Background Infections in patients with systemic lupus erythematosus (SLE) are a leading cause of morbidity and mortality. Preventive measures such as immunizations can reduce this burden. The United States experienced a surge in measles infections with outbreaks totaling more than 1,000 cases in 2019. Most of these outbreaks were associated with communities opposed to vaccination for varying reasons. Little attention has been given to communities at risk for measles due to limited access to health care. Current guidelines for screening prior to immune modulating therapy emphasize tuberculosis, opportunistic infections, viral hepatitis, and HIV, but do not discuss measles. Our clinic serves a majority Hispanic, Central American born population with limited access to healthcare. Measles seroprevalence studies of Central American countries have shown less than optimal rates of immunity. The CDC defines measles immunity as a positive titer, or evidence of 2 measles vaccines, or birth prior to 1957. The population threshold for herd-immunity for measles is generally accepted to be 92–94% immune.

We seek to describe immune status to measles with current immunization status in our cohort of underserved patients with rheumatic diseases.

Methods Cross-sectional with a convenience sample of 95 patients with SLE born after 1957 who were seen in a community-health lupus clinic in 2019. All patients were participants in a natural history study of SLE. Titer for anti-rubeola IgG was requested for each patient with their routine clinical lab draw. Immunization records were requested from primary care providers for patients with negative or equivocal titers.

Results We found evidence of sub-optimal levels of immunity within our cohort. Eleven patients (11.5%) had negative or equivocal titers, and none had records documenting prior measles immunization. Only 2 of the 11 non-immune patients were eligible to receive a live vaccine and both of those patients indicated willingness to receive the MMR vaccine at a future visit. Given the small size of the non-immune group, our study was not sufficiently powered to detect differences across groups by region of birth.

Conclusions Our study shows sub-optimal levels of immunity to measles for our cohort of underserved patients with SLE and particularly highlighting missed opportunity for immunization prior to immunosuppression in patients with SLE. In this manner we expand the public health conversation concerning measles immunization in the United States. Not only is this an issue to address in anti-vaccination communities, but it is an important factor in communities with limited access to healthcare affected by rheumatic diseases such as SLE. Our results suggest that screening titers for measles should be considered by rheumatologists prior to the start of immunosuppression. Given that rheumatology patients living in communities with limited access to healthcare are at further risk due to their dysregulated immune systems and immunosuppressive therapies, both rheumatologists and primary care providers can reduce infection risk in these communities by updating immunizations in patients and family members.

Abstract P60 Table 1 Demographics of patients with lupus tested for measles immunity

<table>
<thead>
<tr>
<th>Region of Birth, %, n</th>
<th>Immune*</th>
<th>Non-immune</th>
</tr>
</thead>
<tbody>
<tr>
<td>USA</td>
<td>19 (22.6%)</td>
<td>4 (36.3%)</td>
</tr>
<tr>
<td>Mexico and Central America</td>
<td>35 (41.6%)</td>
<td>4 (36.3%)</td>
</tr>
<tr>
<td>South America and Caribbean</td>
<td>18 (21.4%)</td>
<td>2 (18.1%)</td>
</tr>
<tr>
<td>Asia</td>
<td>8 (9.5%)</td>
<td>0</td>
</tr>
<tr>
<td>Africa and Middle East</td>
<td>4 (4.7%)</td>
<td>1 (9%)</td>
</tr>
</tbody>
</table>

*Immune as defined as rubeola IgG >30 AU/mL.
transglutaminase (IgA) antibodies. Six patients were DQ2 positive.

After being diagnosed of CD and starting the GFD, SLE patients seem to improve especially the leukopenia, lymphopenia and oral aphthosis, as well as SLEDAI score (shown in attached graphics).

Conclusions SLE patients with CD diagnosis and who started a GFD, showed improvement of leukopenia, lymphopenia, oral aphthosis and even SLEDAI.

In SLE patients with recurrent oral aphthosis and/or gastrointestinal unspecific symptoms, CD should be considered, but since serological screening displays a low sensitivity, HLA testing could be helpful. Gastroscopy should be considered, with biopsy and flow cytometer in uncertain cases. Even though, further studies, especially looking for different clinical profiles and longer observational period are needed.

**P62** POLY-AUTOIMMUNITY FREQUENCY IN SLE PATIENTS FROM A TERTIARY HOSPITAL


Background/Purpose Poly-autoimmunity (PAI) is the presence of more than one Autoimmune Disease (AID) in one patient. The coexistence of Systemic Lupus Erythematosus (SLE) with other AIDs is a clinical challenge due to is one of the issues not yet elucidated in medical practice.

We aimed to determine PAI frequency in the context of SLE patients reported in a tertiary hospital.

Methods Cross-sectional observational study with systematic revision of electronic clinical records of SLE patients with other AIDs (from 2014 to 2018) was performed. Demographic, clinical and immunological data were collected.

Results Of 261 SLE patients, 48 (18.39%) had PAI. Mean age was 51.19 (15.35) years (93.75% were female). 2 patients from the 48 (4.16%) had PAI with three AIDs. The 75% of cases developed SLE as the first AID. The mean age at diagnosis of the first AID was 35.52 (13.33) years and mean age at diagnosis of the second AID was 43.75 (16.31) years. A mean difference of 8.31 (9.24) years between the first and second AIDs debut was observed.

The most frequent AIDs registered that go along with SLE are Antiphospholipid Syndrome (APS)(39.58%), Sjögren Syndrome (SS)(31.25%), and Rheumatoid Arthritis (RA) (16.67%). Moreover, in two cases a third AID was registered: SLE-SS-APS and SLE-APS-autoimmune-thyroiditis.

In the SLE-APS group, SLE was the AID of debut in the 89.47% of cases, instead of SLE-RA group with a 62.5%. The SLE-APS group showed a 47.37% of cases with positive antiphospholipid antibodies and 64.71% positive lupus anticoagulant. In the SLE-RA group a 71.43% and 66.67% positive rheumatoid factor and anti-CCP antibody was reported.

Conclusions 18.39% of patients with PAI in our group of SLE patients was observed, mostly with the SLE as the first AID developed. The most frequent association of AIDs in SLE cases were with APS, SS and RA.

**P63** ABNORMAL DISTRIBUTION OF CD27+ IgD+ UNSWITCHED AND CD27+ IgD- SWITCHED MEMORY B CELLS IN SLE PATIENTS EXPOSED TO ORGANIC SOLVENTS

1Carolina Hurtado, 2Diego Rojas-Guadarrón, 3Elsa María Vázquez-Trespalacios, 4Ricardo Pineda, 5Scott Jenks, 6Gloria Vázquez, 7Fulki Sarz. 1School of Graduate Studies and School of Medicine, CES University, Medellin, Colombia; 2School of Medicine, Ces University, Medellin, Colombia; 3Group of Clinical Information, Antemara IPS, Medellin, Colombia; 4Dept. of Medicine, Division of Rheumatology, Lowance Center for Human Immunology, Emory University, Atlanta, USA; 5Grupo de inmunología celular e inmunogenética, Universidad de Antioquia, Medellin, Colombia

Background Some studies in animal models, support an association between occupational exposure to Organic Solvents (OS) and Systemic Lupus Erythematosus (SLE). The specific physio-pathological changes that these chemicals could induce to accelerate an autoimmune response are not known. Dysregulation of B cells is central in SLE, but very little is known on how OS exposure could influence it. This study aimed to examine the distribution of B cell subsets on Healthy Controls and SLE patients occupationally exposed to OS.

Methods 40 SLE patients who met ACR criteria and 17 Healthy Controls were recruited and classified as occupationally exposed or not to OS. Cryopreserved peripheral lymphocytes were analyzed by multiparametric Flow Cytometry using CD3, CD19, CD27, and IgD markers.

Results SLE patients exposed to OS had increased frequencies of CD27+ Switched Memory (SWM) cells. This change was associated with a specific OS like degreasers and ketones. Additionally, the few HC exposed to OS showed a decrease in Unswitched (USM) cells, with similar frequencies as those seen in SLE patients.

Conclusions Exposure to OS increased SWM cells on SLE patients and decreased USM cells on Healthy Controls. The influence of OS on SWM differentiation may be mediated through T cells. Previous reports of exposure to Trichloroethylene (a common OS), showed increased CD4+ T cell activation and secretion of INF-γ, this causes excessive T follicular helper development and germinal center formation in mice that could induce abnormalities in B cell subsets, and a similar mechanism may operate in OS exposed patients. Further research is needed to verify this hypothesis.

**P66** UTILIZATION OF GEOGRAPHIC INFORMATION SYSTEM (GIS) MAPPING TO ASSESS DISSEMINATION OF A LUPUS COMMUNITY BASED HEALTH AWARENESS MODEL

1Karen Mancera-Cuevas, 2Daniel L Erickson, 1Anh Chung, 2Joan S Chmiel, 3Courtnie Phillip, 3Candace Feldman, 4Patricia Canessa, 3Rosaldin Ramsey-Goldman. 1Medicine/Rheumatology, Northwestern University, Chicago; 2Preventive Medicine, Northwestern University, Chicago; 3Division of Rheumatology, Brigham and Womans Hospital, Boston; 3Illinois Public Health Association, Springfield, USA

Background We used a Popular Opinion Leader (POL) model, which leverages community leaders’ social networks to disseminate health information and change norms in vulnerable communities. We established an academic-community partnerships in Chicago and Boston to increase knowledge about lupus and promote early care-seeking behaviors among African American