APS, pDC and mDC produced BAFF and expressed chemo-
kine receptors.

Conclusion pDC and mDC are differentially affected by IFNα in SLE and APS. IFNα primes pDC for enhanced IFNα pro-
duction which potentiates T-cell activation by mDC, thereby
sustaining the IFN signature in SLE and APS.

CLARIFICATION OF THE ROLE OF DNASE 1 ON THE ONSET OF SYSTEMIC LUPUS ERYTHEMATOSUS IN A MURINE MODEL

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Systemic lupus erythematosus (SLE) is a prototypical autoim-
mune disease resulting in multi-organ damage and a high rate
of morbidity. Onset of SLE is characterised by dysregulated
activation of T and B lymphocytes and the production of
autoantibodies directed against nuclear components. The auto-
antibodies generated during the onset of SLE often recognize
components released by neutrophils during NETosis, a type of
cell death defined by the generation of neutrophil extracellular
traps (NETs). The endonuclease DNase1 has been shown to be
involved in the clearance of NET components. The sera of
SLE patients contain inhibitors of DNase 1 and/or anti-NET
antibodies that block the ability of DNase 1 to degrade NETs.
Thus, whilst NETs are important for clearing infection they
must be tightly regulated and degraded to prevent the onset of
autoimmunity.

In this study we monitored the production of auto-antibod-
ies in the serum of wild type and DNase 1-deficient mice
from the age of 2 to 12 months, along with proteinuria levels
and the development of glomerulonephritis. We show that
DNase 1-deficient mice develop a SLE-like phenotype with
elevated auto-antibody production and kidney damage by 12
months. This model also demonstrates the female bias in SLE
as the female DNase 1-deficient mice had the highest level of
kidney damage. As DNase 1 activity, B cells and aberrant
NETosis are central to progression of SLE understanding their
mechanisms of action are of great therapeutic interest.

MARGINAL-ZONE-LIKE B CELLS DEFICIENCY REPEATEDLY DETECTED IN PERIPHERAL BLOOD AS A POSSIBLE BIOMARKER OF HYPOSPLENISM/ASPLENIA IN SLE

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Objective SLE is a disease associated with a risk of serious
infections, in case of hypospplenism/asplenia especially by
encapsulated bacteria. For opsonization and phagocytosis of
these agents are essential IgM natural Abs, produced only by
B cells of the splenic marginal zone. Significant deficiency of
marginal-zone-like B cell CD19 +CD27+IgM+ cell subpopulation
absolute values x10–6/L in peripheral blood (PB) was demon-
strated in a prospective, comparative, cross-over SLE study; goal of the present study is follow up persistence of this
phenomenon.

Design and method Sixty adult SLE (ACR/1982, update 1997)
pts and 10 age-and sex-matched healthy controls (HC) were
enrolled in month O', and 56 SLE pts also repeatedly after
twelve-month-period, i.e. month 12'; overlap syndromes,
infection, monoclonal gammapathy and renal failure in SLE
under study were excluded. The DuraClone IM panel (Beck-
mann Coulter) was used to identify CD19 +CD27+IgM+B
cell subpopulation in PB samples by flow cytometry Navios
(Beckman Coulter) with software analysis using Kaluza version
1.2: data obtained were expressed in relative% of PB lympho-
cytes and absolute values x10–6/L. Parallel analysis of serologi-
ical SLE biomarkers included C3, C4, ANA/IF (maximal titre),
ANA/ELISA, anti-dsDNA/IFCL (maximal titre), and anti-dsDNA/ELISA
and antinucleosome Abs. Data obtained were statistically
processed using Medcalc-Statistical Software programme.

Results Significant differences (p<0.001) were obtained
between absolute values of CD19 +CD27+IgM+B cells in
HC (median 31.36, 95% CI: 24.49 to 63.35) and SLE month
O' (median 9.82, 95% CI: 6.01 to 14.26), and also SLE month
12' (median 10.09 95% CI: 7.12 to 14.42), but not between
values obtained in SLE month O’ and month 12' (p>0.05); not
significant differences were found in analysis using relative% of PB lymphocytes (p>0.05). In SLE month
O’ was found a slight significant correlation between absolute
values of CD19 +CD27+IgM+B cells and anti-dsDNA/ELISA
Abs (rs= −0.28, p=0.034) without a confirmation in month
12' control (rs= −0.09, p=0.491).

Conclusions The data obtained demonstrated persistent charac-
ter of marginal-zone-like B cells deficiency in peripheral blood,
and are suggesting as possible biomarker of functional hypo-
splenism/asplenia in SLE.

REFERENCE

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CHARACTERISATION OF SLE B CELLS FROM PATIENTS IN REMISSION – PERSISTENT IL-10 SECRETORY DEFECT

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Background SLE is an autoimmune disorder characterised by
polyclonal Bcell activation, the production of anti-double
stranded (ds) DNA autoantibodies and cytokines. Molecular
and clinical studies regarding SLE often address clinically
active patientsand not patients in remission. This study reports
on immunoglobulin, anti-dsDNA-aab and IL-10 secretory
capacity of cultures of CD19 +lymphocytes from SLE patients
in remission in comparison to normal donors. The aim was to
evaluate whether endogenous factors (BAFF, CD40, IL4),
exogenous factors (CpG-ODN-motifs, SAC) or their combina-
tions differentially influence immunoglobulin, cytokine and
anti-dsDNA-aab production in not active SLE patients vs
healthy controls.

Methods Blood samples were obtained from a group of 13 SLE
patients attending clinics at the rheumatology unit at the Heinrich-
Heine University hospital in Düsseldorf and from 5 healthy controls (HC). All patients were randomly collected in clinical remission state (SLEDAI 1,1±1.8). The medication consisted of prednisolone (5/13 patients), mycophenolate mofetil (3/13 patients), azathioprine (1/13 patients), hydroxychloroquine (8/13 patients) or was without immunosuppression.

After 6 days of cell culture levels of IgG, IgM, dsDNA antibodies and interleukin-10 (IL-10) were determined in the supernatants by ELISA. The effect of the stimuli alone or in combination on IgG, IgM, anti-dsDNA-aab, and IL-10 production was analysed.

Results Peripheral B cells from SLE patients in remission or control subjects did not show any difference in IgG, IgM, and anti-dsDNA-aabs to all aforementioned stimuli. The addition of CpG and SAC to cell cultures showed a stimulatory effect on immunoglobulin, cytokine and anti-dsDNA-aab production in SLE B cells and healthy controls alike. The amount of anti-dsDNA IgG-type autoantibodies produced by peripheral B cells was negligible. However, B cells from SLE patients showed diminished capacity to produce IL-10 as compared to B cells from healthy donors (SLE Estimate –40.17, Std.error 17.21, p<0.01).

Conclusion B cells from SLE patients in remission as compared to peripheral B cells from healthy donors have comparable capacity to secrete immunoglobulin including non-IgG anti-dsDNA-aabs whereas their capacity to secrete IL-10 is impaired. This suggests a persisting intrinsic defect of B regulatory cells in SLE.

### Abstracts

**CIRCULATING ANGIOGENIC T-CELLS ARE REDUCED IN PATIENTS WITH SYSTEMIC LUPUS ERYTHEMATOSUS WITH HIGH DISEASE ACTIVITY AND WITHOUT KNOWN CARDIOVASCULAR RISK FACTORS**

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**Background** Recent evidences underlined the central role of T-cells in the pathogenesis of Systemic Lupus Erythematosus (SLE) and in its cardiovascular complications. 1 CD3 +CD31 +CXCR4+angiogenic T-cells (Tang) have been identified as a T-cell subtype involved in the repair of damaged endothelium cooperating with endothelial progenitor cells. 2 Tang were described as selectively expanded in the circulation of systemic sclerosis patients displaying peripheral vascular complications, as a reaction to an inefficient angiogenesis. 3 Not much information is available on Tang in a SLE patients: in a recent study the percentage of circulating CD8 +Tang, but not CD4 +Tang, was higher in SLE than in healthy controls. 4 However, in this study SLE patients with hypertension, dyslipidemia or smoking habit, factors which may influence Tang counts, were not excluded.

The aim of this study was to characterise Tang in a cohort of patients with SLE without known cardiovascular risk factors.

**Methods** Twenty female SLE patients with a recent disease onset (<5 years) and without traditional cardiovascular risk factors or previous events (age: median value=43 [25th-75th percentile=27–54] years) and 18 healthy controls (age: 40 [32–54] years) were enrolled. Phenotypic analysis of peripheral Tang lymphocytes was made by flow-cytometry.

**Disease activity** was evaluated by SLEDAI-2K score.

**Results** SLE patients were divided in two groups according with disease activity. Patients with SLEDAI-2K equal or higher than 6 were defined as patients with high disease activity (n=5). They had a lower percentage of circulating Tang in comparison with healthy controls (10 [8–15] vs 16 [14–23]%) of CD3 +T cells, p=0.04. The result was confirmed in absolute number (83 [60–103] vs 242 [165–328] cell/μl, p=0.04). SLE patients with low disease activity had levels of Tang which were intermediate between, and not significantly different from, healthy controls and patients with high disease activity.

**Conclusions** Tang were reduced in our patients with active SLE, and no known cardiovascular risk factors, suggesting that this reduction was directly explained by disease activity.

**REFERENCES**

### PS5:96 IL-21 DEPENDENT GRANZYME B PRODUCTION OF B-CELLS IS DECREASED IN PATIENTS WITH LUPUS NEPHRITIS

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**Objectives** B-cells play a crucial role in the pathogenesis of lupus nephritis. Recently, a separate subset has been discovered characterised by expression of Granzyme B. The aim of this study is to investigate this subset in patients with systemic lupus erythematosus especially in patients with lupus nephritis.

**Methods** Isolated peripheral blood mononuclear cells of patients with systemic lupus erythematosus (n=30) and healthy controls (n=21) were _in vitro_ stimulated with CPG, IgG +IgM and IL-21. Patients were sub-grouped in patients with and without biopsy proven lupus nephritis. CD19 +B cells were analysed for intracellular Granzyme B expression by flow cytometry. Patients disease activity was assessed by systemic lupus erythematosus disease activity index (SLEDAI).

**Results** The strongest stimulus for Granzyme B secretion of CD19 +B cells was IgG +IgM in presence of IL-21. Patients with systemic lupus erythematosus had a significant decreased percentage of Granzyme B+CD19+B cells. This could be shown in particular for patients with active disease and with lupus nephritis.

**Conclusions** These data demonstrate that CD19 +B-cells of patients with systemic lupus erythematosus are impaired to produce Granzyme B. This may contribute to an imbalanced B-cell regulation towards effector B-cells which might promote the development of lupus nephritis.