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#### ORIGINAL ARTICLE

# Assessment of Association between Duffy Blood Group Phenotypes and Susceptibility to Systemic Lupus Erythematosus and its Severity

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\*Corresponding author: Erfan Taherifard School of Medicin, Zand St., Shiraz, Iran Tel: +98-9172810165 Email: erfantaherifard@gmail.com ABSTRACT

**Background:** Considering the role of Duffy antigenic system expression on RBCs as chemokines for inflammatory cytokines, expression of Duffy antigens as different phenotypes was studied in a group of patients with SLE. The association between different phenotypes of Duffy antigens and occurrence of SLE and its severity was assessed.

**Methods:** This cross-sectional study was carried out on 100 patients diagnosed as SLE using the "Systemic Lupus International Collaborating Clinics 2012 (SLICC) classification" criteria and 100 age-matched healthy subjects as control from April to June 2017. Duffy blood group status was determined in both groups. The patients were categorized into three groups of mild, moderate and severe according to "Systemic Lupus Erythematosus Disease Activity Index 2000 (SLEDAI-2K)" scores.

**Results:** The results of this study showed no significant association between the three different phenotypes of Duffy antigen system and increased risk of SLE or its severity.

**Conclusion:** The results of the current study did not approve of any association between three different Duffy phenotypes and increased risk of SLE or its severity.

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## Introduction

Systemic lupus erythematosus (SLE) is a chronic inflammatory disease characterized by multi organ involvement mediated by tissue-binding autoantibodies and immune complexes (1, 2). This autoimmune disease is associated with various clinical manifestations could result in a complex of complications that may vary significantly among patients (3, 4). It has been documented that both genetic and environmental factors play pivotal roles in the pathogenesis and progression of this disease (5). Various mechanisms have been proposed about the pathogenesis of SLE, but the underlying pathophysiological mechanism is not well understood and several gaps remain to be filled in this area (6). Indeed, all of these proposed mechanisms share some key processes and features; one of them is the reduction in the clearance of immune complexes and apoptotic cells and inflammatory cytokines observed in almost all cases of SLE (7-9). Recent studies have shown that there is a significant relationship between the disease severity and the main signs and symptoms of the disease and serum levels of the cytokines such as interleukin (IL)-1, IL-2, IL-6, IL-8, IFN $\gamma$  (10-13).

Duffy antigen receptor chemokine (DARC); also known as Fy glycoprotein (Fy), is a glycosylated membrane protein which is located on the outer surface of erythroid precursors and red blood cells (RBCs) and the lining surface of the post-capillary venules of several organs such as the spleen, liver and kidney (14-18). DARK antigen system also serves as a nonspecific receptor for most proinflammatory chemokines with high affinity for the CXC class of pro-inflammatory chemokines such as IL-8 and CC chemokines (10, 19, 20). As a result, Duffy antigen system can influence plasma levels of the cytokines and take part in the process of leukocyte recruitment and thus can modulate inflammation (21, 22). In other words, DARK receptors are scavengers for chemokines which acts as a "chemokine sink" (23).

The probable association between SLE and Duffy antigen system has not yet been evaluated. We aimed to assess any possible association between SLE and the presence of different Duffy phenotypes.

#### **Materials and Methods**

In this cross-sectional study which was supported by Shiraz University of Medical Sciences, 100 patients suffering from SLE and equal number of healthy individuals who were among volunteer blood donors referring to the blood transfusion center regularly were enrolled. Due to a relatively large number of this population, the process of age- and sex-matching was conducted precisely. Individuals with positive history of SLE in their first-degree relatives and those who were exposed actively or passively to tobacco were excluded from the control group. The process of participation was completely arbitrary. Patients were employed from those who referred to Hafez Medical Clinic of Shiraz and were diagnosed with SLE during April to June 2017.

Diagnosis of SLE was based upon the "Systemic Lupus International Collaborating Clinics 2012 (SLICC) classification criteria" (24, 25). Then the disease activity was measured using "Systemic Lupus Erythematosus Disease Activity Index 2000 (SLEDAI-2K)" and the patient groups were categorized into three groups of mild, moderate and severe depending on the SLEDAI-2K score (26). Then, the patient and control groups were analysed for the assessment of the Duffy blood group status.

Statistical Package for Social Sciences (SPSS) version 11.5 was used for data analysis. Chi-square and Fisher's exact tests were used to compare the phenotype frequencies between case and control group and to assess if there was any association between individual phenotype and severity of SLE in the patients. P<0.05 was considered statistically significant.

#### Results

As stated earlier, the current study was established to assess the association between three different Duffy positive phenotypes and increased risk of developing SLE or its severity. Distribution of different phenotypes of DARC among both groups is shown in Table 1.

The association between the presence of different Duffy phenotypes and SLE was investigated using the Chi-Square test and no significant association (P=0.426) (Table 2). Then, the patient group was classified into three groups in terms of the severity according to SLEDAI-2K score. The association between the severity of the disease among the patients and the presence of the three different Duffy positive phenotypes was evaluated. The results did not support any significant association (P=0.929).

#### Discussion

Duffy antigen/chemokine receptor (DARC) is a glycosylated membrane protein, serving as a non-specific receptor for a variety of chemokines including angiogenic CXC chemokines (27). There are several known antigens in "Duffy blood group system" including  $Fy^a$ ,  $Fy^b$ , Fy3, Fy4, Fy5, and Fy6 (28). The four most commonly used phenotypes for Duffy blood group system are  $Fy^{(a+/b+)}$ ,  $Fy^{(a-/b+)}$ ,  $Fy^{(a+/b-)}$ , and  $Fy^{(a-/b-)}$ . The Duffy null phenotype is uncommon in white people (29).

DARC plays an important role in inflammatory diseases and malignancies. In a study, DARC expression was showed to be boosted in the early phase of rheumatoid arthritis due to its function of neutrophil recruitment (30). In another study, DARC was found to be upregulated in microvascular endothelial cells of the CNS during active phase of diseases such as immune encephalomyelitis and multiple sclerosis (31). Furthermore, another study showed that rs2814778 polymorphism in the gene encoding DARK has an association with high IgE levels; increasing susceptibility to develop asthma and atopy among certain populations of African descent (32). Besides, in another study, this polymorphism was associated with benign neutropenia among people of African ancestry and specific Middle Eastern ethnic groups (33). To support any role of Duffy antigens in disease processes, enhanced expression of DARK by breast cancer cells was proposed

Table 1: Frequency of diffe	erent phenotypes of Duffy	antigen among patients with SI	E and healthy subjects		
	Duffy antigen Phenotypes				
	Fy <sup>a</sup>	Fy <sup>b</sup>	<b>Double positive Fy</b> <sup>(a+/b+)</sup>		
Control group	24	32	44		
Patients with SLE	31	33	36		
Total	55	65	80		

Table 2: Distribution of different phenotypes of Duffy antigens between different groups of patients with SLE according to disease severity

Severity of the disease	Phenotypes			Total	
	Fy <sup>a</sup>	Fy <sup>b</sup>	Fy <sup>(a+/b+)</sup>		
Mild	11	11	11	33	
Moderate	9	8	12	29	
Severe	11	14	13	38	
Total	31	33	36	100	

as a protection against breast cancer (34). The lack of Duffy antigen expression in patients with sickle cell anemia has been suggested as a poor prognostic factor for end-organ damage in these patients (35).

Moreover, the presence of different Duffy phenotypes (a+/b+, a+/b-, and a-/b+) could determine the severity of various diseases; however, our study did not support this hypothesis. In 2018, Fong and colleagues documented that Plasmodium knowlesi has more affinity for binding to Fy<sup>a+/b+</sup> erythrocytes than to Fy<sup>a+/b-</sup> erythrocytes (36). In another study, double positive Duffy phenotype was more common in patients with multiple myeloma rather than healthy subjects. The authors assumed that some cytokine cascades involved in the pathogenesis of multiple myeloma, act through the DARC pathway (37).

The limitations of our study were small sample size and short period of the study which might influence the results of the study. Meanwhile, to find such associations, conducting epidemiological studies across different geographical regions sound necessary.

## Conclusion

The results of the current study did not approve of any association between three different Duffy phenotypes and increased risk of SLE or its severity. Such studies are recommended to elucidate various pathologic cascades involved in inflammatory diseases, giving us an opportunity to interfere in these pathways.

## Conflict of Interest: None declared.

## References

- D'Cruz DP, Khamashta MA, Hughes GR. Systemic lupus erythematosus. Lancet (London, England). 2007;369(9561):587-96.
- 2. Manson JJ, Rahman A. Systemic lupus erythematosus. Orphanet journal of rare diseases. 2006;1:6.
- Cojocaru M, Cojocaru IM, Silosi I, Vrabie CD. Manifestations of systemic lupus erythematosus. Maedica. 2011;6(4):330-6.
- Cervera R, Khamashta MA, Font J, Sebastiani GD, Gil A, Lavilla P, et al. Systemic lupus erythematosus: clinical and immunologic patterns of disease expression in a cohort of 1,000 patients. The European Working Party on Systemic Lupus Erythematosus. Medicine. 1993;72(2):113-24.
- Kyttaris VC. Systemic lupus erythematosus: from genes to organ damage. Methods in molecular biology (Clifton, NJ). 2010;662:265-83.
- 6. Maidhof W, Hilas O. Lupus: an overview of the disease and management options. P & T : a peer-reviewed journal for formulary management. 2012;37(4):240-9.
- Crispín JC, Liossis SN, Kis-Toth K, Lieberman LA, Kyttaris VC, Juang YT, et al. Pathogenesis of human systemic lupus erythematosus: recent advances. Trends in molecular medicine. 2010;16(2):47-57.
- Robak E, Sysa-Jedrzejowska A, Robak T. [Cytokines in systemic lupus erythematosus]. Przeglad lekarski. 1996;53(8):623-6.
- 9. Jacob N, Stohl W. Cytokine disturbances in systemic

lupus erythematosus. Arthritis research & therapy. 2011;13(4):228.

- Gardner L, Patterson AM, Ashton BA, Stone MA, Middleton J. The human Duffy antigen binds selected inflammatory but not homeostatic chemokines. Biochemical and biophysical research communications. 2004;321(2):306-12.
- Yap DY, Lai KN. The role of cytokines in the pathogenesis of systemic lupus erythematosus from bench to bedside. Nephrology (Carlton, Vic). 2013;18(4):243-55.
- 12. Yap DY, Lai KN. Cytokines and their roles in the pathogenesis of systemic lupus erythematosus: from basics to recent advances. Journal of biomedicine & biotechnology. 2010;2010:365083.
- 13. Hrycek E, Franek A, Blaszczak E, Dworak J, Hrycek A. Serum levels of selected chemokines in systemic lupus erythematosus patients. Rheumatology international. 2013;33(9):2423-7.
- 14. Meny GM. The Duffy blood group system: a review. Immunohematology. 2010;26(2):51-6.
- 15. Neote K, Mak JY, Kolakowski LF, Jr., Schall TJ. Functional and biochemical analysis of the cloned Duffy antigen: identity with the red blood cell chemokine receptor. Blood. 1994;84(1):44-52.
- 16. Hadley TJ, Lu ZH, Wasniowska K, Martin AW, Peiper SC, Hesselgesser J, et al. Postcapillary venule endothelial cells in kidney express a multispecific chemokine receptor that is structurally and functionally identical to the erythroid isoform, which is the Duffy blood group antigen. The Journal of clinical investigation. 1994;94(3):985-91.
- 17. Peiper SC, Wang ZX, Neote K, Martin AW, Showell HJ, Conklyn MJ, et al. The Duffy antigen/receptor for chemokines (DARC) is expressed in endothelial cells of Duffy negative individuals who lack the erythrocyte receptor. The Journal of experimental medicine. 1995;181(4):1311-7.
- Lee JS, Frevert CW, Wurfel MM, Peiper SC, Wong VA, Ballman KK, et al. Duffy antigen facilitates movement of chemokine across the endothelium in vitro and promotes neutrophil transmigration in vitro and in vivo. Journal of immunology (Baltimore, Md: 1950). 2003;170(10):5244-51.
- Langhi DM, Jr., Bordin JO. Duffy blood group and malaria. Hematology (Amsterdam, Netherlands). 2006;11(5):389-98.
- 20. Neote K, Darbonne W, Ogez J, Horuk R, Schall TJ. Identification of a promiscuous inflammatory peptide receptor on the surface of red blood cells. The Journal of biological chemistry. 1993;268(17):12247-9.
- 21. Ntumngia FB, Thomson-Luque R, Pires CV, Adams JH. The role of the human Duffy antigen receptor for chemokines in malaria susceptibility: current opinions and future treatment prospects. Journal of receptor, ligand and channel research. 2016;9:1-11.
- 22. Lukasik E, Wasniowska K. [Duffy blood group antigens: structure, serological properties and function]. Postepy higieny i medycyny doswiadczalnej (Online). 2016;70:143-61.

- 23. Mayr FB, Jilma-Stohlawetz P, Firbas C, Suffredini AF, Derendorf H, Jilma B. Functional Role of the Chemokine Binding Duffy Antigen Receptor Complex (DARC) in Human Inflammation In Vivo. Am Soc Hematology; 2006.
- 24. Petri M, Orbai A-M, Alarcón GS, Gordon C, Merrill JT, Fortin PR, et al. Derivation and validation of the Systemic Lupus International Collaborating Clinics classification criteria for systemic lupus erythematosus. Arthritis and rheumatism. 2012;64(8):2677-86.
- 25. Hartman EAR, van Royen-Kerkhof A, Jacobs JWG, Welsing PMJ, Fritsch-Stork RDE. Performance of the 2012 Systemic Lupus International Collaborating Clinics classification criteria versus the 1997 American College of Rheumatology classification criteria in adult and juvenile systemic lupus erythematosus. A systematic review and meta-analysis. Autoimmunity reviews. 2018;17(3):316-22.
- 26. Gladman DD, Ibanez D, Urowitz MB. Systemic lupus erythematosus disease activity index 2000. The Journal of rheumatology. 2002;29(2):288-91.
- Horne K, Woolley IJ. Shedding light on DARC: the role of the Duffy antigen/receptor for chemokines in inflammation, infection and malignancy. Inflammation research : official journal of the European Histamine Research Society [et al]. 2009;58(8):431-5.
- Cooling L. Blood Groups in Infection and Host Susceptibility. Clinical microbiology reviews. 2015;28(3):801-70.
- 29. Dean L. Blood groups and red cell antigens: National Center for Biotechnology Information; 2005.
- 30. Smith E, McGettrick HM, Stone MA, Shaw JS, Middleton J, Nash GB, et al. Duffy antigen receptor for chemokines and CXCL5 are essential for the recruitment of neutrophils in a multicellular model

of rheumatoid arthritis synovium. Arthritis and rheumatism. 2008;58(7):1968-73.

- Minten C, Alt C, Gentner M, Frei E, Deutsch U, Lyck R, et al. DARC shuttles inflammatory chemokines across the blood-brain barrier during autoimmune central nervous system inflammation. Brain : a journal of neurology. 2014;137(Pt 5):1454-69.
- 32. Vergara C, Tsai YJ, Grant AV, Rafaels N, Gao L, Hand T, et al. Gene encoding Duffy antigen/receptor for chemokines is associated with asthma and IgE in three populations. American journal of respiratory and critical care medicine. 2008;178(10):1017-22.
- 33. Charles BA, Hsieh MM, Adeyemo AA, Shriner D, Ramos E, Chin K, et al. Analyses of genome wide association data, cytokines, and gene expression in African-Americans with benign ethnic neutropenia. PloS one. 2018;13(3):e0194400.
- 34. Wang J, Ou ZL, Hou YF, Luo JM, Shen ZZ, Ding J, et al. Enhanced expression of Duffy antigen receptor for chemokines by breast cancer cells attenuates growth and metastasis potential. Oncogene. 2006;25(54):7201-11.
- 35. Afenyi-Annan A, Kail M, Combs MR, Orringer EP, Ashley-Koch A, Telen MJ. Lack of Duffy antigen expression is associated with organ damage in patients with sickle cell disease. Transfusion. 2008;48(5):917-24.
- 36. Fong MY, Cheong FW, Lau YL. Erythrocytebinding assays reveal higher binding of Plasmodium knowlesi Duffy binding protein to human Fy(a+/b+) erythrocytes than to Fy(a+/b-) erythrocytes. Parasites & vectors. 2018;11(1):527.
- Guler N, Turgut M, Ozatli D, Turgut Y, Gokce AK, Koc S, et al. High ratio of Duffy (a+b+) phenotype in patients with multiple myeloma compared to healthy controls. Hematological oncology. 2009;27(1):50-1.