Lupus arthritis was more common in risk TT-genotype SLE carriers than in other SLE patients ($\chi^2=5.902, p=0.015, \phi=0.027$). We revealed significant increase of CT genotype (RUNX1 rs9979383) in healthy donors vs SLE patients ($\chi^2=4.14, p=0.042$; OR=0.53 ($CI_{95\%} 0.29-0.98$); LR=0.69 ($CI_{95\%} 0.45-0.99$); $LR^+=1.3$ ($CI_{95\%} 1.01-1.56$). SLE arthritis was more common in SLE CT-genotype carriers than in other SLE patients ($\chi^2=4.46, p=0.031; \phi=0.058$). Significant differences in IL6 rs1800795, IL6R rs2228145 and IL6R rs4845618 genotypes distribution between studied groups were not found ($\chi^2=0.42, p=0.559$ and $p=0.407$, correspondingly) but GG-genotype (IL6 rs1800795) carrier-ship in SLE patients was associated with increased APS frequency ($\chi^2=4.45, p=0.033$; OR=0.19 ($CI_{95\%} 0.04-0.99$); $LR^+=0.28$ ($CI_{95\%} 0.07-0.93$; $LR^+=1.41$ ($CI_{95\%} 1.03-1.64$).

**Conclusion:** Our data suggest the susceptibility to SLE in TT genotype of STAT4 rs7574865, protective role of CT genotype of RUNX1 rs9979383 for SLE and association between GG-genotype of IL6 rs1800795 and APS in SLE patients in Belarussian population. Lupus arthritis was associated with TT genotype of STAT4 rs7574865 and CT genotype of RUNX1 rs9979383.

**REFERENCES:**


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**AB0010**

**ASSOCIATION OF SAA1 GENE POLYMORPHISM -13T/C WITH ANKYLOSING SPONDYLITIS IN RUSSIAN POPULATION**

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**Background:** Ankylosing spondylitis is a chronic systemic inflammatory disease. Inflammation and high levels of serum amyloid A (SAA) protein are predisposing factors for secondary AA amyloidosis. The role of SAA1 gene polymorphisms in AS is not well understood.

**Objectives:** To investigate the association of SAA1 gene polymorphism -13T/C (rs12218) with ankylosing spondylitis and to evaluate the influence of this polymorphism on SAA protein concentration.

**Methods:** 123 AS patients (72 males, 51 females; age - M (SD) 37(12.77) years; disease duration - 14.28 (11.22) years; BASDAI - 5.59 (1.13); B27-positive - 111 (90.2%) pts) and 95 gender, age matched healthy individuals (control group) were included in this study. SAA1 gene polymorphism -13T/C was genotyped using allele-specific RT-PCR assay. SAA protein concentration was measured using nephelometry in AS patients.

**Results:** The distribution of genotypes TT, TC and CC differed statistically between AS and control groups (24.4%, 56.1%, 19.5% and 41.1%, 42.4%, 14.7% respectively, $\chi^2=6.9, p=0.03$). The presence of the C allele was associated with the development of AS (OR=1.55 [CI 1.04-2.33], $p=0.03$). The SAA1 -13T/C polymorphism tended to be associated with SAA protein value in AS patients: TT+TC genotypes -13.8 mg/l [4.2; 91.0], CC genotype -7.8 mg/l [1.6; 29.6], p=0.07 ESR, CRP and BASDAI values did not correlated with SAA1 -13T/C polymorphism (p=0.6, p=0.4, p=0.4 respectively).

**Conclusion:** The results of our study demonstrated for the first time that SAA1 gene polymorphism -13T/C (rs12218) is associated with susceptibility to AS. It is also shown that this polymorphism can affect the SAA protein level. Our findings need to be verified in AS patients with high levels of SAA protein in various ethnic and population groups.

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**AB0011**

**EXPRESSION PROFILE AND POTENTIAL FUNCTION OF CIRCINAS IN PERIPHERAL BLOOD MONONUCLEAR CELLS FROM PATIENTS WITH PRIMARY GOUT**

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**Background:** Autoptosis is a phenomenon of "self-phagocytosis" in eukaryotic cells, which maintains cell homeostasis by transporting intracellular materials to lysosomes for degradation and recycling. In recent years, studies have shown that autophagy may be involved in the pathogenesis of rheumatoid arthritis (RA) [1], but its specific mechanism is still unclear.

**Objectives:** The expression levels of autophagy-related genes (ATG) unc-51-like kinase 1 (ULK1), ATG13, ATG17, microtubule-associated protein 1 light chain 3 (LC3), and P62 in peripheral blood mononuclear cells (PBMC) of patients with RA were detected, and their role and clinical significance in the pathogenesis of RA were explored.

**Methods:** Real-time fluorescent quantitative PCR was performed to detect the expression levels of ULK1, ATG13, ATG17, LC3, and P62 in PBMCs of 50 RA patients, 50 healthy controls (HC), and 25 moderate to severe RA patients before and after treatment. Then, 1x2 test, y2 test, Mann-Whitney U test, Pearson test were used for statistical analysis.

**Results:** The levels of hsCRP, white blood cell (WBC), neutrophils (GR), platelet (PLT) and plateletcrit (PCT) in RA group were higher than those in HC group ($P <0.05$). Lymphocytes (LY), red blood cell (RBC), hemoglobin (HGB), hematocrit (HCT), mean corpuscular hemoglobin (MCH), mean red blood cell volume (MCV) and mean red blood cell hemoglobin concentration (MCHC) in RA group were lower than those in HC group ($P <0.05$). The expressions of ULK1, ATG17, and LC3 in RA group were higher than those in HC group, while the expressions of P62 was lower than those in HC group ($P <0.05$) (Figure 1). The correlation analysis suggested that ATG17 was positively correlated with tender joint count (TJC), swollen joint count (SJC), and health assessment questionnaire (HAQ) ($P <0.05$); ULK1 and HAQ were negatively correlated ($P <0.05$).3. Compared with before treatment with TNFi, ATG17, HAQ, DAS-28, ESR, hsCRP, WBC, GR, PLT, and PCT were significantly reduct after treatment ($P <0.05$); the expressions of RBC, HCT, MCV and MCH were significantly increased after treatment, ($P <0.05$); ULK1, ATG13, LC3, P62 and other related clinical and laboratory indicators were not significantly different before and after treatment with TNFi ($P >0.05$).

**Conclusion:** There is abnormal expression of autophagy genes in the peripheral blood of RA patients. ULK1, ATG17, LC3 and P62 may be related to the pathogenesis of RA, among them, ATG17 may regulate the pathogenesis of RA by participating in the TNF-α pathway.

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**AB0012**

**THE EFFECTS OF COMMON THERAPEUTICS ON AIRE METHYLATION STATUS AND THE LEVELS OF IL-16, IL-13, AND IFN-γ IN LPS-INDUCED MACROPHAGE CELLS CAN SULPHASALAZINE'S INFLUENCE ON METHYLATION STATUS OF AIRE EXPLAIN ITS AUTOIMMUNE SIDE EFFECTS?**

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