diagnosed with FMF were referred to the Egyptian National Genetic Diseases Research Clinic	Yes	Unknown
Basaran ¹⁷	2017	Turkey/Ankara	Case-control	Patients did not have any other systemic or inflammatory diseases	Yes	Yalçinkaya–Özen ²⁶
Çakar ³⁰	2017	Turkey/Ankara	Case-control	Patients with other rheumatic diseases, atherosclerotic cardiovascular disease, hypertension, diabetes, and those under 18 y of age were excluded	Yes	Tel Hashomer ²⁵
Daglar ²⁷	2016	Turkey/ Gaziosmanpaşa	Case-control	Women with first pregnancies who had no abnormalities on their ultrasound and no systemic disease other than FMF	No	Tel Hashomer ²⁵
Kelesoglu ¹⁸	2016	Turkey/Istanbul	Case-control	Patients should have complete laboratory and clinical information and should not be treated with anakinra	Yes	Unknown
Celikbilek ²⁸	2015	Turkey/Bezak	Case-control	Patients should not have diabetes, hypertension, liver disease, and acute or chronic infections and should have therapeutic compliance	No	Tel Hashomer ²⁵
Duksal ¹⁹	2015	Turkey/Sivas	Case-control	Patients should not have any infectious or chronic disease and should not receive any treatment other than FMF treatment	Yes	Tel Hashomer ²⁵
Özer ²⁹	2015	Turkey/Tukat	Case-control	Patients should not have infection, pneumonia, diabetes, hypertension, acute or chronic renal failure, obstructive sleep apnea, coronary artery disease, connective tissue disease, inflammatory bowel disease, allergic rhinitis, asthma, or a history of any inflammatory disease	Yes	Tel Hashomer ²⁵
Uluca ²⁰	2014	Turkey/ Diyarbakir	Case-control	Patients included according to Tel Hashomer criteria	Yes	Tel Hashomer ²⁵
Uslu ⁹	2013	Turkey Sivas	Case-control	Patients with diabetes, cardiovascular disease, metabolic syndrome, anemia, acute or chronic infection, other autoimmune diseases, chronic obstructive pulmonary disease, and a history of smoking were excluded. Also, patients should not take any medication other than colchicine	Yes	Tel Hashomer ²⁵
Ahsen ³¹	2013	Turkey/ Afyonkarahisar	Case-control	Patients should not have infection, pneumonia, diabetes, hypertension, acute or chronic renal failure, chronic liver disease, obstructive pulmonary disease, obstructive sleep apnea, coronary artery disease, connective tissue disease, inflammatory bowel disease, allergic rhinitis, asthma, and smoking history	Yes	Unknown

Abbreviation: FMF, familial Mediterranean fever.

FMF patients, 1,605 patients were in the attack-free phase and 232 patients in the attack phase, and 143 patients were assessed both during attack-free and attack phases. The mean age was less than 18 years in seven studies^{16–20,24,29} and more than 18 years in four studies.^{9,27,30,31} One study²⁸ did not report the mean age. The dose of colchicine also varied in different studies and ranged from 0.5 to 2 mg per day. In all studies, except for one that was unclear,¹⁹ patients were treated with colchicine at the time of the study.

Comparison of Neutrophil-to-Lymphocyte Ratio

As shown in **-Table 3**, five studies^{16–18,20,28} made comparisons among three groups (attack, attack-free, and control), while in the rest there were only two study groups (attackfree and control). In all publications with three study groups, NLR was significantly higher in the attack group compared with the attack-free and control groups. Among studies with two-group comparison, only two articles^{24,27} showed no significant differences in NLR between the attack-free and control groups, while in all other studies the amount of NLR

First author	Numb	er of pa	Number of participant	Gender	L					Age of participe	Age of participants (mean \pm SD)	(Age at	Consumption	Dosage of
	AF	A	Control	AF		A		Control	-	AF	A	Control	diagnosis (mean + SD)	of colchicine at the time	colchicine (ma/d)
				Male	Female	Male	Female	Male	Female					of study	
Yorulmaz ¹⁶	143 ^a	143 ^a	143	83	60	83	60	83	60	164.62 ± 51.20 mo	164.62 ± 51.20 mo	164.92 ± 51.10 mo	98.10± 49.11 mo	Yes	0.5–2
Kholoussi ²⁴	42	I	20	23	19	ı	I	11	6	2-7 y	1	2-14 y	Unknown	Yes	Unknown
Basaran ¹⁷	06	70	74	35	55	29	41	38	36	$\begin{array}{c} 12.81 \pm 3.88 \\ y \end{array}$	11.56 ± 4.30 y	$\begin{array}{c} 10.74\pm4.01\\ y\end{array}$	Unknown	Yes	A: 1.03; AF: 1.09
Çakar ³⁰	69	I	35	58	11	I	I	30	5	$\begin{array}{c} 28.30\pm 4.90\\ y\end{array}$	1	$\begin{array}{c} 27.00\pm8.00\\ y\end{array}$	Unknown	Yes	1.1
Daglar ²⁷	33	I	32	0	33	I	I	0	32	$\begin{array}{c} \textbf{25.00} \pm \textbf{4.63} \\ \textbf{y} \end{array}$	I	$\begin{array}{c} 28.50\pm7.20\\ y\end{array}$	Unknown	Yes	1-2
Kelesoglu ¹⁸	509	61	73	239	270	33	28	34	39	$11.20\pm4.46~y$	$9.80\pm4.63~y$	$\begin{array}{c} 10.40\pm5.80\\ y\end{array}$	Unknown	Yes	Unknown
Celikbilek ²⁸	53	26	36	50 ^b	29 ^b	50 ^b	29 ^b	12	24	Unknown	Unknown	Unknown	Unknown	Yes	1.5
Duksal ¹⁹	343	I	199	178	165	I	1	53	146	10.54 ± 3.61 y	I	$\begin{array}{c} 10.21\pm2.98\\ y\end{array}$	Unknown	Unknown	Unknown
Özer ²⁹	153	I	06	78	75	I	I	49	41	12.33 ± 4.41 y	I	$\begin{array}{c} 10.96\pm3.45\\ y\end{array}$	Unknown	Yes	0.5-1.5
Uluca ²⁰	157	75	77	73	84	33	42	38	39	9.10 ± 3.60 y	$\begin{array}{c} \textbf{8.40} \pm \textbf{3.50} \\ \textbf{y} \end{array}$	$\begin{array}{c} 8.30\pm3.50\\ y\end{array}$	Unknown	Yes	0.5–2
Uslu ⁹	94	I	60	30	64	I	I	20	40	$\begin{array}{c} \textbf{29.90} \pm \textbf{12.20} \\ \textbf{y} \end{array}$	I	$\begin{array}{c} 31.30\pm9.40\\ y\end{array}$	$\begin{array}{c} 23.10\pm12.80\\ y\end{array}$	Yes	1.43
Ahsen ³¹	62	I	41	28	34	I	I	14	27	$\begin{array}{c} 32.64 \pm 10.42 \\ y \end{array}$	I	$\begin{array}{c} \textbf{35.56} \pm \textbf{7.46} \\ \textbf{y} \end{array}$	Unknown	Yes	1.5
Abbraviations: A attack AF attack-free	attack.	AF atta	ick-free												

Table 2 Characteristics of the studied population

Abbreviations: A, attack; AF, attack-free. ^aCases in A and AF groups were same. ^bThe gender number was reported for all cases without discriminating between A and AF cases.

First author	Age	e of participants (mean \pm	SD)	p-Value				
	AF	A	Control					
Yorulmaz ¹⁶	2.88±2.9	3.01 ± 1.17	1.59±0.66	A and AF (0.001) A and control (0.001) AF and control (0.431)				
Kholoussi ²⁴	1.23 ± 0.62	-	0.80 ± 0.16	0.10				
Basaran ¹⁷	1.71±0.83	4.1±3.11	1.91±1.86	AF and A (<0.001) A and control (<0.001) AF and control (0.457)				
Çakar ³⁰	2.5 ± 1.6	-	1.9 ± 0.6	0.037				
Daglar ²⁷	1st trimester: 3.6 (1.70–7.06) 2nd trimester: 4.3 (2.0–9.55)	-	1st trimester: 3.25 (1.90–7.22) 2nd trimester: 3.54 (1.97–8.93)	1st trimester (0.23) 2nd trimester (0.19)				
Kelesoglu ¹⁸	1.47	4.1	1.36	AF and A (<0.001) A and control (<0.001) AF and control (0.740)				
Celikbilek ²⁸	1.83 (2.23–1.21)	2.95 (3.46-1.91)	1.63 (2.23–1.41)	0.004				
Duksal ¹⁹	1.55 ± 1.92	-	$\textbf{0.49} \pm \textbf{1.08}$	<0.0001				
Özer ²⁹	1.7 ± 1.99	-	$\textbf{0.45} \pm \textbf{1.26}$	<0.0001				
Uluca ²⁰	0.8±1.6	2.2±2.6	1.3±1.8	AF and A (<0.001) A and control (<0.001) AF and control (0.76)				
Uslu ⁹	0.61 ± 2.06	-	0.42 ± 1.59	<0.0001				
Ahsen ³¹	0.86 ± 2.21	-	$\textbf{0.59} \pm \textbf{1.68}$	<0.0001				

Table 3 Comparison of NLR values between attack, attack-free, and control groups in different studies

Abbreviations: A, attack; AF, attack-free.

in the attack-free group was significantly higher than the control.

Appraisal Results

As all studies were case–control, an appropriate checklist for this type of study was used for quality assessment. As obvious from the results in **~Table 4**, only two studies^{18,24} had notably lower quality compared with the other included articles. However, as all studies had our intended statistical analysis as inclusion criteria, none of them were excluded in this section.

Meta-analysis and Heterogeneity Analysis

- Figs. 2 and **3** show the forest plot of the studies included in the meta-analysis in two subgroups, including FMF attack and FMF attack-free groups. Assessment of these 10 studies showed that the mean NLR in the attack-free group was 0.482 higher than the control group and this difference was reported to be significant (p < 0.0001; standard difference in means = 0.482). Also, the pooled analysis of four studies that compared the NLR during attack period with normal controls demonstrated that the mean NLR in the attack group was 0.853 higher than the control group and this difference was significant (p = 0.001; standard difference in means = 0.483).

The heterogeneity analysis reported an l^2 value of 73.30 (Q = 30.568 and p < 0.0001) for attack-free and normal comparison and an l^2 value of 90.18 (Q = 33.718 and p < 0.0001)

for attack and control comparison. **Fig. 4** also shows the funnel plot of the included study, which shows no evident bias due to the symmetry of the triangle.

Discussion

Our results demonstrated that the NLR values were significantly higher in FMF patients compared with the normal controls; this was consistent for patients both during the attack and during attack-free periods. It implies that NLR may be an indicator of both clinical and subclinical inflammation as it was significantly higher during both disease phases.

The different mutations of the *MEFV* gene pyrin domain cause autoinflammation in FMF patients. The mutation in this domain triggers assembly of inflammasomes^{32,33} and subsequent caspase-1 activation and interleukin 1 (IL-1) release. IL-1, along with interleukin 6 (IL-6) and tumor necrosis factor α , increases the level of acute-phase reactants such as ESR, CRP, and fibrinogen.^{8,29} These acute-phase reactants usually return to their normal value during attack-free period.^{9,29} However, the subclinical inflammation still exists in 30% of the patients during the attack-free phase. The inspection of this undercover inflammation is very important, as it is the cause of different complications of FMF such as amyloidosis, anemia, splenomegaly, and osteopenia.^{16,17} Therefore, studies have tried to propose a suitable marker that can assess this inflammation.^{9,20,29,34,35}

	10. Was riod appropriate ong statistical e analysis used?	7	٨	٨	~	Y	Y	٨	٨	٨	٨	7	Y
	9. Was the exposure period of interest long enough to be meaningful?	~	n	n	Л	Л	n	n	n	n	n	×	n
	8. Were outcomes assessed in a standard, valid, and reliable way for cases and controls?	٨	n	А	٨	А	n	n	n	۸	۸	٨	А
	7. Were strategies to deal with confounding factors stated?	٢	Z	Y	٨	۲	П	Y	Y	Y	Y	٢	Y
	6. Were confounding factors identified?	¥	Z	Y	¥	Y	N	Y	Y	Y	Y	¥	Y
	5. Was exposure measured in the same way for cases and controls?	٢	n	٨	Y	Y	n	٨	٨	٨	٨	٢	n
	 Was exposure measured in a standard, valid, and reliable way? 	Y	n	Х	٨	Х	n	Х	Х	Х	Х	Y	n
}	 Were the same criteria used for identification of cases and controls? 	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
l	2. Were cases and controls matched appropriately?	۲	٢	Z	z	Z	Z	Z	Z	٢	٢	۲	٢
	1. Were the groups comparable other than the presence of disease in cases or the absence of disease in controls?	z	Ν	۲	Y	۲	۲	۲	۲	Ν	Ν	z	Ν
	First author	Yorulmaz ¹⁶	Kholoussi ²⁴	Basaran ¹⁷	Çakar ³⁰	Daglar ²⁷	Kelesoglu ¹⁸	Celikbilek ²⁸	Duksal ¹⁹	Özer ²⁹	Uluca ²⁰	Uslu ⁹	Ahsen ³¹

Abbreviations: N, no; NA, not applicable; U, unclear; Y, yes.

NLR is a cheap, easily accessible, and convenient-to-use tool for detecting inflammation. Most FMF patients have a complete blood cell count with differential analysis, and only with an easy division, the NLR marker can be measured and used for detecting the underlying inflammation. Moreover, this marker provides information about the two immune pathways of adaptive and acquired immune systems. In fact, neutrophils are markers of acute inflammation, lymphocytes show the prolonged part of the inflammatory pathway, and their ratio can demonstrate the state of acute or chronic inflammation. In other words, higher NLR values demonstrate that the acute inflammation persists, as studies have observed in FMF attack cases.^{36,37} Our analysis showed that the NLR level was significantly higher in both FMF attack and FMF attack-free patients compared with that in normal controls. One of the included studies in our systematic review conducted by Özer et al²⁹ reported a strong positive correlation between CRP, as an acute-phase reactant, and NLR. A previous study by Gasparyan et al³⁸ reported that NLR is applicable in rheumatologic diseases with predominant neutrophil inflammations such as Behçet's disease and FMF.

All these findings can be a beacon to the future and offer a suitable cutoff for NLR in the assessment of subclinical inflammatory status in those with inflammation persisting during the attack-free phase. This is particularly beneficial as other inflammatory markers, such as ESR, CRP, and SAA, cannot detect subclinical inflammation in a small portion of FMF patients, and are also more expensive than NLR.²⁹ However, the hypothesis that NLR can be more sensitive than acute-phase reactants remains, and further investigations are needed to compare these markers.

We postulate that patients with high levels of NLR may need higher doses of colchicine, to control their inflammation. However, more studies are needed to assess and confirm these hypotheses. The clinical consequences of an elevated NLR as a marker of subclinical inflammation in FMF need further investigation. For instance, one of the reviewed studies reported that NLR is associated with development of amyloidosis⁹; however, further studies were found to be inconclusive. Another limitation of NLR is that it is a marker of inflammation and thus can be raised in any inflammatory cases, including an infectious disease³⁹; however, acute-phase reactant proteins such as ESR and CRP have this limitation, too.⁴⁰ Moreover, like acute-phase reactants, NLR is dependent on the age of the patients,⁴¹ which should also be considered.

One of the studies was conducted on a population of pregnant women and, due to the probability of imposing bias, was only entered in the systematic review and not in the meta-analysis. It reported no significant difference between those with attack-free FMF and normal controls. It should be considered that pregnancy itself has an immunomodulatory effect and this may affect the underlying inflammation.⁴² Moreover, it is reported that FMF severity remains unchanged or even get better in two-thirds of pregnancies.⁴³

Although our study provided a possible valuable marker for detecting inflammation in FMF patients, there is a further need to identify a set point for detecting inflammation.

Table 4 Quality of studies based on the Joanna Briggs checklist

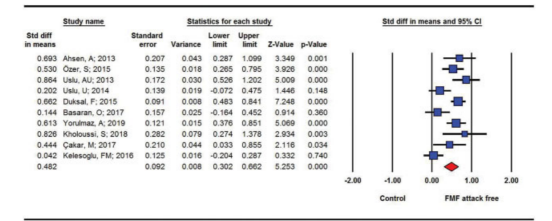


Fig. 2 Forest plot comparing FMF attack-free and control groups. FMF, familial Mediterranean fever.

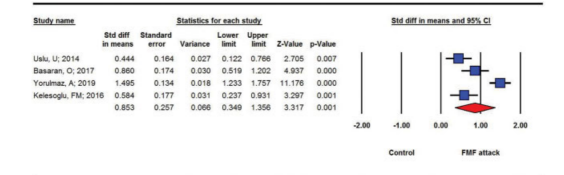
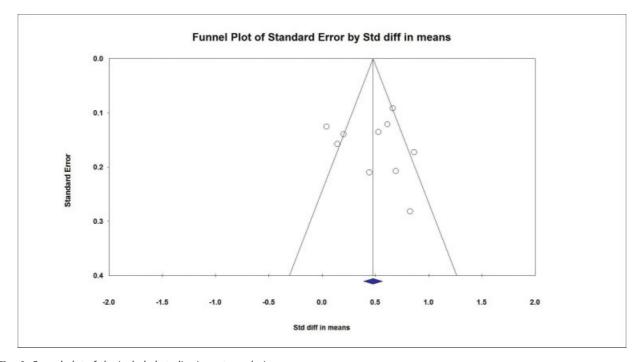


Fig. 3 Forest plot comparing FMF attack and control groups. FMF, familial Mediterranean fever.





Moreover, the available clinical criteria such as Tel Hashomer criteria²⁵—which was used in all studies in our review except one—cannot discriminate between FMF and some other rare autoinflammatory diseases.⁴⁴ Due to this fact, a definite

diagnosis of FMF is made only by genetic detection of its mutations.^{45,46} Two of the reviewed publications in our study lacked genetic confirmation and this can be proposed as a limitation.^{27,28} Different inflammatory status may also

affect the NLR level,⁴⁷ which is another limitation in our analysis—as some studies considered this factor in their inclusion and exclusion criteria, while others did not.

Furthermore, it is proposed that colchicine can halter both clinical and subclinical inflammation.⁴⁸ This may have confounding effects on the NLR values in FMF patients, and may be perceived as another shortcoming of our study. However, it is believed that colchicine mainly compromises neutrophil activity by interfering with activation of the microtubules and their chemotaxis and has no effect on the neutrophil count.^{49,50} This should be considered in future studies, where there may be a need for further adjustment.

Conclusion

Our results showed that NLR values were significantly higher in both attack and attack-free groups compared with the normal controls. As acute-phase reactants are more expensive than NLR and cannot detect subclinical inflammation, NLR can be a valuable and cheap alternative. Further studies are necessary to propose a cutoff for NLR in FMF, taking into consideration the importance of inflammation in their prognosis. Further research in this field to uncover more information is encouraged.

Conflict of Interest None declared.

References

- 1 Özen S. Update on the epidemiology and disease outcome of familial Mediterranean fever. Best Pract Res Clin Rheumatol 2018; 32(02):254–260
- 2 Cekin N, Akyurek ME, Pinarbasi E, Ozen F. MEFV mutations and their relation to major clinical symptoms of familial Mediterranean fever. Gene 2017;626:9–13
- 3 Touitou I. The spectrum of familial Mediterranean fever (FMF) mutations. Eur J Hum Genet 2001;9(07):473–483
- 4 Ozdogan H, Ugurlu S. Familial Mediterranean fever. Presse Med 2019;48(1 Pt 2):e61-e76
- 5 Beheshtian M, Izadi N, Kriegshauser G, et al. Prevalence of common MEFV mutations and carrier frequencies in a large cohort of Iranian populations. J Genet 2016;95(03):667–674
- 6 Sohar E, Gafni J, Pras M, Heller H. Familial Mediterranean fever. A survey of 470 cases and review of the literature. Am J Med 1967; 43(02):227–253
- 7 Barut K, Sahin S, Adrovic A, et al. Familial Mediterranean fever in childhood: a single-center experience. Rheumatol Int 2018;38 (01):67–74
- 8 Yildirim K, Uzkeser H, Keles M, et al. Relationship between serum interleukin-1 β levels and acute phase response proteins in patients with familial Mediterranean fever. Biochem Med (Zagreb) 2012;22(01):109–113
- 9 Uslu AU, Deveci K, Korkmaz S, et al. Is neutrophil/lymphocyte ratio associated with subclinical inflammation and amyloidosis in patients with familial Mediterranean fever? BioMed Res Int 2013; 2013:185317
- 10 Ben-Zvi I, Livneh A. Chronic inflammation in FMF: markers, risk factors, outcomes and therapy. Nat Rev Rheumatol 2011;7(02): 105–112
- 11 Xu S, Ma Y, Wu M, et al. Neutrophil lymphocyte ratio in patients with ankylosing spondylitis: a systematic review and metaanalysis. Mod Rheumatol 2020;30(01):141–148

- 12 Erre GL, Paliogiannis P, Castagna F, et al. Meta-analysis of neutrophil-to-lymphocyte and platelet-to-lymphocyte ratio in rheumatoid arthritis. Eur J Clin Invest 2019;49(01):e13037
- 13 Ma L, Zeng A, Chen B, Chen Y, Zhou R. Neutrophil to lymphocyte ratio and platelet to lymphocyte ratio in patients with systemic lupus erythematosus and their correlation with activity: a metaanalysis. Int Immunopharmacol 2019;76:105949
- 14 Ghodsi A, Mirimoghaddam MM, Sarabi M, et al. Neutrophil-tolymphocyte ratio as a novel and valuable marker for assessing disease severity in ulcerative colitis, multiple sclerosis, and Kawasaki disease: a review. J Basic Res Med Sci 2020;7(03):62–70
- 15 Zhang Z, Su Q, Zhang L, Yang Z, Qiu Y, Mo W. Diagnostic value of hemoglobin and neutrophil-to-lymphocyte ratio in Behcet disease. Medicine (Baltimore) 2019;98(52):e18443
- 16 Yorulmaz A, Akbulut H, Taş SA, Tıraş M, Yahya İ, Peru H. Evaluation of hematological parameters in children with FMF. Clin Rheumatol 2019;38(03):701–707
- 17 Basaran O, Uncu N, Celikel BA, Aydın F, Cakar N. Assessment of neutrophil to lymphocyte ratio and mean platelet volume in pediatric familial Mediterranean fever patients. J Res Med Sci 2017;22:35
- 18 Kelesoglu FM, Aygun E, Okumus NK, et al. Evaluation of subclinical inflammation in familial Mediterranean fever patients: relations with mutation types and attack status: a retrospective study. Clin Rheumatol 2016;35(11):2757–2763
- 19 Duksal F, Alaygut D, Güven AS, et al. Neutrophil-lymphocyte ratio in children with familial Mediterranean fever: original article. Eur J Rheumatol 2015;2(01):20–23
- 20 Uluca Ü, Ece A, Şen V, et al. Usefulness of mean platelet volume and neutrophil-to-lymphocyte ratio for evaluation of children with familial Mediterranean fever. Med Sci Monit 2014; 20:1578–1582
- 21 Moher D, Liberati A, Tetzlaff J, Altman DGPRISMA Group. Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. PLoS Med 2009;6(07):e1000097
- 22 Omranzadeh A, Baradaran A, Ghodsi A, et al. Comparing neutrophil to lymphocyte ratio between a group of familial Mediterranean fever (FMF) and a healthy control group. PROSPERO [Internet] 2018. Available from: https://www.crd.york.ac.uk/prospero/display_record.php?ID=CRD42018092613
- 23 JB Institute. JBI critical appraisal checklist for case control studies. Joanna Briggs Institute Reviewers' Manual 2016
- 24 Kholoussi S, Kholoussi N, Zaki ME, et al. Immunological evaluation in patients with familial Mediterranean fever. Open Access Maced J Med Sci 2018;6(02):310–313
- 25 Livneh A, Langevitz P, Zemer D, et al. Criteria for the diagnosis of familial Mediterranean fever. Arthritis Rheum 1997;40(10): 1879–1885
- 26 Yalçinkaya F, Özen S, Ozçakar ZB, et al. A new set of criteria for the diagnosis of familial Mediterranean fever in childhood. Rheumatology (Oxford) 2009;48(04):395–398
- 27 Daglar K, Kirbas A, Timur H, et al. Subclinical inflammation and simple blood parameters in pregnant with familial Mediterranean fever. J Clin Anal Med 2016;7(05):634–638
- 28 Celikbilek M, Dogan S, Akyol L, et al. Neutrophil-lymphocyte ratio in patients with familial Mediterranean fever. J Clin Lab Anal 2015;29(01):80–83
- 29 Özer S, Yılmaz R, Sönmezgöz E, et al. Simple markers for subclinical inflammation in patients with familial Mediterranean fever. Med Sci Monit 2015;21:298–303
- 30 Çakar M, Akhan M, Doğan T, et al. Investigation of the arterial stiffness and associated factors in patients with familial Mediterranean fever. Anatol J Cardiol 2017;17(02):132–138
- 31 Ahsen A, Ulu MS, Yuksel S, et al. As a new inflammatory marker for familial Mediterranean fever: neutrophil-to-lymphocyte ratio. Inflammation 2013;36(06):1357–1362
- 32 Bilginer Y, Akpolat T, Ozen S. Renal amyloidosis in children. Pediatr Nephrol 2011;26(08):1215-1227

- 33 Savic S, Dickie LJ, Battellino M, McDermott MF. Familial Mediterranean fever and related periodic fever syndromes/autoinflammatory diseases. Curr Opin Rheumatol 2012;24(01):103–112
- 34 Sakallı H, Kal O. Mean platelet volume as a potential predictor of proteinuria and amyloidosis in familial Mediterranean fever. Clin Rheumatol 2013;32(08):1185–1190
- 35 Makay B, Türkyilmaz Z, Ünsal E. Mean platelet volume in children with familial Mediterranean fever. Clin Rheumatol 2009;28(08): 975–978
- 36 Sharaiha RZ, Halazun KJ, Mirza F, et al. Elevated preoperative neutrophil:lymphocyte ratio as a predictor of postoperative disease recurrence in esophageal cancer. Ann Surg Oncol 2011; 18(12):3362–3369
- 37 Azab B, Jaglall N, Atallah JP, et al. Neutrophil-lymphocyte ratio as a predictor of adverse outcomes of acute pancreatitis. Pancreatology 2011;11(04):445–452
- 38 Gasparyan AY, Ayvazyan L, Mukanova U, Yessirkepov M, Kitas GD. The platelet-to-lymphocyte ratio as an inflammatory marker in rheumatic diseases. Ann Lab Med 2019;39(04):345–357
- 39 Forget P, Khalifa C, Defour J-P, Latinne D, Van Pel M-C, De Kock M. What is the normal value of the neutrophil-to-lymphocyte ratio? BMC Res Notes 2017;10(01):12
- 40 Bray C, Bell LN, Liang H, et al. Erythrocyte sedimentation rate and C-reactive protein measurements and their relevance in clinical medicine. WMJ 2016;115(06):317–321
- 41 Li J, Chen Q, Luo X, et al. Neutrophil-to-lymphocyte ratio positively correlates to age in healthy population. J Clin Lab Anal 2015; 29(06):437–443

- 42 Challis JR, Lockwood CJ, Myatt L, Norman JE, Strauss JF III, Petraglia F. Inflammation and pregnancy. Reprod Sci 2009;16(02):206–215
- 43 Eviatar T, Zaks N, Kukuy O, Livneh A, Lidar M. PW01–010–The effect of pregnancy on disease course in FMF. Pediatr Rheumatol Online J 2013;11(Suppl 1):A63
- 44 Karacan İ, Uğurlu S, Tolun A, Tahir Turanlı E, Ozdogan H. Other autoinflammatory disease genes in an FMF-prevalent population: a homozygous MVK mutation and a novel heterozygous TNFRSF1A mutation in two different Turkish families with clinical FMF. Clin Exp Rheumatol 2017;35(06, Suppl 108):75–81
- 45 Samuels J, Aksentijevich I, Torosyan Y, et al. Familial Mediterranean fever at the millennium. Clinical spectrum, ancient mutations, and a survey of 100 American referrals to the National Institutes of Health. Medicine (Baltimore) 1998;77(04):268–297
- 46 Booty MG, Chae JJ, Masters SL, et al. Familial Mediterranean fever with a single MEFV mutation: where is the second hit? Arthritis Rheum 2009;60(06):1851–1861
- 47 Balta S, Demirkol S, Unlu M, Arslan Z, Celik T. Neutrophil to lymphocyte ratio may be predict of mortality in all conditions. Br J Cancer 2013;109(12):3125–3126
- 48 Corsia A, Georgin-Lavialle S, Hentgen V, et al. A survey of resistance to colchicine treatment for French patients with familial Mediterranean fever. Orphanet J Rare Dis 2017;12(01):54
- 49 Soriano A, Manna R. Familial Mediterranean fever: new phenotypes. Autoimmun Rev 2012;12(01):31–37
- 50 Stroka KM, Hayenga HN, Aranda-Espinoza H. Human neutrophil cytoskeletal dynamics and contractility actively contribute to trans-endothelial migration. PLoS One 2013;8(04):e61377