Induction of alternatively activated macrophages enhances pathogenesis during severe acute respiratory syndrome coronavirus infection

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Induction of Alternatively Activated Macrophages Enhances Pathogenesis during Severe Acute Respiratory Syndrome Coronavirus Infection

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Infection with severe acute respiratory syndrome coronavirus (SARS-CoV) causes acute lung injury (ALI) that often leads to severe lung disease. A mouse model of acute SARS-CoV infection has been helpful in understanding the host response to infection; however, there are still unanswered questions concerning SARS-CoV pathogenesis. We have shown that STAT1 plays an important role in the severity of SARS-CoV pathogenesis and that it is independent of the role of STAT1 in interferon signaling. Mice lacking STAT1 have greater weight loss, severe lung pathology with pre-pulmonary-fibrosis-like lesions, and an altered immune response following infection with SARS-CoV. We hypothesized that STAT1 plays a role in the polarization of the immune response, specifically in macrophages, resulting in a worsened outcome. To test this, we created bone marrow chimeras and cell-type-specific knockouts of STAT1 to identify which cell type(s) is critical to protection from severe lung disease after SARS-CoV infection. Bone marrow chimera experiments demonstrated that hematopoietic cells are responsible for the pathogenesis in STAT1−/− mice, and because of an induction of alternatively activated (AA) macrophages after infection, we hypothesized that the AA macrophages were critical for disease severity. Mice with STAT1 in either monocytes and macrophages (LysM/STAT1) or ciliated lung epithelial cells (FoxJ1/STAT1) deleted were created. Following infection, LysM/STAT1 mice display severe lung pathology, while FoxJ1/STAT1 mice display normal lung pathology. We hypothesized that AA macrophages were responsible for this STAT1-dependent pathogenesis and therefore created STAT1/STAT6−/− double-knockout mice. STAT6 is essential for the development of AA macrophages. Infection of the double-knockout mice displayed a lack of lung disease and pre-pulmonary lesions, suggesting that AA macrophage production may be the cause of STAT1-dependent lung disease. We propose that the control of AA macrophages by STAT1 is critical to regulating immune pathologies and for protection from long-term progression to fibrotic lung disease in a mouse model of SARS-CoV infection.
interest in potentially contributing to pulmonary fibrosis development following acute lung injury. In instances where epidermal growth factor (EGF) signaling mediates cell cycle arrest rather than growth, it has been shown that STAT1 is an important signaling component that negatively regulates the EGF pathway (2, 47). It has also been demonstrated that STAT1−/− mice are prone to the development of spontaneous tumors and are more sensitive to varying in vivo models of cancer, including esophageal cancer and melanoma (71, 73). STAT1−/− deficient mice have also been shown to be sensitive to radiation- and bleomycin-induced fibrosis, again illustrating STAT1’s role in control of the regulation of cell cycle progression and proliferation (70).

The involvement of STAT1 in viral infections is associated with its activity in the interferon signaling pathway and the innate immune response (11, 40, 62). In the case of SARS-CoV infection, we have shown that in 129/Sv mice, STAT1 is important in pathways other than the interferon signaling pathway (15). In this model, we found that SARS-CoV infection of mice lacking the alpha/beta interferon receptor (IFNAR1), gamma interferon (IFN-γ) receptor (IFNγR), or interferon lambda receptor (IL28Ra) and IFNAR/IFNGR double mutants display no enhanced pathogenesis, while STAT1−/− mice are highly susceptible and display severe lung disease and pathology (15). STAT1−/− mice in this genetic background have an inability to clear virus by 9 days postinfection and proceed to develop a severe lung disease phenotype that closely replicates the pathology seen in human patients during the SARS-CoV outbreak (31, 49). This includes extensive mixed inflammatory infiltrates, predominantly neutrophils, lymphocytes, and macrophages, as well as pathological features of DAD. The data suggest that a different STAT1-dependent pathway is critical for host protection rather than STAT1’s role in IFN signaling. We have shown that STAT1−/− mice do show an altered immune response to SARS-CoV infection, including an enhanced TH2 profile with a significant increase in genes related to the alternatively activated subset of macrophages (15, 79).

Alternatively activated (AA) macrophages have been implicated in models of lung disease, including asthma and pulmonary fibrosis (16, 51, 75). In contrast to classical macrophages, which become activated and differentiate in response to IFN-γ, AA macrophages differentiate in response to interleukin 4 (IL-4) and IL-13 (66). Induction of the AA macrophage phenotype is dependent upon IL-4R signaling, as well as the presence and activation of STAT6 (7, 44, 67). Their functions also differ from that of classically activated macrophages, as they are associated with wound repair and healing rather than direct pathogen killing (16). In mice, AA macrophages secrete the effector proteins YM1, FIZZ1, and arginase 1 (16). These effector protein functions are not yet fully understood, but their presence has been associated with liver fibrosis (36, 51, 56), as well as allergic airway inflammation (3, 42). In this paper, we utilize a mouse model of SARS-CoV infection to elucidate the role of STAT1 in AI1 as it relates to the induction of AA macrophage subsets in the lung. We demonstrate that the lack of STAT1 in the macrophage and monocyte lineage can shift the immune response and produce significant lung injury, as seen in total STAT1−/− mice. We also show that inhibition of AA macrophage development in STAT1−/− mice, by knocking out STAT6 in concert, protects them from severe lung disease and prefibrotic lesions during SARS-CoV infection. These data demonstrate the role of STAT1 in SARS-CoV pathogenesis, as well as the role that macrophage subtypes play in repair of acute lung injury and development of pulmonary-fibrosis-like diseases.

MATERIALS AND METHODS

Ethics statement. This study was carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. All mice in this study were treated following IACUC guidelines, and procedures were approved by the University of Maryland School of Medicine IACUC. For infection, the mice were pretreated with ketamine and xylazine as an anesthetic. Mice were euthanized if their weight dropped below approved levels or if clinical symptoms warranted it according to our IACUC protocol. Animal housing and care and experimental protocols were in accordance with all Animal Care and Use Committee guidelines.

Viruses and cells. SARS-CoV rMA15 was constructed as previously described (57). All virus stocks were stored at −80°C until they were ready to use. VeroE6 cells were purchased from ATCC (catalog number CRL-1586; Manassas, VA) and were used for growing rMA15 virus, as well as plaque assays to determine the viral load in lung tissue. Cells were grown in minimal essential medium (MEM) (Invitrogen, Carlsbad, CA) with 10% fetal bovine serum (FBS) (Atlanta Biologicals, Lawrenceville, GA) and 1% penicillin/streptomycin (Gemini Bioproducts, West Sacramento, CA).

Mouse breeding. C57BL/6 mice were purchased from Charles River Laboratories (Wilmington, MA). STAT1−/− mice were a gift from David Levy (New York University [NYU]) and were bred and housed in the animal facilities at the University of Maryland, Baltimore, MD (UMB).

For FoxO1−/− crosses, C57BL/6 Ly5.2/Cre mice (catalog number 004781) were purchased from Jackson Laboratory (Bar Harbor, ME), C57BL/6 FoxO1−/− mice were a gift from Michael Holtzman (Washington University in St. Louis), and C57BL/6 STAT1−/− mice were a gift from Lothar Hennighausen (NIDDK, NIH) (24). C57BL/6 STAT1−/− female mice were crossed with male Ly5.1/Cre and FoxO1−/− mice. Initial F1 pup tails were genotyped for the presence of Cre with primers 5'−GCATTAC CGGTGCAATGACAGTCAACCAA and 5'-GAGTGAACGAACCTCGTCCGAAATCGACTGCG-3' and for the STAT1/LoxP allele with primers 5'-CGATGACGTATGACAGTCAACCAA and 5'-ACTGGACGTCAACCAA GCCTG-3'. F1-positive female mice were crossed back to STAT1/LoxP males, and F2 pups were generated. These pups were screened for the presence of Cre and 2 copies of the STAT1/LoxP allele. Dual-positive pups were bred, and the lines were screened by immunohistochemistry (IHC) and PCR for the presence of STAT1 in the affected tissues. The methods of this screen are described below.

For STAT1−/− × STAT6−/− double-knockout production, STAT6−/− mice were purchased from Jackson Laboratory (Bar Harbor, ME). STAT1−/− mice were crossed with STAT1−/− mice, and F1 pups were generated. The F1 pups were crossed, and F2 double-positive pups were produced. These pups were screened for deletion of both genes by PCR, and selected pups were bred for use in the infection experiments.

Histological analysis. Lung sections were fixed in 4% paraformaldehyde (PFA) in phosphate-buffered saline (PBS) for a minimum of 48 hours, after which they were sent to the Histology Core at the University of Maryland, Baltimore, for paraffin embedding and sectioning. Five-micrometer sections were prepared and used for hematoxylin and eosin (H&E) staining by the Histology Core Services (University of Maryland). Stained sections were blinded and analyzed by Lindsay Giocochea at the University of Maryland Medical Center Department of Pathology. Images of histological sections were modified across all sections using Adobe Photoshop CS5 for only minimal modification of images.

Histology scoring. Slides were prepared as 3-μm sections and stained with hematoxylin and eosin. A pathologist (L.G.) was blinded to mouse genetic background, as well as infection status. Fields were examined by light microscopy and analyzed by Lindsay Giocochea (University of Maryland at Baltimore). The degrees of interstitial, peribronchial, and perivascular inflammation was scored from 0 to 3 (Table 1 provides a...
description of the scoring methodology). For each experimental group, Goicochea blindly scored 3 mice, and those scores were tabulated to compare strains, time points, and infections. Other histologic features, such as the presence of reactive bronchial epithelial and pleural changes and the extent of peribronchovascular neutrophilia, were also noted for each group in the text. A combined score, based on the described pathological descriptors (Table 1), was generated for each mouse, and then, 3 mice per group were averaged and the standard deviation for the scoring was computed. These scores are presented in the histological scoring figures.

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| Table 1 Histological scoring criteria |

**YMI and Fizz1 immunohistochemistry.** Five-micrometer paraffin sections of fixed lung tissue were cut by the Histology Core at the University of Maryland, Baltimore. Slides were deparaffinized in xylenes and rehydrated in an ethanol series. Endogenous peroxidase activity was quenched by incubating slides in 0.3% hydrogen peroxide in methanol for 30 min at room temperature (RT). Antigen retrieval was performed by incubating slides for 20 min in IHC select citrate buffer, pH 6.0 (Millipore, Billerica, MA). Tissue sections were then preincubated in 10% goat serum in PBS for 1 h at RT. Rabbit polyclonal Fizz1 antibody (Abcam, Cambridge, MA) was used at a concentration of 1:100 diluted in 10% goat serum in PBS, Rabbit polyclonal YMI1 antibody (StemCell Technologies, Vancouver, BC, Canada) was used at a concentration of 1:200 diluted in 10% goat serum in PBS. Sections were incubated in primary antibody for 30 min at RT. The slides were washed before incubation with biotinylated secondary antibody from the ABC Elite Rabbit kit (Vector Laboratories, Burlingame, CA) for 30 min at RT. The slides were washed again prior to addition of ABC reagent (Vector Laboratories, Burlingame, CA) for 30 min at RT. After 3 changes of PBS, the slides were developed using DAB (3,3-diaminobenzidine) for 2 min per slide. Development was stopped using a 5-min incubation in distilled water (dH₂O). A 10-min incubation in 0.5% methyl green in 0.1 M sodium acetate solution, pH 4.2, was used for counterstaining. The counterstain was cleared using 2 changes of PBS, followed by 2 changes of N-butanol for 2 min each. Slides were dehydrated using 2 5-min incubations in xylenes before being mounted.

**STAT1 immunohistochemistry.** Slides with 5-μm sections of mouse lung were obtained from the Histology Core at the University of Maryland, Baltimore. The slides were deparaffinized using two changes of xylenes for 5 min each change. An ethanol series was then used to rehydrate the sections. The slides were then incubated in MOM mouse IgG blocking reagent from a MOM kit purchased from Vector Laboratories (Burlingame, CA) for 1 h at RT. The slides were washed in PBS and then incubated with MOM protein diluent for 5 min at RT. Anti-STAT1 primary antibody (BD Transduction Laboratories, San Jose, CA) was diluted at 1:100 in MOM protein diluent. The slides were incubated in primary antibody for 30 min at room temperature. The slides were washed in PBS before being incubated in MOM biotinylated anti-mouse IgG antibody provided in the MOM kit for 10 min at RT and washed in 2 changes of PBS. The biotin signal was amplified using a TSA Plus Biotin kit (Perkin Elmer, Waltham, MA). The slides were incubated in working-strength biotin amplification solution made at 1:25 for 5 min at RT and then washed in 2 changes of PBS. An ABC-Alkaline Phosphatase (AP) kit (Vector Laboratories, Burlingame CA) was then used. The slides were incubated in ABC-AP for 30 min at RT and then developed using an alkaline phosphatase kit (Vector Laboratories, Burlingame, CA) by incubating the slides for 60 min in AP. The slides were then counterstained using methyl
green solution before being decolorized, dehydrated in xylenes, and mounted.

Isolation of DNA from intraperitoneal macrophages and peripheral blood. Peritoneal macrophages were obtained by injecting mice with 3 ml of sterile 3% thioglycolate broth (Remel, Lenexa, KS) intraperitoneally. Three days after injection, cells were harvested by lavage of the intraperitoneal cavity. The cells were washed in DMEM and pelleted for further analysis by reverse transcription-PCR. Peripheral blood was collected by terminal cardiac puncture. DNA was isolated using a DNA/RNA Allprep kit from Qiagen Laboratories (Valencia, CA). The DNA was then used for PCR and analyzed on a 1% agarose gel. The template was amplified by reverse transcription-PCR using forward (5'-GCATGACATGATCAGCTTG-3') and reverse (5'-ACTGACGTCACACAGCTG-3') primers.

RESULTS
rMA15 infection is cleared from both C57BL/6 and STAT1−/− mice. To evaluate the role of STAT1 in SARS-CoV infection in mice, 10-week-old WT C57BL/6 and C57BL/6 STAT1−/− mice were infected with 1 × 10^5 PFU of rMA15. rMA15 is a mouse-adapted SARS-CoV containing 6 amino acid mutations from the WT Urbani strain of SARS-CoV that produces lethal disease in BALB/c mice but does not kill C57BL/6 mice (57). The mice were weighed daily over a 9-day course of infection (Fig. 1A), during which lung tissue was harvested at days 2, 5, and 9 for further characterization. WT C57BL/6 mice lost roughly 15% of their body weight during the first 4 days postinfection (p.i.) and then
achieved complete recovery of their starting weight by 9 days p.i. In contrast, STAT1−/− mice continued to lose weight throughout the course of infection, losing more than 20% of their starting weight by 9 days p.i.

Viral titers were analyzed using a plaque assay to compare WT to STAT1−/− mouse infections (Fig. 1B). WT mice displayed peak viral titers at 2 days p.i., with titers of 4.75 × 10^8 PFU/g, after which they decreased by 5 days p.i. to 1.4 × 10^8 PFU/g. Viral titers were below detectable levels by 9 days p.i. STAT1−/− mice displayed a similar trend, with peak titers of 8.75 × 10^7 PFU/g at 2 days p.i. STAT1−/− mice showed a 2-log-unit increase in titer at 5 days p.i. compared to WT mice but still cleared virus by 9 days p.i. to below detectable levels (Fig. 1B). Interestingly, despite the differences in weight loss, we found that WT and STAT1−/− mice on the C57BL/6 background were able to clear rMA15 virus from their lungs during infection, although STAT1−/− mice had higher virus titers at 5 days p.i.

**Induction of prefibrotic lesions in STAT1−/− mice after virus clearance.** Histological analysis was performed on lungs harvested at 2, 5, and 9 days p.i. in C57BL/6 WT and STAT1−/− mice (Fig. 1C and D). As previously reported, WT mice infected with rMA15 show bronchiolar sloughing at 2 days p.i., with significant cell debris in the bronchioles. A peribronchiolar and periarteriolar mixed inflammatory infiltrate consisting of eosinophils and neutrophils was observed. By 5 days p.i., airs spaces were generally clear of cell debris and ciliated epithelial cells were present in the bronchioles. Inflammation was still present in the WT lungs, presenting with main eosinophils and macrophages in focal regions, primarily surrounding small airways. By 9 days p.i., we observed few small focal aggregates predominantly comprised of lymphocytes cuffing around vessels and larger airways; however, in general, inflammatory infiltrates were minimal, and the lungs had returned to their preinfection state.

In C57BL/6 STAT1−/− mice we found a more severe lung pathology that persisted throughout the course of infection. At 2 days p.i., STAT1−/− mice showed lung pathology similar to that seen in WT mice. Lungs showed evidence of bronchiolar sloughing, with mixed inflammatory infiltrates throughout the lung, but most notably displayed cuffing around large airways and blood vessels. At 5 days p.i., STAT1−/− mice did not have the same recovery as WT mice, and we found a continued increase in levels of mixed inflammatory infiltrates comprised of neutrophils and mononuclear cells, macrophages, and lymphocytes. These infiltrates continued to cluster around large airways and vessels and began to fill the alveolar space. By 9 days p.i., there was continued progression in the severity of lung pathology and there was more pronounced neutrophilia and increased macrophage numbers found in the lung interstitium. Lung epithelial cells exhibited squamous metaplasia, which is a characteristic reactive response to damage commonly seen in incidents of ALI, especially in organizing DAD. Interestingly, there was also proliferation of interstitial fibroblasts in both the subpleural and central regions of the lung. This resembled a prefibrotic-like stage sometimes seen following extensive lung damage. H&E-stained slides were blinded and scored by pathologist Lindsay Goicochea (UMB). The slides were scored for perivascular, peribronchiolar, and interstitial inflammation. The scores were tallied as described in Materials and Methods and are graphed in Fig. 1D. While pulmonary inflammation was still found in WT mice at 9 days p.i., we observed that at 9 days p.i. STAT1−/− mice had statistically significant inflammation throughout the lung tissue. Collectively, this lung pathology suggests an inappropriate immune response continuing even after virus had been cleared to below detectable levels.

**Alternatively activated macrophages are induced during rMA15 infection of STAT1−/− mice.** We observed an expansion of total macrophages during infection in our histological sections of STAT1−/− mice after rMA15 infection. Since we had previously reported the induction of AA macrophage phenotypes in STAT1−/− mice (79), immunohistochemistry was performed to look for the presence of proteins commonly associated with AA macrophages. YM1 and FIZZ1 are both expressed from AA macrophages in the lung during diseases such as allergic airway inflammation models and respiratory syncytial virus (RSV) infection and used as the standard marker of AA macrophage induction in mice (54, 55). To identify whether AA macrophages were present in WT and STAT1−/− mice, lung sections were stained with antibodies for YM1 and FIZZ1 to analyze their expression during rMA15 infection (Fig. 2). Wild-type and STAT1−/− mice were found to express FIZZ1 specifically along the lung epithelia in PBS-inoculated controls; however, at 9 days after infection, FIZZ1 protein is highly enriched both in the lung parenchyma and in macrophages of STAT1−/− mice (Fig. 2). This is in contrast to what is observed in WT mice. In WT mice, we found FIZZ1 expression solely associated with the lung epithelium even at 9 days p.i. (Fig. 2). Minimal induction of FIZZ1 was identified at days 2 and 5 postinfection of WT or STAT1−/− mice (Fig. 2). YM1 expression in PBS-inoculated control mice was associated with alveolar macrophages with little to no expression in the ciliated lung epithelium of the large airways in both wild-type and STAT1−/− mice (Fig. 2). However, at 9 days p.i., STAT1−/− mice showed evidence of YM1 still associated with alveolar macrophages, but now they displayed diffuse staining in the lung parenchyma in addition to increased staining in the ciliated lung epithelial cells. Minimal induction of YM1 was identified at days 2 and 5 postinfection of WT or STAT1−/− mice (Fig. 2). This increase in both YM1 and FIZZ1 levels in the lung suggests an increase in AA macrophages in the SARS-CoV-infected STAT1−/− mice.

The induction of FIZZ1 from cell types other than macrophages has been reported in systems of allergic airway inflammation (8). To confirm the presence of macrophages at sites of FIZZ1 and YM1 expression in these mice, we stained with a panmacrophage marker, F4/80, using immunohistochemistry (Fig. 3). We detected F4/80-positive cells at the same sites that stained for FIZZ1 and YM1, which strongly supports the finding that the positive cells are truly AA macrophages.

**Induction of TH2 cytokines, IL-4 and IL-13, during rMA15 infection.** Alternatively activated macrophages differentiate in response to the TH2 cytokines IL-4 and IL-13. We hypothesized that we would be able to detect an increase in IL-4 and IL-13 concomitant with the induction of AA macrophages in response to infection in STAT1−/− mice. RNA from lungs of WT and STAT1−/− mice mock infected with PBS or infected with rMA15 was isolated and used for real-time PCR analysis of IL-4 and IL-13 (Fig. 4). We found that in WT mice, there was little detectable IL-4 or IL-13 message in either PBS-mock-infected or rMA15-infected mice. However, in STAT1−/− mice infected with rMA15, we found a significant increase in IL-4 and IL-13 mRNAs at 5 days p.i., which was then reduced by 9 days p.i. The increase at 5 days p.i. correlates with the induction of AA macrophages seen beginning at 5 days p.i. and increasing through 9 days p.i. (Fig. 2). Once the induction
FIG 2 Alternatively activated macrophages are induced during rMA15 infection of STAT1−/− mice. Immunohistochemistry was performed in lung sections from WT and STAT1−/− mice after either PBS inoculation or rMA15 infection at 2, 5, and 9 days p.i. The sections were stained for the alternatively activated macrophage markers FIZZ1 and YM1. The images showing 10X resolution focused on airway labeling and those at 40X resolution focused on the pulmonary interstitium to highlight the extent of signal.
of AA macrophages occurs, IL-4 and IL-13 are no longer needed to continue the TH2 polarization in STAT1−−/− mice.

**STAT1 is necessary in the hematopoietic lineage to protect from rMA15 pathogenesis.** Our data suggest that STAT1−−/− mice have a dysregulated response to SARS-CoV infection, leading to a more severe disease phenotype and delayed clearance of virus. Immunohistochemical results suggest that part of this response is related to a change in macrophage phenotype and polarity. We next sought to determine if cells from the hematopoietic lineage were responsible for these changes in outcome following infection. In order to test this, we developed bone marrow chimeric mice by lethally irradiating mice and transferring congenitally marked bone marrow; STAT1−−/− bone marrow was injected into wild-type mice, and wild-type bone marrow was injected into STAT1−−/− mice. After screening using the CD45 marker on CD4+ cells to confirm successful reconstitution (Fig. 5A), the mice were infected with 1 × 10⁵ PFU of rMA15 and weighed daily for 9 days. Mouse lungs were harvested at days 2, 5, and 9 postinfection for analysis. Quantification of virus titers was performed on lung tissue to evaluate the ability of each mouse to clear virus. We observed that all groups had peak viral titers at 2 days p.i., with titers beginning to diminish by 5 days p.i. By 9 days p.i., virus was below detectable levels (Fig. 5B). The STAT1−−/− mice receiving wild-type bone marrow displayed a minor delay in clearance at 5 days p.i., while wild-type mice receiving STAT1−−/− bone marrow were indistinguishable from normal wild-type mice.

Lung sections from bone marrow chimera mice were analyzed for the severity of lung pathology after infection (Fig. 5C). At 9 days p.i., the lung pathology was significantly different between the two chimera groups. In STAT1−−/− mice that received wild-type bone marrow, there was damage to the large airways, particularly to the ciliated epithelial cells that line the airways. Cell debris was also still observed in the bronchiolar airspace. These mice also had mixed inflammatory infiltrates that were found cuffing these damaged airways and vessels throughout the lung. Interestingly, these inflammatory infiltrates followed a focal pattern as opposed to the diffuse inflammation seen in the total STAT1−−/− animals.

In contrast, WT mice that received STAT1−−/− bone marrow exhibited less overall damage to the ciliated epithelial cells of the large airways but displayed enhanced inflammatory infiltrates and both perivascular and peribronchial cuffing. There remained foci of reactive squamous metaplasia, but largely, the airways were healthy and intact. We observed a diffuse pattern of inflammation throughout the interstitium comprised of neutrophils and macrophages marked by almost complete collapse of the alveolar space. We also found perivascular neutrophilia, as well as lymphohasmocytic infiltrates, throughout the lung parenchyma. The inflammation seen in the lungs of these wild-type mice receiving STAT1−−/− hematopoietic cells closely resembled that of total STAT1−−/− mice, as did their histological scoring (Fig. 5D).
suggests that one or more cell types from STAT1−/− mice are important contributors to the prolonged lung damage after rMA15 infection.

Deleting STAT1 from macrophages and monocytes produces disease similar to that in complete STAT1−/− mice. After confirming that cells of hematopoietic lineage were predominantly responsible for the increase in pathology in total STAT1−/− mice, we sought to identify potential cell types that could be specifically responsible for this pathology. Together, the hematopoietic link and the increase in AA macrophage markers led us to hypothesize that STAT1 contributes to shaping macrophage responses during SARS-CoV infection and that a change in macrophage polarization was leading to severe lung disease. To test this, we developed 2 mouse strains in which STAT1 is conditionally knocked out. In one strain, we crossed mice containing a STAT1/LoxP integrant with a LysM promoter-driven Cre recombinase to delete STAT1 in monocytes (mice referred to as LysM/STAT1). In the second strain, we crossed mice containing the same STAT1/LoxP integrant with a FoxJ1 promoter-driven Cre recombinase to delete STAT1 in ciliated epithelial cells (77). While we hypothe-
sized that STAT1 was modulating the macrophage response, we created the FoxJ1/STAT1 mouse strain to control for the effect of STAT1 in ciliated epithelial cells, which are the primary target of SARS-CoV in the lung (77). The FoxJ1/Cre mouse has been analyzed and displays specific expression in the ciliated cells of the respiratory epithelium, as well as the choroid plexus, ependyma, oviduct, and testis (77). SARS-CoV specifically infects the respiratory epithelium, so expression in other tissues does not complicate the analysis. To confirm the deletion of STAT1 in monocytes and macrophages in LysM/STAT1 mice, we used thioglycolate to induce intraperitoneal macrophages in WT, STAT1−/−, and LysM/STAT1 mice to compare deletions (Fig. 6A). We confirmed deletion of STAT1 using gene-specific primers and found that in macrophages, a higher-molecular-weight band is found in WT mice than in LysM/STAT1 mice, demonstrating the deletion, and no band is found in the primers used. In blood, where the LysM promoter is not active, we do not see a deletion in the LysM/STAT1 mice, demonstrating that LysM/Cre is specific to monocytes and macrophages. To verify the FoxJ1/STAT1 deletion in ciliated epithelial cells, we used IHC staining for STAT1 to compare WT, STAT1−/−, and FoxJ1/STAT1 mouse airways (Fig. 6B). We found high levels of expression of STAT1 in WT mice and no expression in any lung cells in STAT1−/− mice. In FoxJ1/STAT1 mice, we found STAT1 expres-
sion in the Clara cells of the airways and throughout the alveoli; however, STAT1 expression was lacking in ciliated airway epithelial cells, confirming correct deletion of STAT1 in FoxJ1/Cre-expressing cells.

The 10-week-old WT, STAT1−/−, LysM/STAT1, and FoxJ1/STAT1 mice were inoculated with 1 × 10^5 PFU of rMA15 and were weighed daily for 9 days (Fig. 6C). WT mice displayed a 10% starting weight loss by day 3 postinfection, with the mice regaining 100% of their starting weight by 9 days p.i. STAT1−/− mice lost 10% of their starting weight during the first 3 days and then, as before, lost more than 20% of their starting weight through 9 days p.i. Additionally, FoxJ1/STAT1 mice had weight loss very similar to that of WT mice. We observed 10% weight loss by 3 days p.i. and then complete recovery by 9 days. Interestingly, LysM/STAT1 mice displayed early weight loss that closely followed that of the total STAT1−/− mice and exceeded STAT1−/− mouse weight loss by 4 days p.i.; however, they also proceeded to recover their body weight by 9 days postinfection, albeit more slowly than WT mice.

H&E-stained lung sections were analyzed to determine the degree of lung damage and inflammation in the conditional knockout mice following infection with rMA15. As before, at 9 days p.i., we found minimal residual inflammation in WT mice, while STAT1−/− mice displayed overactive and uncontrolled inflammation, as well as bronchi. Histological scoring demonstrated a reduced score in the WT lung parenchyma at this time point, and FIZZ1 was again found collecting in fluid-filled foci in the interstitium. Conversely, FoxJ1/STAT1 mice displayed very little induced YM1 or FIZZ1 at 9 days p.i. with rMA15 (Fig. 6). Minor induction of YM1 was found after infection; however, it was found at background levels, as seen in WT mice. Collectively, these data suggest that STAT1 plays a role in macrophage polarization during infection with SARS-CoV and that when macrophages lack STAT1, they are increasingly of an alternatively activated phenotype and are induced during rMA15 infection.

**Inhibition of AA macrophages in STAT1−/− mice eliminates severe lung disease.** Our data suggest that AA macrophages are associated with enhanced lung disease after SARS-CoV infection. We hypothesized that these macrophages could be contributing to the exacerbated lung damage, since they have been previously shown to be involved in altered lung repair pathways in fibrosis (48, 69) and asthma (3, 4, 30, 43, 68). Development of AA macrophages is completely dependent on the STAT6 transcription factor (7, 34, 41). In mice lacking STAT6, no AA macrophages are found after treatment with several AA macrophage inducers (7, 34, 41). We created STAT1−/− × STAT6−/− double-knockout mice (called STAT1/6−/−) that contain the STAT1 deletion but lack the ability to produce AA macrophages and thus YM1 and FIZZ1.

As before, we infected these mice and STAT6−/− mice, to control for a phenotype with the loss of STAT6, with rMA15. WT, STAT1−/−, STAT6−/−, and STAT1/6−/− mice were infected with 1 × 10^5 PFU of rMA15 and weighed daily for 9 days. WT and STAT1−/− mice lost more than 20% of their body weight by 9 days p.i. STAT6−/− mice displayed weight loss similar to that of WT mice, with early weight loss by day 3, but then regained weight through 9 days. Interestingly, STAT1/6−/− mice also displayed 10% weight loss by day 3, but unlike STAT1−/− mice, the double-knockout mice regained weight and had returned to their starting weight by 9 days p.i.

Mouse lungs were harvested and analyzed for histological lesions by H&E staining (Fig. 8B). WT and STAT1−/− mice displayed similar lung pathology, as found previously, with minimal residual inflammation in WT mice at 9 days p.i., but STAT1−/− mice displayed severe inflammatory infiltrates around blood vessels and airways and throughout the interstitium. Consistent with the weight loss data, STAT1/6−/− mice displayed minimal inflam-
information and lung damage after rMA15 infection, similar to WT mice (Fig. 8B). Histological scoring of lung sections demonstrated reduced lung damage and inflammation in both STAT6−/− and STAT1/6−/− mice, corroborating the weight loss data (Fig. 8C). Together, these data suggest that elimination of AA macrophages in a STAT1−/− background is able to abrogate induced lung disease and inflammation after rMA15 infection.

To confirm that AA macrophage development was inhibited during infection, we again analyzed YM1 and FIZZ1 induction by IHC as markers of AA macrophage levels in lung tissue (Fig. 9). Lung sections from WT, STAT1−/−, STAT6−/−, and STAT1/6−/− mice inoculated with PBS or rMA15 were analyzed. WT and STAT6−/− mice displayed minimal YM1 and FIZZ1 expression in rMA15-infected mice, while STAT1−/− mice displayed high levels of both YM1 and FIZZ1, as observed previously. Concomitant with weight loss and H&E staining, we found low levels of FIZZ1 and YM1 staining in STAT1/6−/− mice with either PBS inoculation or rMA15 infection. The level of staining was similar to that observed in WT mice after infection. This demonstrates that the STAT1/6−/− mice have decreased numbers of AA macrophages and that this reduction correlates with decreased lung pathology. Together, these data demonstrate that in a STAT1−/− mouse model where AA macrophages are eliminated there is reduced lung pathology and disease after rMA15 infection. We interpret this to mean that AA macrophages contribute to the severe lung disease seen in STAT1−/− mice and that even in the presence of total STAT1 deletion, the removal of AA macrophages inhibits disease progression, prefibrotic lesions, and the enhanced inflammatory infiltrate. We hypothesize that the inhibition of the AA macrophage induction pathway could be a therapeutic target for SARS-CoV infection, as well as that of other highly acute respiratory viruses that may induce similar pathways during infection.

DISCUSSION
The interaction between host and pathogen controls the outcome of an infection. From the virus side, many viral proteins from a wide range of viruses directly affect host cell function during infection. From the host side, the response to infection can exacerbate disease and lead to increased damage, causing more destruction than the viral infection alone. In this series of experiments, we have shown that the host response to SARS-CoV infection is able to exacerbate disease and that the host protein STAT1 is important for controlling that response.

The repair of lung tissue after infection or mechanical trauma normally occurs in a rapid, but controlled series of events: inflammation, growth factor secretion, basement membrane repair, and finally resolution of the injured tissue (58). Under normal circumstances, this process proceeds without any problems; however, when the infectious burden overwhelms the host or there is continuous damage, the wound-healing response can become dys-

FIG 7 Alternatively activated macrophages are induced during rMA15 infection of LysM/STAT1−/− mice but not FoxJ/STAT1−/− mice. IHC was performed in lung sections from LysM/STAT1 and FoxJ/STAT1 mice after either PBS inoculation or rMA15 infection at 9 days p.i. The sections were stained for the alternatively activated macrophage markers FIZZ1 and YM1. The images showing 10X resolution focused on airway labeling, and those at 40X resolution focused on the pulmonary interstitium to highlight the extent of signal.
regulated, resulting in scarring and fibrosis (39, 61). When scarring develops into ARDS and pulmonary fibrosis, lung function is reduced, which can ultimately lead to death. Both ARDS and pulmonary fibrosis are traditionally treated with corticosteroids, which act to dampen the immune response; however, this course of treatment has not been shown to be particularly effective (38, 53). In cases where the etiology is an infectious agent, such a treatment can actually be harmful to the patient, as it delays the clearance of the pathogen (6, 17, 76).

After injury occurs, there is a cascade of events that progress from initiation of the wound-healing response to recruiting inflammatory cells and fibroblasts to resolution of the wound. Al-

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**FIG 8** Pathogenesis of rMA15 in STAT1/6−/− mice. (A) Weight loss curves of WT and STAT1/6−/− mice infected with rMA15 (n = 5 for each time point). The error bars indicate standard deviations. (B) H&E staining of WT, STAT1−/−, STAT6−/−, and STAT1/6−/− mice either mock infected with PBS or infected with rMA15. (C) Histological scoring of H&E-stained mouse lungs for WT, STAT1−/−, STAT6−/−, and STAT1/6−/− mice at 9 days p.i.
FIG 9 Immunohistochemical staining of lung sections from WT, STAT1−/−, STAT6−/−, and STAT1/6−/− mice for AA macrophage markers YM1 and FIZZ1. Brown staining is positive staining with antibody. All sections are from either PBS-inoculated or rMA15-inoculated mice at 9 days p.i.
tivation in any of these cascades can drive the host to induce development of fibrosis in the tissue that is in need of repair. Several factors have been shown to be involved in pushing a profibrotic spectrum in animal models; however, these targets have yet to be therapeutically validated in human studies (20, 52, 74). A novel pathway in this process is the STAT1/EGFR signaling axis. Mutations in STAT1 have been found in many cancers throughout the body, especially the lung (21, 71). STAT1 has been shown to activate cell cycle arrest and apoptosis and has demonstrated the properties consistent with a tumor suppressor. Additionally, STAT1−/− mouse embryonic fibroblasts (MEFs) are highly susceptible to uncontrolled cell growth compared to WT MEFs (70), and STAT1−/− mice display increased sensitivity to radiation (12, 22) and bleomycin-induced fibrosis (70).

In viral models of chronic airway infection using Sendai virus as the inducer, STAT1 expression in airway epithelial cells has been shown to be key to protecting against severe lung disease (63). In this model, STAT1−/− mice are highly susceptible to Sendai virus, and using bone marrow chimeras of WT and STAT1−/− mice, it was found that in STAT1−/− mice reconstituted with WT bone marrow, the enhanced susceptibility remained, but not with the converse reconstitution. This result is the opposite of what we found for SARS-CoV infection in the above-mentioned bone marrow chimera experiment and suggests that the intimate and virus-specific interactions between the virus and host can modify the inflammatory and disease states. Using the same model of Sendai virus-induced chronic airway infection, Kim et al. found that even after viral clearance, induction of natural killer T (NKT) cells activates macrophages in the lung (23). These macrophages and NKT cells secrete IL-13, which polarizes macrophages into an alternatively activated macrophage subtype. This activation produces a positive-feedback loop to amplify IL-13 production and alternative activation of macrophages (23). Importantly, the authors found that it is the persistent activation of the AA macrophage pathway that is critical to the continued lung damage and inflammation.

In our previous study, we showed the result of infection by rMA15 in 129/Sv STAT1−/− mice, which display a phenotype different in important ways from that of C57BL/6 STAT1−/− mice (15). We found that 129/Sv wild-type mice infected with a mouse-adapted SARS-CoV (rMA15) lose around 15% of their starting weight through 4 days of infection that the mice recovered to their starting weight by 9 days p.i. Importantly, in WT C57BL/6 mice, the rMA15 virus is cleared by 9 days p.i., and it is also cleared in the STAT1−/− mice, unlike what is seen in 129/Sv STAT1−/− mice (15). Even with virus clearance, the STAT1−/− mice still developed pre-pulmonary-fibrosis-like lesions in the lungs by 9 days p.i. This suggests that, as we hypothesized, the STAT1 protein has a role in the development of lung pathology in addition to its role in the innate immune response to infection.

Based on previous data, we predicted that STAT1 was playing a role either in the lung ciliated epithelial cells, which are the only cells in the lung that are infected by SARS-CoV (64), or in macrophages, since we found an altered macrophage response in total STAT1−/− mice infected with rMA15 (79). Using this C57BL/6 model, we produced mice that lacked STAT1 in only specific cell types, either FoxJ1/STAT1 mice (77) or LysM/STAT1 mice (5). FoxJ1/STAT1 mice have STAT1 deleted in the ciliated epithelial cells, and LysM/STAT1 mice have STAT1 deleted in monocytes and macrophages. FoxJ1/STAT1 mice infected with rMA15 displayed lung pathology, weight loss, and virus growth kinetics similar to those of WT C57BL/6 mice. LysM/STAT1 mice displayed severe lung pathology and a cytokine response similar to that of total STAT1−/− mice. Interestingly, the weight loss was not significantly different from that of WT mice, suggesting that weight loss and lung pathology may be separate responses during infection. While the lung pathologies and the AA macrophage induction in STAT1−/− and LysM/STAT1 mice are very similar, the weight loss is not, with LysM/STAT1-infected mice regaining their weight through the experiments. We hypothesize that this difference may be due to the role of STAT1 in cytokine induction from host cells and that this induction involves unidentified host response factors affecting morbidity and subclinical symptoms of mice. We are currently using transcriptomics and mathematical modeling to identify pathways contributing to weight loss and pathogenesis in these mice.

The LysM/STAT1 mice developed lung pathology very similar to that of the total STAT1−/− mice. When assayed for the induction of alternatively activated macrophages, which we suspect are critical for the development of these lesions, we found significant upregulation of this macrophage population compared to WT mice and to levels very similar to those in total STAT1−/− mice. Recently, it has been shown that macrophages isolated from patients with idiopathic pulmonary fibrosis (IPF) have an increased number of alternatively activated macrophages, as shown by the human AA macrophage markers of increased CD163 and decreased inducible nitric oxide synthase (iNOS) expression (60). In order to evaluate the role of the AA macrophages more closely, we crossed STAT6−/− mice, which are known to have a blocked AA macrophage phenotype (59), with our STAT1−/− mice. These STAT1/6−/− mice developed disease in a manner that replicated that seen in wild-type mice and also lacked induction of AA mac-
ropahge-associated proteins YM1 and Fizz1. This further supports our hypothesis that AA macrophage induction in STAT1−/− mice is responsible for the enhanced disease phenotype.

Together, our findings demonstrate that control of the AA macrophage response is critical during the response to acute lung injury, potentially from a variety of etiologies, and that STAT1 is an important mediator of the development of AA macrophages during infection. We are actively investigating the role of this pathway in other models of chronic inflammation in the lung to identify the role of AA macrophages, STAT1, and the inflammatory response in the induction and maintenance of lung disease. We hypothesize that if we can modulate the host response during infection to skew the classical versus alternatively activated macrophage induction pathways, we may be able to alter the disease outcome in cases of ALI and limit the development of fibrotic-like disease.

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