ANTI-ATP SYNTHASE AUTOANTIBODIES ALTER ENDOTHELIAL CELL (EC) SURVIVAL BY DISRUPTING PH REGULATION IN VASCULIT

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Background
B-cell clusters represent genuine germinal centres (GCs) in the SGs of SS patients, but their majority display transitional and marginal zone (MZ)-like B cell characteristics. We thus asked the question as to whether TLR9 stimulation that occurs during inflammatory reaction could influence such an aberrant B-cell differentiation.

Methods
Following laser microdissection, TLR expression was measured in B-cell aggregates using quantitative RT-PCR. As controls, transitional and mature B cells from cord blood and tonsils were FACS sorted and stimulated with CpG-ODN. CFSE enabled to evaluate their proliferation, they were also analysed by flow cytometry to determine their activation status. Differentiation into Ig-secreting cells was demonstrated by ELISA.

Results
Transitional B-cell aggregates of SGs expressed high level of TLRs whereas real GC lacked TLR expression. In vitro, TLR9 stimulation induced proliferation of transitional B cells, activation and maturation as MZ B cells (Notch2high, CD21high, IgMhigh, IgDlow, CD23low, CD27−). Mature B cells proliferated also but differentiated toward a follicular pathway (Notch2low, IgMlow, IgDlow, CD23low, CD27+). Production of Ig was observed in both B-cell supernatants.

Conclusion
High TLR level in transitional SG B-cell aggregates indicates how highly sensitive to inflammatory stimuli they are. Immature B cells can differentiate into Ig-secreting cells following TLR9 stimulation through a MZ-like maturation pathway without entering the GCs. In contrast, mature B cells differentiate towards a classical follicular pathway. Whether one or both differentiation pathway generate autoreactive B cells in SGs remains to be established in SS patients.
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*Ann Rheum Dis* 2010 69: A10
doi: 10.1136/ard.2010.129577y

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