are positive for anti-ds DNA, anti-Sm, and/or anti-nuclear antibodies. Exclusion criteria include: active, clinically significant, interstitial lung disease/pulmonary arterial hypertension; proteinuria >4 g/day and/or eGFR <45 mL/min/1.73 m²; recent acutely worsened renal function; and use of oral corticosteroids >30 mg/day prednisone equivalent, injectable corticosteroids, or change in dose of corticosteroids within 2 weeks prior to or during screening. The primary efficacy endpoint family comprises response based on SLE Responder Index (SRI)-4 among all patients and SRI-6 in patients with high disease activity (baseline SLEDAI-2K ≥10) at Week 52. Success on either endpoint will support a conclusion of efficacy. Primary safety endpoints include nature, severity, and incidence of adverse events (AEs) and serious AEs. Secondary endpoints include the time to first severe flare up to Week 52, SRI-4 and SRI-6 response at Week 52 in the serologically active subgroup, and disease activity over time, including attaining low disease activity, and change from baseline in Cutaneous Lupus Erythematosus Disease Area and Severity Index (CLASI)-A at each study visit. The primary analysis is planned when all patients have completed the safety follow up for the 52-week blinded portion of the study, entered the OLE, or have discontinued prematurely from the study. Results Recruitment is ongoing. Target enrolment is 432 to 468 participants. The first patient was randomized on 20 January 2017; study completion is expected end of 2019. Conclusions This study will provide clinical proof of concept of the efficacy and safety of evobrutinib in SLE. Funding Source(s): Merck KGaA, Darmstadt, Germany

Abstract 213

Assocation between age of SLE diagnosis and disease damage differs across racial/ethnic groups: results from the California Lupus Epidemiology Study

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Background Earlier age of SLE onset is associated with greater disease damage, even after taking into account the effects of current age and disease duration. We sought to determine if this association was consistent across racial and ethnic groups, given the differences in disease severity among these groups.

Methods Data derive from the baseline visit of the California Lupus Epidemiology Study (CLUES), an ongoing cohort of patients in the San Francisco Bay Area with confirmed SLE diagnoses, drawn from a variety of clinical sources and prior SLE studies. Participants provided access to medical records and had a visit with a study physician in which clinical labs were drawn. Disease damage was measured using the SLICC/ACR Damage Index (SDI), calculated at the study visit. Age of diagnosis was ascertained by the study physician or from the medical records. Race/ethnicity (White, African American, Hispanic of any race, and Asian) and educational attainment (high school or less, some college, college graduate) were determined by patient report. Due to the small sample size, patients from other racial groups were excluded from this analysis (n=5). Using multiple linear regression, we estimated a model of SDI as a function of race/ethnicity and age of diagnosis, plus terms for interaction between the variables. The model controlled for sex, current age, and education.

Abstract 213 Figure 1

SDI by race/ethnicity and age of diagnosis
Results Among 323 participants, 89% were female, 39% Asian, 11% African American, 22% Hispanic of any race, and 29% White. Mean age was 45±14; mean age at diagnosis 29±12. Nearly half of respondents had a college degree. SDI at the baseline study visit ranged from 0 to 10 points, mean 1.8±2.0; 70% of the cohort had SDI>0. The regression model showed strong evidence (p=0.01) for interaction of age of diagnosis with race/ethnicity. As seen in the figure 1, SDI scores in racial/ethnic minorities were much higher among those diagnosed at younger ages; this relationship was not seen among whites.

Conclusions In this multi-ethnic cohort of SLE patients, the association of diagnosis age and disease damage varied according to race/ethnicity, with whites diagnosed at younger ages accumulating less damage than those in other racial/ethnic groups diagnosed at comparable ages. Future research should examine if these differences are due to phenotypic differences among the groups, diagnostic delays, or other access to care issues.

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214 RELATIONSHIP BETWEEN SERUM LEVEL OF RENALASE AND LUPUS NEPHRITIS ACTIVITY

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Background Lupus nephritis is a major risk for overall morbidity and mortality in SLE (Systemic lupus erythematosi- sis), and despite potent anti-inflammatory and immunosuppres- sive therapies still ends in Chronic Kidney Disease (CKD) or End Stage Renal Disease (ESRD) for too many patients. Renalase is a novel, kidney secreted cytokine-like protein that promotes cell survival.

Aim of the work studying the relationship between level of Human Serum Renalase (RNLS) with LN and its role in the disease activity and progression.

Methods For the current cross-sectional study 23 healthy controls and 48 patients with LN were screened and 30 subjects were selected. These patients were subdivided into two equal groups according to disease activity measured by SLEDAI (SLE Disease Activity Index). Human Serum Renalase (RNLS) concentration was measured by a highly sensitive, commercial sandwich enzyme immunoassay which uses (RNLS) antibody to capture Renalase from serum. Assessment before and after treatment was done for 17 patients who received prednisone and immunosuppressive therapy were recruited and followed up for three months to evaluate the serum renalase levels before and after treatment.

Results The level of renalase was significantly higher in LN patients compared to healthy controls, (P value<0.001). Moreover, patients with active LN had higher serum renalase levels compared to patients with inactive LN (P value<0.005) Serum renalase levels were positively correlated with SLEDAI, 24 hour urine protein excretion, ds-DNA and ESR and CRP but inversely correlated with serum C3 and the class (espe- cially in proliferative type (Class III, IV, more than class V). Renalase amounts decreased significantly after three-months of standard therapy. Also we found there is insignificant differ- ence of renalase level according to treatment by MMF (myco- phenolate mofetil) and Cyclophosphamide during and after activity (P value=0.655, 0.550)

Conclusions Serum renalase levels were correlated with disease activity in LN. Serum renalase might serve as a potential indicator for disease activity in LN.

215 FIRST-IN-MAN STUDY EVALUATING THE SAFETY, TOLERABILITY, PHARMACOKINETICS AND CONCENTRATION QT ANALYSIS OF THE NOVEL BTK INHIBITOR EVOBRTINIB (M2951) IN HEALTHY VOLUNTEERS

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Background Brutons tyrosine kinase (BTK) regulates B-cell receptor signaling and is a rational target for B-cell driven pathologies, including SLE. Evobrutinib (M2951) is a highly specific, oral inhibitor of BTK that demonstrated efficacy in preclinical models of autoimmune disease. This first-in-man study investigated the safety, tolerability, and pharmacokinetics (PK) of evobrutinib in healthy volunteers and examined the relationship between evobrutinib exposure and changes in QT interval.

Methods This was a single-center, Phase I, double-blind, placebo-controlled trial. In Part 1, 48 healthy participants in six successive dose cohorts (25, 50, 100, 200, 350, 500 mg) were randomized (6:2) to a single oral dose of evobrutinib or placebo. In Part 2, 36 participants in three ascending dose cohorts (25, 75, 200 mg/day) were randomized (9:3) to evobrutinib or placebo once daily for 14 days. Safety and tolerability were assessed following single and multiple dosing, and PK parameters determined by non-compartmental methods. Change from baseline in QT interval was obtained from 12-lead electrocardiogram recordings and corrected for heart rate by Fridericias method (QTcF).

Results Treatment-emergent adverse events (TEAEs) with evobrutinib were mostly mild, occurring in 25% of participants after single dosing, and 48% after multiple dosing. The nature and incidence of TEAEs were similar among evobrutinib and placebo groups, with no apparent dose relationship regarding the frequency or type of TEAEs. Evobrutinib showed rapid absorption (tmax ~0.5 hour), short half-life (~2 hour), and dose-proportional PK following single and multiple dosing, with no accumulation or time dependency on repeat dosing. Concentration QTcF analyses revealed no significant exposure-effect relationship. Based on a linear mixed-effects model for change from baseline in QTcF (QTcF), the slope of the relationship between mean placebo-adjusted QTcF (QTcF) and concentration was negative and close to zero (0.00027 ms/ng/ml; p=0.86; figure 1). The predicted QTcF effect at geometric mean Cmax for the highest dose (1512 ng/mL) was 1.16 ms, with an upper limit of 3.26 ms for the 90% two-