**291**

**IDENTIFICATION OF THE LONG NONCODING RNA ENST00000602652 AS A NOVEL INFLAMMATORY REGULATOR ACTING THROUGH TYPE I INTERFERON PATHWAY IN SYSTEMIC LUPUS ERYTHEMATOSUS**

Z Liao1, 1Shanghai Institute of Rheumatology, Rheumatology, Shanghai, China

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**Background and aims** Long noncoding RNAs (lncRNAs) have recently been identified to be tightly linked to diverse human diseases. Systemic lupus erythematosus is an autoimmune disease and renal involvement is the most frequent complication. Inflammatory cytokines produced by renal mesangial cells (RMCs) play a vital role in lupus nephritis (LN). In the present study we investigated the contribution of the lncRNA Enst00000602652 to the pathogenesis to LN.

**Methods** The high throughput RNAseq data between LN and healthy control was used to screen for candidate lncRNA. SYBR Green quantitative RT–PCR (RT-qPCR) was used to detect the expression of lncRNA and individual interferon-stimulated genes (ISGs). Western blotting and luciferase was used to confirm the regulatory function of lncRNA.

**Results** LncRNA Enst00000602652 expression was abnormally increased in LN patients and correlated to degree of renal damage. Additionally, Expression of lncRNA Enst00000602652 was induced by stimulation of type I interferon. Silencing Enst00000602652 significantly reduced the expression of a group of chemokines and cytokines, including IFIT1, oas1, etc., which were induced by type I interferon. Furthermore, LncRNA Enst00000602652 affects IFN receptor I and phosphorylation of Jak1 and Stat1.

**Conclusions** Long noncoding RNA Enst00000602652 is a positive regulator of the IFN signalling pathway in LN. LncRNA Enst00000602652 may contributes to the pathogenesis of LN and provides potentially novel target for therapeutic intervention.

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**293**

**EFFECT OF HIGH DOSE CHOLECALCIFEROL DURING SLE FLARES IN MIR-146A EXPRESSION, REGULATORY T-CELLS AND IL-17A EXPRESSION**

1A Marinho*, 2C Carvalho, 2D Boleia, 2A Bettencourt, 3B Martins da Silva, 1C Vasconcelos.
1Centro Hospitalar Do Porto, Internal Medicine, Porto, Portugal; 2ICBAS, Immunogenetics, Porto, Portugal

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**Background and aims** A few microRNAs have known gene expression regulatory roles in innate immunity. The miR-146a, seems to be a negative regulator of innate immunity. Interestingly, miR-146a has been reported to be downregulated in PBMCs of SLE patients, being negatively correlated with clinical disease activity and with IFN levels. The ability of vitamin D to regulate miRs and their emerging relationship have been proposed through several experimental approaches. The aim of this study was to determine the Vitamin D effect in miR-146a expression and in T-Reg and TCD4+ IL-17A producing cells, in SLE.

**Methods** An interventional study with 3 weeks follow-up of SLE patients with a high dose vitamin D supplementation (50,000 UI or 100,000 UI/Week) was done. We assessed four female patients who had a SLEDAI >6 and at least one BILAG A. At screening, relevant data were compiled: SLEDAI-2K, BILAG score, concomitant therapy, previous SLE manifestations, 25(OH)D levels, T-Reg/IL-17A ratio and miR-146a expression. At Week 3: 25(OH)D levels, T-Reg/IL-17A ratio, miR-146a expression, SLEDAI 2 K, BILAG and concomitant therapy.

**Results** No significant difference were found, regarding Vitamin D levels, before and after supplementation. Regarding Tregs/IL-17A ratio before and after supplementation, no benefits, regarding enhancing of T-reg or decrease of IL 17-A producing cells. No significative differences were found in miR-146a expression between controls and SLE active patients before and after vitamin D supplementation.

**Conclusions** Severe SLE activity may cause resistance to Vitamin D therapeutic effects, including enhancing of Vitamin D levels and immunogenetic effects.