CD11b activation suppresses TLR-dependent inflammation and autoimmunity in systemic lupus erythematosus

Mohd Hafeez Faridi, …, Mariana J. Kaplan, Vineet Gupta


Genetic variations in the *ITGAM* gene (encoding CD11b) strongly associate with risk for systemic lupus erythematosus (SLE). Here we have shown that 3 nonsynonymous *ITGAM* variants that produce defective CD11b associate with elevated levels of type I interferon (IFN-I) in lupus, suggesting a direct link between reduced CD11b activity and the chronically increased inflammatory status in patients. Treatment with the small-molecule CD11b agonist LA1 led to partial integrin activation, reduced IFN-I responses in WT but not CD11b-deficient mice, and protected lupus-prone MRL/Lpr mice from end-organ injury. CD11b activation reduced TLR-dependent proinflammatory signaling in leukocytes and suppressed IFN-I signaling via an AKT/FOXO3/IFN regulatory factor 3/7 pathway. TLR-stimulated macrophages from *CD11B* SNP carriers showed increased basal expression of IFN regulatory factor 7 (IRF7) and IFN-β, as well as increased nuclear exclusion of FOXO3, which was suppressed by LA1-dependent activation of CD11b. This suggests that pharmacologic activation of CD11b could be a potential mechanism for developing SLE therapeutics.

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CD11b activation suppresses TLR-dependent inflammation and autoimmunity in systemic lupus erythematosus

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Genetic variations in the ITGAM gene (encoding CD11b) strongly associate with risk for systemic lupus erythematosus (SLE). Here we have shown that 3 nonsynonymous ITGAM variants that produce defective CD11b associate with elevated levels of type I interferon (IFN-I) in lupus, suggesting a direct link between reduced CD11b activity and the chronically increased inflammatory status in patients. Treatment with the small-molecule CD11b agonist LA1 led to partial integrin activation, reduced IFN-I responses in WT but not CD11b-deficient mice, and protected lupus-prone MRL/Lpr mice from end-organ injury. CD11b activation reduced TLR-dependent proinflammatory signaling in leukocytes and suppressed IFN-I signaling via an AKT/FOXO3/IFN regulatory factor 3/7 pathway. TLR-stimulated macrophages from CD11B SNP carriers showed increased basal expression of IFN regulatory factor 7 (IRF7) and IFN-β, as well as increased nuclear exclusion of FOXO3, which was suppressed by LA1-dependent activation of CD11b. This suggests that pharmacologic activation of CD11b could be a potential mechanism for developing SLE therapeutics.

Introduction
Systemic lupus erythematosus (SLE, lupus) is a debilitating autoimmune disease that is characterized by hyperactive immune cells, serum autoantibodies, immune complex deposition, multiorgan damage, and accelerated vascular disease (1). Pathways downstream of various pattern recognition receptors — in particular, Toll-like receptors (TLRs) — are central to the aberrant immune responses contributing to SLE and result in elevated levels of inflammatory cytokines, such as type I interferons (IFN-I), IL-1β, IL-6, TNF-α, and IL-17, in patients (2). Elevated levels of IFN-I in circulation are a heritable risk factor for SLE and play a pathogenic role (3). Although the underlying mechanisms influencing IFN-I levels in SLE subjects and conferring a strong predisposition to disease remain to be fully characterized, genetic variations, in combination with environmental stressors, play a key role. Genome-wide association studies have identified single-nucleotide polymorphisms (SNPs) in a number of immune response genes as key contributors to disease activity, and SNPs in ITGAM (coding for CD11b or αM, the α chain of the β2 integrin) are strongly associated with SLE risk (4). These include genetic variants that produce a decreased affinity for the β2 integrin partner, CD11c, resulting in reduced receptor function (5). However, the link between genetic variation in ITGAM and disease pathogenesis is not fully understood.

Here we report that genetic variants in the ITGAM gene strongly correlate with elevations in type I interferon (IFN-I) in SLE and have a pathogenic role in lupus disease pathogenesis. 3 nonsynonymous variants in ITGAM produce defective CD11b and lead to elevated IFN-I in lupus. Pharmacologic activation of CD11b via a small molecule reduces IFN-I signaling and proinflammatory activity in lupus.

Authorship note: M.H. Faridi, S.Q. Khan, W. Zhao, H.W. Lee, and M.M. Altintas contributed equally to this work.

Conflict of interest: V. Gupta is an inventor of pending patents related to this study (WO2013159082) and has the potential for financial benefit from their future commercialization. V. Gupta is also a cofounder and a member of the scientific advisory board of Adhaere Pharmaceuticals Inc., a company developing novel therapeutics for commercialization. V. Gupta is an inventor of pending patents related to this study.
kocytic integrin heterodimer CD11b/CD18) show highly significant correlation with SLE (4–6). Three ITGAM coding region SNPs (7, 8) (rs1143678, rs1143679, and rs1143683), which result in missense mutations P1146S (a C>T substitution), R77H (a G>A substitution), and A858V (a C>T substitution), respectively, in the protein, have strong correlation with the incidence of SLE and with SLE subphenotypes including lupus nephritis, discoid rash, and immunologic manifestations (reviewed in ref. 9). How these nonsynonymous mutations confer SLE risk is not clear, but they appear to reduce CD11b function, including integrin activation, ligand binding and cell adhesion, phagocytosis, and catch-bond formation (7, 10–13). Recent studies also show that CD11b acts as a negative regulator of TLR signaling pathways and of B cell autoreactivity (14, 15), implying that the ITGAM SNPs found in SLE patients may affect disease activity via a reduction in CD11b’s normal, antiinflammatory signaling activities in leukocytes, although the exact mechanism behind how these variants contribute to autoimmunity is unclear. In murine systems, while CD11b deficiency was shown to reduce neutrophil accumulation and glomerular injury in an acute model (16), it has also been shown to increase susceptibility to chronic inflammatory and autoimmune diseases (refs. 14, 17–21, and reviewed in ref. 22), including increased tissue infiltration of leukocytes and immune-mediated injury in lupus-prone mice (19, 20). These data further suggest that CD11b plays a protective role in SLE. Here, we show that SLE patients carrying SNPs in the ITGAM gene present with elevated IFN-I activity, molecularly linking reduced CD11b function with increased disease risk. Ex vivo, cells homozygous for ITGAM SNPs showed a basal increase in IFNB and IRF7 expression and a decreased level of nuclear FOXO3, which regulates their expression, suggesting reduced suppression of TLR signaling at a basal level. Pharmacologic activation of CD11b, protein product of ITGAM, reduced IFN-I responses ex vivo and in animals via suppression of TLR-dependent FOXO3/IRF7 pathway in a CD11b-dependent fashion. It also similarly protected lupus-prone mice from end-organ injury. Suppressing overactive immune responses is a key strategy for treating SLE, and our data suggest that allosteric activation of CD11b with small molecules could be a novel therapeutic strategy.

Results

**SLE subjects carrying ITGAM SNPs have significantly elevated serum IFN-I activity.** To test for a direct link between ITGAM SNPs and the IFN-I pathway in SLE, we measured the serum IFN-I activity (3) in 171 SLE subjects of European ancestry and determined their ITGAM genotype (Supplemental Table 1; supplemental material available online with this article; https://doi.org/10.1172/JCI88442DS1). This IFN assay was previously normalized using 100 healthy controls to establish normal values for serum IFN in healthy individuals (3). We found that the ITGAM risk alleles showed strong association with high IFN-I serum activity in patients (Figure 1A). Furthermore, analysis showed that the SNPs rs1143678 and rs1143683 were in complete linkage disequilibrium, forming a haplotype (not shown). Regression analysis demonstrated that alleles in rs1143679 and rs1143678/rs1143683 showed independent evidence for association with elevated IFN-I serum activity. Surprisingly, we found that a subset of these subjects (21/171) carried a haplotype containing the common SNP in rs1143679 (G) along with the rare SNP in rs1143683 (T). This GT haplotype was also associated with significantly elevated IFN-I activity (odds ratio = 3.57, \( P = 0.0068 \)) (Figure 1B). To examine whether there was increased disease activity or flare in the ITGAM risk allele carriers at the time of blood draw that might account for this association, we examined their SLE Disease Activity Index (SLEDAI) scores, which were available for approximately 80% of the SLE patients at the time of blood draw. The average SLEDAI score in the overall cohort was 3.16, indicating mild to moderate average disease activity, typical of an outpatient SLE cohort. Next, we examined the SLEDAI scores in the same categories as those in Figure 1, A and B (Supplemental Figure 1). While there seems to be a trend toward higher disease activity, we did not find any significant increase in SLEDAI in the ITGAM risk allele carriers versus noncarriers. Furthermore, we examined it in a multivariate model, by performing a logistic regression analysis to assess the impact of SLEDAI score on the relationship between the ITGAM SNPs and IFN-I levels. SNPs were entered as predictor variables, with IFN as the outcome, and the model was assessed with and without inclusion of SLEDAI score. The \( \beta \) coefficients and odds ratios for the associations between SNP genotype and IFN did not change significantly between the 2 models (<10%), and the SLEDAI variable did not show significant association with the outcome in the overall model (odds ratio = 1.04), further supporting the idea that SLEDAI was not confounding the relationship observed between SNP genotype and IFN-I.

An issue that commonly arises in the population cytokine data is that some of the data appear to fit a bimodal pattern, and the serum IFN levels in patients with the ITGAM GT genotype appear to show a similar behavior. To address this, we have previously reported an independent methodology for data analyses that uses...
Figure 2. LA1 induces intermediate-affinity conformation in CD11bA and reduces TLR-stimulated proinflammatory cytokines. (A) IL-1β, IL-6, TNF-α, and MCP-1 levels in primary mouse macrophage supernatants treated with vehicle DMSO (C), LA1 (20 μM), LPS (50 ng/ml), or LPS+LA1 for 8 hours (MCP-1) or 12 hours (IL-1β, IL-6, and TNF-α). Bars show mean ± SEM (n = 3) from 1 of at least 3 independent experiments. (B) IL-6 and TNF-α levels in supernatants of human macrophages treated with vehicle DMSO (C), LPS (50 ng/ml), or LPS+LA1 for 12 hours. Bars show mean ± SEM (n = 3) from 1 of at least 2 independent experiments. (C) IL-6 and TNF-α levels in supernatants of human macrophages treated with vehicle DMSO (C), R848, or R848+LA1 for 12 hours. Bars show mean ± SEM (n = 3) from 1 of 2 independent experiments. (D) IFN-β levels in the sera of WT or CD11b–/– mice treated with DMSO (C), LPS (100 ng/ml), or LPS+LA1 for 4 hours. Bars show mean ± SEM (n = 3). (E) IFN-β levels in supernatants of primary WT or CD11b–/– macrophages treated with DMSO (C), LA1 (20 μM), LPS (100 ng/ml), or LPS+LA1 for 12 hours. Bars show mean ± SEM (n = 3) from a representative experiment. *Not detectable. (F) Survival of C57BL/6 mice (n = 12) subjected to CLP and treated with vehicle (1% DMSO in saline) or LA1 (2 mg/kg/d). (G) IL-1β, IL-6, and TNF-α in plasma obtained 24 hours after C57BL/6 mice were subjected to either sham surgery (C) or CLP and treated with either vehicle (CLP+Veh) or LA1 (CLP+LA1). Bars show mean ± SEM. (H) Body weight loss (percentage of the initial weight) over 14 days in C57BL/6 mice (n = 20 per group) infected with WT H1N1 virus and treated daily with either vehicle (H1N1 + Vehicle) or LA1 (H1N1 + LA1). Weight loss in noninfected animals (control) and noninfected animals treated with LA1 (Control + LA1) is also shown. Each data point represents mean ± SEM (*P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001; A–E, 1-way ANOVA, Tukey’s test; F–H, Student’s t test).
binning of the data into high and low bins followed by categorical statistical analyses (3), in addition to the rank-order nonparametric \(t\) test of the quantitative data. Additionally, in such analyses, we have previously successfully used a value of 2.0 as a cutoff for high versus low, which corresponds to 2 SD above the mean of healthy donors. Applying this cutoff to the data in Figure 1, A and B, and using categorical analyses, we arrive at results similar to those of the nonparametric \(t\) test (Figure 1A, \(P = 0.037\) by \(\chi^2\); odds ratio = 2.01; Figure 1B, \(P = 0.0083\) by Fisher’s exact test, odds ratio = 3.49). These analyses, for the first time to our knowledge, suggest a direct link between reduced CD11b function as a result of these SNPs and the increased IFN-\(\alpha\) activity in SLE. Indeed, recent studies with macrophages from C\(D\)\(k\)\(b\)\(\alpha\) mice show significantly higher IFN-\(\beta\) production upon stimulation with the TLR4 agonist LPS in comparison with WT macrophages (14), further supporting the idea that functional deficit in CD11b is directly linked to increased IFN-\(\alpha\) activity in SLE.

**CD11b agonist LA1 suppresses proinflammatory cytokine secretion by TLR-activated leukocytes.** CD11b/CD18 ligation and clustering mediate proinflammatory signaling in leukocytes, including synergistic potentiation of signaling by other receptors (23–31). Conversely, they have also been shown to negatively regulate signaling by TLRs, TNF receptors, and cytokine receptors (14, 17, 23, 24, 32–34), which could suppress downstream IFN-\(\beta\) pathways. To test whether pharmacologic activation of CD11b could suppress TLR-dependent responses, and thus be a therapeutic strategy in SLE, we used our recently discovered small-molecule CD11b agonist LA1 as a tool (35, 36). LA1 binds to the CD11b/\(\alpha\) domain (CD11b/\(\alpha\)) in an allosteric pocket (Supplemental Figure 1), enhancing ligand binding. Molecular dynamics and isothermal calorimetry studies with CD11b/\(\alpha\) and fluid shear stress studies with cell surface–expressed full-length integrin CD11b/CD18 (Supplemental Methods; Supplemental Tables, 2 and 3; and Supplemental Figures 2–4) further established that LA1 binding promoted partial CD11b/\(\alpha\) activation, inducing an intermediate conformation in CD11b/\(\alpha\) (37) and no large conformational changes in the integrin heterodimer (38), defining this as its preferred binding mode. Given that the SLE-associated CD11b mutations are located outside of the ligand-binding \(\alpha\) domain (Supplemental Figure 2, A and B), it is unclear whether they would affect LA1 binding to the \(\alpha\) domain, and thus affect LA1’s activity. To assess, we generated cells stably expressing either full-length WT integrin CD11b (and its partner CD18) or the CD11b R77H mutant and used them in adhesion assay. The results show that both cell types displayed a similar level of LA1-mediated adhesion to immobilized Fg (Supplemental Figure 5), with similar EC\(_{50}\) values, suggesting that LA1 is able to activate the WT and the mutant integrin equally well and its binding is unaffected by the ITGAM SNP. The results also corroborate a recently published study, which also showed that the functional defects in CD11b R77H are rescued by integrin activation (13). Next, we treated primary murine macrophages and neutrophils with LPS in the absence or presence of LA1 and measured secretion of IL-1\(\beta\), IL-6, TNF-\(\alpha\), and monocyte chemoattractant protein-1 (MCP-1) in cellular supernatants. LA1 significantly suppressed levels of all 4 in macrophages (Figure 2A) and neutrophils (Supplemental Figure 6A) in a dose-dependent manner (Supplemental Figure 6B). Similarly, TNF-\(\alpha\)-stimulated IL-6 production was also significantly reduced (Supplemental Figure 6C). LA1 treatment also suppressed levels of IL-6 and TNF-\(\alpha\) in cell culture supernatants of human macrophages stimulated with the TLR agonist LPS (Figure 2B) or R848 (Figure 2C), further validating the role of LA1-mediated integrin activation as suppressor of proinflammatory signaling by heterologous receptors in leukocytes. As TLR activation also induces IFN-\(\alpha\) and downstream IFN-\(\beta\) signaling pathways (2), we tested the effects of TLR stimulation on IFN-\(\beta\) generation in vivo. LPS administration significantly upregulated IFN-\(\beta\) levels, with the level of increase significantly higher in the CD11b\(\alpha\) animals as compared with the WT animals (Figure 2D), suggesting that CD11b activation–dependent modulation of TLR-signaling extends to IFN-\(\beta\) pathways. Importantly, we found that CD11b activation with LA1 significantly suppressed this upregulation of IFN-\(\beta\) in WT mice, but not in the mice lacking CD11b (CD11b\(\alpha\); Figure 2D), suggesting that the in vivo effects of LA1 here are mediated via CD11b. In vitro, LPS treatment also resulted in significantly increased production of IFN-\(\beta\) by CD11b\(\alpha\) macrophages, as compared with the WT cells (Figure 2E), in accordance with the published studies (14). Furthermore, while LA1 significantly reduced the LPS-mediated upregulation in IFN-\(\beta\) levels in the cultured macrophages from WT animals, it failed to suppress IFN-\(\beta\) upregulation in the CD11b\(\alpha\) cells (Figure 2E). These data, in combination with our previously published results showing that LA1 (and other analogs) significantly reduces peritoneal infiltration of CD11b\(\alpha\) leukocytes in response to thioglycollate-induced peritonitis in WT but not in CD11b\(\alpha\) animals (35), strongly suggest that LA1 is primarily mediating its effects here via CD11b.

**LA1 treatment suppresses inflammatory cytokines in vivo and protects mice from severe sepsis.** To study the effects of CD11b activation with LA1 on TLR-stimulated cytokine generation in vivo, we subjected WT C57BL/6 mice to cecal ligation and puncture (CLP) (39), a model of polymicrobial sepsis, and treated them with LA1 or vehicle control. Among the control, vehicle-treated mice, 75% died within 120 hours (Figure 2F), and more than 90% (11/12) of them died within 240 hours. Surprisingly, more than 60% of LA1-treated mice were alive for 120 hours, and 50% of them survived more than 240 hours. Vehicle-treated mice showed high serum levels of the 3 hallmark cytokines (IL-1\(\beta\), IL-6, and TNF-\(\alpha\)) that play a significant role in cytokine storm–mediated septic shock (40), whereas LA1-treated animals showed a significant reduction in these cytokines (Figure 2G). Significantly, serum bacterial count 24 hours after CLP showed few remaining bacteria in LA1-treated mice as compared with high levels in the vehicle-treated group (Supplemental Figure 7), which suggests that LA1 treatment also suppresses bacterial load and correlates with previous data that LA1 enhances phagocytosis (35). Viral infections also lead to exacerbated leukocyte activation and cytokine storm, which prospectively predicts morbidity and mortality in patients (41). To determine whether LA1-mediated integrin activation in leukocytes would similarly suppress excessive immune response upon severe viral infection, we infected C57BL/6 mice via intranasal administration of influenza A virus (42) and monitored the transient loss in body weight, which is a primary man-
manifestation of infection in these animals (41, 43). While the vehicle-treated animals exhibited significant weight loss within 3 days after viral infection (Figure 2H), reaching a maximum weight loss around day 6, before regaining it, LA1-treated animals did not show any loss of body weight after infection. These data suggest that LA1 blunts cytokine storm in vivo, improving survival.

LA1-mediated CD11b activation suppresses proinflammatory TLR and IFN-I pathways via MyD88/NF-κB and AKT/FOXO3/IRF3/7 axes. To identify TLR signaling components that are regulated by LA1-mediated CD11b activation, we performed genome-wide mRNA expression profiling of human macrophages stimulated with LPS (50 ng/ml) in the absence or presence of LA1 (Figure 3A). Our analysis showed that 416 genes were significantly affected by LA1 treatment (Supplemental Table 4) and that LA1 normalized a majority of the LPS-regulated genes (Figure 3B). In addition to the known genes downstream of TLR engagement (MYD88, IL6, IL1A, CXCL9), pathway analysis showed that the genes in the TLR-associated canonical antiviral response (P < 1.3 × 10⁻¹², overlap 13/52) and IFN-I pathways (P < 2.5 × 10⁻⁹, overlap 8/24) were highly enriched (Supplemental Figure 8). This includes CXCL9,
CXCL10, IDO1, CCL5, IFIT1, IRF7, and ISG15. We used a multiplex mRNA counting method to measure the expression of a number of these genes in macrophages treated with LPS or LPS plus LA1 over time (Figure 3C), which validated the expression level changes induced by LA1 in the genes in the TLR and IFN-I pathways. Curiously, the microarray data also showed a concomitant increase in FOXO3 transcription, which was recently shown to suppress IFN-I pathway by limiting IFN regulatory factor 7 (IRF7) autoregulation (44). Additional known direct targets of FOXO3, including IFIH1, DDX58, RSAD2, MX2, and CMPK2, were also reduced by LA1. This suggests that LA1-mediated CD11b activation suppresses TLR signaling via MyD88-dependent NF-κB pathway and downstream IFN-I signaling via the FOXO3/IRF pathway.

Western blot (WB) analysis confirmed that LA1 suppressed the LPS-stimulated NF-κB pathway (reduced levels of phosphorylated IKKα/β and p65) (Figure 3D), which is also in line with previous studies showing that CD11b negatively regulates TLR-dependent NF-κB pathway, by increasing degradation of MyD88, and that deficiency of CD11b exacerbates this TLR-dependent proinflammatory signaling by failing to reduce the MyD88 levels (14). Furthermore, active CD11b down-regulates MyD88 protein levels by inducing Src and CBL phosphorylation (14), and the WB analyses show increased pSrc and pCBL by LA1 (Figure 3E), suggesting that this CD11b agonist may use a similar mechanism. Densitometric quantification of WBs for the level of MyD88, the most upstream TLR signaling event, confirmed that LPS treatment increased MyD88 levels that were significantly reduced by LA1-mediated CD11b activation (Supplemental Figure 9), suggesting it to be a key node. Transcriptional induction of IFN-I is mediated by transcription factors IRF3 and IRF7 that require phosphorylation for nuclear entry and activity (45). Under basal conditions, nuclear FOXO3 acts as a negative regulator of IFN-I pathway via suppression of IRF7 (44). Upon TLR stimulation, FOXO3 is excluded from the nucleus and degraded, via its AKT-mediated phosphorylation, thereby derepressing IRF7 and IRF7-dependent genes, including the IFN-I pathway. As TLR stimulation in macrophages induces MyD88-dependent AKT phosphorylation (46), we hypothesized that LA1-mediated CD11b activation mechanistically suppresses IFN-I pathway by reducing AKT phosphorylation, thereby reducing FOXO3 phosphorylation and, thus, enhancing nuclear FOXO3. Indeed, WB analysis showed that CD11b activation with LA1 suppressed the LPS-stimulated upregulation of pAKT and pFOXO3 in macrophages (Figure 3F). Reduced pFOXO3 levels also correlated with reduced phosphorylation of IRF3 and IRF7, suggesting that CD11b activation blunts IFN-I signaling via the AKT/FOXO3/IRF3/7 axis. To further confirm that CD11b regulates these pathways, we measured the levels of these proteins in LPS-stimulated primary macrophages from WT and CD11b–/– animals using WB (Figure 3G). Whereas the WT cells showed an expected time-dependent increase in the levels of MyD88, pAKT, pFOXO3, pIRF3, and pIRF7, the CD11b–/– cells showed a larger increase in the levels of these proteins at baseline and upon LPS stimulation, suggesting that the AKT/FOXO3/IRF3/7 signaling pathway is exacerbated in the CD11b-deficient macrophages in a manner similar to the previously reported exacerbation of the MyD88/IKK/NF-κB pathway in these cells (14).
Cells stimulated with TLR3 agonist pI:C also showed that LA1 reduced upregulation of pAKT and pFOXO3 in macrophages (Supplemental Figure 10), and reduced pIRF3. Given that TLR3 is MyD88-independent and is known to regulate the IFN-I pathway via IRF3 (47, 48), this suggests that LA1 suppresses TLR-dependent signaling via IRF3 in the MyD88-independent arm. Cells stimulated with other TLR agonists (R848 or CpG) also showed similar results (not shown), suggesting that this might be a general mechanism of downmodulating TLR signaling by CD11b. Since the downstream transcriptional activity of FOXP3 is mediated via its nuclear retention (49) and because LA1-mediated CD11b activation affects FOXP3 phosphorylation, we predicted that LA1 treatment would affect FOXP3’s subcellular localization. Confocal microscopy–based imaging showed that FOXP3 was mostly inside the nucleus under basal conditions in RAW macrophages (Figure 4A). As a control, we stained for NF-κB, which expectedly showed a mostly cytoplasmic localization. Upon stimulation with LPS, FOXP3 expectedly translocated out of the nucleus and was primarily in the cytoplasm (49), whereas NF-κB showed the opposite pattern. Strikingly, cotreatment with LA1 suppressed the translocation of FOXP3 from the nucleus into the cytoplasm, suggesting that LA1-mediated CD11b activation reduces IFN-I signaling pathway via protection of FOXP3 levels in the nucleus. In contrast, NF-κB showed the exact opposite results, suggesting that partial activation of CD11b is sufficient to limit the TLR-dependent IFN-I pathway in cells via an AKT/FOXO3/IRF3/7 circuit (Figure 4B).

LA1 suppresses inflammatory injury and end-organ pathology in lupus-prone mice. To test the efficacy of LA1 in a lupus model, we used the MRL/Lpr mice that develop IFN-I–dependent multiorgan lupus, which is similar to human lupus, including skin, vascular, and renal inflammation (50), and involves a leukocyte-mediated inflammatory process (51). There was no evidence of overt toxicity in these animals, with no differences in total body weight throughout treatment (not shown). Vehicle-treated MRL/Lpr mice (but not the haplotype-matched MRL/Mpj) lost whiskers and developed skin lesions and alopecia, identifiable within the intrascapular area and on the ears and face (muzzle) by 14 weeks of age (refs. 52, 53, and Figure 5, A–C). Increased leukocyte infiltration was identified in the dermis of these patches, along with enhanced epidermal thickness (Figure 5B). Treatment with LA1 significantly reduced alopecia, leukocyte infiltration, and epidermal thickness (Figure 5, A–C). In addition, vehicle-treated MRL/Lpr mice developed significant features consistent with lupus glomerulonephritis, including increased urine albumin/creatinine ratio and IgG renal immune complex deposition, which were significantly reduced with treatment with LA1 (Figure 5, D and E). LA1-treated animals also showed decreased glomerular damage (Figure 5E and Supplemental Figure 11). Consistent with this amelioration in lupus phenotype, LA1 treatment resulted in significant amelioration in levels of serum anti-dsDNA antibodies and total IgG, various inflammatory cytokines and chemokines, and spleen weight, in comparison with vehicle-treated mice (Figure 5, F–H), which correlates with reduced inflammation in LA1-treated mice. While the differences in numbers of splenic double-negative T cells (not shown), activated T and B cells, or plasma cells, DCs, macrophages, and neutrophils did not achieve statistical significance between LA1-treated and vehicle-treated mice (Supplemental Figures 12 and 13), the trends showed decreases in splenic macrophages, activated B cells, and plasma cells, which may explain the differences in autoantibody synthesis, as previously described (54, 55). To investigate whether the efficacy of LA1 in vivo was associated with modulation of the IFN-I pathway, we isolated mRNA from MRL/Lpr splenocytes and quantified relative expression of several candidate genes. Indeed, cells from LA1-treated animals showed significantly lower expression of MCP-1, IIf7, IgL5, and Idha (Figure 5I) and, while not statistically significant, a trend for increased Foxo3 mRNA levels (Figure 5J). Consistent with the hypothesis that increased phosphorylation of FOXO3 drives IFN-I signaling, bone marrow monocytes from LA1-treated animals showed significantly reduced levels of pFOXO3 (Figure 5K). These data suggest that LA1-mediated CD11b activation mitigates lupus phenotype and end-organ damage in association with significant suppression of the AKT/FOXO3/IRF7 pathway and modulation of IFN-I activity. Since aberrant leukocyte activation and enhanced IFN-I activity contribute to the development of endothelial dysfunction in SLE (56), we asked whether the endothelial dysfunction inherent to MRL/Lpr mice might also be reduced following LA1 treatment. Aortic rings were isolated from 19-week-old MRL/Lpr mice after 10-week treatment with LA1 or vehicle, and acetylcholine-dependent relaxation triggered after phenylephrine precontraction was determined as described in Methods. Indeed, treatment with LA1 significantly improved endothelium-dependent vasorelaxation, indicating a vasoprotective effect on the lupus vasculature (Figure 6A). Examination of vascular reendothelialization in balloon-injured Fisher male rats also showed significant reendothelialization at the site of balloon injury with LA1 treatment (Figure 6, B–D, and Supplemental Methods), suggesting that LA1 may accelerate the process of endothelial regeneration (57). Increased neutrophil adhesion via LA1 to HUVECs did not show any damage to the HUVEC monolayer (Supplemental Figure 14, A–C, and Supplemental Methods). Together, these results show that LA1 treatment does not harm the endothelium, similarly to previous studies using knock-in animals that express constitutively active mutants of integrins CD11a/CD18 (58, 59) or αβ, (60) that also did not report any signs of vascular injury in the various experimental models, suggesting that integrin activation, per se, does not harm the vasculature or have any harmful consequences for animals under normal conditions.

Cells homozygous for CD11b SNPs show basal level increase in IFNB and IRF7 expression and increased FOXP3 nuclear exclusion that is suppressed by LA1 treatment. To examine the molecular link between ITGAM SNPs and IFNB in human cells, we used quantitative reverse transcriptase PCR to measure the expression level of IFNB and IRF7 in peripheral blood mononuclear cells (PBMCs) from donors carrying either major or minor ITGAM alleles at the rs1143679 or rs1143683 loci and found that the cells carrying a minor allele consistently showed increased expression of both transcripts (Figure 7A). Given that their expression is modulated by nuclear levels of FOXP3, we also quantified the levels of FOXP3 in the nucleus versus cytoplasm and found that, under basal conditions, the cells carrying a minor allele showed significantly reduced levels of FOXP3 in the nucleus, as compared with the cells carrying the common allele (Figure 7B). Recent reports show that
**A** MRL/Lpr Vehicle LA1 MRL/Mpj control

**B** MRL/Lpr Vehicle LA1 MRL/Mpj control

**C**

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<tbody>
<tr>
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<td>Veh</td>
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**E** MRL/Lpr

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<th>LA1</th>
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**F**

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<th>Anti-dsDNA IgG (KU/ml)</th>
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<td>LA1</td>
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**G**

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<th>IP-10 (pg/ml)</th>
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<td>LA1</td>
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**H**

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**I**

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**J**

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<th>Foxo3 expression (fold change)</th>
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<tr>
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<td>Veh</td>
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</table>

**K**

|        | p-FOXO3a | FOXO3a |

* denotes statistical significance.
TLR pathway is basally activated in SLE (61), and the data presented here suggest that ITGAM SNPs likely reduce a CD11b-dependent “tonic” suppression of TLR signaling in SLE, due to a failure of the mutant CD11b in maintaining levels of nuclear FOXC3, resulting in reduced suppression of the IRF7 (44) and increased IFN-I. Future research will further clarify this connection. We also found that LPS treatment of cells resulted in rapid movement of FOXO3 across various ethnic groups. However, how these polymorphisms link between an integrin mutation and a pathogenic circulating factor that may predispose to SLE and promote immune dysregulation characteristic of this disease. We found that LA1 enhanced CD11b-dependent ligand binding and cell adhesion by inducing an intermediate affinity conformation in the ligand-binding CD11bA domain. Together with our previously published studies that showed that LA1 induces increased binding of mAb24 (38, 64, 65) (which detects an activation-dependent neoepitope in the βA domain), but not KIM127 (66) (which detects an integrin extension-dependent neoepitope), this further suggests that LA1 binding does not induce an extended integrin conformation (38). Such LA1-mediated integrin activation significantly blunted generation of TLR-stimulated proinflammatory cytokines and chemokines in leukocytes, suppressing TLR-dependent intracellular signaling pathways in vitro and in vivo, and further explaining the observed strong antiinflammatory effects of LA1 in vivo. Importantly, in addition to suppressing the MyD88-dependent NF-κB pathways in leukocytes, LA1 also dampened the IFN-I signaling, indicating an important mechanism by which it may modulate immune dysregulation in SLE. Detailed molecular analysis showed that LA1 reduced phosphorylation of AKT and its substrate FOXO3α, thereby increasing the levels of FOXO3α in the nucleus that suppresses IRF3/7-dependent gene expression (44). IRF3 and IRF7 are members of the IFN regulatory transcription factor family, a master regulator of the IFN-I-inducible pathway and a key player in innate immunity (67). As such, our findings directly link integrin biology with crucial pathways in IFN-I responses that may predispose to SLE and promote immune dysregulation characteristic of this disease.

Given that the TLR and IFN-I pathways play a significant role in the pathogenesis of SLE, the significance of these findings was tested in murine lupus. LA1 reduced IFN-I signaling and protected animals from end-organ injury and vascular dysfunction, by similarly modulating the CD11b/FOXO3 axis in leukocytes. Importantly, the relevance of these findings is also translated into humans in that the subjects carrying missense ITGAM SNPs were found to have dysregulation of FOXO3 in leukocytes and higher IFN-I activity. This, for the first time to our knowledge, suggests a direct link between an integrin mutation and a pathogenic circulating factor. As the binding site for LA1 on CD11b does not overlap with the SLE-associated mutations in CD11b, cell-based studies also showed that LA1 is able to induce cell adhesion via the WT and the mutant R77H integrin equally well. Indeed, LA1-mediated CD11b be fully characterized. Integrin CD11b combines with the β2 chain (ITGB2, CD18) to form Mac-1, an integrin heterodimer with crucial effects on myeloid cell adherence to stimulated endothelium. In addition to its role in inflammation-induced leukocyte tissue recruitment, prior studies have also highlighted a protective role of CD11b (reviewed in ref. 22), including maintenance of autoreactive B cell tolerance (15), suppression of DC maturation and function, and restriction of Th17 differentiation (62). CD11b (along with ITAM receptors) has also been shown to negatively regulate TLR4-triggered inflammatory responses (14, 23, 63). However, it has remained unclear whether aberrant function of CD11b in individuals with ITGAM polymorphisms can promote IFN-I responses and whether modulation of CD11b in SLE, by restoring integrin activation, can modify IFN-I signaling and modulate autoimmunity. Here, we have uncovered a direct link between an integrin mutation and a pathogenic circulating factor that may predispose to SLE and promote immune dysregulation characteristic of this disease. We found that LA1 enhanced CD11b-dependent ligand binding and cell adhesion by inducing an intermediate affinity conformation in the ligand-binding CD11bA domain. Together with our previously published studies that showed that LA1 induces increased binding of mAb24 (38, 64, 65) (which detects an activation-dependent neoepitope in the βA domain), but not KIM127 (66) (which detects an integrin extension-dependent neoepitope), this further suggests that LA1 binding does not induce an extended integrin conformation (38). Such LA1-mediated integrin activation significantly blunted generation of TLR-stimulated proinflammatory cytokines and chemokines in leukocytes, suppressing TLR-dependent intracellular signaling pathways in vitro and in vivo, and further explaining the observed strong antiinflammatory effects of LA1 in vivo. Importantly, in addition to suppressing the MyD88-dependent NF-κB pathways in leukocytes, LA1 also dampened the IFN-I signaling, indicating an important mechanism by which it may modulate immune dysregulation in SLE. Detailed molecular analysis showed that LA1 reduced phosphorylation of AKT and its substrate FOXO3α, thereby increasing the levels of FOXO3α in the nucleus that suppresses IRF3/7-dependent gene expression (44). IRF3 and IRF7 are members of the IFN regulatory transcription factor family, a master regulator of the IFN-I-inducible pathway and a key player in innate immunity (67). As such, our findings directly link integrin biology with crucial pathways in IFN-I responses that may predispose to SLE and promote immune dysregulation characteristic of this disease.

Discussion

The association between specific ITGAM SNPs and SLE risk is one of the strongest and most specific lupus genetic risk factors across various ethnic groups. However, how these polymorphisms in ITGAM that result in decreased function of CD11b functionally predispose to lupus and its associated organ damage remained to
activation in ex vivo donor leukocytes suppressed TLR-stimulated cytokines and IFN-I levels, by correcting the signaling change in the AKT/FOXO3/IRF3/7 axis. Published findings and these observations suggest that CD11b plays an important role in SLE and that CD11b activation should be explored as a potential novel therapeutic target in this autoimmune disease, particularly as a personalized approach in patients identified as carriers of specific genetic polymorphisms.

Methods

Full experimental details can be found in the Supplemental Methods. Human samples and measurement of serum IFN-I activity. We analyzed 171 SLE cases from multiple study centers consisting of patients of self-reported European ancestry that had serum available for IFN-I analysis (Supplemental Table 1). Informed consent was obtained from all patients in both cohorts included in this study, and the study was approved by the institutional review boards at the respective institutions. Patients were genotyped at coding change SNPs in ITGAM (rs1143678, rs1143679, and rs1143683) using custom-designed Applied Biosystems TaqMan primers and probes on an Applied Biosystems 7900HT PCR machine with greater than 98% genotyping success. Additional genotyped human PBMCs from donors carrying ITGAM α according to the company’s protocol. Levels of IL-1, IL-6, and TNF-α were measured using murine ELISA kits (R&D Systems) according to the company’s protocol. Primary macrophage assays. Human PBMCs were isolated from fresh blood collected from healthy volunteers under an IRB-approved protocol and using RosetteSep (human monocyte Enrichment Cocktail, catalog 15068, Stemcell Technologies) followed by Ficoll-Hypaque density gradient centrifugation according to published protocols (73). Cells adherent to the tissue culture flask were cultured for 5–7 days at 37°C in 5% CO₂, to promote their full differentiation into monocyte-derived macrophages (74, 75).

Figure 6. LA1 improves vasorelaxation in mice. (A) Quantification of acetylcholine-dependent (Ach-dependent) relaxation after phenylephrine (PE) precontraction in aortic rings from 19-week-old MRL/Lpr mice after 10-week treatment with LA1 or vehicle. Control data are from MRL/Mpj. Data are mean ± SEM (n = 7 LA1-treated, n = 8 vehicle-treated). (*P < 0.05, **P < 0.01, ***P < 0.001, comparing LA1 vs. controls.) Curves were first analyzed using an asymmetric (5 parameters) logistic equation; significance was determined by 2-way ANOVA. (B) Representative photomicrographs of balloon-injured arteries from rats treated with vehicle or LA1. Images of arteries stained with Evans blue vital staining in situ before sacrifice are shown at the top. Photomicrograph of cross section of arteries stained with anti-vWF antibody to label the endothelium (arrow) is presented below. Scale bars: 50 μm. (C) Quantification of the amount of Evans blue extracted from the balloon-injured arteries from rats treated with vehicle or LA1 (as in B). Data are mean ± SEM (n = 7 vehicle-treated, n = 8 LA1-treated). (D) Quantification of the extent of reendothelialization 7 days after balloon injury from the balloon-injured arteries from rats treated with vehicle or LA1 after staining with anti-vWF antibody (as in B). Data shown are mean ± SEM (n = 7 vehicle-treated, n = 8 LA1-treated) (*P < 0.05, Student’s t test).
monocytes as described in the literature (77). For 5–7 days, macrophages were generated from bone marrow–derived cells plated in 6-well plates in RPMI containing 10% FBS and were cultured for 4 hours as previously described (35). Cells were washed with ice-cold PBS and then lysed in TRizol (Thermo Fisher Scientific) for total RNA isolation. RNA sample quality was analyzed using a 2100 Bioanalyzer (Agilent). Whole-genome transcriptional profiling was performed using Illumina HT-12 human gene expression arrays and analyzed using GeneSpring software version 13. Only significant genes (P ≤ 0.05 and fold change ≥ 2) were considered for pathway analysis. We deposited all microarray data in the Gene Expression Omnibus database in the series GSE76802. Expression levels of selected genes in human macrophages stimulated with LPS (50 ng/ml) for various amounts of time (0, 0.5, 2, 4, 8, and 24 hours) in the absence or presence of LA1 (20 μM) were measured using an inflammation panel set and the nCounter Digital Analyzer (Nanostring Technologies) as previously described (76).

Isolation of murine primary neutrophils and macrophages. Thiglycolate-elicited neutrophils and macrophages were isolated from B6 WT mice or B6 CD11b−/− mice (Jackson Laboratory), as described previously (35). Neutrophils were collected 4 hours after thioglycolate injection followed by 1-hour adherence purification before collection of nonadherent cells. The macrophages were collected 4 days after thioglycolate injection and adherence-purified for 1 hour, followed by a wash with PBS to remove nonadherent cells. For the bone marrow–derived macrophages, 2 × 10^6 cells per well of harvested cells were plated in 6-well plates in RPMI containing 10% FBS and were cultured for 5–7 days. Macrophages were generated from bone marrow–derived monocytes as described in the literature (77).

Statistical analysis. For microarray analyses, human macrophages (2 × 10^6) were cultured in 6-well tissue culture plates and were treated with LPS (50 ng/ml) in the absence or presence of LA1 (20 μM) for 4 hours as previously described (35). Cells were washed with ice-cold PBS and then lysed in TRizol (Thermo Fisher Scientific) for total RNA isolation. RNA sample quality was analyzed using a 2100 Bioanalyzer (Agilent). Whole-genome transcriptional profiling was performed using Illumina HT-12 human gene expression arrays and analyzed using GeneSpring software version 13. Only significant genes (P ≤ 0.05 and fold change ≥ 2) were considered for pathway analysis. We deposited all microarray data in the Gene Expression Omnibus database in the series GSE76802. Expression levels of selected genes in human macrophages stimulated with LPS (50 ng/ml) for various amounts of time (0, 0.5, 2, 4, 8, and 24 hours) in the absence or presence of LA1 (20 μM) were measured using an inflammation panel set and the nCounter Digital Analyzer (Nanostring Technologies) as previously described (76).

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Statistics. To calculate statistical significance, a 2-tailed student’s t test was used, unless otherwise specified. The differences in the cytokine concentrations were statistically analyzed using the Mann-Whitney test, and plotted with the GraphPad Prism software package. For multiple comparisons, 1-way ANOVA with Tukey’s test was used. A value of P less than 0.05 was considered statistically significant. For endothelium-dependent vasorelaxation, curves were first analyzed using an asymmetric (5 parameters) logistic equation, and significance of each individual data point was determined by 2-way ANOVA. Unless otherwise specified, results are represented as mean ± SEM.

Study approval. The studies on human samples were reviewed and approved by the Mayo Clinic IRB in Rochester, Minnesota, USA. All subjects provided appropriate prior informed consent to participate in the study. The respective studies on C57BL/6 WT and CD11b−/− animals were reviewed and approved by the IACUC at Rush University Medical Center or the IACUC at University of Miami School of Medicine, or in accordance with Portuguese guidelines and regulations after approval by the local ethics committee at Instituto Gulbenkian de Ciência, Oeiras, and the national regulating agency (Direcção Geral de Alimentação e Veterinária). All experiments using MRL/Lpr and MRL/Mpj mice were approved by the Animal Ethics Committee (ACUC) at the National Institute of Arthritis and Musculoskeletal and Skin Diseases/NIH, Bethesda, Maryland, USA.

Author contributions. MHF, SQK, and WZ designed and performed in vitro and in vivo experiments. EM and CCR performed animal experiments. KZ, VK, ARA, RD, DS, VH, SJK, PB, A.M. Santander, MTK, MMA, XL, SD, KHK, HWL, JSR, JR, and JFC contributed to in vitro experiments. JMD, A.M. Schnaith, YGP, MJ, EMB, JSR, BS, SKN, and TBN contributed to human sample analyses. TZ and RP contributed to molecular dynamics simulations and data analysis. BI analyzed the microarray data and helped build pathways. LFM and AZ contributed to CLP and virus infection experiments, respectively. MP, JIC, and CCR performed immunophenotyping of lupus mice and protein FOXO3 assessments in lupus. VH performed scoring of lupus murine skin. DJC performed scoring of lupus murine kidneys. PSY performed HPLC analyses for mouse creatinine. WLT performed cytokine work in lupus mice. VG, MJK, and RIVP codesigned and cowrote the paper.

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