Mycophenolate mofetil reduces STAT3 phosphorylation in systemic lupus erythematosus patients

Samantha Slight-Webb, … , Paul J. Utz, Judith A. James


Systemic lupus erythematosus (SLE) is a highly variable autoimmune disease that can involve severe organ-threatening symptoms, such as lupus nephritis. Certain drugs, such as mycophenolate mofetil (MMF), are effective at reducing morbidity associated with nephritis; however, the immune pathways associated with disease suppression are poorly defined. Here, we provide evidence that MMF inhibits phosphorylation of STAT3 and other associated immune pathways. Using mass cytometry and bead-based or ELISA assays, the systemic phenotype of SLE patients not taking (MMF–) or taking (MMF+) MMF were studied. MMF+ SLE patients had significant reductions in total numbers of transitional B cells, plasmablasts, and T cells, specifically CD4+ Th17-type and CD4+ Treg-type cells, compared with MMF– patients. Plasma soluble mediators were decreased in MMF+ patients including chemokines (MIG/CXCL9 and SDF-1α/CXCL12) and growth factors (VEGF-A and PDGF-BB). Soluble mediators and cell subsets grouped by functional properties revealed significant modifications associated with STAT3 and B cell pathways. Further, healthy PBMCs treated with IL-6 revealed a reduction in p-STAT3 following the addition of mycophenolic acid (the active metabolite of MMF). In conclusion, the inhibition of STAT3 phosphorylation by MMF may explain the effectiveness of this treatment in SLE patients, since increased levels of p-STAT3 are associated with disease pathology.
Mycophenolate mofetil reduces STAT3 phosphorylation in systemic lupus erythematosus patients

Samantha Slight-Webb, Joel M. Guthridge, Eliza F. Chakravarty, Hua Chen, Rufei Lu, Susan Macwana, Krista Bean, Holden T. Maecker, Paul J. Utz, and Judith A. James

Department of Arthritis and Clinical Immunology, Oklahoma Medical Research Foundation, Oklahoma City, Oklahoma, USA. Departments of Medicine and Pathology, University of Oklahoma Health Sciences Center, Oklahoma City, Oklahoma, USA. Institute for Immunity, Transplantation and Infection, and Division of Immunology and Rheumatology, Stanford University School of Medicine, Stanford, California, USA.

Systemic lupus erythematosus (SLE) is a highly variable autoimmune disease that can involve severe organ-threatening symptoms, such as lupus nephritis. Certain drugs, such as mycophenolate mofetil (MMF), are effective at reducing morbidity associated with nephritis; however, the immune pathways associated with disease suppression are poorly defined. Here, we provide evidence that MMF inhibits phosphorylation of STAT3 and other associated immune pathways. Using mass cytometry and bead-based or ELISA assays, the systemic phenotype of SLE patients not taking (MMF–) or taking (MMF+) MMF were studied. MMF+ SLE patients had significant reductions in total numbers of transitional B cells, plasmablasts, and T cells, specifically CD4+ Th17-type and CD4+ Treg-type cells, compared with MMF– patients. Plasma soluble mediators were decreased in MMF+ patients including chemokines (MIG/CXCL9 and SDF-1α/CXCL12) and growth factors (VEGF-A and PDGF-BB). Soluble mediators and cell subsets grouped by functional properties revealed significant modifications associated with STAT3 and B cell pathways. Further, healthy PBMCs treated with IL-6 revealed a reduction in p-STAT3 following the addition of mycophenolic acid (the active metabolite of MMF). In conclusion, the inhibition of STAT3 phosphorylation by MMF may explain the effectiveness of this treatment in SLE patients, since increased levels of p-STAT3 are associated with disease pathology.

Introduction

Systemic lupus erythematosus (SLE) is a chronic, systemic autoimmune disease, characterized by a heterogeneous clinical presentation driven by autoreactive lymphocytes and proinflammatory cytokine production. Lupus inflammation can often lead to renal injury and multiple organ damage even when treated (1). Mycophenolate mofetil (MMF) is an immunosuppressive agent that was first shown to be effective in preventing organ rejection in renal transplant patients, but more recently was found to be efficacious in lupus, especially in patients with lupus nephritis (2, 3). Compared with cyclophosphamide, MMF was found to elicit a similar reduction in disease signs and symptoms in patients with severe lupus, but with a safer side-effect profile (4, 5). MMF is now considered a standard treatment for lupus nephritis, while further investigation into its usefulness for nonrenal manifestations is underway (6). Despite the known safety and utility of MMF in lupus treatment, the systemic impact of MMF treatment on cellular subsets, cell activation, and soluble mediator pathways in SLE remains poorly defined.

MMF is the prodrug of mycophenolic acid (MPA), which reversibly inhibits inosine monophosphate dehydrogenase (IMPDH), and most strongly inhibits the IMPDH II isoform (7). IMPDH is a rate-limiting enzyme in the de novo synthesis of guanine nucleotides and 2-deoxyribonucleotides. The type II isoform is primarily expressed in activated lymphocytes, while the type I isoform is expressed in almost all cell types. While it is recognized that MMF can affect many cell types, the cytostatic effect on lymphocytes is the principal mechanism of action by which immunosuppression is thought to occur (7). In lupus, patients taking MMF were found to have elevated frequencies and numbers of B cells compared with SLE patients not taking immunosuppressants; however, plasmablast numbers and frequencies were decreased in patients
taking MMF compared with controls (8, 9). In a separate study, SLE patient samples taken at baseline and 4 weeks after MMF treatment had elevated frequencies of CD4+ T cells and NKT cells and decreased frequencies of CD8+ T cells and NK cells after 4 weeks, but no significant differences in B cells were observed and total numbers by cell type were not reported (10). In addition to phenotypic changes, MMF was found to induce epigenetic modifications in lupus CD4+ T cells by enhancing H4 acetylation and H3K4 trimethylation that increased regulatory miRNA levels and decreased expression of CD40L (11, 12). Variable results, and the absence of data on dendritic cell (DC) and monocyte alterations in SLE patients taking MMF, leave much to be determined for the cellular mechanisms of MMF in SLE pathogenesis.

Although some work has been done to identify phenotypic changes in SLE patients taking MMF, little to no data exist about the effect of MMF on the systemic cytokine milieu. SLE patients are characterized by elevated plasma levels of Th1, Th2, Th17, and myeloid-derived proinflammatory cytokines and chemokines such as interleukin 1α (IL-1α), IL-2, IL-5, IL-10, IL-13, IL-17, interferon-γ (IFN-γ), IP-10, G-CSF, and TNF-α (13). Many SLE patients are also distinguished from healthy controls by elevated blood levels of type I IFN accompanied with upregulation of IFN-inducible genes (13, 14). After 48 hours in vitro, MMF reduced the production of IL-2, IL-3, IL-4, IL-5, IL-6, IL-10, IFN-γ, and TNF-α in human lymphocytes (7). In lupus patients assessed for systemic IFNs, type I IFN was decreased in patients taking MMF compared with patients not taking MMF, but no significant difference was observed in IFN-γ production (15). In CD4+ T cells isolated from lupus patients and treated with either MPA or methanol, IL-17F mRNA levels were downregulated in cells treated with MPA, while no differences were observed in IL-2, IL-17A, or IFN-γ mRNA levels (11).

Using observational data collected from peripheral blood mononuclear cells (PBMCs) of SLE patients taking MMF (MMF+) and control SLE patients matched by age, gender, disease activity, other medication use, and ethnicity that were not taking MMF (MMF−), we assessed differences in cell numbers and activation for all major peripheral cell subsets using a comprehensive mass cytometry staining panel. In addition, 38 plasma soluble mediators were measured to assess the effect of MMF on systemic lupus inflammation and to identify the involvement of particular cellular pathways in MMF-associated immunomodulatory pathways in SLE. To examine STAT3 regulation following MMF treatment, in vitro studies using IL-6–stimulated PBMCs treated with MPA were analyzed by phosphoepitope flow cytometry.

Results

Phenotype evaluation by mass cytometry in MMF+ SLE patients. To understand the systemic immune changes that occur in MMF+ SLE patients, we utilized a mass cytometry panel to 33 heavy metal–labeled antibodies to identify major and minor immune cell subsets in patients taking or not taking MMF. The panel included a broad range of markers to allow segregation of T cells (CD4+ and CD8+ memory, naive, and effector cell subsets and TCRγδ+), B cells (naive, transitional, memory, plasmablasts, double-negative, and nonswitched memory), monocytes (classical, nonclassical, and intermediate), DCs, NK cells (CD8+/-, CD57+/-, CD16+/-, CD94+/-), and NKT cells (Supplemental Tables 1 and 2; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.124575DS1). Cell activation and chemokine markers were also included to assess activation and functional properties of cell subsets. To evaluate cellular profiles, a high-dimensional method of analysis using t-distribution stochastic neighbor embedding (t-SNE) and a traditional bivariate plot analysis were both utilized. t-SNE creates an unsupervised 2-dimensional map by using the collected cellular data to draw pairwise comparisons and plot similar cells near each other.

In our analysis, t-SNE reduced 33 parameters into 2 dimensions creating 22 distinct clusters (Figure 1A). The median intensity of each cellular marker within a cluster was assessed by gating and projection into a heatmap to classify cell subsets (Figure 1B). Following gating, cell frequencies were determined and total cell numbers/ml were back-calculated. MMF+ SLE patients trended lower in the total number of circulating cells/ml, but was not statistically significant (Figure 2A). However, 3 populations were found to have significantly fewer numbers in MMF+ patients by t-SNE, namely, T cells (CD3+CD56−), Th17-type cells (CD3+CD4+CCR6+), and plasmablasts (CD19+IgD CD27+CD38+) (Supplemental Table 1).

Using a standard biaxial gating strategy, we confirmed t-SNE results and identified additional cell subtypes such as T effector cells (Th1-, Th2-, Th17-, Treg-, and Thh-type cells), CXCR3+ DCs, and NK cell subsets for a total of 55 cell subsets (Supplemental Figure 1). We identified differences in cell numbers of 5 cell types using bivariate plots, all significantly lower in MMF+ patients. T cells, specifically Th17-type cells (CCR6+CD161+ and Treg-type cells (CD25+CD127+), transitional B cells (IgD+CD24+CD38+CD27+), and CD8+CD56+ NK cells (Figure 2, B–F) were significantly reduced in MMF+ patients (Supplemental Table
2). CXCR3+ DCs, plasmacytoid DCs, and CD94− and CD57− NK cells also trended lower in numbers, but did not reach statistical significance (Supplemental Table 2). Based on these observations, MMF targets T cell, B cell, and NK cell subsets in SLE patients, primarily CD4+ T effector cells and CD38hi B cells.

Cell subset activation status and MMF use. To better define the activation status of our defined cell populations we analyzed the frequencies of activation (CD27, CD28, CD38, CD86, HLA-DR, and ICOS), regulatory (CD85j, CD94, CD33, and PD-1), and chemokine (CCR6, CCR7, CXCR3, and CXCR5) receptor expression on each gated cell subset (Supplemental Table 3). Although MMF+ patients had no observed differences in naive, memory, or double-negative B cell numbers compared with MMF− patients (Supplemental Tables 1 and 2), the expression of CD38, CD86, and CCR6 on naive B cells and CD86 on memory B cells was at significantly lower frequencies on peripheral blood cells of MMF+ patients compared with MMF− patients (Figure 3, A–E). Plasmablasts, which were decreased systemically in MMF+ patients by t-SNE analysis, also had lower expression of CCR6, a chemokine receptor upregulated after activation in B cells and important for the recall response to antigen (16, 17) (Figure 3, D and F). Despite differences in memory B cells, IgD−CD27− double-negative B cells, also known as nonconventional memory B cells, only trended lower in CCR6, CD38, and CD86 expression (Figure 3, G–I).

B cell activation markers and CCR6 expression were significantly lower in MMF+ patients. In addition, T cells also had lower expression of CCR6, specifically on CD8+ memory T cells, in MMF+ patients compared with MMF− patients (Figure 3, D, J, and K). CD4+ T cells and CXCR3+ DCs had lower expression of the activation marker CD38 in MMF-treated SLE patients (Figure 3, L and M), suggesting disease suppression by MMF may occur by hindering both adaptive and innate cell pathways.
Plasma soluble mediator levels negatively correlate with MMF use. As activation marker frequency was diminished in MMF+ patients, we expected specific proinflammatory cytokines, growth factors, and chemokines to also be downregulated in MMF+ SLE patients. To assess this, 38 soluble mediators were assayed in plasma by multiplex assays and ELISA assays. The majority of soluble mediators were lower in the plasma of MMF+ SLE patients including IL-5, IL-7, IP-10/CXCL10, and MCP-1 compared with MMF– patients (Figure 4A and Supplemental Table 4). In addition, vascular endothelial growth factor (VEGF) and platelet-derived growth factor (PDGF-BB) and the chemokines stromal cell–derived factor 1 (SDF-1α/CXCL12) and IFN-γ–inducible MIG/CXCL9, were all downregulated in MMF+ SLE patients (Figure 4, B–E). The most significantly depressed soluble mediator was VEGF (P = 0.0007), which drives angiogenesis, capillary permeability, and endothelial cell proliferation and chemotaxis.

STAT3 phosphorylation and associated immune pathways are downregulated by MMF. To determine the cellular pathways most affected by MMF treatment in SLE patients, cytokines, chemokines, and significant cell populations were grouped by functional properties (Figure 5A) or biomarkers/variables impacted by STAT signaling (Figure 5B) using the PANTHER classification system, which categorized our variables into the following groups: apoptosis signaling, cell growth, angiogenesis, cell differentiation, cell migration, B cell activation, T cell activation, innate cell activation, adipocyte activation, regulatory and adhesion groups, or STAT1/2/3 associated, STAT1/3/5 associated, STAT3 associated, STAT5 associated, and STAT6 associated. Group medians of each variable were converted to Z scores and plotted using a radar plot (Figure 5, A and B). STAT3-associated variables, namely VEGF, PDGF-BB, CD4+ Th17-type cells, and CD4+ Treg-type cells, were most significantly reduced in MMF+ SLE patients, followed closely by diminished activation in B cell pathways, where STAT3 is also required to form long-lived antibody responses (18).

To investigate whether MMF reduced STAT3 phosphorylation, PBMCs from healthy individuals were stimulated with IL-6 and incubated with MPA for 3 hours at varying concentrations. Phosphoepitope flow analysis revealed a significant reduction in phosphorylated STAT3 following incubation with 10 μM MPA, a concentration equivalent to therapeutic levels found in SLE patients (19, 20) (Figure 6A), with no differences observed in phosphorylated STAT5 (Figure 6B) or cell apoptosis (Figure 6C) after 3 hours. Furthermore, reductions in STAT3 phosphorylation were similar across all major cell populations (Figure 6D). Together, these data suggest that p-STAT3 reduction is a principal mechanism of MMF-induced SLE disease suppression.
# Discussion

One of the more severe manifestations of SLE, lupus nephritis, occurs in up to 60\% of patients during the course of the disease (21). MMF is becoming a mainstay of treatment of lupus nephritis (4, 5). In general, the use of MMF in severe SLE patients leads to significant disease suppression, and maintenance therapy with MMF can prevent relapse of nephritis. The therapeutic efficacy of MMF in reducing disease activity is undisputed; however, the cellular alterations and systemic inflammatory pathways responsible for the successful actions of MMF have been unclear. In the current study, in-depth phenotyping of 55 cell subsets and 38 plasma soluble mediators, and in vitro p-STAT3 assessment, provide pivotal insight into the mechanism of MMF in SLE.

STAT3 is a critical regulator of numerous cellular processes including pathways that drive cell proliferation, differentiation, migration, inflammation, and apoptosis (22). The significance of STAT3 in biological actions is revealed in the disruption of the STAT3 gene in mice, which results in early embryonic death (23). STAT3 is the only STAT gene that is developmentally lethal. Conditional deletions of STAT3 have revealed its importance in CD4\(^+\) T cell differentiation, and effects on Th17, Treg, and Th9 pathways (24–26). Further, STAT3 is required for memory B cell development and T-dependent antibody responses (27).

In SLE, total and phosphorylated levels of STAT3 are elevated and correlate with increased Th17 cells, organ involvement, and SLE disease activity of patients (28–30). Our study is the first to our knowledge to demonstrate that MMF reduces phosphorylation of IL-6–driven STAT3 in healthy PBMCs and may be a primary method for limiting disease activity in SLE.

Further support that MMF hinders the STAT3 pathway was observed in systemic reductions of Th17-type cells (CCR6\(^+\)/CD161\(^+\)) (31) in MMF+ SLE patients. Th17 cells and IL-17A levels are elevated during active SLE (32, 33). CCR6 and CD161 expression distinguish CD4\(^+\) T cell populations that produce IL-17...
insight.jci.org   https://doi.org/10.1172/jci.insight.124575

FIGURE 4. Growth factor and chemokine downregulation distinguish MMF+ SLE patients from MMF− patients. Proinflammatory soluble mediators (38 total) were measured by multiplex or ELISA. Overall trends in relative plasma soluble mediator levels are depicted by (A) radar plot using the group median Z scores. MMF− SLE patients (n = 10) are highlighted in blue and MMF+ SLE patients (n = 5) are in red. Soluble mediators are represented by dot plots and include (B) VEGF, (C) PDGF-BB, (D) SDF-1α/CXCL12, and (E) MIG/CXCL9. *P < 0.05, ***P < 0.001 by Kruskal-Wallis test with Dunn’s multiple comparisons test. Median ± standard deviation shown.

(32), and reductions were observed in Th17-type cells of SLE patients treated with MMF. In vitro studies with CD4+ lupus T cells found decreased IL-17F mRNA expression in MPA-treated cells compared with methanol-treated cells, but no differences in IL-17A mRNA levels (11). Interestingly, IL-17+ CD4+ T cells have been found to be targeted by MMF compared with IFN-γ CD4+ T cells in a mouse diabetes model (34). The decrease in Th17 cells of MMF+ lupus patients was also associated with a reduction in Treg-type cells (CD25hiCD127–). Although this may seem counterintuitive as the number and function of Treg cells negatively correlate with lupus disease activity (33), MMF has been found to decrease CD4+CD25+Foxp3+ Treg cells in transgenic T cell, graft-versus-host, and experimental ischemia-reperfusion mouse models (31, 35, 36). Further, STAT3 is required for differentiation and expression of FoxP3 in Tregs (37). Our data suggest that MMF may have a more profound impact on Th17 and Treg cells in SLE patients that may be important in disease suppression; however, the differential effects on induction and maintenance of Th17 and Treg cells in a longitudinal study looking at IL-17 and FoxP3 by single-cell analysis should be considered.

The results of the current study indicate that B cells are a major target of MMF, which had a significant effect on almost all B cell subsets, except IgD Ig27- double-negative B cells. Decreases in cell activation were observed in naive B cells, transitional B cells, memory B cells, nonswitched memory B cells, and plasmablasts in MMF+ patients. Furthermore, a decrease in CD38hi B cell subsets (plasmablasts and transitional B cells), as found either by t-SNE or manual gating, were found in patients taking MMF. A reduction in CD38hi B cells in transplant patients and a reduction of plasmablasts in MMF-treated SLE patients has been previously observed; thus, our data confirm the impact of MMF on activated B cell populations in SLE (8, 9, 38, 39).
In addition, MPA inhibited expansion, activation, and immunoglobulin production in primary B cells in vitro from both healthy individuals and rheumatoid arthritis patients; however, MPA had little effect on the expansion of terminally differentiated plasma cells (40). This suggests that reductions in plasmablast numbers in MMF-treated SLE patients may largely be due to blocking of plasma cell differentiation. Decreases in transitional B cells also suggest a possible impact of MMF on B cell maturation out of the bone marrow, and further in vitro investigation into cell subset–specific maturation and expansion in SLE patients is of interest. Despite the known effects of MMF on B cell proliferation (40), double-negative B cells were not reduced in patients taking MMF in this study. The lack of differences may be due to the known exhausted or anergic phenotype of double-negative B cells, also known as nonconventional memory cells, which may result in a reduced ability to expand and respond to stimuli (41). Our data suggest that MMF-induced STAT3 inhibition profoundly effects B cells, most specifically later-stage CD38hi B cells to suppress lupus disease.

Finally, 38 systemic cytokines were assessed in SLE patients on MMF with significant differences found in only 4 key cytokines, VEGF, PDGF-BB, SDF-1α/CXCL12, and MIG/CXCL9, which were all reduced in patients taking MMF. VEGF and PDGF-BB, both of which are angiogenesis factors, were most significantly affected by MMF treatment. SLE is known to be associated with immune-mediated vascular damage and angiogenesis. Elevated VEGF levels correlate with vascular damage in SLE patients, which positively associates with disease activity, suggesting VEGF is a potential driver of SLE endothelial dysfunction (42). VEGF has also been found to be downregulated following treatment with MMF in several murine models (43, 44). The VEGF gene is regulated directly by the STAT3 protein, and decreases in VEGF may directly contribute to decreases in peripheral B cell and T cell numbers (45–48). Further, we also found significant reductions in 2 chemokines, MIG/CXCL9 and SDF-1α/CXCL12, in patients taking MMF. MIG is largely produced by activated B cells, monocytes, and DCs, and reduction in MIG may be due to decreases in B cell activation and numbers seen in
MMF-treated patients. In addition, SDF-1α is critical for hematopoiesis including B lymphocyte development within the bone marrow niche (49). Reductions in SDF-1α likely contribute to lower B cell numbers and alterations in the B cell compartment, including decreased transitional B cells, seen in MMF+ patients.

STAT inhibitors are a promising family of therapeutic targets for lupus. Indeed, STAT3 inhibition in MRL/lpr lupus-prone mice inhibited the onset of lupus nephritis, decreased anti-dsDNA antibodies, and reduced inflammatory cytokine production (50). We have identified inhibition of activated STAT3 as a putative mechanism of disease suppression by MMF in patients with SLE. These data hold promising information for drug therapy development utilizing MMF and associated pathway inhibitors. Future studies utilizing a larger cohort assessed longitudinally for STAT3 phosphorylation and upstream proteins, such as the mammalian target of rapamycin pathway and the mitogen-activated protein kinase pathway, following MMF treatment are of particular interest and necessary in determining the involvement of MMF in STAT3 inhibition.

Methods
Study population. PBMCs and plasma samples from 10 SLE patients not taking MMF and 5 SLE patients taking MMF were studied. Patients were recruited at the Oklahoma Medical Research Foundation (OMRF). SLE patients fulfilled the American College of Rheumatology (ACR) criteria for SLE classification. Subjects were matched by gender, ethnicity, age ± 5 years, medication use, and disease activity by SELENA-SLEDAI (Supplemental Table 5) (51). Similar frequencies of SLE patients with active disease, as defined by SLEDAI ≥ 4, are found in both groups, with 50% of MMF– patients and 60% of MMF+ patients. PBMCs were isolated using Lymphocyte Separation Medium (Mediatech, Inc.) and stored in freezing media (20% human serum and 10% DMSO in RPMI) at –80°C until use. Plasma was also collected and stored at –80°C until use.

CyTOF immunophenotyping. Assays were performed in the Human Immune Monitoring Center at Stanford University. Briefly, PBMCs were stained with a surface antibody cocktail specific for CCR6, CCR7, CD11b, CD11c, CD123, CD127, CD14, CD16, CD161, CD19, CD161, CD19, CD20, CD24, CD25, CD27, CD28, CD3, CD33, CD38, CD4, CD45RA, CD56, CD57, CD8, CD85j, CD86, CD94, CXCR3, CXCR5, HLA-DR, ICOS, IgD, PD-1, and TCRγδ (see Supplemental Table 6 for clone list, supplier, and metal isotope utilized) for 60 minutes on ice. PBMCs were then stained with DOTA-maleimide (Macrocyclics) containing natural-abundance indium (viability marker). The cells were resuspended in iridium-containing DNA intercalator and incubated at room temperature for 20 minutes. The cells were washed twice and diluted before injection into the CyTOF (DVS Sciences). Data analysis was performed using Cytobank by gating on intact cells based on the iridium isotopes of the intercalator, and then on singlets by Ir-191 versus cell length, and then on live cells (indium-Live/Dead–negative population), followed by cell subset–specific gating or viSNE analysis.

Figure 6. Mycophenolic acid reduced STAT3 phosphorylation in vitro. Healthy control PBMCs from 6 individuals were treated with IL-6 (100 ng/ml) for 15 minutes prior to incubation with varying levels of MPA (0 μM, 1 μM, or 10 μM) for 3 hours. Phosphoepitope flow assessment of (A) p-STAT3 MFI, (B) p-STAT5 MFI, and (C) active caspase 3/7 MFI were determined following fixation and staining of cells immediately following the 3-hour incubation. Major cell populations were assessed for differences in (D) p-STAT3 inhibition including T cells, B cells, monocytes, NK cells, and dendritic cells compared between 0 μM MPA and 10 μM MPA. *P < 0.05, ***P < 0.001 by Kruskal-Wallis test with Dunn’s multiple comparisons test. Median ± standard deviation shown.
Soluble mediator measurement. Plasma of SLE patients was analyzed for the concentration of soluble mediators using a multiplex bead-based assay (eBioscience/Affymetrix) to measure sCD40L, CXCL13, eotaxin, G-CSF, GROα, ICAM-1, IFN-α, IFN-β, IFN-γ, IL-1α, IL-1β, IL-1RA, IL-10, IL-12p70, IL-13, IL-15, IL-17α, IL-2, IL-21, IL-23, IL-2RA, IL-4, IL-5, IL-6, IL-7, IL-8, IP-10, leptin, LIF, MCP-1, MCP-3, MIG, MIP-1β, MIP-1α, NGF-β, PAI-1, PDGF-BB, RANTES, resistin, SCF/c-kit ligand, SDF-1α/CXCL12, sEslectin, sFasL, TGF-β, TNF-α, TNFR1, TNFR2, TRAIL, VCAM-1, and VEGF and run on a Bio-Plex 200 suspension array reader (Bio-Rad). A standard control serum was included on each plate (Cellgro human AB serum, Mediatech). Soluble mediator concentrations (pg/ml) were calculated using Bio-Plex manager software using 5-parameter logistic curve fitting. Soluble mediators with >60% of concentration values reported as out of range in both treatment groups were dropped from the analysis to minimize data loss from row-based deletion of missing values. These mediators included IFN-α, IFN-γ, IL-1α, IL-1β, IL-10, IL-12p70, IL-13, IL-15, IL-17α, IL-23, IL-4, NGF-β, and TGF-β. The 38 soluble mediators that passed quality control were normalized and values were used for subsequent analyses. Plasma levels of BLyS were assessed using enzyme-linked immunosorbent assay (ELISA) per the manufacturer’s protocol (R&D Systems).

Pathway analyses. Functional properties and STAT pathways for all cytokines and relevant cell receptors from significant cell populations were assessed using the PANTHER classification system (version 13.1) (52), which is used to classify protein family/subfamilies, biological process, molecular function, and signaling pathways. All proteins were assessed for pathways and functional properties within PANTHER, and are reported in Supplemental Table 7. Cytokines and cell subsets were grouped according to function or STAT-associated pathway outputs, and radar plots were created as mentioned in the data analysis section below.

STAT3 phosphorylation by phosphoepitope flow. Previously frozen PBMCs were thawed, washed, and resuspended in supplemented RPMI 1640 at 1 × 10^6 cells/well in a 24-well plate. Cells were rested 1 hour, and then stimulated with IL-6 (100 ng/ml) for 15 minutes prior to addition of MPA, the active metabolite of MMF, at varying concentrations (0 μM, 1 μM, or 10 μM). Cells were incubated with MPA for either 0 or 3 hours, after which cells were washed with cold PBS and stained with near-IR Live/Dead stain (Thermo Fisher Scientific). PBMCs were fixed in paraformaldehyde, washed, and Fc receptor blocked. Cells were then permeabilized with saponin and stained for lineage markers (CD3, CD4, CD8, CD20, CD14, CD11c, and CD56) followed by staining for intracellular markers for p-STAT3 and p-STAT5 (BD Biosciences). FLICA caspase 3/7 stain (ImmunoChemistry Technologies, LLC) was added 1 hour prior to the end of MPA incubation. Cells were resuspended in FACS buffer and immediately run on a FACSCelesta (BD Biosciences).

Data analysis. t-SNE analysis were performed using Cytobank’s viSNE (53). t-SNE was performed after gating off live, intact singlet cells. For frequency analysis, 24,000 events were used per sample. Cell clusters were gated and frequencies exported for cell number calculations and analysis. Concatenated files of 110,000 cells per group were used to generate representative t-SNE images. Manual gating was also performed on individual subject files for B cells, monocytes, NK cells, T cells, and DCs in Cytobank (Supplemental Figure 1, A–D).

Statistics. Dot plots, heatmaps, statistical analyses, and multiple testing corrections were performed using GraphPad Prism 6.0 for Windows or TIBCO Spotfire 6.0.1 (TIBCO Software Inc.). Radar plots were generated by using the group median of modified Z scores in Microsoft Excel. P values less than 0.05 were considered significant (*P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001). All error bars indicate the standard deviation.

Study approval. All work contained in this publication was approved by the Institutional Review Board at the Oklahoma Medical Research Foundation. Blood donors gave written informed consent prior to study inclusion in accordance with the ethics principles stated in the Helsinki Declaration.

Author contributions
SSW analyzed data, wrote the manuscript, and designed and conducted experiments. JMG and JAJ designed research studies and experiments, and wrote the manuscript. EFC designed research studies. HC, RL, and KB analyzed data. SM designed and conducted experiments and acquired data. HTM and PJU designed and conducted experiments.

Acknowledgments
We would like to thank Virginia Roberts and Wade DeJager for technical assistance. In addition, we would like to thank the Stanford Human Immune Monitoring Center (HIMC) for performing the CyTOF phenotyping assays. Research reported was supported by NIAID, NIAMS, and NIGMS of the NIH (U19AI082714, U19AI082719, U54GM104938, RC1AR058554, P30GM103510, U01AI101934, U10AI143822).
P30AR053483, T32AI007633 [to SSW], and S10RR026735) and was conceived through interactions of the NIAID Autoimmunity Center of Excellence. The contents are solely the responsibility of the authors and do not necessarily represent the official views of the NIH or its institutes.

Address correspondence to: Judith A. James, Arthritis and Clinical Immunology, Oklahoma Medical Research Foundation, 825 N.E. 13th Street, Oklahoma City, Oklahoma 73104, USA. Phone: 405.271.4987; Email: Judith-James@omrf.org.


