not differ by low complements, anti-dsDNA, or anti-Ro antibodies. Stepwise multivariable linear regression analysis demonstrated that the association between extrarenal disease and lower PROMIS-29 summary scores was primarily driven by arthritis and independent of potential confounders (tables 2 and 3).

Conclusion The majority of patients had isolated renal disease and report a QOL similar to that of the general population. In contrast, those with extrarenal manifestations report significantly worse QOL outcomes. These results reinforce the critical importance of routine laboratory surveillance and medication compliance for nephritis even in patients with seemingly quiescent clinical disease since lupus nephritis is often asymptomatic.

SLE Diagnosis

**POSITIVE ANA TESTING IN AN ACADEMIC MEDICAL CENTER: IMPACT ON DIAGNOSIS**

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Background Anti-nuclear antibodies (ANA) are common in systemic rheumatic diseases, non-rheumatic autoimmune diseases and the general population. In clinical practice, testing for ANA can inform further clinical diagnoses even in the absence of symptoms to suggest SLE or connective tissue disease. This study was undertaken to evaluate whether ANA testing result informed clinical diagnosis.

Methods The UT Southwestern Medical Center IRB approved this study. Data were obtained by SQL queries of the Epic electronic health record. The study population included all patients for whom an ANA was ordered at UT Southwestern Medical Center January 1, 2010 to June 30, 2017. The titer and pattern of the ANA as well as the results of ENA or anti-dsDNA testing were obtained. Patient characteristics included age and sex. Encounter characteristics included the date of testing, frequency of testing, primary encounter diagnosis, and provider specialty.

Results During the study period, a total of 33,270 ANA were ordered in 28,659 unique patients. Twenty two thousand, five hundred and twenty nine of the ANAs were tested in outpatients, representing 0.7% of all office visits during this time. 3,505 patients (11.9%) had ANA tested multiple times (range: 2-25 times). In 31% of the patients having multiple ANA, the result was repeatedly negative; 41% stayed repeatedly positive; 16% of patients had an initial negative ANA and a subsequent positive; 12% had an initial positive and a subsequent negative ANA.

Forty-nine percent of the ANA tests were positive at a titer of 1:80 or greater; slightly more women (51.1%) had a positive ANA (≥1:80) than did men (43.0%). Fifty-four and a half percent of the positive ANA in women were 1:320 or greater vs. 1:80 or greater; slightly more women (51%) had a positive ANA; 12% had an initial positive and a subsequent negative ANA.

Conclusions ANA testing in the inpatient and outpatient setting is common. Diagnoses precipitating testing are most often non-rheumatic conditions. A positive ANA result changed the clinical diagnosis in a small percentage of patients.

**CAPSTONE: AN OBSERVATIONAL RETROSPECTIVE ANALYSIS COMPARING A COMPLEMENT ACTIVATION MULTIANALYTE PANEL VS. STANDARD ANA TESTING: SYSTEMIC LUPUS ERYTHEMATOSUS DIAGNOSIS, TREATMENT OUTCOMES AND ECONOMIC IMPACT**

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Background Systemic Lupus Erythematosus (SLE) is a complex clinical disease that often takes years and repeated testing to adequately diagnose. Standard biomarkers have low specificity (ANA) or low sensitivity (anti-dsDNA, anti-Smith). The advent of a multianalyte assay panel (MAP) incorporating innovative cell-bound complement activation markers warrants comparison of its clinical utility to conventional autoantibodies for the diagnosis and treatment of SLE.

Objective This study compares the likelihood of SLE diagnosis, SLE treatment initiation, and the downstream impact on healthcare utilization on over 40,000 patients tested with either MAP (AVISE Lupus) versus standard of care lab testing with traditional anti-nuclear antibody (ANA) testing strategy cohort (tANA).

Methods Using electronic health record (EHR) data from the Illumination Health registry (an integrated EHR record database encompassing records from over 300 US rheumatologists), an observational retrospective cohort study was performed. Health records from 01/2016 to 12/2020 and administrative claims with cost data for a subset of patients linkable to the HealthCore Integrated Research Database (HIRD) and Medicare data were analyzed. Two cohorts were established: MAP testing strategy and the tANA approach. Each test result was classified as positive, negative, or indeterminate and outcomes were stratified based on test results. Multivariable logistic regression was used to estimate odds ratios (OR) comparing the likelihood of SLE medication initiation and SLE diagnosis between the MAP and tANA cohorts. Test impact on SLE diagnosis, treatment initiation, patterns of repeat testing, and downstream healthcare utilization were analyzed.

Results 21,827 MAP testing episodes and 22,778 tANA testing episodes were included in the main cohort. Findings include: 2,437 (11.2%) patients tested positive by MAP compared to 5,364 (23.6%) of tANA(+) patients. MAP(+) patients were over 5-fold more likely to be diagnosed with SLE as compared to the tANA patients, 31% vs. 8% (OR = 5.11, 95% CI 4.43-5.89). Similarly, the new patient cohort was 6-fold more likely to achieve an SLE diagnosis when MAP is used, 30% vs. 6% (OR = 6.34, 95% CI 5.12-7.86).

Among patients with no baseline prescription for SLE medication(s), MAP(+) patients were more likely to initiate SLE medications compared to tANA(+) (43% vs. 32%, OR = 1.57, 95% CI 1.41-1.76). In patients new to the practice within the preceding year, the treatment effect was even larger 55% vs. 33% (adjusted OR = 2.77, 95% CI 2.31-3.32).