of smaller marker panels, we have developed anti-MVP into prototypic bead-based ELISA format.

Results Discovery and validation experiments using the NavigAID SLE array showed that anti-MVP antibodies occurred with frequencies of 15%-30% in three different SLE cohorts at a specificity of 97%. Exploratory testing of multi-marker panels consisting of anti-MVP in combination with anti-dsDNA, anti-ribosomal P and anti-SmD yielded a 6% increase in sensitivity at 98% without loss of specificity. Multivariate data projection methods revealed that anti-MVP is detected in a subset of SLE patients with little overlap to established marker. A bead-based ELISA was developed for measuring anti-MVP antibodies and showed good correlation with Luminex data (R=0.88) indicating successful platform transfer.

Conclusions Anti-MVP autoantibodies represent a useful marker in SLE and, in combination with established markers, optimises the strategy for autoantibody testing. Furthermore, although more studies are needed, our findings suggest a previously undescribed linkage of type I IFN and autoantibody targets in SLE.

269 SINGLE NUCLEOTIDE POLYMORPHISMS (SNPS) OF INTEGRIN-ALPHA-M (ITGAM) ARE ASSOCIATED WITH LUPUS NEPHRITIS (LN) IN AN ASIAN SYSTEMIC LUPUS ERYTHEMATOSUS (SLE) COHORT

M Chan*, WG Law, TY Lian, KO Kong, CY Yu, YW Song, HH Chng, Leung, TS Tan, HS Howe, Tan Tock Seng Hospital, Rheumatology and Allergy, Singapore; Singapore; *The Research Institute at Nationwide Children’s Hospital, Centre for Molecular and Human Genetics, Columbus, USA; †Seoul National University, Rheumatology, Seoul, Republic of Korea; ‡University of California Los Angeles, Rheumatology, Los Angeles, USA.

Background The Integrin-alpha-M (ITGAM) rs1143679 SNP has been associated with susceptibility to SLE and lupus nephritis (LN) in oriental Chinese and Thai populations. We previously found 13 ITGAM SNPs in linkage disequilibrium (LD) that were associated with susceptibility to SLE, but found no association with rs1143679.

Aim To determine associations of ITGAM SNPs with SLE subphenotypes and autoantibodies.

Methods We studied 248 patients fulfilling the 1997 ACR revised criteria for SLE. SLE-associated ITGAM SNP alleles were identified using custom-designed Immunochip arrays and gPLINK 1.062 software, with Bonferroni corrections for multiple comparisons. Associations of SLE-related ITGAM SNPs with SLE subphenotypes (malar or discoid rash, serositis, mouth ulcers, arthritis, haematological, renal or neurological involvement) and autoantibodies to dsDNA, Ro, RNP or Sm were determined with chi-square and Fisher’s tests and logistic regression.

Results All 13 SLE susceptibility ITGAM SNPs as well as the uncommon rs1143679 SNP (n=11) were associated with LN (Table 1). The strongest association was with rs2359661 (p=0.002, uncorrected). Subjects with these SNPs were less likely to have discoid rash. There was a trend towards an association with anti-Sm. Logistic regression models for 11 SNPs retained the factors LN, discoid rash and anti-Sm, suggesting strong LD for these SNPs.

Conclusions This study demonstrated novel ITGAM SNP associations with LN and confirmed the association of rs1143679 with LN. Most associated SNPs were in the regulatory region of ITGAM bearing promoter/enhancer histone marks and have been associated with expression levels in several cell types, suggesting modulation of levels of ITGAM expression to impact these subphenotypes.

270 ASSOCIATION OF TLR2 (23BP INS/DEL) POLYMORPHISM WITH SYSTEMIC LUPUS ERYTHEMATOSUS (SLE) AND P. FALCIPARUM MALARIA: A STUDY IN MALARIA ENDEMIC AREA OF ODISHA, INDIA

B Das*, TR Tripathy, SP Pattanaik, A Panda, S.C.B. Medical College, Medicine, Cuttack, India; S.C.B. Medical College, Biochemistry, Cuttack, India; Central University of Jharkhand, Centre for Life Sciences, Ranchi, India.

Background and aims Human toll-like receptors (TLRs) participate in the innate response and signal the activation of adaptive immunity. TLRs play a vital role in sensing infection. A common 23 bp insertion/deletion polymorphism at 5’UTR of TLR2 gene has been shown to affect TLR2 expression and plasma levels of pro-inflammatory molecules. We hypothesised...