Irgm1 coordinately regulates autoimmunity and host defense at select mucosal surfaces

Kathleen M. Azzam,1 Jennifer H. Madenspacher,1 Derek W. Cain,2 Lihua Lai,1 Kymberly M. Gowdy,3 Prashant Rai,1 Kyathanahalli Janardhan,4,5 Natasha Clayton,4 Willie Cunningham,4 Heather Jensen,4 Preeyam S. Patel,4 John F. Kearney,6 Gregory A. Taylor,7 and Michael B. Fessler1

1Immunity, Inflammation and Disease Laboratory and 2Signal Transduction Laboratory, National Institute of Environmental Health Sciences, Research Triangle Park, North Carolina, USA. 3Department of Pharmacology and Toxicology, Brody School of Medicine, East Carolina University, Greenville, North Carolina, USA. 4Cellular & Molecular Pathology Branch, National Toxicology Program, National Institute of Environmental Health Sciences, Research Triangle Park, North Carolina, USA. 5Integrated Laboratory Systems, Inc., Research Triangle Park, North Carolina, USA. 6Department of Microbiology, University of Alabama at Birmingham, Birmingham, Alabama, USA. 7Geriatric Research, Education, and Clinical Center, Durham VA Medical Center, Durham, North Carolina, USA.

The pathogenesis of primary Sjogren’s syndrome (SS), an autoimmune disease that targets the mucosa of exocrine tissues, is poorly understood. Although several mouse models have been developed that display features of SS, most of these are within the larger context of a lupus-like presentation. Immunity-related GTPase family M protein 1 (Irgm1) is an interferon-inducible cytoplasmic GTPase that is reported to regulate autophagy and mitochondrial homeostasis. Here, we report that naive Irgm1−/− mice display lymphocytic infiltration of multiple mucosal tissues including the lung in a manner reminiscent of SS, together with IgA class–predominant autoantibodies including anti-Ro and anti-La. This phenotype persists in the germ-free state, but is abolished by deletion of Irgm3. Irgm1−/− mice have increased local production in the lung of TECP15-idiotype IgA, a natural antibody with dual reactivity against host and pneumococcal phosphorylcholine. Associated with this, Irgm1−/− mice display enhanced opsonization and clearance of Streptococcus pneumoniae from the lung and increased survival from pneumococcal pneumonia. Taken together, our results identify Irgm1 as a master regulator of mucosal immunity that dually modulates evolutionarily conserved self- and other-directed immune responses at the interface of host with environment.

Introduction

Primary Sjogren’s syndrome (SS) is an autoimmune disorder characterized by lymphocytic infiltration of exocrine tissues including salivary glands, lacrimal glands, and other mucosal organs such as the lungs and pancreas, in the absence of evidence of another underlying multisystem autoimmune disorder (e.g., systemic lupus erythematosus [SLE]). Among autoimmune diseases, primary SS is second in prevalence only to rheumatoid arthritis (1). Several mouse models have been developed that spontaneously display features of SS (2). While these models have advanced insight into the pathogenesis of SS, fundamental gaps persist in our understanding of the unifying mechanisms that target immune-mediated injury specifically to the epithelium of select mucosal organs.

Approximately 10%–20% of SS patients have clinical pulmonary involvement, but it is speculated that a much larger proportion have subclinical lung disease involving abnormal expansion of bronchus-associated lymphoid tissue (BALT) (3). BALT, a network of peribronchovascular tertiary lymphoid tissue composed of follicular DCs and organized B and T cell zones, is found in some species, but is typically absent in mice and humans except in the setting of infection, chronic lung disease, and autoimmunity (4). BALT-derived antibodies may contribute locally to progression of lung disease and also to host defense against pathogens (5, 6). Identification of the molecular determinants of submucosal lymphoid hyperplasia in the lung and other organs (i.e., mucosa-associated lymphoid tissue) may offer new therapeutic avenues for human disease.
Immunity-related GTPase family M protein 1 (Irgm1) is a member of a family of interferon-inducible (IFN-inducible) cytoplasmic immunity-related GTPases (IRG) encoded by approximately 20 genes in the mouse (7, 8). Irgm1-null mice display dramatic susceptibility to intracellular pathogens that is thought to arise from a combination of deficient cell-autonomous resistance and infection-induced hematopoietic failure (9). Although the mechanism of action remains somewhat obscure, Irgm1 is thought to promote autophagy and mitochondrial homeostasis in part through regulating the organellar localization and activation of other IRGs (10–12). Polymorphisms in the human homolog, IRGM, have been associated with Crohn’s disease (13), suggesting a possible role in restraint of spontaneous mucosal immunity against host and/or microbiota. To date, however, no role has been demonstrated for Irgm1 in regulation of spontaneous autoimmunity.

Here, we report for the first time to our knowledge that naive Irgm1−/− mice display lymphocytic infiltration of multiple mucosal tissues including the lung in a manner reminiscent of SS, together with IgA class–dependent autoantibodies including TEPC15-idiotype (T15-idiotype) IgA, a natural antibody with dual reactivity against host and pneumococcal phosphorylcholine (PC) (14). Associated with this, Irgm1−/− mice display enhanced opsonization and clearance of Streptococcus pneumoniae from the lung. Irgm1 deletion thus reveals coordinate immune targeting of evolutionarily conserved host and pathogen epitopes at the environmental interface. Taken together, our results suggest that Irgm1 is a key regulator of mucosal immunity.

Results

Irgm1−/− mice have spontaneous peribronchovascular B and T cell infiltration. Irgm1−/− mice have defective host defense against several intracellular pathogens (9, 15). Cellular functions that have been identified as Irgm1 dependent including autophagy (13) and migration (16) also govern steady-state immune cell constitution of the lung and other organs. Given this, we examined the lungs of naive Irgm1−/− mice and littermate controls. Remarkably, we found that 8- to 15-week-old Irgm1−/− mice of both sexes housed under specific pathogen–free conditions had multifocal, well-formed lymphocytic aggregates in their lungs in a predominantly peribronchovascular pattern (Figure 1A). Increased lymphocytes were also found in the airway lumen of naive Irgm1−/− animals, as revealed by bronchoalveolar lavage (Supplemental Figure 1; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.91914DS1). Immunohistochemical (IHC) staining revealed the lung parenchymal lesions to be B cell (Pax5+) predominant, with fairly discrete B and T cell (CD3+) zones (Figure 1B), reminiscent of tertiary lymphoid tissue, i.e., BALT (4). IHC verified the presence of both CD4+ and CD8+ T cells in the lesions (Figure 1C). Flow cytometry confirmed increased B cells, CD4+ T cells, and CD8+ T cells in digests of Irgm1−/− lungs (Figure 1D and Supplemental Figure 2), and moreover revealed that there was an increase in the Irgm1−/− lung of both conventional B2 (CD19+CD5-CD11b+IgMlo) cells as well as B1a (CD19+CD5+CD11b+IgMhi) and B1b (CD19+CD5-CD11b+IgMlo) cells (Figure 1E and Supplemental Figure 3), innate-like B cells known to produce polyreactive natural (germline-encoded) antibodies that mediate host defense during acute infection (17).

Lymphocyte infiltration into the lung raised the possibility of a systemic lymphoproliferative disorder. However, consistent with a prior report of defective hematopoietic stem cell function in Irgm1−/− mice (18), we found reduced numbers of lymphocytes (Figure 2, A and B, and Supplemental Figure 4A) as well as erythrocytes and platelets (Supplemental Figure 4B) in the circulation, and deficiencies in B cell (Supplemental Figure 4, C and D) and myeloid (data not shown) lineages in the bone marrow. The peritoneal cavity (PerC) is the primary anatomic site for B1 B cells, from which location these cells are thought to emigrate to the lungs and other organs in response to inflammatory insults (19). Flow cytometric characterization of cell populations in the PerC of naive Irgm1−/− mice revealed a significant reduction in B1a cells, no change in B1b cells, and an increase in B2 cells (Figure 2C). Collectively, these results suggested dysregulation of B cell lineage development and/or trafficking in Irgm1−/− mice.

B cells accumulate in BALT and other tertiary lymphoid organs in response to CXCL13, where local growth factors induce formation of germinal centers, sites of active B cell proliferation, differentiation, and class switching (4, 5). We detected increased CXCL13 in the Irgm1−/− lung that was restricted to the cellular aggregates (Figure 3, A and B). Positive staining was also noted for B cell–activating factor (BAFF), a key B cell prosurvival and differentiation factor (1) (Figure 3, C and D). Upon more extensive profiling, we found that, although some additional cytokines were modestly increased in Irgm1−/− lung homogenates, several cytokines that promote B cell proliferation/differentiation and/or BALT formation, including IL-2, IL-5, IL-6, IL-9, and IL-10 were not increased (Figure 3E). Nonetheless, consistent with active local cell proliferation,
moderate numbers of lymphocytes in the lung lesions displayed positive staining for Ki-67 (Figure 3F). In addition, positive staining was noted with peanut agglutinin, a marker of activated germinal center B cells (Figure 3G), and an increased frequency and number of CD4^+^CD8^+^ T cells (CD4^+^CD8^+^ CD45^+^B220^+^GL7^+^CD38^–^) was also noted in Irgm1^{–/–} lung digests by flow cytometry (Figure 3H and Supplemental Figure 5).

Irgm1 deletion induces lymphocytic infiltration of multiple exocrine tissues. Given the striking pulmonary phenotype, which suggested increased trafficking and tissue persistence of B cells, we extended our histopathologic screen of naive Irgm1^{–/–} mice to more than 40 tissues (Supplemental Table 1). Lymphocytic infiltration was also noted in the submandibular and parotid glands, extraocular lacrimal glands, and pancreas of both sexes, with IHC confirming B and T cells in all cases (Figure 4, A–F). Associated acinar atrophy of the submandibular glands was present. In the pancreas, extensive exocrine pancreatic atrophy with adipocyte replacement and scattered infiltration of lymphocytes was present (Figure 4E). By contrast, the islets and ducts were not affected. Consistent with functional compromise of the exocrine acini of the pancreas and/or salivary glands, serum amylase was reduced in Irgm1^{–/–} mice (Supplemental Figure 6A). All Irgm1^{–/–} mice also exhibited fur loss around the eyelid margin of at least one eye (Supplemental Figure 6B), suggestive of lacrimal dysfunction.

A subset of 8- to 15-week-old (7 of 12 males, 1 of 4 females) Irgm1^{–/–} mice also displayed mild chronic active inflammation in the liver. No other tissues, however, displayed any notable pathology. Examination of 9- to 14-month-old Irgm1^{–/–} mice of both sexes revealed marked progression of the aforementioned lung
lesions with age (Supplemental Figure 7), but no involvement of additional tissues beyond those noted in younger mice. Collectively, this selective pattern of exocrine/mucosal organ involvement in the absence of disease in organs typically involved in SLE and other autoimmune disorders (i.e., kidneys, skin, lymph nodes, central nervous system, joints) is reminiscent of primary SS in humans.

**Tissue lesions of Irgm1−/− mice persist in the germ-free state.** Given that Irgm1−/− mice have compromised immunity and BALT can be induced by infection (4, 5), we performed an extensive screen for bacterial, viral, and fungal infection. However, no pathogens were detected by serology (Supplemental Table 2), nor was *Pneumocystis murina* detected in the Irgm1−/− lung by PCR (data not shown). Moreover, similar lesions were found in the lungs, salivary glands, and pancreas of germ-free Irgm1−/− mice (Supplemental Figure 8), confirming that the SS-like tissue phenotype was autonomous from microbiota.

**Tissue lesions of Irgm1−/− mice are abolished by Irgm3 deletion.** The alternate IRG subfamily M member, Irgm3, has been proposed to have complex functional cooperativity with Irgm1 in which it coregulates the organellar deposition and activation of other IRG proteins (12). Several of the phenotypes of Irgm1−/− mice reportedly revert upon deletion of Irgm3 (i.e., in Irgm1−/− Irgm3−/− mice), including defective host defense, macrophage motility, autophagy, and stress hematopoiesis (8, 10). It has been proposed that Irgm3 deletion relieves the cytopathic effects of Irgm1 deletion by promoting intracellular clearance/relocalization of aggregated IRG proteins away from lysosomes (10, 12). Remarkably, we found that lymphocytic infiltration in lung, salivary glands, lacrimal glands, and pancreas of germ-free Irgm1−/− mice (Supplemental Figure 8), confirming that the SS-like tissue phenotype was autonomous from microbiota.

**Irgm1 deletion induces spontaneous IgA-predominant autoimmunity.** The lymphocytic infiltration of several exocrine tissues in Irgm1−/− mice was reminiscent of human autoimmune disease, SS in particular. Given this, we tested Irgm1−/− serum for the presence of anti-nuclear antibodies (ANAs) using HEP-2 cells as substrate, a classical indirect immunofluorescence procedure used in clinical medicine (and mouse models) to screen for autoantibodies targeting nuclear antigens (20). Of interest, although IgG class ANAs were not detected in Irgm1−/− (or WT) serum, an abnormal increase in IgA ANA was evident, yielding nuclear staining in a coarse speckled pattern (Figure 5A).

In patients, a coarse speckled pattern of ANA is specific for autoimmune disease (21). Given this, we used ELISA to quantify specific nucleus-targeting autoantibodies, testing serum from Irgm1−/− mice of 2 ages (8–12 weeks, 1 year) (Figure 5, B and C). Anti–double stranded DNA (anti-dsDNA) is a commonly used confirmatory autoantibody in human autoimmune disease. Anti-Ro and anti-La, ribonucleoprotein-specific autoantibodies commonly found in SS, have been proposed to arise from autoreactive targeting of mucosal epithelia (1, 22). Consistent with the ANA finding, anti-dsDNA IgA was dramatically increased in the serum of Irgm1−/− mice of both ages. By contrast, anti-dsDNA IgG was modestly increased only in young Irgm1−/− mice and anti-dsDNA IgM was not elevated. Sexual dimorphism was evident for IgA levels of anti-Ro (both Ro-52/tripartite motif–containing protein 21 [Trim21] and Ro-60 antigens) and anti-La, with elevation only in female Irgm1−/− mice, and a dramatic age-dependent further increase in titer (Figure 5C). Anti-Ro IgG and anti-La IgG levels were normal (data not shown).

Autoimmune disorders commonly exhibit systemic induction of cytokines. In SS, excess type I IFNs induce autoantigens and enhance B cell function and IgA class switching via the IFN-stimulated gene (ISG) BAFF, leading to a vicious cycle (1, 22). A wide range of cytokines including BAFF and CXCL13

---

**Figure 2. Dysregulated lymphocyte populations in Irgm1−/− mice.** (A) Counts of circulating total leukocytes (WBC) and leucocyte subtypes were enumerated in naive Irgm1−/− mice and controls (n = 10/genotype). (B) Circulating B cells (CD3−B220+) and T cells (CD3+B220−) in naive Irgm1−/− mice and controls (n = 4/genotype) were quantified by flow cytometry. (C) Peritoneal B1a (CD19+CD5–CD11b+IgM−/−CD23+), B1b (CD19+CD5+CD11b+IgM+CD23+), and B2 (CD19+CD5+CD11b–IgM–CD23+) cells in naive Irgm1−/− mice and controls (n = 7/genotype) were quantified by flow cytometry. Data are the mean ± SEM and are representative of at least 2–3 independent experiments.

*P* < 0.05, **P** < 0.01, ***P*** < 0.001 by unpaired 2-tailed Student’s *t* test. ND, not detected; PMN, neutrophil; Lymph, lymphocyte; Mono, monocyte; EO, eosinophil; Baso, basophil.
were elevated in the serum of naive 8- to 12-week-old Irgm1 –/– mice (Figure 5, D and E). BAFF plus several additional ISGs, including Irf7, Eif2ak2, and Mx2 were also upregulated in naive Irgm1 –/– lung tissue (Figure 3D and Figure 6A). Suggesting increased local IFN effect specifically in pulmonary epithelial cells, we found increased expression of Irf7, Eif2ak, and Mx2 in the lungs of naive Irgm1 –/– mice were stained by IHC for Ki-67 (original magnification, ×20) and (G) peanut agglutinin (PNA; original magnification, ×40). (H) CD45+ B220+ GL7+ cell counts were quantified in lung digests of naive Irgm1 –/– mice and controls (n = 4-5/genotype). Data are the mean ± SEM and are representative of at least 3 independent experiments. IHC images are representative of at least 3–4 mice. *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001 by unpaired 2-tailed Student’s t test.

Figure 3. Lung lesions of Irgm1 –/– mice display active germinal centers. (A–D) Lungs of naive Irgm1 –/– mice and controls were evaluated by IHC and ELISA for CXCL13 (A and B) and B cell–activating factor (BAFF) (C and D). Original magnification for panels A and C, ×20. For ELISA, 4–8 mice/genotype were used. (E) Cytokines were quantified in lung homogenates from naive Irgm1 –/– mice and controls (n = 7/genotype) using multiplex technology. (F and G) Lung lesions of Irgm1 –/– mice were stained by IHC for (F) Ki-67 (original magnification, ×20) and (G) peanut agglutinin (PNA; original magnification, ×40). (H) CD45+ B220+ GL7+ cell counts were quantified in lung digests of naive Irgm1 –/– mice and controls (n = 4-5/genotype). Data are the mean ± SEM and are representative of at least 3 independent experiments. IHC images are representative of at least 3–4 mice. *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001 by unpaired 2-tailed Student’s t test.
Irgm1-null lymphocytic lung lesions produce antibody locally. Human autoimmune disorders including SS commonly exhibit polyclonal hyperglobulinemia. Given the reduction in circulating and bone marrow B cells in the face of increased B cells in lung and other tissues of Irgm1−/− mice, it was of interest to define systemic and tissue antibody levels. Irgm1−/− mice 8–12 weeks old had a marked increase in serum total IgG

Figure 4. Lymphocytic infiltration of multiple exocrine tissues in Irgm1−/− mice. (A and B) Submandibular salivary glands from naive Irgm1+/+ mice and controls were stained with hematoxylin and eosin (H&E; original magnification, ×1.4) (A), and lymphocytic lesions detected in Irgm1−/− tissue (shown by arrows in A) were then IHC stained for T cells (CD3) and B cells (Pax5) (original magnification, ×40) (B). (C and D) Extraorbital lacrimal glands from naive Irgm1−/− mice and controls were stained with H&E (original magnification, ×4) (C), and lymphocytic lesions detected in Irgm1−/− tissue were then IHC stained for T cells (CD3) and B cells (Pax5) (original magnification, ×20) (D). (E and F) Pancreas from naive Irgm1−/− mice and controls was stained with H&E (original magnification, ×4) (E). Arrows indicate ducts and islets, which are unaffected; asterisks indicate surrounding adipose replacement of exocrine regions. Rare lymphocytic infiltrates in Irgm1−/− tissue were IHC stained for T cells (CD3) and B cells (Pax5) (original magnification, ×20) (F). (G–J) Lungs (G), submandibular salivary glands (H), extraorbital lacrimal glands (I), and pancreas (J) were evaluated by H&E stain in naive Irgm1+/+ Irgm3−/− mice. Original magnification, ×0.9 for G and J, ×1.4 for H, and ×4 for I. Tissues in all cases are representative of at least 3–4 mice per genotype.
Increased IgG and, in particular, marked elevation of IgA, was also noted in BALF of naive Irgm1–/– mice (Figure 7B), whereas the concentration of albumin, a serum protein, was not increased (data not shown). This, together with past reports that airspace IgA derives from local synthesis (24), suggested that the increase in airway immunoglobulin was likely arising, at least in part, from increased local antibody production in the lung. Dimeric IgA made by submucosal B cells is secreted across mucosal epithelia in complex with polymeric immunoglobulin receptor (pIgR) after the latter is proteolytically cleaved into so-called secretory component (SC) (25). Of interest, we found elevated expression of pIgR in Irgm1–/– lungs (Figure 7C). Consistent with increased local release of secretory IgA into the airspace, we found elevated SC in Irgm1–/– BALF (Figure 7D), as well as increased high molecular weight (>250 kDa) IgA signal on nonreducing immunoblot of Irgm1–/– BALF (Figure 7E).

IHC of the Irgm1–/– lung confirmed that B cells in the center of the follicular lesions were predominantly IgM+, but that IgG+ and IgA+ cells were also present (Figure 7F), suggesting that class switch recombination was occurring locally. In order to quantify local antibody release from B cells in various tissues, ELISPOT was performed (Figure 7, G and H). In bone marrow and spleen of Irgm1–/– mice, no change in frequency and IgA, but not IgM (Figure 7A). Increased IgG and, in particular, marked elevation of IgA, was also noted in BALF of naive Irgm1–/– mice (Figure 7B), whereas the concentration of albumin, a serum protein, was not increased (data not shown). This, together with past reports that airspace IgA derives from local synthesis (24), suggested that the increase in airway immunoglobulin was likely arising, at least in part, from increased local antibody production in the lung. Dimeric IgA made by submucosal B cells is secreted across mucosal epithelia in complex with polymeric immunoglobulin receptor (pIgR) after the latter is proteolytically cleaved into so-called secretory component (SC) (25). Of interest, we found elevated expression of pIgR in Irgm1–/– lungs (Figure 7C). Consistent with increased local release of secretory IgA into the airspace, we found elevated SC in Irgm1–/– BALF (Figure 7D), as well as increased high molecular weight (>250 kDa) IgA signal on nonreducing immunoblot of Irgm1–/– BALF (Figure 7E).

IHC of the Irgm1–/– lung confirmed that B cells in the center of the follicular lesions were predominantly IgM+, but that IgG+ and IgA+ cells were also present (Figure 7F), suggesting that class switch recombination was occurring locally. In order to quantify local antibody release from B cells in various tissues, ELISPOT was performed (Figure 7, G and H). In bone marrow and spleen of Irgm1–/– mice, no change in frequency and IgA, but not IgM (Figure 7A). Increased IgG and, in particular, marked elevation of IgA, was also noted in BALF of naive Irgm1–/– mice (Figure 7B), whereas the concentration of albumin, a serum protein, was not increased (data not shown). This, together with past reports that airspace IgA derives from local synthesis (24), suggested that the increase in airway immunoglobulin was likely arising, at least in part, from increased local antibody production in the lung. Dimeric IgA made by submucosal B cells is secreted across mucosal epithelia in complex with polymeric immunoglobulin receptor (pIgR) after the latter is proteolytically cleaved into so-called secretory component (SC) (25). Of interest, we found elevated expression of pIgR in Irgm1–/– lungs (Figure 7C). Consistent with increased local release of secretory IgA into the airspace, we found elevated SC in Irgm1–/– BALF (Figure 7D), as well as increased high molecular weight (>250 kDa) IgA signal on nonreducing immunoblot of Irgm1–/– BALF (Figure 7E).

IHC of the Irgm1–/– lung confirmed that B cells in the center of the follicular lesions were predominantly IgM+, but that IgG+ and IgA+ cells were also present (Figure 7F), suggesting that class switch recombination was occurring locally. In order to quantify local antibody release from B cells in various tissues, ELISPOT was performed (Figure 7, G and H). In bone marrow and spleen of Irgm1–/– mice, no change in frequency...
lipid moieties common to host and pathogens.

sites, with increased local production of IgA-predominant antibodies targeting evolutionarily conserved

deletion confers a coordinate disinhibition of autoimmunity and innate immunity at selected mucosal

Irgm1 deletion augments successful pathogen clearance. Taken together, these findings suggest that Irgm1 deletion

or absolute number of IgG antibody-secreting cells (ASCs) was detected compared with WT counterparts. By contrast, although not statistically significant, a reciprocal decrease in IgM and increase in IgA ASCs was detected in both tissues, suggesting increased IgA class switching. In the lung, ASCs of all 3 classes were markedly increased, suggesting that the lung, and almost certainly the B cell–rich foci specifically, is a site of robust antibody production in Irgm1-deleted animals. Consistent with increased local antibody production, total spot area (i.e., integrated Ig+ area surrounding all ASCs) on ELISPOT was markedly increased for all 3 antibody classes in lung tissue (Supplemental Figure 10). In addition, ELISA of saline-perfused Irgm1−/− lung tissue confirmed increased IgM, IgG, and IgA, with most marked elevation in the latter (Figure 7I).

Irgm1-null lung overproduces T15-idiotype anti-PC IgA. Natural antibodies are germline-encoded, innate (i.e., spontaneous), polyreactive immunoglobulins, largely produced by B1a B cells, that are thought to have been evolutionarily selected in response to important antigens (26). Among natural antibodies are the well-studied T15-idiotype anti-PC autoantibodies, which are reactive against oxidized phospholipid (i.e., oxidized 1-palmitoyl-2-arachidonoyl-sn-phosphatidylcholine) in apoptotic cells, as well as against PC exposed on the surface of S. pneumoniae (14). Given that B1a cells and anti-PC play complex roles in autoimmunity (27, 28) and accumulate in the lung in chronic inflammatory conditions (29), we surveyed for dysregulated levels of anti-PC immunoglobulins in the lung and other tissues of Irgm1−/− mice.

Irgm1−/− mice had a reciprocal decrease in PC-specific IgM and increase in PC-specific IgA in the serum, whereas PC-specific IgG levels were equivalent to WT (Figure 8A), together suggesting increased IgA class switching, perhaps due to increased anti-PC production at a mucosal site. ELISPOT revealed an increase in frequency and number of anti-PC IgA ASCs in Irgm1−/− lung tissue (Figure 8B). This was associated with increased PC-specific IgA in BALF of Irgm1−/− mice (Figure 8C). An increase in T15-idiotype anti-PC IgA was also detected in the BALF, lung tissue, and serum of Irgm1−/− mice (Figure 8D). TC68 and ABI-2 are anti–T15 idiotype antibodies; the former detects the VhS107.1 region of the T15 idiotype, and the latter, the CDR3 region (30). Consistent with increased local production of T15 IgA in the Irgm1−/− lung, we detected increased frequency and numbers of B220−PC-BSA−TC68−ABI-2− cells in lungs of Irgm1−/− mice (Figure 8E and Supplemental Figure 11). These findings suggest that, even in the face of reduced circulating and bone marrow B cells, Irgm1 deletion favors trafficking of B1a cells to the lung, at which site local conditions including BAFF overexpression may favor both enhanced survival and IgA class switching of T15-idiotype cells.

Augmented innate Pneumococcal host defense in the Irgm1-null lung. Exogenous treatment with T15-idiotype antibody improves survival in mice infected by S. pneumoniae, likely due to enhanced PC-targeted binding (14). Given this, we speculated that the local increase in native airway T15-idiotype antibody in Irgm1−/− mice might be associated with augmented natural host defense. To test this, Irgm1−/− mice and controls were infected intratracheally with PKH26-labeled S. pneumoniae. Analysis of BALF collected 30 minutes after infection revealed increased IgM and IgA opsonization of S. pneumoniae in the Irgm1−/− airway (i.e., an increased percentage of IgM+ and IgA+ PKH26+ bacteria) (Figure 9, A and B). This augmented recognition by natural antibody was associated with enhanced clearance of S. pneumoniae from the lung 48 hours after infection (Figure 9C), and also with improved survival of Irgm1−/− mice following S. pneumoniae infection (Figure 9D). With S. pneumoniae, we thus extend the purview of Irgm1 in host defense to extracellular bacteria, showing in this case, contrary to that of intracellular pathogens, that Irgm1 deletion augments successful pathogen clearance. Taken together, these findings suggest that Irgm1 deletion confers a coordinate disinhibition of autoimmunity and innate immunity at selected mucosal sites, with increased local production of IgA-predominant antibodies targeting evolutionarily conserved lipid moieties common to host and pathogens.

Figure 6. Spontaneous induction of interferon-stimulated genes in the Irgm1+/− lung. mRNA expression (normalized to GAPDH) of the indicated targets was quantified in whole lung (A), pulmonary epithelial (CD45 CD31 CD34 EpCAM+) cells sorted by FACS (B), and pulmonary hematopoietic (CD45+) cells purified by column (C) from naive Irgm1+/+ and Irgm1−/− mice (n = 3–7/genotype). Data are the mean ± SEM and are representative of at least 3 independent experiments. *P < 0.05, **P < 0.01, ***P < 0.001 by unpaired 2-tailed Student’s t test.
Discussion

Primary SS is the second most common autoimmune disease, with an estimated prevalence of 0.9–6 per 1,000 individuals, involving a 9:1 female predominance (1). Roles have been identified for IFNs, CXCL13, BAFF, and B cells, all of which are proposed to impinge upon epithelial cells, leading to a vicious cycle of autoantibody generation (1). Autoantigens such as Ro externalized on apoptotic cell blebs may trigger autoantibody production and germinal center formation by infiltrating B cells at mucosal sites (22). While mouse models have been developed that recapitulate features of SS, several of these including (NZB/NZW)F1, MRL/lpr, BAFF-transgenic, IL-12–transgenic, IL-14–transgenic, and TGF-β1−/− mice are dominated by multiorgan involvement in an SLE-like pattern (i.e., glomerulonephritis, skin disease, lymphadenopathy), while others are confounded by comorbidities such as diabetes mellitus (2). Thus, many questions remain about the disease mechanisms specific to SS pathogenesis, and there is a need for additional, well-defined murine models that faithfully mimic the human disorder.
Here, we report for the first time to our knowledge that Irgm1–/– mice spontaneously display several features reminiscent of SS by approximately 8 weeks of age, including lymphocytic infiltration of lacrimal and salivary glands, hyperglobulinemia, and elevated BAFF, ANA, and anti-Ro and -La antibodies. We found some evidence for gender dimorphism, in particular, elevated anti-Ro and -La autoantibodies in females. Associated with these hallmark tissue lesions was selective atrophy of the exocrine pancreas consistent with autoimmune pancreatitis, marked expansion of peribroncholar lymphoid tissue, and, in a subset of mice, lesions potentially consistent with autoimmune hepatitis. All of these are tissue complications that have been associated with human SS (22, 31). In addition to representing a novel model of SS with extraglandular manifestations, the Irgm1–/– mouse may thus also facilitate new molecular insights into BALT and autoimmune pancreatitis.

Of interest, the Irgm1–/– mouse exhibits increased serum IgG and IgA, but normal IgM. Among antibody-producing tissues surveyed, bone marrow and spleen of the Irgm1–/– mouse had normal/reduced IgM/IgG ASCs but increased IgA ASCs, whereas lung exhibited a dramatic increase in ASCs for all 3 isotypes. Collectively, these findings suggest increased IgA class switching in Irgm1–/– tissues, and also the interesting possibility that the prominent lung lesions may serve as a significant source of systemic immunoglobulin in the Irgm1–/– mouse. Increased autoantibody levels in the Irgm1–/– mouse were restricted to the IgA class, perhaps pointing to an affected mucosal site as a source of origin. Consistent with this, we confirmed the lung as a source of T15-idiotype anti-PC IgA, and identified increased pulmonary B1a cells as a likely source (26). Whereas several prior publications have identified macrophage and T cell abnormalities in the setting of Irgm1 deletion (9, 15, 16), we are unaware of past reports of B cell dysregulation in Irgm1–/– mice.

B1a cells have previously been implicated in autoimmunity (28), in which context they may be attracted to affected target organs by CXCL13 (32), a chemokine we found to be elevated in the follicular lesions of the Irgm1–/– lung (Figure 3A), as well as in the salivary and lacrimal gland lesions of Irgm1–/– mice (data not shown). CXCL13 is reportedly required for B1 cell production of T15 anti-PC antibodies (33). B1a cells have been identified as major sources of IgA in vivo (34), and also shown to class switch to IgA preferentially, especially in response to BAFF (19), which we also found to be increased in Irgm1–/– lung lesions. Aberrant expression of select cytokines may thus underlie abnormal homing of B1a cells to the Irgm1–/– lung as well as programming of B1a cells at that site to release IgA class natural antibodies. Suggesting human relevance, BAFF and CXCL13 are also elevated in human SS and 80% or more of primary SS patients have elevated anti-Ro and -La IgA (1, 35, 36).

Irgm1 is an IFN-stimulated p47 GTPase that has been proposed to support host defense against intracellular pathogens by promoting autophagy and mitochondrial homeostasis (7, 8, 10). It is widely expressed in murine immune cells, including B1 and B2 cells, CD4+ T cells, CD8+ T cells, conventional B cells, and natural killer cells (7, 8). Irgm1 is also expressed in non-immune cells such as epithelial cells, astrocytes, and neurons, raising the possibility that Irgm1 may represent a conserved molecule involved in a variety of cell types that may be important for the regulation of immune responses.

Figure 8. Production of T15-idiotype anti-phosphorylcholine IgA in the Irgm1–/– lung. (A) Anti-phosphorylcholine (anti-PC) IgM, IgA, and IgG was quantified by ELISA in serum of naive Irgm1–/– mice and controls. (B) Anti-PC IgA antibody-secreting cells (ASCs) were quantified in perfused lung of naive Irgm1–/– mice and controls. Data were verified by BSA negative control run in parallel and background subtracted. (C) Anti-PC IgA was quantified by ELISA in bronchoalveolar lavage fluid (BALF) of naive Irgm1–/– mice and controls. (D) T15-idiotype IgA was quantified by ELISA in BALF, lung, and serum of naive Irgm1–/– mice and controls. (E) T15-idiotype B cells were quantified by flow cytometry in lung of naive Irgm1–/– mice and controls. Data are the mean ± SEM and are representative of at least 3 independent experiments involving n = 4–6/genotype. *P < 0.05, **P < 0.01, ***P < 0.001 by unpaired 2-tailed Student’s t test.
Irgm1−/− mice (43). Given that elevated type I IFNs reportedly drive increased epithelial cell death and turnover in the pancreas and salivary gland of Irgm1−/− mice (43), we further speculate that IFN overproduction may target autoimmune disease (37). However, experimental autoimmune encephalomyelitis and similar artificial models, involving vaccination and acute challenge with autoantigens, almost certainly diverge from the pathogenesis of native autoimmune disease.

Irgm1−/− leukocytes reportedly display enhanced TLR signaling through the adaptor protein myeloid differentiation program protein 88 (MyD88) (38). In other settings, MyD88 has been shown to support TLR7-dependent induction of type I IFNs by pDCs (39), BAFF production by and signaling in B cells (40, 41), as well as several B cell functions including autoantibody production (42). Hyperactivated MyD88-dependent signaling in DCs is sufficient to cause autoimmune disease in mice (20). Taking these results together, we thus propose that enhanced MyD88-dependent signaling in select interacting cell types of the Irgm1−/− mouse may promote an axis of IFN overproduction, downstream BAFF induction, and BAFF hypersensitivity, leading to autoimmunity. Using ELISA, we found only very low levels of IFN-α in BALF of Irgm1−/− (and Irgm1+/+) mice that were near the limit of detection (data not shown); however, a recent report using a highly sensitive viral suppression bioassay did confirm elevated type I IFNs in serum of Irgm1−/− mice (43). Given that elevated type I IFNs reportedly drive increased epithelial cell death and turnover in the pancreas and salivary gland of Irgm1−/− mice (43), we further speculate that IFN overproduction may target autoreactive B cells to the Irgm1−/− epithelium. Interestingly, MyD88 may possibly contribute to autoimmune targeting of Irgm1−/− mucosa given that it reportedly underlies autoimmune targeting of the lung and other mucosal surfaces in Foxp3-deficient mice (44).

As has been reported for the host defense defects of the Irgm1−/− mouse (12), we found that the SS-like phenotype was abolished by Irgm3 deletion. Irgm3 deletion has been proposed to rescue cell-autonomous defects in Irgm1-null cells by restoring dysregulated subcellular localization and activation of IRG proteins (12). Our results thus suggest that Irgm1-disinhibited IRGs in one or more interacting cell types may be the proximal cause of autoimmune pathogenesis. This could arise, for example, from damage-associated molecular patterns released by Irgm1-deficient cells inducing IFN production in pDCs or other cells through TLR/MyD88 signaling. Identification of the Irgm1-deficient cell types that drive the in vivo SS-like phenotype will have to await generation of tissue conditional Irgm1−/− mice.

Irgm1−/− cells have deficient autophagic flux (11). However, mice with deficient canonical autophagy do not display spontaneous autoimmune disease (45). Moreover, B cell autophagy is required for the SLE-like phenotypes of TLR7-transgenic mice and Lpr mice (46, 47), as well as for differentiation and survival of plasma cells (48). There is evidence for increased autophagic flux in T cells in human SLE and murine SLE models (49). We thus speculate that deficient autophagy is unlikely to be the underlying mechanism of spontaneous autoimmunity in Irgm1−/− mice, and may possibly mitigate against it.
Future studies will be required to better define the mechanisms underlying the enhanced anti-Pneumococcal host defense of the Irgm1–/– lung. The degree to which locally produced secretory IgA versus IgM or other antibody isotypes contribute is uncertain. T cells and other cells of the local BALT-like aggregates may also collaborate in the immune response. Finally, the possibility also remains that altered recruitment of circulating immune cells (e.g., neutrophils) to the infected Irgm1–/– airspace may contribute to enhanced bacterial clearance. Detailed immune profiling of the infected lung, as well as cell-specific Irgm1 deletion may ultimately be required to properly dissect the role(s) of Irgm1 in this context.

Taken together, our studies identify Irgm1 as a unifying regulator of self- and other-directed immune responses at selected mucosal sites. Phosphatidylcholine is the predominant phospholipid of pulmonary surfactant, in which location it has remarkably high concentrations not found elsewhere in vivo (50). It is intriguing to consider that this PC-headgroup phospholipid, perhaps oxidized by the inflammatory milieu and surfactant, in which location it has remarkably high concentrations not found elsewhere in vivo (50). It is intriguing to consider that this PC-headgroup phospholipid, perhaps oxidized by the inflammatory milieu of the Irgm1–/– lung, may serve as the inciting autoantigen for local expansion of T15-idiotype B cells in the lung. We propose that IRGM polymorphisms should be examined for their possible association with BALT, B cell homeostasis, and autoimmune disease in humans, and also that investigation of IRGM as a potential therapeutic target in autoimmunity and pulmonary host defense is warranted.

Methods
Mice. Irgm1-deficient mice have been described (9) and were crossed more than 8 generations onto the C57BL/6 background. Age- and gender-matched WT littermate controls were used. Germ-free mice were housed in sterile flexible film Trexler isolators with sterilized bedding and sterilized chow and water (ad libitum), within the National Gnotobiotic Rodent Resource Center, UNC/Chapel Hill. Sterility was confirmed on a monthly basis by fecal Gram stain, aerobic and anaerobic cultures, and 16S rRNA PCR analysis of feces and bedding. Irgm1–/– Irgm3–/– mice were obtained from G.A. Taylor.

Flow cytometry. Lung tissues were perfused, digested, and processed to single-cell suspension as previously reported (51). Cells were blocked with anti–mouse CD16/32 (Biolegend, 101302) and stained for 30 minutes with the following anti-mouse antibodies for individual plots: CD45-eFluor 450 (eBiosciences, 48-0451-82); CD3e PE Cy7 (Biolegend, 100320); CD19-FITC (Biolegend, 115507); CD45.2-PE (Biolegend, 109808); CD4-FITC (eBiosciences, 11-0041-81); CD8a APC (Biolegend, 100711); CD45R/B220-eFluor 450 (eBiosciences, 48-0452-82); CD5 PE (Biolegend, 100608); CD11b V500 (BD Horizon, 562128); IgM APC (Biolegend, 406515); CD43 PerCP-Cy5.5 clone S7 (BD Biosciences, 562865); CD3-AF488 (Biolegend, 100321); IgM-PE (Biolegend, 406508); CD43 clone 1B11 APC/Cy7 (Biolegend, 121220); GL7-FITC (Biolegend, 144603); CD38 PE (Biolegend, 250505); IgA-FITC (Southern Biotech, 1165-02); and IgM AF647 (Life Technologies, A21238). S107 (PC-specific IgA) cells, provided by the Kearney laboratory, were used as a positive control. PC-BSA AF488 was generated with the Alexa Fluor 488 Monoclonal Antibody Labeling kit (Invitrogen). Purified monoclonal antibodies for TC68 Pacific Blue and AB1-2 AF555 were provided by the Kearney laboratory. PC-BSA AF488 was generated with the Alexa Fluor 488 Monoclonal Antibody Labeling kit (Invitrogen). CD45R/B220 PerCP was from BD Biosciences (561086). All flow cytometric analyses were performed on the LSR II (BD Biosciences) and analyzed using FACSDiva (BD Biosciences) and FlowJo software (Tree Star).

Clinical blood assays. Blood cell counts were analyzed using the HEMAVET 1700 hematology analyzer (Drew Scientific, Inc.). Manual leukocyte differential counts were performed, and smear estimates were used to confirm values. Serum amylase was measured with an Olympus AU400e (Beckman Coulter) using reagents from the manufacturer.

Immunoglobulin and albumin ELISAs. Mouse IgA, IgM, IgG, and albumin ELISA quantitation sets (Bethyl Laboratories) were used with the following dilutions: serum 1:10,000 (IgA, IgM) or 1:50,000 (IgG); tissue ELISAs neat or 1:4; BALF (immunoglobulins) 1:2; and BALF (albumin) 1:1,000.

ANA assay. An indirect fluorescent antibody assay utilizing HEp-2 cells as a substrate (MBL Bion) following the manufacturer’s specifications used mouse serum at an initial concentration of 1:40 for screening and then diluted up to 1:640 for endpoint analysis. FITC-conjugated goat anti-mouse IgG (γ-chain specific), IgM (μ-chain specific), or IgA (α-chain specific) (Sigma-Aldrich) was added to wells at 1:200 in PBS. For MetaMorph (Molecular Devices) analysis, negative controls (PBS) were used to determine a threshold of 35 and most of the data had intensities of 38–39.

Autoantibody ELISAs. Mouse anti-SSB (La), anti-SSA (Ro-52), and anti-SSA (Ro-60) ELISAs (Signosis) were used per manufacturer’s specifications for IgG detection, and utilizing HRP-conjugated IgA (Bethyl...
Lectin Laboratories) at a dilution of 1:1,000. Serum samples were diluted 1:50. For anti-dsDNA ELISA, calf thymus DNA (Sigma-Aldrich) stock was resuspended at 2 mg/ml in 1× SSC buffer overnight at 4°C, sheared the next day by running through an 18-gauge needle, and then diluted to 1 μg/ml in carbonate buffer for coating of plates overnight at 4°C. The next day, wells were washed with 0.5× PBS/0.5% BSA, and blocked with PBS/3% BSA (room temperature, 2 hours). Serum samples (1:50 dilution) and anti-dsDNA antibody (Abcam, clone HYB331-01) used for standard for IgG isotype were diluted using 0.5× PBS/0.5% BSA buffer, and plates were incubated at 37°C for 1 hour, and then 4°C for 1 hour. After washing, HRP-conjugated anti-mouse IgG, IgA, or IgM (Bethyl Laboratories) was added to the plate at 1:5,000 dilution in 0.5× PBS/0.5% BSA and wells were incubated at room temperature for 1 hour. Wells were then washed with 0.5× PBS/0.5% BSA, developed with TMB substrate (Biolegend), stopped with 2N H2SO4, and read at OD450 (Bio-Tek Synergy 2 microplate reader).

**Quantitative RT-PCR.** Total RNA was extracted from cells and tissues with RNeasy kits (QIAGEN) using the manufacturer’s protocol. For tissue-extracted RNA, tissues were homogenized with a TissueLyser (Qiagen). Predeveloped, validated primer/probe sets were used (Applied Biosystems). Reactions were run in duplicate on an ABI Prism 7900HT with thermal cycling parameters specific for 1-step RT-PCR. The efficiency (slopes) of the target amplification and the efficiency of the reference endogenous control (GAPDH) amplification were 100% (±10%). For sorted lung epithelial cells, the RNAqueous-Micro Total RNA Isolation kit (Thermo Fisher Scientific) was used with elution in a 20-μl volume; 500 ng RNA was converted to cDNA.

**Mouse exposures and harvests.** Mice were intratracheally administered *S. pneumoniae*, serotype 3 (5 × 10^4 to 2 × 10^6 CFU, or 5 × 10^5 CFU for survival studies) (ATCC). Lung was homogenized in PBS, and serial dilutions plated on blood agar for bacterial quantification. BALF was spun at 300 g for 6 minutes. Erythrocytes were lysed in hypotonic buffer on ice for 1 minute. Cells were then resuspended in PBS and counted. Cytospins were stained and differentials counted by light microscopy.

**Cell sorting.** Lungs were perfused, inflated, and digested with elastase (Worthington Biochemicals) for 45 minutes at 37°C. Minced lung was then digested with DNase (Sigma-Aldrich) for 15 minutes. Cells were strained (70-μm filter), deplated of CD45+ cells using AutoMACS column (Miltenyi Biotec), and then blocked with anti–mouse CD16/CD32 (2.4G2) and normal mouse and rat serum (Jackson Immunoresearch). Cells were incubated with one or more of the following antibodies/markers: CD34 (clone MEC14.7), CD45 (clone 104), EpCAM/CD326 (clone G8.8), CD31 (clone 390), CD103 (clone 2E7), 7-AAD (all Biolegend). Cells were then sorted (7-AAD CD31 CD34 CD45 EpCAM+) on an ARIA-II flow cytometer (BD Biosciences) with purity greater than 92%–97%.

**Cytokine analysis.** Cytokines were quantified by multiplex assay (Bio-Plex; Bio-Rad) or ELISA (eBioscience). BAFF (MBLYSO) and CXCL13 (MCX130) ELISAs (R&D Systems) were used as directed. Lung homogenate was processed using the Bioplex Cell Lysis kit (Bio-Rad), and equivalent protein was used for each sample, as quantified by BCA.

**BCA protein assay.** Manufacturer’s instructions (Pierce) were followed, with colorimetric reading at OD562 on a Bio-Tek Synergy 2 microplate reader.

**Immunoblotting.** Saline-perfused lungs were homogenized in 0.1% Triton X-100 supplemented with protease inhibitors and then clarified by centrifugation. An equal protein mass of lung or BALF (first concentrated using a 3-kDa MW cutoff [Amicon Ultra-0.5 centrifugal filter unit with ultracel-3 membrane; MilliporeSigma]) was subjected to SDS-polyacrylamide gel electrophoresis and nitrocellulose membrane transfer as previously reported (51). For BALF IgA immunoblot, reducing agents were omitted from sample buffer. Membranes were probed with mouse anti–β-actin-peroxidase (1:25,000; Sigma-Aldrich, A3854), goat anti-IgA HRP (1:1,000; Abcam, ab97235), or goat anti-pIgR (1:1,000; R&D Systems/Bio-technne, AF2800). Membranes were then washed and the latter target exposed (60 minutes) to 1:5,000 HRP-conjugated secondary antibody (GE Healthcare) in 5% milk-buffer. After further washes, signal was detected with ECL Western Blot detection reagents (GE Healthcare), followed by film exposure (GE Healthcare).

**Anti-PC and anti–T15 idiotype ELISAs.** High-binding enzyme immunoassay/RIA plates (Costar) were coated with 2 μg/ml PC-BSA (Biosearch Tech) or 2 μg/ml AB1-2 (anti-Vk22 antibody) in PBS and incubated overnight at 4°C. Wells were washed and blocked in 1% BSA in PBS for 1 hour at room temperature. Purified monoclonal unlabeled antibodies provided by the Kearney laboratory (BH8 [anti-PC IgM] or S107 [anti-PC IgA]) were used as standards in some experiments where values are expressed as ng/ml. After
sample and standard addition, plates were incubated overnight at 4°C. The following day, the plates were washed and anti-mouse isotype antibody conjugated to alkaline phosphatase (Southern Biotechnology) was added at 1:1,000 dilution for 1 hour. Plates were then developed with BCIP-NBT (Sigma-Aldrich) and read at 405 nm on a Bio-Tek Synergy 2 microplate reader.

**Statistics.** Analysis was performed using GraphPad Prism software. Data are represented as the mean ± SEM. Two-tailed Student’s *t* test was applied for comparisons of 2 groups and ANOVA for comparisons of more than 2 groups. Survival was evaluated by log-rank test. For all tests, *P* less than 0.05 was considered significant.

**Study approval.** All experiments were performed in accordance with the Animal Welfare Act and the U.S. Public Health Service Policy on Humane Care and Use of Laboratory Animals after review by the National Institute of Environmental Health Sciences Animal Care and Use Committee.

**Author contributions**
KA, KG, PR, and HJ designed, conducted, and analyzed experiments and contributed to the writing of the manuscript. NC, WC, JM, and LL contributed to tissue harvest, processing, and/or evaluation. KJ contributed to IHC, pathology evaluation, interpretation/report, and manuscript review. DC, MF, NC, PP, JK, and GT provided critical reagents, contributed to experimental design and interpretation, and contributed to the writing of the manuscript.

**Acknowledgments**
The authors thank Ligon Perrow for assistance with breeding; Tiwanda Masinde, Yvette Rebolloso, Geoffrey Hurbert, and David Olson (NIEHS IHC core); the NIEHS necropsy and histology core; Cristina Evans and David Goulding (colony support); Deborah King for blood cell count analysis; Ralph Wilson for clinical chemistry analysis; Carl Bortner and Maria Sifre for flow cytometry assistance; Jeff Tucker and Agnes Janoszahi from the Fluorescence Microscopy and Imaging Center for metamorphic analysis; Emily Mesev for assistance with immunoblotting; and David Kurtz and Jacqueline Locklear from the NIEHS QA Laboratory. We thank Marilyn Diaz and Chuancang Jiang for technical support with ANA and immunoglobulin ELISAs. We are indebted to the National Gnotobiotic Rodent Resource Center, UNC (5-P39-DK034987, 5-P40-OD010995). This research was supported by the Intramural Research Program of the NIH, National Institute of Environmental Health Sciences (Z01 ES102005), and by NIH National Institute of Allergy and Infectious Diseases grants AI14782 and AI124072 (to JFK).

Address correspondence to: Michael B. Fessler, National Institute of Environmental Health Sciences, 111 T.W. Alexander Drive, P.O. Box 12233, MD D2-01, Research Triangle Park, North Carolina 27709, USA. Phone: 919.541.3701; Email: fesslerm@niehs.nih.gov.
predicting the dramatic susceptibility of Irgm1-deficient mice to infection. *BMC Biol.* 2016;14:33.


