Bim suppresses the development of SLE by limiting myeloid inflammatory responses

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The introduction of Bcl-2 family is considered the guardian of the mitochondrial apoptotic pathway. We demonstrate that Bim acts as a molecular rheostat by controlling macrophage function not only in lymphoid organs but also in end organs, thereby preventing the break in tolerance. Mice lacking Bim in myeloid cells (LysMcreBimfl/fl) develop a systemic lupus erythematosus (SLE)-like disease that mirrors aged Bim−/− mice, including loss of marginal zone macrophages, splenomegaly, lymphadenopathy, autoantibodies (including anti-DNA IgG), and a type I interferon signature. LysMcreBimfl/fl mice exhibit increased mortality attributed to glomerulonephritis (GN). Moreover, the toll-like receptor signaling adaptor protein TRIF (TIR-domain–containing adapter–inducing interferon-β) is essential for GN, but not systemic autoimmunity in LysMcreBimfl/fl mice. Bim-deleted kidney macrophages exhibit a novel transcriptional lupus signature that is conserved within the gene expression profiles from whole kidney biopsies of patients with SLE. Collectively, these data suggest that the Bim may be a novel therapeutic target in the treatment of SLE.

INTRODUCTION

Systemic lupus erythematosus (SLE) is a multifactorial, multigenetic autoimmune disease of unknown etiology that is characterized by the presence of autoantibodies and severe end-organ damage (Shirai and Hirose, 2006). The origin of the break in tolerance leading to the development of systemic autoimmunity and production of autoantibodies is unknown. However, studies have suggested that a failure to process apoptotic body antigens by marginal zone macrophages (MZMs) may be required for the activation of lymphocytes in SLE-like disease (McGaha and Karlsson, 2016). Monocytes and macrophages are mononuclear phagocytes that are crucial for maintaining homeostasis (Ginhoux and Jung, 2014). Macrophages are highly plastic and are therefore credited with essential roles in inflammation as well as tissue injury and repair (Ginhoux and Jung, 2014). Recent studies have shown that, similar to peripheral blood monocytes, renal macrophages from SLE patients are increased in number and exhibit elevated expression of activation markers (Katsiari et al., 2010). Further, the numbers of glomerular macrophages, tubular luminal macrophages and/or CD16+ macrophages in the kidney correlate with clinical activity and outcome in patients with SLE (Hill et al., 2001). Studies in murine models also support the importance of monocytes and macrophages in the pathogenesis of SLE-like disease (Hutcheson et al., 2008; Katsiari et al., 2010). Collectively, these data suggest a pivotal role for monocytes and macrophages in the pathogenesis of SLE and SLE-like disease, but the factors that control their state of activation and function are unknown.

Apoptosis or programmed cell death is necessary for immune cell development and homeostasis. Cells undergo apoptosis through two distinct pathways: an extrinsic path-
way of apoptosis and an intrinsic pathway of apoptosis. Specifically, the intrinsic pathway is regulated by the Bcl-2 (B cell lymphoma 2) protein family and proceeds through a mitochondrial-dependent mechanism. Antiapoptotic proteins of the Bcl-2 protein family include Bcl-2, Bcl-xL, Bcl-w, Mcl-1, and A1. Proapoptotic proteins of the Bcl-2 protein family consist of two types: those with multiple Bcl-2 homology (BH) domains, including Bak, Bax, Bok, and Bcl-x5 and those containing only a single BH3 domain, including Bim, Bad, Bid, Noxa, and Puma. Studies using BH3 peptides reveal that Bim, Bid, and Puma may function as direct activators of apoptosis, whereas Bad and Noxa exist as indirect activators of cell death (Billard, 2013). However, only mice deficient in Bim develop spontaneous systemic autoimmunity (Bouillet et al., 1999). Given the role of Bim as a mediator of cell death and the lymphocyte-centric hypothesis of SLE development, significant attention has understandably been paid to the role that Bim plays in eliminating self-reactive lymphocytes. However, Bim deficiency also impacts innate immune cell populations (Hutcheson et al., 2008). Little is known about the role of Bim on innate immune cells or their relative contribution to systemic autoimmune. In this study, we demonstrate that myeloid cells are central initiators of SLE-like disease in Bim−/− mice and potentially dispute the conventional dogma that the central role of Bim in autoimmune disease is to prevent the escape of autoreactive lymphocytes from apoptosis. Novel strategies that target Bim may be useful for the treatment of systemic autoimmunity.

RESULTS

Mice deficient for Bim in macrophages develop SLE-like disease

We and others have reported that Bim−/− mice develop systemic autoimmunity and end-stage glomerulonephritis (GN; Bouillet et al., 1999; Hutcheson et al., 2008). To determine whether Bim might prevent systemic autoimmunity via its function in myeloid cells, we generated mice with conditional deletion of Bim in the myeloid cell compartment on a mixed background (LysMCreBimfl/fl) and compared them to conventional deletion of Bim in the myeloid cell compartment on the C57BL/6 genetic background (Fig. S1 A and Table S1). 8-mo-old female LysMCreBimfl/fl mice showed disrupted splenic architecture (Fig. 1 A) and developed severe GN (Fig. 1, B–D). Mice deficient for Bim in macrophages (LysMCreBimfl/fl) mice showed enhanced splenic topography as compared to age- and sex-matched control mice (LysM−/−Bimfl/fl, LysM−/−Bim−/−, CD19CreBimfl/fl, and CD4CreBimfl/fl). At 6 mo of age, female LysMCreBimfl/fl mice showed disrupted splenic architecture (Fig. 1 A) and developed severe GN (Fig. 1, B–D). LysMCreBimfl/fl mice also showed reduced splenic architecture (Fig. 1 A) and lower kidney scores than control mice (Fig. 1 C). In contrast, mice deficient for Bim in B or T cells did not exhibit any features of systemic autoimmunity, GN (Fig. 1, A–D), or enhanced mortality (Fig. 1 E).

Mice on a 129 background have a predisposition to lupus disease (Obata et al., 1979); therefore, we backcrossed LysMCreBimfl/fl mice onto a C57BL/6 background. Genomic screening of 150 single-nucleotide polymorphism (SNP) markers confirmed that LysMCreBimfl/fl mice are 98% on the C57BL/6 genetic background (Fig. S1 A and Table S1). 8-mo-old female LysMCreBimfl/fl mice exhibited severe splenomegaly and lymphadenopathy compared with controls (Fig. 2, A–E). Enlargement of the spleen was associated with abnormal splenic architecture marked by distention of red pulp area and lymphoid hyperplasia in white pulp area (Fig. 2 C). There was a substantial increase in antinuclear autoantibody (ANA) titers in LysMCreBimfl/fl mice (Fig. 2 F), including anti–single-stranded DNA (anti-ssDNA) and anti–double-stranded DNA (anti-dsDNA) IgG ANA. LysMCreBimfl/fl mice also showed increased IgM ANA directed against ssDNA, dsDNA, histones, and nucleosomes (Fig. 2 F). Additionally, Ig isotypes, in particular the pathogenic IgG2a and IgG2b subclasses, were elevated in LysMCreBimfl/fl mice (Fig. 2 G). Moreover, the serum levels of IL-12p70, IL-17, IFN-β, and IFN-γ were higher in LysMCreBimfl/fl mice (Fig. 2, H–J). LysMCreBimfl/fl mice also had evidence of IgG- and IgM-immune complex deposition in kidneys (Fig. 2 K), increased proteinuria (Fig. 2 L), and severe GN (Fig. 2, M and N). Further, deleting Bim did not affect the balance of the Bcl-2 family members in splenic macrophages (Fig. S1, B and C). Although the expression levels of Bcl-2 and Bcl-xL were lower in 3- and 8-mo-old LysMCreBimfl/fl kidney macrophages, the other Bcl-2 family members, including Bcl-w, Mcl-1, Bax, and Bak, were unaffected (Fig. S1, D and E).

LysMCreBimfl/fl mice showed altered splenic cellularity

We next examined the topographical localization of macrophages in spleen of LysMCreBimfl/fl mice. It has been well established that Siglec-1 (CD169; Sialoadhesin) and DC-SIGN (CD209b) delineate marginal metallophilic macrophages (MMs) and MZMs, respectively (McGaha and Karlsson, 2016). MZMs were markedly reduced in LysMCreBimfl/fl mice as early as 3 mo of age, and this loss dramatically progressed with age (Fig. 3 A). In contrast, MMTM topography were maintained in the spleens of young LysMCreBimfl/fl mice, although there was a slight decrease as LysMCreBimfl/fl aged (Fig. 3 A). We then used multiparameter flow cytometry to quantify the numbers of MZMs and MMs in spleen. MZMs and MMs were gated via the exclusion of doublets, dead cells, red-pulp macrophages (RPMs), lineage− and CD19+ /B220+ cells, and CD209b+ and CD169+ cells, respectively (Fig. S2, A and B). The expression of CD11b further divided MZMs and MMs into two subtypes: the CD11b+ and CD11b− populations. MZMs were larger in size as measured by diameter (CD11b+ MZMs: 29.03 ± 1.58 µm; CD11b− MZMs: 25.87 ± 1.48 µm) than MMs (CD11b+ MMs: 25.00 ± 1.22 µm; CD11b− MMs: 21.61 ± 0.42 µm; Fig. S2, C and D; CD11b−/− MZMs vs. CD11b−/− MMs, P < 0.05). The number of CD11b+ MZMs was significantly lower in LysMCreBimfl/fl spleens than in control mice at 3 mo of age (Fig. 3 B). The number of CD11b+ MZMs and CD11b− MZMs further decreased in LysMCreBimfl/fl spleens at 8 mo (Fig. 3, B and C). In contrast, there was no significant difference in the number of CD11b+ MMs between control and LysMCreBimfl/fl mice (Fig. 3 D). However, CD11b− MMs numbers increased in 8-mo-old LysMCreBimfl/fl mice (Fig. 3 E). LysMCreBimfl/fl MZMs and MMs also displayed enhanced costimulatory signals, including CD80 and CD86 as compared...
with controls (Fig. S2, E–H). Because marginal zone B (MZB) cells maintain close contact with MZMs under steady-state condition, we observed a thinner layer of CD1d+ MZB cells in the splenic MZ region of LysMCreBimfl/fl mice (Fig. S2, I and J). We then performed RNA sequencing (RNA-seq) on the CD11b+ and CD11b− populations of MZMs in order to assess transcriptional changes caused by Bim deletion. The presence or absence of CD11b elicited more variation in gene expression than the deletion of Bim (Fig. 3 F). CD11b+ MZMs and CD11b− MZMs exhibited 20 and 64 differentially expressed genes, respectively, when Bim was deleted, including Lyz2, Hjurp, and Ms4a6d (Fig. 3, G and H).
Figure 2. **Loss of Bim in myeloid cells, including macrophages, leads to a break in tolerance and the development of SLE-like disease.** 8-mo-old female CTRL and LysM$^{Cre}$Bim$^{fl/fl}$ (C57BL/6) mice were examined for systemic autoimmune disease phenotypes. (A) Spleen weights for CTRL ($n = 51$) and LysM$^{Cre}$Bim$^{fl/fl}$ ($n = 44$) mice. (B) Total number of live spleen cells from CTRL ($n = 24$) and LysM$^{Cre}$Bim$^{fl/fl}$ ($n = 19$) mice. (C) Representative photomicrographs of H&E-stained spleen sections. Bars, 200 μm. (D) Cervical and inguinal lymph node weights for CTRL ($n = 26$) and LysM$^{Cre}$Bim$^{fl/fl}$ ($n = 20$) mice. (E) Total

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**Primary Language**: English

**Section**: Loss of Bim in myeloid cells, including macrophages, leads to a break in tolerance and the development of SLE-like disease.

**Description**: The figure illustrates the effects of loss of Bim in myeloid cells on autoimmune disease. It shows graphs and images comparing spleen and lymph node weights, cell counts, and immunohistochemical staining for IgG and IgM in CTRL and LysM$^{Cre}$Bim$^{fl/fl}$ mice. The graphs depict significant differences in spleen weight, spleen cell counts, and lymph node weights between the two groups, with a notable increase in autoimmune disease markers in the mice lacking Bim.

**Citation**: Tsai et al.

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**Figure Breakdown**:

- **A**: Spleen weight comparison between CTRL and LysM$^{Cre}$Bim$^{fl/fl}$ mice.
- **B**: Live spleen cell count comparison.
- **C**: Representative H&E-stained spleen sections.
- **D**: Lymph node weight comparison.
- **E**: Total number of live spleen cells.
- **G**: Serum immunoglobulin levels (IgA, IgG1, IgG2a, IgG2b, IgM).
- **H**: Cytokine concentrations (IL-12, IL-17, IL-1β, TGF-β).
- **I**: IFN-α and IFN-β levels.
- **J**: IFN-γ levels.
- **K**: Representative immunohistochemical staining for IgG and IgM.
- **L**: Proteinuria level comparison.
- **M**: Kidney histology comparison.

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**Additional Commentary**: The data suggest a role for Bim in maintaining immune tolerance and preventing the development of autoimmune diseases such as SLE.
We further expanded our studies to examine the splenic cellular composition of LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice using the flow gating strategy shown in Fig. 4 A. The numbers of monocytes, RPMs (Fig. 4, B–D), plasmacytoid dendritic cells (pDCs; Fig. 4 E) and CD11b\textsuperscript{+} dendritic cells (DCs; Fig. 4 F) were elevated in LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice. Further, both CD4 and CD8 naive, memory, and effector T cells (Fig. 4, G and H) and Foxp3\textsuperscript{+} regulatory T cells (Fig. 4 I) showed higher numbers in LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice than controls. All B cell types with the exception of MZ and T2 B cells were also greater in number (Fig. 4 J) in LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice. The enlargement of the lymphocyte compartment was associated with increased activation of monocytes, RPMs, and DCs, as indicated by elevated levels of CD80, MHCI, and CD40 (Fig. 4, K–R). Furthermore, T cells were hyperactivated, as indicated by the significant increase in CD69 and CD279 expression (Fig. 4, S–Y). T cell proliferation was significantly higher in macrophages and DCs isolated from LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice than in those isolated from controls in a mixed leukocyte reaction assay (Fig. 4 Z).

**Bim-deleted myeloid cells are sufficient to cause systemic autoimmunity**

Previous studies have suggested that the LysM\textsuperscript{Cre} (lysozyme M) transgenic mouse may be promiscuous with respect to its promoter (Miyamoto et al., 2002; Ye et al., 2003). We also observed that hematopoietic cells from LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice displayed variable reduction of Bim (Fig. 5 A). Therefore, we adaptively transferred WT CD45.1 CD4\textsuperscript{+}CD8\textsuperscript{−} Bim\textsuperscript{fl/fl} mice that underwent adoptive transfer (Fig. 5, E and F). Moreover, there was no difference in the loss of EdU between LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} and Rag\textsuperscript{−/−} mice once a month (Fig. 5 G). Only Rag\textsuperscript{−/−}LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice that received WT lymphocytes developed splenomegaly (Fig. 5, C and D) and increased lymph node weights (not depicted). The numbers of CD11b\textsuperscript{+} MZMs and CD11b\textsuperscript{−} MZMs were reduced in the spleens of Rag\textsuperscript{−/−}LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice that underwent adoptive transfer (Fig. 5, E and F). In contrast, the numbers of MMMs were mildly affected by the loss of Bim in the Rag\textsuperscript{−/−} mice (Fig. 5, E and F). Moreover, the significantly expanded lymphocyte pool (Fig. 5 G) was associated with increased activation of monocytes, RPMs and DCs in Rag\textsuperscript{−/−}LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice that underwent adoptive transfer as compared with controls (Fig. S3, A–H). This enhanced activation status was also consistent with LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice (Fig. 4, K–R). Serum levels of histone-reactive IgG ANA and ssDNA-reactive IgM ANA after adoptive transfer were significantly elevated in Rag\textsuperscript{−/−}LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice than in Rag\textsuperscript{−/−} mice (Fig. 5 H). Importantly, in the adoptive transfer group, Rag\textsuperscript{−/−}LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice developed GN, as indicated by higher pathology scores (Fig. 5 I), distorted kidney structure (Fig. 5 J), and increased infiltrating CD45\textsuperscript{+} cells and F4/80\textsuperscript{+} macrophages in the glomeruli as compared with Rag\textsuperscript{−/−} mice (Fig. 5 K). Moreover, analysis of Bim expression in Rag\textsuperscript{−/−}LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice after adoptive transfer revealed that the transferred lymphocytes had intact Bim expression, whereas the myeloid cells had consistent loss of Bim (Fig. 5 L). Further, CD4\textsuperscript{+}Bim\textsuperscript{fl/fl} and dLck\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice on a C57BL/6 background (Herold et al., 2014; Li et al., 2017) did not display splenomegaly (Fig. S3, I and J) or MZM loss (Fig. S3, K–N). CD4\textsuperscript{+}Bim\textsuperscript{fl/fl} and dLck\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice did not develop GN (Fig. S3 O) and showed no significant difference in the numbers of kidney monocytes/macrophages compared with controls (Fig. S3, P and Q). These data demonstrate that the deficiency of Bim in myeloid cells is sufficient to drive the SLE pathology in LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice. Further, we support a potential cell intrinsic effect of Bim deficiency in myeloid cells, we generated mixed bone marrow chimeras by injecting LSK (Lin\textsuperscript{−}/Sca1\textsuperscript{+}/c-kit\textsuperscript{+}) cells from B6.CD45.1 (CTRL) or/and LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} (CD45.1) donor mice into host B6.CD45.1/2 mice (Fig. 5 M). These chimeric mice developed a mild systemic autoimmune disease at a ratio of 1:1 CTRL and LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} LSK cells (Fig. 5, N–Q).

**Bim mediates systemic autoimmunity through its BH3 domain**

Because the central function of Bim involves the apoptotic pathway, we examined the effect of Bim deletion on the lifespan of myeloid cells in LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice by monitoring EdU (5-ethyl-2'-deoxyuridine) incorporation and loss over time (Fig. 6 A). Monocytes and neutrophils from various tissues (bone marrow, blood, spleen, and kidney) shared a similar pattern of EdU incorporation at days 35 and 49 (Fig. 6, B–E). Moreover, there was no difference in the loss of EdU between control and LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} monocytes and neutrophils at day 63 (Fig. 6, B–E). pDCs and B cells from the spleen of LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} mice exhibited a lower retention of EdU, whereas little to no difference was observed in control and LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} DCs (Fig. 6 D). EdU-labeling kinetics of macrophages were different in spleen and kidney (Fig. 6, D and E). At day 63, a substantial percentage of LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} RPMs were persistently labeled with EdU (Fig. 6 D), whereas at the same time, LysM\textsuperscript{Cre}Bim\textsuperscript{fl/fl} kidney macrophages lost EdU labeling.
To further identify a potential apoptotic function for Bim, TUNEL analysis was performed. The number of TUNEL+ cells in 3- and 8-mo control and LysMCreBimfl/fl spleens was comparable (Fig. S4, A, C, and D). Similarly, there was no difference in the number of TUNEL+ cells in the tubular regions of 3- and 8-mo LysMCreBimfl/fl kidneys (Fig. S4, B, E, and F). In contrast, loss of Bim significantly affected the survival of bone marrow–derived macrophages (BMDMs) after growth factor withdrawal for 96 h (Fig. S4, G and H). These data suggest that Bim may have apoptotic function in an in vitro cell model, but its in vivo apoptotic function may be less evident.

The BH3 domain of Bim has been implicated in the induction of apoptosis. A previous study created mice where the BH3 domain of Bim was replaced with the domain from other BH3-only proteins such as Bad (BimBad) and Puma (BimPuma; Mérino et al., 2009). To examine a potential nonapoptotic function for the BH3 domain, BMDMs were treated with TLR agonists. LysMCreBimfl/fl and BimBad BMDMs produced significantly higher levels of IL-12p40 in response to LPS or CpG treatment than controls (Fig. 6, F and G). Furthermore, BimBad mice exhibited higher spleen weight and developed worse kidney pathology than controls (Fig. S4, I and J), whereas BimPuma mice were similar to con-
trols (Fig. 6, F and G; and Fig. S4, I and J). We then generated mice that express one Bim-BH3 mutant allele and one floxed Bim allele by crossing Bim<sup>fl<sup/><sub>puma</sub></sup>, Bim<sup>fl<sup/><sub>Bad</sub></sup> or Bim<sup>fl<sup/><sub>Noxa</sub></sup> mice to LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> mice. LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> and LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/puma</sub></sup> mice exhibited splenic weight and cell numbers (Fig. 6, H and I), ANA (Fig. 6, J and K), and kidney pathology (Fig. 6, N and O) comparable with controls. In contrast, LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> and LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/Noxa</sub></sup> mice displayed sphenomegaly, increased ANA, and severe GN similar to LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> mice (Fig. 6, H, I, and L–O).

### Deletion of TRIF ameliorates GN in LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> mice

The TLR pathway has been implicated in the development of SLE-like disease in mice (Baccala et al., 2007; Kim et al., 2009; Nickerson et al., 2010). MyD88 is the adaptor protein for most TLRs, except TLR3 and, in some cases, TLR4, which signal through TRIR-domain–containing adapter–inducing IFN-β (TRIF). Therefore, we deleted MyD88 in myeloid cells or TRIF in all cells of LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> mice to generate MyD88<sup>fl<sup/><sub>/fl</sub></sup>LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> mice and TRIF<sup>−/−</sup>LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> mice, respectively. Both MyD88<sup>fl<sup/><sub>/fl</sub></sup>LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> and TRIF<sup>−/−</sup>LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> mice exhibited sphenomegaly (Fig. 7, A and B) and loss of MZMs (Fig. 7 C). Further, MyD88<sup>fl<sup/><sub>Bad</sub></sup>LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> (Fig. 7 D) and TRIF<sup>−/−</sup>LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> mice (Fig. 7 E) showed higher titers of IgG ANA to ssDNA, dsDNA, histones, and nucleosomes and evidence of IgG-immune complex deposition in glomeruli than controls (Fig. 7 F).

Deletion of TRIF, but not MyD88, prevented GN in LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> mice (Fig. 7, G and H). We then analyzed the immune cell composition of the kidney from normal mice under steady-state conditions using multiparameter flow cytometry. Macrophages in the kidneys were denoted by CD11b<sup>+</sup>CD64<sup>+</sup>F4/80<sup>+</sup>MHCI<sup>II</sup>− (Fig. 7 I). There were significantly higher numbers of Ly6C<sup>hi</sup> monocytes and macrophages in the kidneys of LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> mice than in control kidneys (Fig. 7, K and L). However, the number of Ly6C<sup>lo</sup> monocytes and macrophages was significantly lower in TRIF<sup>−/−</sup>LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> mice than LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> mice (Fig. 7, J and L). To further identify whether TRIF signaling is enhanced in LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> mice, we treated BMDMs with poly(I:C). There was a dramatic increase of IRF3 and TBK1 phosphorylation/activation in LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> BMDMs compared with controls (Fig. S5 A). In contrast, LPS stimulation had no effect on IRF3/TBK1 activation (Fig. S5 A) but induced IL-12 and TNF production (Fig. S5 B).

### LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> mice display an SLE-like specific transcriptional signature

We performed RNA-seq of kidney macrophages (CD11b<sup>+</sup>CD64<sup>+</sup>F4/80<sup>+</sup>MHCI<sup>II</sup>−) from control and LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> mice to study the transcriptional response in kidney macrophages. We found 87 significantly up-regulated and 69 significantly down-regulated genes in LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> kidney macrophages compared with controls (Fig. 8 A). GO enrichment analysis revealed that the up-regulated genes, such as Nod1, Cd84, and Cd274, are involved in “regulation of immune system process” (false discovery rate [FDR] adjusted P = 6.66 × 10<sup>−2</sup>; Fig. 8 B), whereas the down-regulated genes, such as Hdc9, Cd36, and Adrb2, are enriched for “regulation of vasculature development” (Fig. 8 B). K-means clustering (K = 4; Fig. 8 C) of the 5,694 differentially expressed genes (fold change ≥1.5, compared across any two of the strains) yielded three main categories: genes that were regulated by both Bim and TRIF are denoted in cluster I (lupus nephritis [LN] signature), genes that were affected by Bim independent of TRIF are indicated in cluster II (Bim-regulated signature), and genes that were regulated by TRIF independent of Bim are expressed in clusters III and IV (TRIF-regulated genes). Next, we verified that the overlap of genes regulated by both Bim and TRIF (cluster I, LN signature) did not occur by chance (Fig. 8 D). We found that the number of genes demonstrating reverse regulation between control to LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> kidney macrophages and LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> to TRIF<sup>−/−</sup>LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> kidney macrophages was higher than expected (χ<sup>2</sup> test, P < 10<sup>−110</sup>; Fig. 8 E).

### LysM<sup>Cre</sup>Bim<sup>fl<sup/><sub>/fl</sub></sup> kidney macrophages share a common transcriptional signature with Bim-deleted MZMs, NZB/W kidney macrophages, and human LN kidneys

We then calculated the number of differential genes as a result of Bim deletion across the affected tissues by ex-
aminsing LysM<sup>cre</sup>Bim<sup>fl/fl</sup> CD11b<sup>+</sup> MZMs, CD11b<sup>−</sup> MZMs, and kidney macrophages (Fig. 3, G and H; and Fig. 8 A). Significant numbers of genes from CD11b<sup>+</sup> MZMs (two genes, P < 10<sup>−5</sup>) and CD11b<sup>−</sup> MZMs (three genes, P < 0.05) overlapped with kidney macrophages (Fig. 9 A and Table S2). Additionally, CD11b<sup>−</sup> MZMs shared a significant number of genes with CD11b<sup>+</sup> MZMs (two genes, P < 10<sup>−5</sup>; Fig. 9 A and Table S2). We also compared our results to a microarray dataset from a previously published mouse model of lupus (Berthier et al., 2012), which examined kidney macrophages from young and old NZB/W mice. We identified a strong positive correlation (R = 0.63) in gene expression profiles between LysM<sup>cre</sup>Bim<sup>fl/fl</sup> kidney macrophages and NZB/W kidney macrophages (Fig. 9 B). Additionally, we compared our results to a microarray dataset from whole kidney biopsy specimens from human LN patients separated into tubulointerstitial or glomerular compartments (Berthier et al., 2012). LysM<sup>cre</sup>Bim<sup>fl/fl</sup> kidney macrophages and tubulointerstitial and/or glomerular tissue of human LN demonstrated a low but positive correlation (R = 0.1; Fig. 9, C and D). There were 33 genes (22 up- and 11 down-regulated; Table S3) in common between LysM<sup>cre</sup>Bim<sup>fl/fl</sup> kidney macrophages and NZB/W kidney macrophages (P < 10<sup>−4</sup>; Fig 9 E). These genes were enriched for GO processes such as “adaptive immune response” (FDR adjusted P = 6.03 × 10<sup>−2</sup>), “positive regulation of mononuclear cell migration” (FDR adjusted P = 1.13 × 10<sup>−1</sup>), and “negative regulation of IL-10 production” (FDR adjusted P = 9.63 × 10<sup>−5</sup>). The kidney macrophages from LysM<sup>cre</sup>Bim<sup>fl/fl</sup> mice shared a significant number of genes with the tubulointerstitial area (seven genes, P < 10<sup>−2</sup>; Fig. 9 F and Table S3) but shared only five genes with glomerular tissue (Fig. 9 F). The five shared genes from tubulointerstitial and glomerular area of human LN included ISG15, Ly6E, LYZ, MS4A6A, and IFITM3 (Table S3). Collectively, these data suggest that the loss of Bim in myeloid cells leads to systemic autoimmunity and end-stage GN. Thus, LysM<sup>cre</sup>Bim<sup>fl/fl</sup> mice represent a new model of SLE that shares characteristics with established murine models of SLE-like disease and LN patients.

**DISCUSSION**

The origin of the break in tolerance leading to development of SLE and subsequent GN is unknown. Previous studies suggested that the Bcl-2 family members may be critical for this process. Bim<sup>−/−</sup> and mice lacking Bim in all hematopoietic cells (Vav<sup><sub>Bim<sup>-/−</sub></sup></sub>) develop systemic autoimmunity and end-stage LN (Bouillet et al., 2002; Herold et al., 2014). Only CD19<sup><sub>Bam<sup>-/−</sub></sup></sub>Bax<sup><sub>Bak<sup>-/−</sub></sup></sub> chimeras and vav-Bcl-2tg mice exhibit a systemic autoimmune disease phenotype similar to Bim<sup>−/−</sup> mice (Takeuchi et al., 2005; Mason et al., 2013). Here, we examine the impact of deleting Bim in the myeloid cell compartment using LysM<sup>cre</sup>Bim<sup>fl/fl</sup> mice. LysM<sup>cre</sup>Bim<sup>fl/fl</sup> mice display hallmarks of SLE-like disease, including loss of MZMs, splenomegaly, lymphadenopathy, autoantibody production, IFN signature, and increased mortality. However, further analysis reveals promiscuity in the LysM promoter, consistent with previous studies that suggested LysM may be expressed not only in myeloid cells but also in developing hematopoietic cells that differentiate into B and T cells (Miyamoto et al., 2002; Ye et al., 2003). Interestingly, Fas and caspase-8 deletion is intact in lymphocytes from LysM<sup>cre</sup>Fas<sup>−/−</sup> and LysM<sup>cre</sup>caspase-8<sup>−/−</sup> mice, respectively (Cuda et al., 2012, 2015). These data suggest that the leakiness of LysM may be more dependent on the accessibility of the floxed gene. Nonetheless, studies using Rag<sup>−/−</sup>LysM<sup>cre</sup>Bim<sup>fl/fl</sup> mice that received adoptively transferred WT lymphocytes eliminates any possibility that the susceptibility to SLE-like disease in LysM<sup>cre</sup>Bim<sup>−/−</sup> may be derived from Bim-deleted B and T cells. However, our data do not rule out an effect of the loss of Bim in other myeloid populations such as DCs and neutrophils. As opposed to CD19<sup><sub>Bam<sup>-/−</sub></sup></sub>Bax<sup><sub>Bak<sup>-/−</sub></sup></sub> mice (Takeuchi et al., 2005), CD4<sup><sub>Bam<sup>-/−</sub></sup></sub> and CD19<sup><sub>Bam<sup>-/−</sub></sup></sub>Bim<sup>−/−</sup> mice fail to develop an SLE-like disease phenotype, even though they are on an autoimmune prone background. These data suggest that Bim plays a role in autoimmunity that does not parallel other Bcl-2 proteins; furthermore, deletion of Bim in myeloid cells, including macrophages, is sufficient to induce the development of autoimmunity in mice.

Previous studies have suggested that a failure to process apoptotic bodies by MZMs, which normally induces toler-
Figure 6. The systemic autoimmunity in LysM\textsuperscript{Cre}\textsuperscript{Bim\textsuperscript{f/f}} mice may not require enhanced survival of monocytes/macrophages. (A) Representation of experimental design for EdU study as described in Materials and methods. (B–E) FACS analysis of percentage of EdU incorporation in multiple immune cell populations from CTRL and LysM\textsuperscript{Cre}\textsuperscript{Bim\textsuperscript{f/f}} mice at day 35 (n = 4), 49 (n ≥ 8), and 63 (n = 6) in bone marrow (B), blood (C), spleen (D), and kidney (E). The experiment was performed one time. Data are presented as mean ± SEM and were compared by Mann–Whitney test. Asterisks indicate a significant difference between LysM\textsuperscript{Cre}\textsuperscript{Bim\textsuperscript{f/f}} and CTRL (*, P < 0.05; **, P < 0.01; ***, P < 0.001). (F and G) Bim\textsuperscript{Bad} GM-CSF BMDMs were hyperresponsive to TLR activation. BMDMs were stimulated with either LPS (F) or CpG (G) for 6 h, and supernatants were collected and evaluated for IL-12p40 by multiplex immunoassay as
ance, may be necessary for the activation of lymphocytes in SLE-like disease (McGaha and Karlsson, 2016). We observe that the MZMs are absent as early as 1.5 mo of age in LysMCre Bim−/− mice, whereas the numbers or topography of the MMMs are minimally affected. Similarly, Rag−/−/LysMCre Bim−/− mice reconstituted with WT lymphocytes also exhibit a reduced number of MZMs, whereas the MZMs in Rag−/− mice transferred with WT lymphocytes are normal. The increased numbers of pDCs in the spleen are associated with elevated serum levels of type I IFN in LysMCre Bim−/− mice. These data are consistent with a recent study, which suggested that the lack of MZMs may be because of the recruitment of type I IFN–expressing pDCs. Further, type I IFN alters the location of MZB cells, thereby disrupting the lympho toxin signaling pathway in MZMs (Li et al., 2015). In line with this, we detected a change in the localization of MZB cells but not in their total number in LysMCre Bim−/− mice. However, we did not observe any difference in the gene expression of the members of lympho toxin pathway, including megakaryoblastic leukemia 1, megakaryoblastic leukemia 2, and serum response factor (Li et al., 2015). Although our study has some limitations, the differences in lympho toxin pathway gene expression between the previous model and LysMCre Bim−/− mice may be because of the numbers of replicates or the lupus models being investigated.

Although Bim is crucial for maintaining hematopoietic cell homeostasis through the induction of apoptotic cell death, our data suggest that Bim may also play a nonapoptotic role that affects the development of SLE-like disease. These data are also consistent with previous studies suggesting that the Bcl-2 pathway, including Bim, may have nonapoptotic functions in macrophages and DCs (Nopora and Brocker, 2002; Hou and Van Parijs, 2004; Chen et al., 2007; Gautier et al., 2008; Hutcheson and Perlman, 2008; Hutcheson et al., 2008; Scatizzi et al., 2010; Yeretsian et al., 2011). We show that Bim–deleted macrophages display increased markers of activation and secrete elevated levels of cytokines, and these results are similar to those of our previous study (Scatizzi et al., 2010). Loss of Bim has no effect on the turnover of MZMs or even in development of MZMs in the Rag−/− background. Additionally, pulse–chase EdU studies suggest that Bim either reduces the rate of proliferation or has only a minor effect on the cell death of myeloid cells. However, because we do not observe a difference in the rate of EdU uptake, these data would suggest that the rate of proliferation is the same in monocytes and macrophages regardless of the presence of Bim. We cannot rule out that there is a marginal effect on apoptosis, because Bim-deleted BMDMs have lower levels of apoptosis than controls.

Previous studies have shown that the BH3 protein family is divided into direct and indirect activators of cell death. Bid, Bim, and Puma are considered activators of apoptosis, as BH3 peptides from these proteins are capable of inducing apoptosis (Billard, 2013). Macrophages from BimBad and LysMCre Bim−/−, but not BimPuma mice, exhibit enhanced cytokine production in response to TLR stimulation that is independent of cell death. Further, replacement of one of the Bim alleles in LysMCre Bim−/− mice with that of Bad−/− or Noxa−/Bim leads to an SLE-like disease that mirrors LysMCre Bim−/− mice. In contrast, replacement of Bim−/Bim with Puma−/Puma prevents SLE-like disease and resembles WT or LysMCre Bim−/− mice. These data are consistent with the notion that the BH3 domain of Puma may function in a manner similar to the BH3 domain of Bim (Mérino et al., 2009). Our data partially agree with the study demonstrating that BimBad or BimNoxa rescues the lethal disease seen in Bcl-2−/− mice (Mérino et al., 2009). Differences between the studies may reflect the mechanism of disease (i.e., Bcl-2−/− mice develop hematopoietic deficiency and PKD, whereas LysMCre Bim−/− mice develop systemic autoimmunity and LN). Collectively, these data suggest that the BH3 domain of Bim in myeloid cells is required to suppress systemic autoimmunity. Moreover, Bim may have multiple functions in myeloid cells that can contribute to SLE-like disease. Future studies are required to definitively show that Bim has a direct nonapoptotic function.

Our cell culture data using BMDMs suggest that the loss of Bim may enhance TLR signaling. However, deletion of MyD88 or TRIF in LysMCre Bim−/− mice has no effect on systemic autoimmunity. In contrast, loss of TRIF in LysMCre Bim−/− mice is sufficient to suppress the progression to full-blown GN. Although TRIF−/−/LysMCre Bim−/− mice display immune complex deposition, there is limited macrophage infiltrate and reduced GN. These data suggest that macrophage infiltration is necessary for the destruction of the kidney, consistent with human studies (Hill et al., 2001). The deletion of TRIF in LysMCre Bim−/− mice clearly delineates systemic autoimmunity from end-organ disease, consistent with a previous study (Ge et al., 2013). Our cell culture studies using BMDMs suggest that loss of Bim leads to enhanced TBK1 activation in response to TLR3 signaling, which occurs...
through TRIF. Future studies may require generating LysM^{Cre} Bim^{fl/fl} mice that also lack TBK1. However, we cannot rule out that TRIF-deleted kidney stromal cells or other nonmyeloid cells influence macrophage recruitment. Although loss of Myd88 exacerباتеs the development of LN in LysM^{Cre} Bim^{fl/fl} mice, this altered phenotype may be caused by overcompensation by the TRIF signaling pathway. Because both Myd88 and TRIF are involved in downstream signaling for TLR4, future studies are necessary to examine the role that TLR4 pathway plays in disease. Our data are novel and are in contrast with other studies showing that Myd88 deletion is required in B cells and DCs to prevent systemic autoimmunity in Lyn^{−/−} mice (Ban et al., 2016) and for the expansion of lymphocytes and end-organ damage in MRL/lpr mice (Teichmann et al., 2010). Differences between our data and those using Lyn^{−/−} and MRL/lpr mice may be attributed to background of the mice and the affected cells caused by global gene deficiency.

Numerous studies have used gene expression profiling of peripheral blood mononuclear cells and whole kidney tissue from SLE patients to gain insights into the pathobiology of disease or identification of a potential biomarker (Tsokos et al., 2016). Nonetheless few studies isolated individual cell populations to understand their specific roles in SLE disease pathogenesis. Our global gene expression studies demonstrate a lupus-specific as well as a TRIF- and Bim-specific signature in kidney macrophages from control, LysM^{Cre} Bim^{+/+} and TRIF^{+/−}LysM^{Cre} Bim^{+/−} mice. For example, the expression level of CD36, a receptor known to be involved in phagocytosis of apoptotic bodies, is lower in LysM^{Cre} Bim^{+/−} kidney macrophages than control kidney macrophages. Expanding our analysis to MZMs from LysM^{Cre} Bim^{+/−} mice yields some commonalities. We also identify similarities between kidney macrophage transcriptional signatures in LysM^{Cre} Bim^{+/−} mice and those from old NZB/W mice, even though these cells are from two different backgrounds. Of note, both Cd274 (PD-L1; Dai et al., 2014) and Ccl2 (Tesch et al., 1999), known contributors to SLE, are enhanced in these cells. Additionally, we detected a significant number of overlapping genes between tubulointerstitial tissue of LN patients and kidney macrophages from LysM^{Cre} Bim^{+/−} mice. Despite the difficulties of comparing transcriptional signatures across species and techniques (RNA-seq vs. microarray), seven genes are shared between LysM^{Cre} Bim^{+/−} kidney macrophages and tubulointerstitial tissue, including IFITM2 (Rai et al., 2016), LY6E, and ISG15 (Feng et al., 2006), genes known to be highly expressed in SLE patients. However, given that the patient data represent whole tissue, it is unknown whether the increased expression in these genes from the lupus patient datasets is macrophage specific. Future studies require a direct comparison of isolated macrophages from kidneys of SLE patients with those from murine studies. Collectively, we have established a novel model of SLE-like disease that is mediated by loss of Bim in myeloid cells. Further, our data demonstrate that Bim is a new therapeutic target for SLE treatment.

**MATERIALS AND METHODS**

**Mice**

Mice homologous for loxP-flanked Bim allele (CTRL: LysM^{+/−}Bim^{+/−}, C57BL/6:129 mixed background) were a gift from the late S. Korsmeyer and J. Opferman (Takeuchi et al., 2005; Steimer et al., 2009) and were mated with mice expressing Cre under the control of LysM promoter (LysM^{Cre}; The Jackson Laboratory), CD19-Cre (CD19^{Cre}) knock-in mice, or CD4-Cre (CD4^{Cre}; Taconic Biosciences, Inc.) transgenic mice. In addition, Bim^{+/−} were backcrossed to C57BL/6 by The Jackson Laboratory using speed congenics to achieve 98% C57BL/6. The mice were then backcrossed to C57BL/6 for two additional generations to generate congenic Bim^{+/−} mice. After backcross, Bim^{+/−} mice were mated with LysM^{Cre} mice to generate LysM^{Cre} Bim^{+/−}. The fidelity of deleting Bim in LysM^{Cre} Bim^{+/−} was previously published and confirmed by us (Takeuchi et al., 2005; Steimer et al., 2009). Rag^{−/−} mice (The Jackson Laboratory), Myd88^{−/−} mice (The Jackson Laboratory), and TRIF^{−/−} mice (The Jackson Laboratory) were crossed to LysM^{Cre} Bim^{+/−} to generate Rag^{−/−}LysM^{Cre} Bim^{+/−}, LysM^{Cre} Bim^{+/−}Myd88^{−/−} mice, and TRIF^{−/−}LysM^{Cre} Bim^{+/−} mice. Bim^{−/−}, Bim^{−/−}Noxa, and Bim^{−/−}Puma mice were provided by P. Bouillet (Mérino et al., 2009). Bim^{−/−}Bim^{−/−}Noxa, and Bim^{−/−}Puma mice were crossed with LysM^{Cre} Bim^{−/−} to generate LysM^{Cre} Bim^{lox/−}Bim^{−/−}, LysM^{Cre} Bim^{lox/−}Noxa, and LysM^{Cre} Bim^{lox/−}Puma mice. CD4^{−/−} Bim^{−/−} and dLck^{Cre} Bim^{−/−} mice were generated on a C57BL/6 background as previously described (Herold et al., 2014; Li et al., 2017; provided by D.A. Hildeman, University of Cincinnati, Cincinnati, OH). B6 CD45.1/2 mice were generated by crossing B6 mice to B6.CD45.1/2. All mice were housed and bred at a barrier- and specific-pathogen-free facility at the Center for Comparative Medicine at...
Figure 8. Bim cooperates with TRIF to regulate transcriptional profiles of kidney macrophages. (A) Volcano plot of differentially expressed genes in LysM<sup>Cre<sup>Bim<sup>fl/fl</sup></sup> kidney macrophages compared with CTRL kidney macrophages. Significantly regulated genes are indicated in red (n = 2). (B) Bar graphs indicating the normalized gene expression of individual genes in CTRL and Cre<sup>LysM<sup>Bim<sup>fl/fl</sup></sup> kidney macrophages. Data are presented as mean ± SEM. (C) Heat map of K-means (K = 4) clustering on 5,694 differential genes and GO processes from each cluster. (D) Scatterplot showing fold change of gene expression...
Northwestern University. Genotyping of the mice was done by Transnetyx. Female mice were used for all experiments. Proteinuria was measured with Uristix reagent strips (Siemens Healthcare Diagnostics). All housing conditions and animal experiments were approved by the Northwestern University Institutional Animal Care and Use Committee.

**Histopathology and immunohistochemistry**

Spleen and kidney were isolated and fixed in 2% paraformaldehyde. 4-µm paraffin-embedded kidney and spleen sections were stained with periodic acid–Schiff base (PAS), H&E, F4/80, or CD45 as previously described (Hutcheson et al., 2008). The PAS-stained kidney sections were scored by a pathologist (G.K. Haines) blinded to the study. The kidney scores were determined by the addition of the values from glomerulosclerosis, interstitial fibrosis, and different morphological indexes based on the previous published scoring system (Hill et al., 2000). Photomicrographs were taken on an Olympus BX41 microscope equipped with an Olympus DP21 camera.

**Serological tests**

Serum was obtained from 8-mo-old CTRL (Bim fl/fl) and LysM¿⁄¿Bim¿⁄¿ mice and was diluted 50 times in 0.1% gelatin supplemented with 2% BSA (Sigma-Aldrich), 3 mM EDTA, and 0.05% (vol/vol) Tween-20 (Sigma-Aldrich). Immulon 2 HB 96-well flat bottom plates (Immunology Technology) were precoated with methylated BSA followed by immobilization of antigens including dsDNA, ssDNA, histones, or dsDNA and histones (nucleosomes). Plates were washed with DPBS (Lonza) after incubation, and serum samples were added into the wells. Unbound serum samples were washed with DPBS containing 0.05% (vol/vol) Tween-20 (Sigma-Aldrich), and serum samples were conjugated to goat anti–mouse IgG or IgM antibodies conjugated with alkaline phosphatase (SouthernBiotech). pNPP (p-nitrophenyl phosphate disodium salt; Sigma-Aldrich) was used for colorimetric detection, and the plates were read at OD 405 nm on a BioTek Synergy HT Multi Mode plate reader. For measurement of serum isotyping antibodies, serum was diluted at 1:25,000, and a mouse immunoglobulin isotyping 96-well plate assay kit (EMD Millipore) was used according to the instructions of the manufacturer. The results were read on a Luminex 200 instrument running on an xPONENT software. Serum cytokines were evaluated using Luminex bead-based assays (Affymetrix).

**Immunofluorescent microscopy**

Kidneys were processed analysis for immune complex deposition as previously described by us (Hutcheson et al., 2008; Cuda et al., 2014). For splenic MZMs, MMMs, and MZB cells, spleens were isolated and snap-frozen in Tissue-Tek OCT and sectioned at 8 µm thickness followed by fixing in cold acetone and blocking with Fc Block (BD Biosciences) and normal rat serum (Jackson ImmunoResearch Laboratories). Finally, the spleen sections were stained with anti–CD209b–APC (clone REA125; Milteny Biotech), anti–CD169–PE (clone REA197; Milteny Biotech), anti–CD169–FITC (clone MOMA-1; AbD Serotec), anti-B220–Alexa Fluor 488 (clone RA3-6B2; BioLegend), anti-B220–Alexa Fluor 647 (clone: RA3-6B2; BioLegend), anti–CD11d–PE (clone 1B1; BioLegend), or anti–CD4–Alexa Fluor 647 (clone GK1.5; BioLegend). Images were acquired using a 10× objective on a Nikon TE2000E2-PFS running NIS-Elements software.

**Tissue preparation and flow cytometry**

Preparation of single-cell suspension from blood and spleen was performed as previously described (Rose et al., 2012). Kidneys were collected into cold MACS buffer (Miltenyi Biotech), carefully decapsulated, weighed, cut into small fragments (1–2 mm), and subjected to mechanically disruption using C-tubes (Miltenyi Biotech) and GentleMACS dissociator (Miltenyi Biotech) with the program m_liver_01.02. The disrupted kidney tissue was continuously rotated at 37°C for 30 min and subjected to GentleMACS dissociator (Miltenyi Biotech) using the program m_liver_02.02. Each kidney was digested with 2.5 ml in-house digestion buffer consisting of 0.76 U/ml Liberase TL (Sigma-Aldrich) and 2mg/ml DNase I (Roche) dissolved in HBSS with Ca²⁺ and Mg²⁺ (Mediatech Inc.). Cells released during enzymatic digestion were filtered through Falcon 40-µm cell strainers (Corning), and red blood cells were lysed with Pharm Lyse Buffer (BD Biosciences). Isolated kidney cells were labeled with CD45 MicroBeads (Miltenyi Biotech) and separated on MultiMACS Cell24 Separator Plus (Miltenyi Biotech). Total numbers of live cells were counted by Countess Automated cell counter (Invitrogen), and dead cells were distinguished by Trypan Blue stain. Cells were stained with Live/Dead Aqua fluorescent dye (Invitrogen), incubated with Fc Block and then stained for 30 min with the fluorochrome-conjugated antibodies. Data were acquired on an LSR II flow cytometer (BD Biosciences). Compensation and data were analyzed using FlowJo software (Tree Star). FMO (fluorescence minus one) controls were used to gate MZM and MMM populations or when necessary. Splenic MZMs, MMMs, and kidney macrophages were sorted on a BD FACSARia SORP instrument (BD Biosciences) at the Northwestern University Robert H. Lurie Comprehensive Cancer Center Flow Cytometry Core Facility.
Mixed leukocyte reaction assay
Isolation of single-cell suspension from the spleens was performed as previously described (Rose et al., 2012). In brief, splenic CD11b+ cells were purified by immune-magnetic positive selection. Splenocytes were incubated with anti-CD11b MACS beads (Miltenyi Biotec), and CD11b+ cells were puri-
fied with a MACS column. Purity of macrophages was ~90% as determined by FACS analyses. Purified CD11b⁺ splenic cells were pulsed with 10 μg/ml OVA peptide (aa 323–339) for 60 min at 37°C in 5% CO₂. The pulsed cells were washed two times with RPMI. OVA-specific CD4 T cells were isolated from the spleens of OT-II/Rag⁻/⁻ mice using the CD4 T cell isolation kit (Miltenyi Biotec) according to the manufacturer's protocol. The resulting CD4 T cells were >90% purified as determined by flow cytometry. Thereafter, CD4 T cells were labeled with 500 nM CFSE (Invitrogen) for 12 min 37°C in 5% CO₂. 2.5 × 10⁴ OVA-pulsed CD11b⁺ splenic cells were incubated with 20 × 10⁶ CFSE-labeled CD4 T cells in the presence or absence of 5 μg/ml CpG-B (ODN 1668) in a 96-well flat-bottom plate for 3 d at 37°C in 5% CO₂. All cultures were set up in triplicate. To harvest the cells, 7.5 mM EDTA was added to the cells for 15 min. Cells were then washed with MACS buffer and stained with anti-CD4 (BD Biosciences). A constant number of Calibrite APC beads (BD Biosciences) was added to allow the acquisition of equal parts per culture. 0.25 mg 7-aminoactinomycin D per test (BD Biosciences) was used to exclude the dead cells. For data acquisition, a constant number of Calibrite APC beads was counted. Live CD4 T cells (CD4⁺ 7-aminoactinomycin D⁻) were gated. The number of dividing cells (i.e., those showing less than the maximal CFSE fluorescence intensity) was determined as previously described (Cuda et al., 2015).

Isolation and adoptive transfer of lymphocytes
Isolation of single-cell suspension from the spleens of 8-wk-old B6.CD45.1 mice was performed as previously described (Rose et al., 2012). Splenic lymphocytes were purified by negative immune-magnetic selection. In brief, spleen cells were treated with Fc Block and incubated with PE-conjugated myeloid-lineage antibodies including anti-Ter119 (clone TER119; eBioscience), anti-CD11b (clone M1/70; BD Biosciences), anti-F4/80 (clone BM8; BioLegend), anti-CD11c (clone HL3; BD Biosciences), anti-mPDCA-1 (clone JF5-1C2.4.1; Miltenyi Biotec), anti-SiglecF (clone CD117; BD Biosciences), and depletion on autoMACS Pro Separator (Miltenyi Biotec) with the “depletes” program. The resulting lymphocyte preparation was >80% purity as determined by flow cytometry. Each CD45.2. Rag⁻/⁻ or CD45.2. Rag⁻/⁻/LysMCreBim⁰/⁺ mouse received ~4.0 × 10⁷ to 5.0 × 10⁷ donor lymphocytes through retro-orbital injections every month over an 8-mo period and bled every month to ensure the efficacy of adoptive transfer. The recipient CD45.2. Rag⁻/⁻ and CD45.2. Rag⁻/⁻/LysMCreBim⁰/⁺ mice were then sacrificed after 8 mo of adoptive transfer to assess SLE-like disease symptoms.

Mixed bone marrow chimeras
Bone marrow was isolated from the femurs and tibias from 6-wk-old healthy donors from CTRL (Bim⁰/⁺) and LysMCre-Bim⁰/⁺ mice. Red blood cells were lysed with Pharm Lyse Buffer (BD Biosciences). Enrichment of LSK (Lin⁻Sac-1⁻ c-kit⁻) cells was performed via labeling with CD117 (c-kit) MicroBeads (Miltenyi Biotec) and positively selecting the c-kit⁺ cells on an autoMACS Pro Separator. The positively selected fraction was then incubated with Fc Block and labeled with fluorochrome-conjugated antibodies. LSK cells were obtained by FACS using a BD FACSAria SORP instrument (BD Biosciences) at the Northwestern University Robert H. Lurie Comprehensive Cancer Center Flow Cytometry Core Facility. 3-mo-old recipient B6.CD45.1/2 mice were subjected to a single dose of 1,000 cGy g-irradiation using a Cs-137–based Gammatrac 40 irradiator (Best Therantronsics). 12 h after irradiation, recipients were injected 2 × 10⁵ LSK cells from CTRL mice, 2 × 10⁵ LSK cells from LysMCre-Bim⁰/⁺ or 10⁵ of LSK cells from each of CTRL and LysMCre-Bim⁰/⁺ intravenously. Chimeric mice were given trimethoprim/sulfamethoxazole (40 mg/5 mg, respectively; Hi-Tech Pharmacal/Akorn) through autoclaved drinking water for 1 mo and then switched back to normal drinking water. Chimeras were assessed for SLE-like disease symptoms after 8 mo.

EdU study
CTRL (Bim⁰/⁺) and LysMCre-Bim⁰/⁺ mice were given 2 mg EdU (Santa Cruz Biotechnology, Inc.) through subcutaneous injections daily for 49 d. To examine turnover rate and proliferation in tissue-resident monocytes/macrophages, mice were sacrificed at several time points (days 35, 49, and 63). Single-cell suspensions from bone marrow, blood, spleen, and kidney were stained with fluorochrome-conjugated antibodies. EdU staining was assayed with intracellular flow cytometry using a Click-iT Plus EdU Alexa Fluor647 Flow Cytometry Assay kit (Invitrogen) according to the manufacturer's protocol. Each tissue came along with an FMO control to identify the EdU-positive cells. Tissues were acquired on an LSRII flow cytometer. Compensation and data were analyzed using FlowJo software (Tree Star).

TUNEL analysis
Frozen spleen and kidney sections were cut at 4 μm thickness and fixed with 1% paraformaldehyde in PBS, pH 7.4, for 10 min at room temperature. Mouse thymus was used as a positive control for TUNEL. Slides were then postfixed in precooled ethanol/acetate 2:1 for 5 min at −20°C. TdT-enzyme and dUTP-conjugated fluorescein cocktail were added according to manufacturer’s protocol (ApopTag Peroxidase In Situ Apoptosis Detection kit; Millipore). Slides were mounted with mounting medium containing DAPI nuclei staining.

In vitro assays: Cell culture, cytokine measurements, and Western blot analysis
Bone marrow was prepared from the femurs and tibias, and erythrocytes were lysed. Bone marrow was then cultured in DMEM (Mediatech Inc.) supplemented with 10% heat-inactivated fetal bovine serum (ATLAS Biologicals), 2 mM l-glut...
tamine (Mediatech Inc.), 100 IU/ml penicillin, 100 μg/ml streptomycin (Mediatech Inc.), and 1 mM sodium pyruvate (Mediatech Inc.) in the presence of 25 ng/ml recombinant murine GM-CSF (PeproTech, Inc.) or M-CSF (PeproTech, Inc.) at 37°C in 5% CO2. Culture medium was changed on day 3 and day 5. (G)M-CSF BMDMs were seeded on day 7 at a concentration of 1.0–1.8 × 10^6 cells/ml in the 6-well plates or 12 × 75 mm culture tubes. For the cell death assay, growth factors were removed or kept in M-CSF BMDMs for 96 h at 37°C in 5% CO2 and stained with Annexin V and Aqua Live/Dead. To measure cytokine fluctuations in response to TLR agonists, GM-CSF BMDMs were stimulated with or without 5 μg/ml imiquimod (InvivoGen) or 5 μg/ml CpG (InvivoGen) for 6 h, and supernatants were harvested for cytokine measurements using ProcartaPlex Multiplex Immunoassay kit (Affymetrix). For Western blot analysis, GM-CSF BMDMs were seeded at 3.2 × 10^5 cells per well in a 24-well plate for 24 h before treatment. For poly(I:C) stimulation, GM-CSF BMDMs were mock transfected for 1 h or transfected with 0.5 μg LMW-poly(I:C) (InvivoGen) per well at a final concentration of 1 μg/ml for 1 or 2 h using 2.5 μl Lipofectamine 2000 Reagent per 1 μg poly(I:C). For LPS stimulation, GM-CSF BMDMs were treated with crude LPS (Sigma–Aldrich) at a final concentration of 100 ng/ml for 1 h. GM-CSF BMDMs were harvested in 150 μl Laemmli, and 30 μl of the protein lysate was separated by SDS-PAGE followed by immunoblot with phospho–TBK1, TBK1, phospho–IRF3 Bim or GAPDH (Cell Signaling Technology) at a dilution of 1:1,000. HRP-conjugated anti–rabbit IgG secondary antibody (GE Healthcare) was used at 1:6,000.

RNA-seq

Cells were sorted as described in “Tissue Preparation and Flow Cytometry” and immediately lysed with RLT Buffer from the Qiagen RNeasy Plus Mini kit or RNA extraction buffer from PicoPure RNA isolation kit (Arcturus Bioscience, Inc.). Cell lysates were stored at −80°C until RNA was extracted. RNA isolations were performed using the Qiagen RNeasy Plus Mini kit or PicoPure RNA isolation kit. RNA quality and quantity were measured using the Agilent Technologies High Sensitivity RNA ScreenTape System. SMA RT-Seq v4 Ultra Low Input RNA kit (Clontech Laboratories, Inc.) was used for full-length cDNA synthesis, and the Nextera XT DNA sample preparation kit (Illumina Inc.) was used for library preparation. DNA libraries were sequenced on an Illumina NextSeq 500 instrument with a target read depth of ~10 million aligned reads per sample. The pool was denatured and diluted, resulting in a 2.5 pM DNA solution. PhiX control was spiked at 1%, and the pool was sequenced by 1 × 75 cycles using the NextSeq 500 High Output reagent kit (Illumina Inc.).

RNA-seq analysis

RNA-seq data were first demultiplexed using Bcl2fastq, and RNA-seq reads were aligned to the mouse reference genome (NCBI, mm10) using TopHat and Bowtie aligners. Normalized gene counts were calculated using HT-seq. For the RNA-seq analysis, we focused on the set of highly expressed genes (log2 normalized expression >4). For visualization, GENE-E (https://software.broadinstitute.org/GENE-E/) was used to generate pairwise correlation matrix (Pearson’s correlation) and to perform K-means clustering shown in the heat map. Volcano plots were generated using the log2 fold change of normalized gene counts between control and LysMCreBim+/− cell populations (CD11b+/−MZMs or kidney macrophages) on the x axis and p-values (−log10) on the y axis, p-values were calculated with a two-tailed unpaired t test. The fold change scatterplots in Fig. 8 were constructed using log2 fold change of normalized gene counts in control and LysMCreBim+/− kidney macrophages versus LysMCreBim+/− and TRIF−/−LysMCreBim+/− kidney macrophages. Additional fold change scatterplots were generated in a similar fashion with NZB/W and LN patient data. GO associations and the related p-values were determined by GOrilla (Eden et al., 2009). Lists of differential genes, as in the Venn diagrams, were defined by log2 fold change >1 and P < 0.01. The significance of the enrichment for overlapping genes in the Venn diagram was calculated using a hypergeometric distribution.

Microarray data comparison

Normalized gene expression files for NZB/W kidney macrophages and human LN were obtained through the Gene Expression Omnibus under accession numbers GSE32583 and GSE32591. The mouse genes were converted to the corresponding human orthologues using NCBI homologue (Build 68). The fold-change scatterplots in Fig. 9 (B–D) were constructed as described (see previous paragraph). Differential genes were defined using log2 fold change >1 and P < 0.01, and Venn diagrams were used to compare the differential genes from RNA-seq analysis.

Statistical analysis

The data are presented as mean ± SEM or mean ± SD, and differences between groups were compared using the Mann–Whitney test or Student’s t test with GraphPad Prism 6.0 software.

Online supplemental material

Fig. S1 (related to Fig. 2) shows genomic confirmation of LysMCreBim+/− mice; LysMCreBim+/− mice are 98% on a C57BL/6J background, and the lack of Bim does not affect the balance between total expression levels of proapoptotic Bcl-2 members and antiapoptotic Bcl-2 members. Fig. S2 (related to Fig. 3) shows the alteration of the location of MZB cells in the spleen of LysMCreBim+/− mice. Fig. S3 (related to Fig. 5) shows that the deletion of Bim in lymphocytes is not sufficient to cause SLE-like disease. Fig. S4 (related to Fig. 6) shows Bim-BH3 has an antiinflammatory function. Fig. S5 (related to Fig. 7) shows that Bim-deleted BMDMs are more responsive to TLR agonists. Table S1 (related to Fig. S1 A) lists 150 SNP markers that are tested in the genomic
scan for LysM<sup>Cre<sub>Bim<sup>/fl</sup></sub> mice. Table S2 (related to Fig. 9 A) lists the overlapping genes shared by LysM<sup>Cre<sub>Bim<sup>/fl</sub></sub> kidney macrophages, CD11b<sup>+</sup>MZMs, and CD11b<sup>-</sup>MZMs. Table S3 (related to Fig. 9, E and F) lists the overlapping genes shared by LysM<sup>Cre<sub>Bim<sup>/fl</sub></sub> kidney macrophages and NZB/W kidney macrophages and LysM<sup>Cre<sub>Bim<sup>/fl</sub></sub> kidney macrophages and kidney tissue of LN patients.

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