Pseudomonas aeruginosa Protease IV Exacerbates Pneumococcal Pneumonia and Systemic Disease

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ABSTRACT  Pneumonia is a pulmonary disease affecting people of all ages and is consistently a leading cause of childhood mortality and adult hospitalizations. Streptococcus pneumoniae and Pseudomonas aeruginosa are major lung pathogens commonly associated with community-acquired and nosocomial pneumonia. Additionally, mixed lung infections involving these bacterial pathogens are increasing in prevalence and are frequently more severe than single infections. The cooperative interactions of these two pathogens that impact pulmonary disease severity are understudied. A major secreted virulence factor of P. aeruginosa, protease IV (PIV), cleaves interleukin 22 (IL-22), a cytokine essential for maintaining innate mucosal defenses against extracellular pathogens. Here, we investigate the ability of PIV to augment the virulence of a pneumococcal strain with limited virulence, S. pneumoniae EF3030, in a C57BL/6 murine model of pneumonia. We demonstrate that pulmonary coinfection involving P. aeruginosa 103-29 and S. pneumoniae EF3030 results in pneumococcal bacteremia that is abrogated during pneumococcal coinfection with a PIV-deficient strain. Furthermore, intratracheal administration of exogenous PIV and EF3030 resulted in abundant immune cell infiltration into the lung with large abscess formation, as well as severe bacteremia leading to 100% mortality. Heat-inactivated PIV did not worsen pneumonia or reliably induce bacteremia, suggesting that the specific activity of PIV is required. Our studies also show that PIV depletes IL-22 in vivo. Moreover, PIV-mediated enhancement of pneumonia and disease severity was dependent on the expression of pneumolysin (Ply), a prominent virulence factor of S. pneumoniae. Altogether, we reveal that PIV and Ply additively potentiate pneumonia in a murine model of lung infection.

IMPORTANCE S. pneumoniae remains the leading cause of bacterial pneumonia despite widespread use of pneumococcal vaccines, forcing the necessity for appropriate treatment to control pneumococcal infections. Coinfections involving S. pneumoniae with other bacterial pathogens threaten antibiotic treatment strategies and disease outcomes. Currently, there is not an effective treatment for alveolar-capillary barrier dysfunction that precedes bacteremia. An understanding of the dynamics of host-pathogen interactions during single and mixed pulmonary infections could elucidate proper treatment strategies needed to prevent or reduce invasive disease. Antibiotic treatment decreases bacterial burden in the lung but also increases acute pathology due to cytotoxins released via antibiotic-induced bacterial lysis. Therefore, targeted therapeutics that inhibit or counteract the effects of bacterial proteases and toxins are needed in order to limit pathology and disease progression. This study identifies the cooperative effect of PIV and Ply, products of separate lung pathogens that additively alter the lung environment and facilitate invasive disease.

KEYWORDS Pseudomonas aeruginosa, Streptococcus pneumoniae, coinfection, invasive disease, pneumococcus, pneumonia, proteases
Pneumonia is the leading cause of childhood mortality worldwide and is also a primary reason for hospitalizations associated with high morbidity and financial burden in the United States (1, 2). The causative agent in cases of pneumonia has classically been diagnosed and treated as a single infecting microorganism, and the principal etiological agents identified during episodes of bacterial pneumonia include Streptococcus pneumoniae, Haemophilus influenzae, Staphylococcus aureus, and Pseudomonas aeruginosa (3). However, increases in concurrent lung infections have been reported during both nosocomial and community-acquired pneumonia (CAP), and these mixed bacterial infections have resulted in more severe disease presentations than single infections (4–8). Specifically, S. pneumoniae is the most common causative agent of CAP and is also the most prevalent bacterium during mixed CAP infections (5–7).

In healthy individuals, the environment of the lower respiratory system is nutrient poor and unfavorable to invading pathogens (9). During acute lung injury or chronic lung disease, the alveolar environment favors bacterial growth through nutrient-rich fluid accumulation, temperature gradients, development of oxygen pockets, and a host inflammatory response that selectively favors potential pathogens (10–13). Contrary to the previous assumption that the lung is a sterile environment, a heterogeneous microbiome has been identified in the lungs of healthy individuals (9). Furthermore, the lung microbiome shapes the ecological forces that influence the colonization or elimination of potential pathogens in the lower airways (10). Lung microbiome dysbiosis occurs during acute lung injury such as viral infection and favors the growth of P. aeruginosa and S. pneumoniae (10). Moreover, lung coinfections linking P. aeruginosa and S. pneumoniae have been reported (7, 14–16). This coexistence implies that these two pathogens are cooperatively interacting during pneumonia. These interactions may be a result of secreted virulence factors that alter the lung environment, select for outgrowth of these pathogens, and thus enhance persistence of both pathogens.

Here, we investigated the ability of a secreted Pseudomonas protease, protease IV (PIV), to augment an active lung infection caused by S. pneumoniae EF3030, a strain of pneumococcus with limited virulence in a murine model (17). PIV has been shown to alter the lung mucosal defense by proteolytically degrading host immune effectors, such as interleukin 22 (IL-22) and surfactant proteins (18, 19). IL-22 is a critical facilitator of mucosal innate immunity whose effects maintain mucosal barrier integrity through specific targeting of mucosal epithelial cells to induce tissue repair and secretion of antimicrobial peptides (20). Surfactant proteins are innate immune effectors that bind, aggregate, and increase phagocytic uptake of bacterial pathogens in the lung (21). Furthermore, IL-22 and surfactant proteins have been shown to be essential for limiting pneumococcal pneumonia, and EF3030 is a pneumococcal strain that does not readily induce bacteremia (22, 23). Thus, a lung environment altered by PIV activity may favor the outgrowth of pneumococci, leading to enhanced lung pathology driven by pneumococcal virulence factors. Pneumolysin (Ply) is a cytotoxin and prominent virulence factor of pneumococci that has been shown to be essential during pulmonary infection and dissemination into the bloodstream (24–27). Based on these prior studies, we hypothesize that depletion of IL-22 by PIV in the localized lung environment exacerbates pneumococcal pneumonia and permits invasive disease mediated by Ply.

RESULTS

Pulmonary coinfection involving S. pneumoniae and P. aeruginosa expressing PIV results in pneumococcal bacteremia. EF3030 is a pneumococcal strain with limited virulence that does not readily induce bacteremia during pneumonia in a murine model of infection. We performed a coinfection of S. pneumoniae EF3030 with either the P. aeruginosa 103-29 parent strain (designated PA103-29) or a piv deletion mutant of PA103-29 to determine if secreted virulence factors of P. aeruginosa could stimulate pneumococcal bacteremia. In mice coinfected with EF3030 and PA103-29, there was no significant difference in pneumococcal densities in the lungs compared to mice infected with EF3030 alone (Fig. 1). However, pneumococci were isolated from the
blood of all mice coinfected with EF3030 and PA103-29. A deletion of *piv* in PA103-29 significantly reduced (*P* < 0.001) pneumococcal loads in murine lungs and abrogated pneumococcal bacteremia (Fig. 1).

Protease IV is active in murine lungs during infection. To further examine PIV-specific effects on pneumococcal pulmonary infections, we measured protease activity in lung homogenates of mice intratracheally administered phosphate-buffered saline (PBS), PA103-29, or PIV alone to determine a physiologically relevant concentration of PIV to administer (Fig. 2). When evaluating the secretome of *P. aeruginosa*, the Chromozym PL assay has been previously shown to be specific for measuring PIV activity (28). Compared to mock-treated mice, significantly higher (*P* < 0.05) proteolytic activity was measured in the lungs of mice during PA103-29 infection. Administration of 10 μg PIV resulted in a nonsignificant 0.25-fold decrease in proteolytic activity compared to proteolytic activity in the lungs of mice during active infection (Fig. 2).

PIV exacerbates pneumococcal pneumonia that develops into highly fatal bacteremia in a murine model of infection. We investigated whether the proteolytic effects of PIV could augment the virulence of EF3030 in a C57BL/6 murine pneumonia model. LasB is a *Pseudomonas* protease that cleaves a number of host proteins, including surfactants, but this specific protease does not cleave IL-22 (18). Therefore, LasB was used as a negative control for IL-22 cleavage. EF3030 combined with PIV

![Figure 1](http://msphere.asm.org/)

**FIG 1** *P. aeruginosa* expressing protease IV (PIV) during murine lung coinfection with *S. pneumoniae* EF3030 stimulates pneumococcal bacteremia. C57BL/6 mice were intratracheally inoculated with 10^6 CFU of *P. aeruginosa* 103-29 (PA103-29) or PA103-29 piv deletion mutant followed by intratracheal inoculation of 10^6 CFU of *S. pneumoniae* EF3030 at 24 h after *Pseudomonas* infection. Pneumococcal burden in the lungs and blood was enumerated by plating on selective blood agar 48 h after pneumococcal infection. Deletion of *piv* in PA103-29 significantly reduced pneumococcal load in the lung and abrogated pneumococcal bacteremia. Each symbol represents the value for a single mouse, and each bar represents the mean for that group of mice. Data are representative of two independent studies. *, *P* < 0.05; **, *P* < 0.01.

![Figure 2](http://msphere.asm.org/)

**FIG 2** Protease activity in murine lungs at 24 h after intratracheal administration. C57BL/6 mice were intratracheally inoculated with phosphate-buffered saline (PBS; control), 10^6 CFU of *P. aeruginosa* 103-29 (PA103-29), or 10 μg protease IV (PIV). At 24 h after intratracheal administration, protease activity in murine lungs was measured on a spectrophotometer using the chromogenic substrate Chromozym PL. Optical density readings at 405 nm were converted into proteolytic units per milliliter as described by the manufacturer. Protease activity significantly increased during PA103-29 infection, and there was no significant difference between protease activity measured during active infection and that measured during administration of PIV alone. Data represent two independent studies performed in triplicate. Error bars represent standard errors of the means. *, *P* < 0.05; ns, no significant difference.
significantly increased ($P < 0.01$) bacterial burden during pneumococcal pneumonia (Fig. 3A) and allowed the infection to advance to significant bacteremia ($P < 0.05$) compared to EF3030 alone (Fig. 3B). Heat-inactivated PIV or LasB did not significantly increase bacterial burden during pneumococcal pneumonia compared to EF3030 alone (Fig. 3A). However, there was a strong trend ($P = 0.0625$) toward increased bacteremia in mice inoculated with LasB compared to EF3030 alone (Fig. 3B). PIV and EF3030 combined inoculation resulted in 100% mortality, while all other inoculations caused very low or no mortality by 2 days postinfection (see Fig. 5).

PIV increases host susceptibility to Ply. Pneumolysin (Ply) is a virulence factor of pneumococcus that compromises alveolar-capillary barrier function, leading to permeability and flooding of alveoli (29). Since PIV enhanced pneumococcal pneumonia in a murine model (Fig. 3), we examined whether this increase in pathology was a result of additive Ply activity. Whether PIV was administered or not, there was no significant difference in bacterial load in the lungs (Fig. 4A) or blood (Fig. 4B) of mice during experimental pneumonia when infected with the Ply-deficient strain JLB12. Higher CFU were recovered from mice infected with JLB12 alone (Fig. 4A) than from mice infected with EF3030 alone (Fig. 3A), with some mice infected with JLB12 alone becoming bacteremic (Fig. 4B). Nonetheless, all mice infected with either EF3030 or JLB12 alone survived through 2 days postinfection (Fig. 5). Moreover, PIV administered with EF3030 resulted in higher bacterial burdens in murine lungs and blood (Fig. 3) as well as significantly increased mortality (Fig. 5) compared to combined PIV and JLB12-infected mice (Fig. 4 and 5).

PIV does not impact pneumococcal adherence to human pulmonary epithelial cells. Pneumococcal adherence to human pulmonary epithelial cells was examined to determine if PIV alters *S. pneumoniae* EF3030 interactions with cells relevant to bacterial pneumonia. PIV coadministered with bacterial cells did not impact adherence to epithelial cells in either EF3030 or JLB12, a ply mutant strain of EF3030 (Fig. 6). However, a deletion of *ply* significantly increased ($P < 0.05$) pneumococcal adherence to pulmonary epithelial cells (Fig. 6). EF3030 and JLB12 were not able to invade epithelial cells with or without the presence of PIV (data not shown).
PIV and Ply induce neutrophil (polymorphonuclear leukocyte [PMN]) recruitment and lung damage during active pneumonia. We used histology to visualize relative lung pathology during active pneumonia. Intratracheal administration of EF3030 and PIV led to massive infiltration of immune cells and shrinkage of alveolar spaces with evidence of necrosis and abscess formation (Fig. 7, panels 3 and 4). No indication of pneumonia was observed when mice were inoculated with PBS or with EF3030 combined with heat-inactivated PIV (Fig. 7, panels 1 and 2 and panels 5 and 6, respectively). Mice inoculated with a combination of active PIV and the Ply-deficient strain JLB12 displayed evidence of focal, mild pneumonia characterized by restricted infiltration of immune cells and attenuated shrinking of alveolar air spaces (Fig. 7, panels 7 and 8).

PIV depletes IL-22 levels in vivo during active pneumococcal pneumonia. IL-22 is required for host control of pneumococcal pneumonia, and IL-22 is cleaved by Pseudomonas PIV but not LasB (18, 22). To assess if IL-22 cleavage was occurring in vivo, we measured IL-22 levels in murine lungs during infection. IL-22 levels were signifi-

FIG 4 Pneumolysin (Ply) is required for PIV-mediated enhancement of pneumococcal pneumonia in a murine model. Mice were intratracheally inoculated with 10⁶ CFU of an EF3030 ply mutant (JLB12) with or without 10 μg of Pseudomonas PIV. Mouse lungs and blood were collected 2 days following lung infection, and pneumococcal CFU were enumerated by plating on BA. PIV did not significantly enhance pneumococcal burden in the lungs (A) or blood (B) of mice infected with a pneumolysin-deficient mutant. Data shown are representative of two independent studies (n = 10). Error bars represent standard errors of the means. ns, no significant difference.

FIG 5 Pseudomonas PIV and pneumococcal pneumolysin contribute to murine mortality during experimental pneumonia. Murine survival was monitored for 2 days following pulmonary infection. Survival rates are expressed as percentages (PIV, n = 8; EF3030, n = 12; EF3030 + PIV, n = 8; EF3030 + PIV³, n = 10; EF3030 + LasB, n = 6; JLB12, n = 10; JLB12 + PIV, n = 11). Mice infected simultaneously with Pseudomonas PIV and S. pneumoniae EF3030 expressing pneumolysin experienced 100% mortality, while all other inoculations resulted in very little to no mortality by 2 days postinfection. Data are representative of two independent experiments. Survival curves were determined to be significantly different (P < 0.0001) by log rank (Mantel-Cox) test.
cantly reduced ($P < 0.05$) when PIV was administered in combination with either EF3030 (Fig. 8A) or JLB12 (Fig. 8B) compared to IL-22 levels in mice infected with EF3030 alone or JLB12 alone, respectively. When mice were inoculated with heat-inactivated PIV or LasB, there was no significant difference in IL-22 levels compared to inoculation with EF3030 alone (Fig. 8A). IL-22 levels were also higher in mice infected with EF3030 (Fig. 8A) than in mice infected with the Ply-deficient strain JLB12 (Fig. 8B).

**DISCUSSION**

In this study, we demonstrate that a *Pseudomonas* protease IV (PIV) augments the virulence of an *S. pneumoniae* strain in a murine model of acute lung infection. Our findings reveal that PIV and Ply, virulence factors exclusively expressed by two separate pathogens, additively potentiate an active lung infection. The coadministration of active PIV and Ply-expressing strain EF3030 resulted in the highest bacterial loads and severe bacteremia (Fig. 3), as well as 100% mortality (Fig. 5) and enhanced immune cell infiltration leading to large abscess formation (Fig. 7, panel 4). This severe pathology must be attributed to a protease function that is not shared by PIV and LasB, as LasB coadministration with EF3030 did not result in significantly increased pulmonary bacterial loads, bacteremia, or mortality (Fig. 3 and 5). Since PIV did not impact interactions of *S. pneumoniae* EF3030 with pulmonary epithelial cells (Fig. 6), it is likely that altered immune signaling events are responsible for increased pathology within our murine model of lung infection. Both PIV and LasB cleave host surfactant proteins, but PIV exclusively cleaves IL-22 (18). To our knowledge, the activity of exogenously administered PIV has not been previously investigated in the lung. Our findings were able to quantify PIV activity (Fig. 2) and confirm the depletion of IL-22 by PIV *in vivo* (Fig. 8). This observation during an active murine lung infection is consistent with previous findings of IL-22 degradation in tracheal aspirates of *P. aeruginosa*-infected patients (18). Therefore, it is likely that IL-22 depletion in the lung is allowing for enhanced pneumococcal burden. PIV is a very weak immunogen (30), so increased pathology was not an effect of PIV immunogenic properties perpetuating deleterious inflammation. Accordingly, specific activity of PIV was required, as heat inactivation of PIV attenuated pathological findings (Fig. 3, 5, and 7). Although IL-22 played a large role in severity of pneumonia in our studies, the role of surfactants cannot be overlooked. LasB coadministration with EF3030 did enhance bacteremia, albeit not to statistical significance (Fig. 3B). Since surfactants have been shown to be important in restricting pneumococcal pneumonia (23), it is likely that the compounded effects of both
Nonetheless, increased pathology was not solely dependent on the activity of PIV. PIV coadministration with Ply-deficient strain JLB12 did not enhance pneumococcal pneumonia or bacteremia (Fig. 4) and resulted in restricted, focal evidence of pneumonia (Fig. 7). Surprisingly, we recovered higher pneumococcal CFU from the lungs of PIV and Ply-deficient strain JLB12 coadministration mice compared to Ply-positive strain EF3030 and Ply-positive control strain EF3030 Δply mice. Preserved airspaces and healthy tissues were observed in the inactivated PIV (panels 5 and 6) and PBS (panels 1 and 2)-treated mice. Arrows indicate areas of acute inflammatory cells within the lung parenchyma, with the largest abscess formation seen within panel 4.

FIG 7 P. aeruginosa PIV and S. pneumoniae producing pneumolysin induce massive neutrophil recruitment and lung damage during active murine pneumonia. Histological analysis of murine lung sections at magnifications of ×4 (left) and ×40 (right). Data are representative of two biological replicates of mice inoculated with PBS (panels 1 and 2), S. pneumoniae EF3030 and PIV (panels 3 and 4), EF3030 and heat-inactivated PIV (PIVIN; panels 5 and 6), and JLB12 (EF3030 Δply) and PIV (panels 7 and 8). EF3030 and PIV combined inoculation resulted in severe, diffuse pneumonia (panels 3 and 4) compared to restricted, focal pneumonia during JLB12 infection (panels 7 and 8). Preserved airspaces and healthy tissues were observed in the inactivated PIV (panels 5 and 6) and PBS (panels 1 and 2)-treated mice. Arrows indicate areas of acute inflammatory cells within the lung parenchyma, with the largest abscess formation seen within panel 4.
mice infected with JLB12 alone than from mice infected with EF3030 alone (Fig. 3A and 4A), and 50% of mice inoculated with JLB12 alone were bacteremic at 2 days postinfection (Fig. 4B). However, increased adherence to pulmonary cells (Fig. 6) and decreased induction of protective IL-22 (Fig. 8) afforded by JLB12 compared to EF3030 may have allowed for the increased pneumococcal burden in the lung and breakdown of epithelial barrier integrity. The major cellular sources of IL-22 during pneumococcal pneumonia are innate-like cells (ILCs) and thymocyte cells (T cells) expressing γδ T-cell receptors, which are stimulated by IL-23 secreted from activated dendritic cells and macrophages in the lung (22, 31). Ply has been shown to activate macrophages through Toll-like receptor 4 (TLR-4) signaling (32). Thus, decreased activation of macrophages responsible for producing proinflammatory cytokines is likely contributing to the decreased induction of IL-22 and enhanced bacteremia in JLB12-infected mice (Fig. 4B and 8).

Overall, innate mucosal barriers to infection play an important role in preventing pneumococcal invasive disease (PID). Altered lung environments provided by comorbidities or coinfection that lead to microbiome dysbiosis and pathogen outgrowth result in intimate interactions between invading pathogens. Specifically, our study reveals that a secreted *Pseudomonas* virulence factor dampens the host immune response and intensifies disease severity during *S. pneumoniae* pulmonary infections. Our findings also show that *Pseudomonas* PIV and *Streptococcus* Ply production have a combinatorial effect on disease severity. Unfortunately, the effects of these combined virulence factors cannot be alleviated with antibiotic usage. Since Ply is not secreted and is released upon bacterial lysis (33, 34), bactericidal antibiotics that cause a rapid release of Ply could result in major lung damage. In an altered lung such as the environment created by PIV, increased damage mediated by rapid Ply release could be catastrophic. Thus, studies examining targeted therapeutics that minimize damage created by proteases and cytotoxins are necessary in order to develop proper treatment strategies for patients with critical lung infections.

**MATERIALS AND METHODS**

**Materials.** Recombinant *P. aeruginosa* protease IV (PIV) was purified and assayed using Chromozym PL (Roche) as previously described (35). PIV was heat inactivated at 60°C for 1 h, and inactivity was verified by gelatinase zymography. *P. aeruginosa* protease LasB was purchased from Elastin Products Company, Inc. Human pulmonary epithelial cell line A549 (ATCC CCL-185) was obtained from Stephen Stray (University of Mississippi Medical Center, Jackson, MS) and authenticated by the American Type Culture Collection on 23 October 2017. Mouse IL-22 Ready-Set-Go! was purchased from Invitrogen.
CS7BL/6 (B6) mice were bred at the University of Mississippi Medical Center or purchased from The Jackson Laboratory.

**Bacterial strains and growth conditions.** *S. pneumoniae* strains were grown at 37°C with 5% CO₂ in Todd-Hewitt broth supplemented with 0.5% yeast extract (THY) or on sheep blood agar (BA) containing 0.5 μg/ml gentamicin. EF3030 is a serotype 19F *S. pneumoniae* strain (36). An EF3030 isogenic pneumolysin (ply) deletion mutant, JLB12, was created by allelic replacement of ply with an erythromycin resistance cassette amplified from ΔPLY2 as previously described (37). JLB12 was cultured using selective medium containing 3 μg/ml erythromycin. *P. aeruginosa* strains were grown at 37°C on MacConkey agar (MA) or in Luria-Bertani (LB) broth with shaking at 180 rpm. *P. aeruginosa* 103-29 and the isogenic protease IV (piv) deletion mutant have been previously characterized (38). During murine pulmonary coinfections, lung and blood samples were plated on BA containing 200 μg/ml polymyxin B to select for *S. pneumoniae* growth and MA without antibiotics to select for *P. aeruginosa* growth.

**Murine pulmonary infections.** Adult B6 mice (10 to 12 weeks old, 20 to 25 g) were anesthetized with an intraperitoneal injection of 500 μl Avertin (6.25 mg tribromoethanol dissolved in 2.5% amylene hydrate solution) and intratracheally inoculated with 50 μl containing 10⁶ CFU pneumococci or 10 μg of PIV. For bacterial coinfections, mice were inoculated with 50 μl containing 10⁶ CFU of *Pseudomonas* followed by 50 μl containing 10⁶ CFU of pneumococci at 24 h after *Pseudomonas* infection. For mixed exogenous administration, mice were inoculated with 50 μl containing 10⁶ CFU of pneumococci and 10 μg of either PIV, heat-inactivated PIV, or LasB. The Chromozym PL assay (Roche) was used according to the manufacturer’s instructions to measure the proteolytic activity in fresh lung homogenates of mice administered PIV or infected with 10⁶ CFU of *P. aeruginosa* 103-29. Mice were monitored for survival and euthanized at 2 days postinfection. Blood was collected by retroorbital bleed or from the heart. Lungs were removed and homogenized in 2 ml PBS. Blood and lung samples were plated on selective BA or MA to enumerate CFU per milliliter recovered from each mouse. Mouse studies were approved by the University of Mississippi Medical Center Institutional Animal Care and Use Committee in accordance with National Institutes of Health laboratory animal use guidelines.

**Pneumococcal adherence to human pulmonary epithelial cells.** Adhesion assays were performed as previously described (39). Briefly, human A549 pulmonary epithelial cells that had reached at least 90% confluence in 24-well plates were washed three times with PBS and incubated with approximately 10⁶ CFU of pneumococci in each well. Pneumococci were allowed to adhere for 30 min before epithelial cells were washed three times with PBS, lifted with 0.25% trypsin-EDTA, suspended in PBS, and plated on BA to enumerate adherent pneumococcal CFU. Recovered pneumococcal CFU were adjusted by standardizing inocula to 10⁶ CFU.

**Histology of murine lungs.** At 24 h postinfection, mice were euthanized and all five lobes were fixed in modified Davidson’s fixative (20% ethanol, 6% acetic acid, 10% formaldehyde dissolved in water). Fixed tissues were processed and stained with hematoxylin and eosin (H&E) by Excalibur Pathology Inc. Stained slides were examined by a reviewer blind to the specimen source. Pneumonia was characterized by exudate of fibrin and neutrophils within the airspaces in order to determine inflammation loci (focal versus diffuse) and severity of pneumonia (severe versus mild).

**IL-22 levels in murine lungs.** Murine lungs were homogenized in PBS containing 1× Halt protease inhibitor cocktail (Thermo Scientific). Lung homogenates were centrifuged at 1,500 × g, and supernatants were collected for use in a sandwich enzyme-linked immunosorbent assay (ELISA). IL-22 levels in supernatants were quantified using the mouse IL-22 Ready-Set-Go! ELISA kit according to the manufacturer’s instructions.

**Statistical analysis.** Results were analyzed using the InStat program (Prism 4 software; GraphPad, San Diego, CA). The Mann-Whitney test was used to test significant differences in the means of two groups. The Kruskal-Wallis test with Dunn’s post hoc comparisons was used for analysis of differences in means for three or more groups. Survival curves were analyzed with a log rank test. A P value of <0.05 was considered statistically significant.

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J.L.B., A.R.C., R.J.O., and L.S.M. conceived the experiments. J.L.B., A.R.C., M.A.B., A.T., R.J.O., and L.S.M. designed the experiments. J.L.B., A.R.C., M.A.B., and H.R.P. performed the experiments. J.L.B. and K.V.A. analyzed the data. J.L.B. wrote the original manuscript. All authors reviewed and edited the manuscript.

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