Methods: Fifteen severe, refractory SLE patients were included and followed for 2 years. Patients received RTX at week 0 and 2 and BLM at week 4, 6, 8 and then 24-weekly until week 104. Clinical response was assessed by achievement of low level of disease activity (LLDAS). By using specific antibody assays and high sensitivity flowcytometry (HS-FACS), we longitudinally followed, respectively, levels of SLE-specific ANAs and B-cell subsets.

Results: Ten patients (67%) showed a good clinical response after 24 weeks, referred to as ‘responders’. Two of these patients (13%) switched treatment after 24 weeks due to a pregnancy wish and 8 patients (53%) continued study treatment throughout the complete 2 years of follow-up. Five patients (33%) discontinued treatment due to persistent LN (n=2), major flare (n=2) or relapsing minor flare (n=1), together referred to as ‘non-responders’. Responders achieved LLDAS at a median of 24 weeks [range 12-36] and remained in LLDAS for 76 weeks [56-92] out of 104 weeks of follow-up. In 7 patients with active LN, 6 attained a complete renal response. In responders, ANAs showed significant and specific reduction throughout 2 years with achievement of seronegative anti-dsDNA immunofluorescence in 6 out of 6 anti-dsDNA positive patients at baseline while total IgG, anti-tetanus and anti-rubella antibodies remained stable. By using HS-FACS, a median decrease of 97% [99;±35] CD19+ B-cell depletion was achieved at 24 weeks. Long-term follow-up showed that B-cell repopulation was inhibited throughout 2 years with a persistent median decrease of 84% [92;±22] compared to baseline. Further analysis of B-cell subsets revealed that in the responder group, double-negative (DN) B cells (CD27- IgD-) reached maximum depletion at 4 weeks (median 1.09*10^6 cells/liter [range 0.23*10^6-4.31*10^6]), which lasted up to week 72 with a median of 1.26*10^6 cells/liter [0.79*10^6-4.11*10^6] at week 72, similar to values at nadir. This was in contrast to non-responders, where maximum depletion of DN B cells was reached at 12 weeks (0.48*10^6 [0.17*10^6-5.86*10^6]), after which these cells increased to 2.29*10^6 [0.49*10^6-4.39*10^6] at 24 weeks.

Conclusion: Over 2 years follow-up, RTX+BLM for severe, refractory SLE patients prevented complete B-cell repopulation with persistent and specific reduction of ANAs. Clinical response was observed in 67% of patients and treatment discontinuation due to high disease activity was associated with early repopulation of DN B-cells. These data warrant further studies on clinical and immunological benefits of combination treatment RTX+BLM.

REFERENCE:


OP0043
WITHDRAWAL OF A LOW DOSE (5 MG) OF CORTICOSTEROIDS IN SYSTEMIC LUPUS IN REMISSION FOR MORE THAN A YEAR IS AT RISK OF RELAPSE – THE CORTICOLUP TRIAL
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Background: Glucocorticoids (GCs) are a mainstream of treatment for patients with SLE but are associated with significant adverse effects. Some SLE patients are maintained for a long-term under a low dose of prednisone to prevent relapse. There is no scientific data to sustain this therapeutic strategy.

Objectives: We hypothesized that maintaining a daily dose of 5 mg prednisone in patients with an inactive SLE for at least a year would prevent the risk of relapse.

Methods: The CORTICOLUP study (NCT02356517) was a prospective randomised, open-labeled, controlled, monocentric study sought to compare maintenance vs withdrawing of low-dose of prednisone (5mg) to reduce SLE flares, conducted from January 2014 to March 2017. Inclusion criteria were SLE patients who during the year preceding the inclusion had 1 or inactive SLE defined by a SLEDAI-2K score ≤ 4, a BILAG-2004 index C, D or E in all systems and a PGA <0 and 2/4 stable SLE treatment including prednisone 5 mg daily. The primary end point was the number of patients with flares during 12 months of follow-up defined by the revised-SELENA SLEDAI Flare Index (rSFI). Secondary outcomes were occurrence of a BILAG scores A or B ≥1, clinical SLEDAI-2k ≥0, PGA >0.5 and increase of the SLICC damage index (SDI). All patients were included in the intention-to-treat analysis.

Results: A total of 124 patients (61 in the maintenance group and 63 in the withdrawal group) were included. No patients were lost to follow up. At baseline There were no significant differences between the maintenance and the withdrawal group with respect to: duration of SLE (mean [standard deviation] duration of 11.8 (±0.9) years vs 13.1 (±1.5) years), duration of remission (55.7 (±5.6) vs 67.5 (±8.8) months), HCO treatment [96.2% vs 100.0%], immunosuppressive drugs [27.9% vs 25.4%], previous renal involvement [34.4% vs 41.3%], low C3 [16.4% vs 15.9%], positive Farr test [47.5% vs 46.0%], and SDI [mean index of 0.5 (±0.1) vs 0.7 (±0.2)].

There were significantly more flares in the withdrawal group compared to the maintenance group (17 flares versus 4, p=0.0034 using the Fisher’s exact test). Mild or moderate flares were more frequent in the withdrawal group compared to the maintenance group (12 vs 3, p=0.029). The occurrence of severe flare was not significantly different between the two groups (5 vs 1, p = 0.208). More than two-thirds of the flares in the withdrawal group occurred within the first six months. Within the withdrawal group, using forest plot analysis, no significant association was found between the occurrence of a flare and age, sex, duration of SLE, duration of GCs treatment, immunosuppressants and serological SLE activity at randomization. Four patients in the withdrawal group and none in the maintenance group experienced damage: 2 osteoporosis bone fractures, 1 hydroxychloroquine retinopathy and 1 cataract.

Conclusion: Withdrawal of low dose of steroids in patients with inactive SLE and stable therapeutic regimen for more than a year is associated with a high risk of relapse.

Disclosure of Interests: None declared

OP0044
A SYSTEMATIC REVIEW AND META-ANALYSIS OF THE GONADOTOXIC EFFECTS OF CYCLOPHOSPHAMIDE AND BENEFITS OF GONADOTROPIN RELEASING HORMONE ANALOGUES IN WOMEN OF CHILD-BEARING AGE WITH AUTOIMMUNE RHEUMATIC DISEASE
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Background: Premature ovarian insufficiency (POI) is a side effect of intravenous cyclophosphamide (i.v.CYC), which is the treatment of choice for many severe manifestations of autoimmune rheumatic disease (ARD) (1). The risk of POI post-CYC appears to be dependent on cumulative dose and patient age but has not been precisely quantified. Gonadotropin Releasing Hormone Analogues (GnRHa) may reduce this risk (2). Studies however, on GnRHa and CYC use in ARD patients have been limited in size and/or by combined analysis with patients receiving CYC for malignancies.

Objectives: In order to more precisely categorise the risk of POI in ARD patients, and determine whether GnRHa are effective in reducing this risk, we performed a systematic review to assess the risk of sustained amenorrhea with i.v.CYC in ARD and a meta-analysis of the efficacy of GnRHa in reducing this risk.

Methods: Using Preferred Reporting Items for Systematic Reviews and Meta Analyses principles, we searched English language papers using PubMed, MEDLINE, EMBASE and the Cochrane Library to April 2018. We included all studies...