P22 ANTI-CARBAMYLATED PROTEIN ANTIBODIES IN SYSTEMIC LUPUS ERYTHEMATOSUS: CLINICAL AND SEROLOGICAL ASSOCIATIONS

Micaela Fredi, Ilaria Cavazzana, Francesco Poiaati, Michele Boldrini, Silvia Piantoni, Rajesh Kumar, Roberta Ottaviani, Angela Tincani, Franco Franceschini, Rheumatology and Clinical Immunology Unit, ASST Spedali Civili, Brescia; Dept. of Clinical and Experimental Sciences, University of Brescia, Brescia, Italy

Background/Purpose Anti-carbamylated protein antibodies (anti-CarbP) have been described not only in Rheumatoid arthritis but in other systemic autoimmune diseases. Recently, they have been reported in different cohorts of Systemic Lupus Erythematosus (SLE) with a prevalence of 9–28%. In patients selected with arthritis/arthralgias. Anti-CarbP have been proposed as a marker of erosive arthritis in SLE. The aim was to assess the prevalence of anti-CarbP in SLE patients from a single center cohort and their association to clinical and laboratory data.

Methods Serum anti-CarbP levels were evaluated using a homologous ELISA (n=384 AU/mL). Clinical data were obtained from clinical charts.

Results Complete clinical and serological data were available for 314 consecutive patients: 85 (27%) positive and 229 (73%) negative. No association was found among CarbP+ and arthritis/arthralgias. CarbP+ patients presented an earlier disease onset compared with CarbP− (mean ±11±21±14.7, p=0.001), a trend towards a higher prevalence of xerophthalmia (36% vs 26.5%, p=0.075; OR: 1.62, 95% CI: 0.95–2.75) and extractable nuclear antigen positivity (67% vs 54%, p=0.064; OR: 1.65, 95% CI: 0.97–2.8). Interestingly, patients anti-CarbP+ less frequently experienced class IV glomerulonephritis (12% vs 21.8%, p=0.05; OR: 0.53, 95% CI: 0.26–1.08). Fifty-six patients evaluated were treated with anti-Blys therapy and longitudinally sera were available (T0, T6, T12). At baseline anti-CarbP were positive in 10 (17.8%) andtitre significantly decreased at T6 (p=0.006) and T12 (p=0.01). Negative seroconversion was observed in 710 sera.

Conclusions The prevalence of anti-CarbP antibodies found in our unselected cohort is in line with what previously reported. In our hands, anti-CarbP antibodies seems to identify a less severe form of SLE, with less kidney involvement and probably in overlap with Sjögren disease. Further studies are needed in order to be able to identify a possible role for this autoantibody.

REFERENCES

P23 LONGITUDINAL ANTINUCLEAR ANTIBODY (ANA) SEROCONVERSION IN SYSTEMIC LUPUS ERYTHEMATOSUS: A PROSPECTIVE STUDY OF SWEDISH CASES WITH RECENT-ONSET DISEASE

Martina Fredlund, Jonas Wetterö, Charlotte Dahlé, Orjan Dahlström, Thomas Skogh, Johan Rönndel, Christopher Skövål, Rheumatology/Division of Neuro and Inflammation Sciences, Dept. of Clinical and Experimental Medicine, Linköping University, Linköping; Clinical Immunology/Division of Neuro and Inflammation Sciences, Dept. of Clinical and Experimental Medicine, Linköping University, Linköping; Swedish Institute for Disability Research, Dept. of Behavioural Sciences and Learning, Linköping University, Linköping; Dept. of Immunology, Genetics and Pathology, Uppsala University, Uppsala, Sweden

Background Immunoglobulin G (IgG) anti-nuclear antibodies (ANA) detected by indirect immunofluorescence (IF) microscopy remain a hallmark of systemic lupus erythematosus (SLE). Since it is controversial if IF-ANA status varies over time, we designed a prospective study with longitudinal follow-up of recent-onset patients with SLE.

Methods The study population consisted of 56 newly diagnosed SLE cases, all meeting the 1982 ACR and/or the 2012 SLICC criteria. Clinical follow-up data, including disease activity and organ damage, and serum were collected from SLE onset and onwards, in most cases yearly (0 to 96 months). IF-ANA on HEP-2 cells was analysed and categorized regarding staining patterns. Using an addressable laser bead assay (ALBIA; FIDIS™ Connective profile), we measured IgG-ANA fine specificities to Ro52/SSA, Ro60/SSA, La/SSB, Sm, Sm/RNP, U1RNP, dsDNA, ribosomal P protein and histone.

Results At baseline, all patients were judged IF-ANA positive at an abnormal titre corresponding to the 95th percentile of healthy blood donors, but seven of 54 patients (13%) lost IF-ANA-positivity over time, see figure 1. Homogenous (46%) and speckled (31%) were the most frequently observed staining patterns at inclusion, whereas 7% switched their pattern at least once during follow-up, see figure 1. Established associations between ANA fine-specificities and clinical data were confirmed. Levels of anti-Sm/RNP, but not of anti-dsDNA, correlated with clinical SLE disease activity (mSLEDAI-2K).

Conclusion A considerable proportion of Swedish patients with SLE lose IF-ANA positivity over time. Consistent staining patterns were frequent. The clinical and mechanistic relevance of ANA seroconversion remains uncertain. Further prospective evaluations in larger SLE populations with diverse ethnicities are warranted.

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Abstract P23 Figure 1 Frequencies of ANA staining patterns over time