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Association of Rs763780 Polymorphism of Gene IL17F with the Risk of Developing Rheumatoid Arthritis in Uzbekistan 2

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Abstract 6

As a result of the study conducted, it was witnessed that the G allele and the heterozygous A / G genotype of the IL17F gene (rs763780) among patients with RA are significantly higher than in the control group. In particular, the most significant discrepancies were registered in 9 patients with articular-visceral form of the disease whom the G allele exceeded the proportion 10 of carriage in the control statistically significantly 2.58 times (2 = 4.512; P = 0.037; OR = 11 2.58;9512

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Index terms— rheumatoid arthritis (RA), rs763780 polymorphism of gene IL17F, carriage, allele, genotype, 14 developent risk. 15

Introduction 1 16

mong all the variety of inflammatory diseases of joints, rheumatoid arthritis (RA) is amid the most prevalent 17 nosology which affects about 1% of adult population worldwide [5,7]. Along with this, within diverse populations 18 of the world, large epidemiological studies have established differences in the prevalence of the disease [6,7,14]. 19 Hence, on a frequently basis, RA occurs in American Indians (up to 7%) while among other nationalities the 20 incidence of the disease is in the range of 0.2-0.4% [14]. Pathogenic aspect of RA development remains poorly 21 22 perceived. However, it is a fact that in the implementation of the pathological process that gives a rise to the 23 disease, a connection is observed in conformity with a number of factors such as the impact of the environment, bad habits, microbial and viral agents, genetic polymorphisms, etc. [2,8,9,16]. Inflammation, being the basis 24 for the development of RA, come to light with transformations in the articular tissue. The progression of 25 inflammation in the subsequent passes to the bone tissue inducing its destruction [3]. The bulk of factors are 26 involved in the regulation of inflammatory processes among which the leading role is played by polymorphic 27 variants of a number of pro-inflammatory cytokines (IL17F, etc.) [10,12]. Meanwhile, the results of studies on 28 the assessment of participation in increasing the risk of developing RA are ambiguous [4,11,15,10,12,18]. Thus, 29 researchers C. N. Carvalho (2015) did not find a correlation between the IL-17F (7488T / C) gene polymorphism 30 and the development and severity of RA [4]. Similar results with no differences between the IL17F gene and 31 the development of articular and extraarticular forms of the disease were obtained by A. Pawlik (2016) when 32 33 investigating Polish patients with RA (n = 422) [15]. S. Louahchi (2016) also did not find an association of 34 IL17F (rs763780, rs2397084) with susceptibility to RA among Algerians (n = 343) [15]. Nevertheless, the results 35 of studies by Y. H. Lee, S.C. Bae, (2017), O. S. Marwa (2017), M. Shao (2020) confirm the role of the IL17F gene in the development of RA [10,12,18]. The resulting disagreements are possibly related to the traits of the 36 studied populations. Corollary analysis of the studies performed delineates ambiguous conclusions regarding the 37 contribution of the IL-17F gene to the mechanisms of RA onset. In this regard, it is of significant magnitude 38 to conduct supplementary examinations to assess the relationship of this gene with the development of RA. 39 Furthermore, the data obtained will assist to better conceive and explain the degree of participation of the 40 41

IL-17F gene in the formation of this complex disease.

42 **2** II.

⁴³ **3** Material and Methods

The study encompassed 106 adults (combined general group) of unrelated patients living in the Republic of 44 Uzbekistan with a diagnosis of RA verified taking into account the ACR / EULAR criteria (2010) [1]. All 45 patients, in the period of 2018-2021, were examined and hospitalized at 3 clinics of the Tashkent Medical Academy 46 (Uzbekistan, Tashkent), which, depending on the form of the disease, were stratified into two subgroups 1A (n 47 48 = 74) -patients with articular RA and 1B (n = 32) -patients with articular-visceral form of RA. When it comes 49 to control, conditionally healthy individuals (n = 109) without a history of autoimmune diseases, comparable in 50 sex, age and living in the territory of the re-public, were examined. In order to comply with ethical standards, informed consent was resulted from all individuals included in the study. For molecular genetic studies, DNA was 51 isolated from venous blood leukocytes using the "AmpliPrime RIBOprep, Russia" kit according to the standard 52 method [13]. Detection of rs763780 polymorphism of the IL17F gene (SYNTOL, Russia) was carried out by 53 SNP-PCR (Applied Biosystems, thermocycler 2720 (USA)) with verification of the specificity and number of 54 amplified fragments by electrophoretic method in agarose gel. The obtained data were statistically processed 55 using the "OpenEpi 2009, Version 9.3" software package. 56

57 III.

58 4 Results and Discussion

Table ??: Analysis of allelic distribution and genotypic frequencies of the IL 17F (rs763780) gene polymorphism
 in the studied groups

If the escalation in the proportion of allele G carriage of the polymorphic variant of the IL-17F gene (rs763780) 61 in the 1st combined group of patients with RA and in the "1A" subgroup of patients with the articular form 62 of the disease tended to amplify the risk of developing RA by almost twice ($2^2 = 3.344$; P = 0.07; OR = 63 1.919; 95% CI: 0.954-3.859) and 1.65 times (?2 = 1.57; P = 0.211; OR = 1.65; 95% CI: 0.756-3.594). Then 64 in the subgroup of patients "1B" with the articularvisceral form of RA, the risk of developing the disease was 65 statistically significantly boosted by 2.58 times (2 = 4.512; P = 0.037; OR = 2.58; 95% CI: 1.076-6.188) (Table 66 2). Genotype A / A carriage proportion of the polymorphic variant of the IL-17F gene (rs763780) in all groups 67 enabled particular differences: in the combined group of RA patients it was 79.3%, in subgroups "1A" and 68 "1B" -81.1% and 75.0%, respectively, and in the control group -88.1%. Along with this, the frequency of the 69 heterozygous genotype A / G had a clear discrepancy in the groups of patients (combined group RA -19.8%; 70 "1A" subgroup -18.9%, "1B" -21.9%) compared with the control (11.9%). In addition, it is important to note 71 72 that the mutant G / G genotype was recorded only among patients with the articular-visceral form of the disease (subgroup 1B), the proportion of which was 3.1%. The decrease in the frequency of the wild A / A genotype 73 among patients in contrast to the control did not differ statistically (in the combined group of RA patients -?2 74 = 3.073; P = 0.084; OR = 0.517; 95% CI: 0.247-1.081; in the "1A" subgroup -?2 = 1.713; ? = 0.193; OR = 0.084; O 75 0.58; 95% CI: 0.257-1.311 and in subgroup "1B" -?2 = 3.336; ? = 0.072; OR = 0.406; 95% CI: 0.154-1.068) 76 (Table 2). The distribution of genotypes of the polymorphic variant of the IL-17F gene (rs763780) in the studied 77 groups did not deviate from the Hardy-Weinberg equilibrium (P > 0.05). In particular, genotypes A / A, A / G, 78 79 and G / G in the combined group of RA patients were 0.79%, 0.2%, and 0.01% while in the control group their 80 values amounted to 0.88%, 0.12%, and 0.0%, respectively. Analysis of allele distribution frequencies represented a 81 greater registration of the proportion of carriers of the G allele among RA patients in the general group compared to controls (10.8% versus 6.0%). There was ascendance in the frequency of this indicator due to an increase in 82 their share in both subgroups of patients which reached 9.5% in subgroup 1A of patients with articular RA 83 and 14.1% in subgroup 1B of patients with articular-visceral RA (Table ??). Meanwhile, the differences in the 84 proportion of the heterozygous genotype A / G carriage in the groups of RA patients compared with the controls 85 turned out to be more paramount. So, if in the 1st combined group of RA patients this genotype boosted 1.8 86 times (?2 = 2.509; P = 0.119; OR = 1.824; 95% CI: 0.867-3.837); then in "1A" subgroup 1.72 times (?2 = 1.713; 87 ? = 0.193; OR = 1.723; 95% CI: 0.763-3.892) and in subgroup "1B" more than twice (?2 = 2.011; ? = 0.165; 88 OR = 2.068; 95% CI: 0758-5.645). The obtained differences indicate the presence of a clear tendency towards 89 an increased risk of RA formation in carriers of the A / G genotype. Perhaps, with a larger coverage of the 90 91 sample under study, disparities could be reliably significant. Consequently, differences that we established in the 92 frequency of distribution of the G allele and the A / G genotype among RA patients compared to controls allow 93 us to determine their role in proliferating the risk of developing the disease, especially the articular-visceral form.

94 5 Group n

⁹⁵ 6 Allele frequency Genotype distribution frequency

96 A G A /A A/G G /G n % n % n % n % n % First combined

97 **7** IV.

98 8 Conclusion

Rheumatoid arthritis is induced by a complex autoimmune disease, the origin of which is complicated by the 99 lack of pathological mechanisms [14]. However, the results of modern studies emphasize the special role of 100 genetic polymorphisms of genes of proinflammatory cytokines which are involved not only in increasing the risk 101 of developing RA, but also in the severity of its course [7]. IL17F is considered as one of these genes, which 102 can serve as a potential candidate gene leading to the development of RA [10,12]. Meanwhile, in relation to this 103 point of view, the views of researchers differ. So, if C. N. Carvalho (2015), S. Louahchi (2016), A. Pawlik (2016) 104 [4,11,15] did not find an association between the IL17F gene and the onset of RA in their studies, the results of 105 later works by Y. H. Lee, S.C. Bae, (2017), O. S. Marwa (2017), M. Shao (2020) indicate the participation of 106 the IL17F gene in the mechanisms of RA formation [10,12,18]. Taking into account the existing disagreements in 107 this regard, we found it interesting to assess the participation degree of the IL17F (rs763780) gene polymorphism 108 in the risk of developing RA among the population of the Republic of Uzbekistan. As a result of our studies, we 109 have encountered in that the G allele and the heterozygous A / G genotype of the IL17F gene (rs763780) among 110 patients with RA are significantly higher than in the control group. In particular, the most significant differences 111 112 were found in patients with articular-visceral form of the disease in which the G allele exceeded the proportion of carriage in the control statistically significantly 2.58 times (?2 = 4.512; P = 0.037; OR = 2.58; 95% CI: 1.076 113 -6.188) and on the part of the heterozygous genotype A / G there was a clear tendency to increase its frequency 114 by more than twice (2 = 2.011; P = 0.165; OR = 2.068; 95% CI: 0758-5.645) which in turn indicates the possible 115 participation of this polymorphism in the pathogenesis of the disease. Moreover, only among patients with this 116 form of RA was the carriage of the mutant genotype G / G (3.1%); 2 = 2.011; P = 0.165; OR = 2.068; 95% CI: 117 0758-5.645). The obtained data emphasize the role of the polymorphic variant of the IL17F gene (rs763780) in 118 the development of RA among the population of Uzbekistan. In addition, these results contribute to a deeper 119 understanding of the pathogenetic mechanisms of RA formation which is overly consequential in predicting the 120 development of RA and searching for the most effective methods of treating the disease.

 $\mathbf{2}$

Groups scrutiny	under	Alleles and	OR	Statistical difference compared to control 95% CI: ? 2		
		geno-				
		Δ	3 344	0.072	0 521	0 259 -1 048
Group 1	\mathbf{R}^{2}	G	3,344	0,072	1 010	0.954 - 3.859
(n=106) pat	ients	u	3,311	0,012	1,010	0,001-0,000
(11 100) par	101100	?/?	3,073	0,084	0,517	0,247 - 1,081
		?′/G	2,509	0,119	1,824	0,867 -3,837
Subgroup 1 $(n=74)$?, R?	Á	1,577	0,211	0,607	0,278 -1,323
articular form		G	1,577	0.211	1,648	0.756 - 3.594
		?/?	1,713	0,193	0,580	0,257 -1,311
		?′/G	1,713	0,193	1,723	0,763 -3,892
Subgroup articular-vis- form of RA	1B, ceral (n =	Á	4,512	0,037	0,388	0,162 -0,929
32)	× ·					
,		G^*	4,512	0,037	2,580	1,076 -6,186
		?/?	3,336	0,072	0,406	0,154 -1,068
		?/G	2,011	0,165	2,068	0,758 - 5,645

?

Figure 1: Table 2 :

121

8 CONCLUSION

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