Abstracts

**PO.4.81** VARIABILITY OF XANTHINE OXIDOREDUCTASE ACTIVITY PATTERNS IN SYSTEMIC LUPUS ERYTHEMATOSUS AND RHEUMATOID ARTHRITIS

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**Purpose** Characterization of activity patterns of several mutually converting xanthine oxidoreductase subtypes, xanthine oxidase (XO) and xanthine dehydrogenase (XDG), within plasma and the blood cell compartments in systemic lupus erythematosus as compared with rheumatoid arthritis patients.

**Methods** The research was carried out in agreement with the WMA Declaration of Helsinki principles. 56 SLE patients and 77 RA patients were enrolled in this study. Diagnosis of SLE was verified using the ACR criteria (1997). RA was verified according to ACR/EULAR criteria (2010). Disease activity was assessed according to SLEDAI-2K and DAS28 indices, respectively. The reference group consisted of 35 healthy controls. Lymphocytes and erythrocytes were separated by means of density gradient centrifugation (1077 g/ml). XO and XDG activities were measured in plasma, lysed lymphocytes and lysed RBC using previously published kinetic techniques. Results were expressed as median and quartiles. Correlations were analyzed using Spearman’s correlation coefficient. Differences were considered significant when p<0.05.

**Results** Mean age of SLE patients was 35 (31; 42) years, mean duration of disease was 8 (5; 11) years. Mean age of RA patients was 45 (37; 49) years, mean RA duration was 8 (6; 10) years. 15 (26.8%) SLE patients had mild disease activity, 26 (46.4%) had moderate activity, and 15 (26.8%) had high activity. 16 (20.8%) RA patients had mild disease activity, 49 (63.6%) had moderate activity, and 12 (15.6%) had high activity. Both xanthine oxidoreductase subtypes had various activity shifts in plasma and lysed blood cells in RA as well as in SLE. Both SLE and RA patients had high plasma XO activity in combination with low plasma XDG activity (all p<0.05), while low XO and XDG activities were demonstrated in lysed lymphocytes for these two groups (all p<0.001). Lysed red blood cells in RA had high XO activity in combination with low XDG activity (all p<0.001). SLE patients were revealed low XDG activity without significant shift of XO activity in red blood cells. When comparing SLE and RA, SLE patients had lower plasma XDG (p=0.012), higher lymphocyte XO (p<0.001), lower erythrocyte XO (p<0.001), and lower erythrocyte XDG (p<0.001) activities. There was positive correlation between plasma XO activity and the disease activity index as well as negative correlations between plasma XDG activity, lymphocyte XO activity, lymphocyte XDG activity and the disease activity index both in SLE and RA (all p<0.001). Red blood cells in SLE had negative XO correlation and positive XDG correlation with disease activity; such correlation pattern in RA was inverse (all p<0.001).

**Conclusion** The imbalance between oxidase and dehydrogenase subtypes of xanthine oxidoreductase in SLE was expressed in higher levels of circulating XO activity that is responsible more for free radicals generation. A decrease of lymphocytic XO and XDG activities could be an indirect evidence of purine metabolism disturbance in SLE and RA. Increase of XO/XDG ratio in erythrocytes may affect the lifespan of these cells both in SLE and RA.

**PO.4.82** CORRELATION BETWEEN ANTI DS-DNA AND SYSTEMIC LUPUS ERYTHEMATOSUS DISEASE ACTIVITY
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**Objectives** To determine the correlation between serum anti-dsDNA titer and Systemic Lupus Erythematosus Disease Activity

**Introduction** Lupus is a chronic autoimmune inflammatory disease which affects more women with a peak in incidence of reproductive age. Serological tests are used to determine the activity of the disease and predict the precipitation of the disease. Anti ds-DNA antibodies are found in high levels, on patients serum with LES flare, but not in normal person serum and in patients with other autoimmune disease such as discoid lupus erythematosus, chronic hepatitis or rheumatoid arthritis. Usually during the activation of the disease we have a decrease of complement and increase of anti ds-DNA.

**Methods** The patients included in the study are 42, with an average age of 25-35 years and with the highest percentage of patients being female (40 females and 2 males). The double stand ds-DNA antibody serum was measured. The disease activity was evaluated by SLEDAI evaluation. The SLEDAI is a global index that was developed and introduced in 1985 as a clinical index for the assessment of lupus disease activity in the preceding 10 days. It consists of 24 weighted clinical and laboratory variables of nine organ systems.

**Results** The growth of anti-ds DNA was observed in all patients with a flare of disease. Positive correlation was observed between SLEDAI and anti-dsDNA levels. It was observed that this correlation was significant between the anti-ds-DNA titer and the SLEDAI activity r = 0.52 (p<0.0001).

**Conclusion** The implementation of SLEDAI is a clinically important tool for evaluating patients with LES. Serial measurement of anti-ds DNA can help us diagnose lupus flare and make the right therapeutic decision for patients with SLEDAI high points.

**PO.4.83** ANEMIA AS AN OBTAINABLE MARKER OF SYSTEMATIC LUPUS ERYTHEMATOSUS ACTIVITY ASSESSMENT
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**Purpose** Anemia is frequent manifestation of SLE. Numerous reasons for anemia are well recognized including chronic disease anemia, iron deficiency anemia, autoimmune hemolytic anemia, anemia due to chronic renal failure, anemia caused by the treatment and others. On the other hand, correlation of anemia severity with SLE disease activity and with particular clinical manifestations is not yet satisfactory examined. The aim of this study is to investigate anemia grade in patients with SLE and propose it as a potential severity and prognostic marker.
Infections in Infants Born to Mothers with Systemic Lupus Erythematosus and the Role of Preterm Birth

Abstract

Methods 56 patients with a mean age of 33 ± 6.3 years and with 7.5 ± 3.2 years' duration of SLE were enrolled in this study. All patients fulfilled with EULAR/ACR 2019 classification criteria and were evaluated with SLEDAI-2K. Anemia was graded as mild (91–110 g/l), moderate (71–91 g/l) and severe (51–70 g/l). Data was analyzed with SPSS statistical software. Results Out of all examined patients 85% were females and 15% males. Mean SLEDAI-2K was 11.26 ± 7.2. 68.4% of patients presented with anemia, in particular 27 patients (47.4%) had mild, 7 (12.3%) moderate, 4 (7.0%) severe anemia. According to the study lower values of anemia significantly correlate with high SLEDAI-2K (p < 0.01) and presence of Lupus nephritis (p < 0.042). Diversely, significant association between ESR, CRP and aPL titers and anemia levels was not found. 78.9% of patients with anemia had mucocutaneous manifestations, 57.9% arthritides, 47.4% hepatitis, 44.7% pleuritis, 42.1% pericarditis, 36.8% lupus nephritis, 28.9% cerebral vasculitis. Conclusion Anemia could be used as an additional marker for the evaluation of SLE activity status. In purpose to determine, if specific anemia type is more connected with disease activity, further assessment of hepcidin, ferritin and coombs test is required.

Purpose To investigate the risk of infections in the first year of life in infants born to mothers with systemic lupus erythematosus (SLE) compared to infants born to general population comparators, and to examine the role of preterm birth as a mediator of the association.

Methods Liveborn singletons born to mothers with SLE and general population comparators were identified in the Medical Birth Register (MBR; 2006–2012), sampled from the Swedish Lupus Linkage (SLINK) cohort (1987–2012). SLE was defined by ≥ 2 International Classification of Diseases (ICD)-coded visits in the National Patient Register (NPR) and MBR, with ≥ 1 visit before pregnancy. Infections were defined as any ICD-coded visit listing an infection diagnosis in the in- and outpatient records of the NPR or dispensed antibiotic prescriptions in the Prescribed Drug Register. Hospitalized infections were defined as a primary ICD-coded hospitalization. Modified Poisson regression models estimated risk ratios and 95% confidence intervals (RR; 95% CI) of infant infection associated with maternal SLE adjusted for maternal age, first-trimester smoking, and calendar year. Causal mediation analysis estimated the percentage of the total effect explained by preterm birth (< 37 weeks).

Results Of 441 SLE offspring, 17% had an infection diagnosis in the first three months of life compared to 12% of 4,485 non-SLE offspring, with a corresponding RR of 1.4 [95% CI 1.1–1.7] which was greater for infants born preterm (1.8 [95% CI 1.2–2.9]). Hospitalized infections occurred in 5% of SLE offspring and 3% of non-SLE offspring in the first three months (RR 2.1 [95% CI 1.3–3.3]), and in 9% of SLE offspring and 5% of non-SLE offspring in the first year (RR of 1.6 [95% CI 1.1–2.2]), regardless of being born preterm. Fifty-two percent of the total effect of maternal SLE on any infection in the first three months was mediated through preterm birth.

Conclusions When looking at the first three months, maternal SLE was associated with a nearly 40% higher risk of any infection and a two-fold higher risk of hospitalized infections. Preterm birth may explain half of the association between maternal SLE and any infection in the first three months of life.

WHAT ARE THE SEX DIFFERENCES IN PATIENTS WITH SYSTEMIC LUPUS ERYTHEMATOSUS?

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Purpose Systemic lupus erythematosus (SLE) is a chronic, multiorgan, systemic autoimmune disease that is more common in women than men. Men with SLE diagnosed often have a more aggressive clinical course with rapid accrual of organ damage, resulting in a poorer prognosis compared with women with SLE.

Methods This cross-sectional study included 227 pts: 69% (n=156) females, aged 35.4±8.0 years and 31% (n=71) males, aged 36.1±1.3 years with SLE according to SLICC/ACR 2012 criteria, attending a routine visit at our Clinic between February 2015 and February 2020. Regarding SLE-specific evaluations, patients were assessed for age at onset, disease duration, cumulative organ involvement, cumulative serology and ongoing therapy, SLE Disease Activity Index (SLEDAI-2K) and the Systemic Lupus International Collaborating Clinics damage index (SDI).

Results There were no sex differences in: mean age, age at onset (24.5±9.4 years in females and 25.6±11.0 years in males), disease duration (133.7±9.6 and 131.4±12.9 months), SLEDAI-2Kscore (9.5±0.59 and 9.15±0.98), SDI score (1.9±0.5 and 1.85±0.2), steroid use duration (99.8±8.3 and 89.8±10.4 months), taking immunosuppressants in the disease course (49% and 53%). SLE men compared to SLE women had a higher incidence of discoid cutaneous lupus – 23% versus 7%, p<0.001, venous thromboses – 27% vs 15%, p<0.05 and cumulative prednisolone dose -80.1±6.6 vs 39.1±3.4 grams, p<0.001. Oral ulcers (28% vs 46%, p<0.05), Sjögren syndrome (1% vs 11%, p<0.00001) and taking hydroxychloroquine (30% vs 63%, p<0.00001) appears to be less common in men than in women. No differences were found in the frequency of other clinical and serological disorders (women and men): acute cutaneous lupus – in 63% vs 62%, joint involvement – 90% vs 96%, serositis involvement – 42% vs 54%, nephritis – 53% vs 62%, neuropsychiatric involvement – 30% vs 31%, haematological involvement – 60% vs 46%, ANA – 85% vs 82%, other serological disorders (anti-dsDNA or anti-Sm, or low complement) – 81% vs 69% (p=0.05), antiphospholipid antibodies – 55% vs 41%, antiphospholipid syndrome – 21% vs 30%, arterial thromboses – 11% vs 8%, fever – 25% vs 30%, Raynaud phenomenon – in 25% vs 30% of patients (p≥0.05 in all cases).