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Depletion of Alveolar Macrophages during Influenza Infection Facilitates Bacterial Superinfections

Hazem E. Ghoneim,*†‡ Paul G. Thomas,§ and Jonathan A. McCullers*†

Viruses such as influenza suppress host immune function by a variety of methods. This may result in significant morbidity through several pathways, including facilitation of secondary bacterial pneumonia from pathogens such as Streptococcus pneumoniae. PKH26-phagocytic cell labeling dye was administered intranasally to label resident alveolar macrophages (AMs) in a well-established murine model before influenza infection to determine turnover kinetics during the course of infection. More than 90% of resident AMs were lost in the first week after influenza, whereas the remaining cells had a necrotic phenotype. To establish the impact of this innate immune defect, influenza-infected mice were challenged with S. pneumoniae. Early AM-mediated bacterial clearance was significantly impaired in influenza-infected mice: ~50% of the initial bacterial inoculum could be harvested from the alveolar airspace 3 h later. In mock-infected mice, by contrast, >95% of inocula up to 50-fold higher was efficiently cleared. Coinfection during the AM depletion phase caused significant body weight loss and mortality. Two weeks after influenza, the AM population was fully replenished with successful re-establishment of early innate host protection. Local GM-CSF treatment partially restored the impaired early bacterial clearance with efficient protection against secondary pneumococcal pneumonia. We conclude that resident AM depletion occurs during influenza infection. Among other potential effects, this establishes a niche for secondary pneumococcal infection by altering early cellular innate immunity in the lungs, resulting in pneumococcal outgrowth and lethal pneumonia. This novel mechanism will inform development of novel therapeutic approaches to restore lung innate immunity against bacterial superinfections. The Journal of Immunology, 2013, 191: 1250–1259.
on pulmonary innate immune cells, particularly resident macrophages. We reasoned that early escape from the first line of defense in the lung could have profound effects on immunity to a variety of pathogens, including secondary bacterial invaders. Enhanced bacterial growth and replication through this mechanism could allow enhanced expression of virulence factors and the resulting inflammatory response. We used standard murine influenza virus infection and coinfection models (20), in which BALB/c female mice are infected intranasally by a sublethal dose of influenza A virus; this primary infection may then be followed by a sublethal dose of \textit{S. pneumoniae} at different time points after influenza infection. However, we improved on this model by introducing more specific gating strategies for distinguishing resident macrophage subtypes, coupled with in vivo labeling of lung-resident macrophages before influenza infection. This allowed accurate tracking of resident macrophages during the course of influenza infection. We demonstrate that influenza virus depletes AMs. This innate defect impairs early bacterial clearance, supporting development of secondary bacterial pneumonia. These data have implications for understanding virus-induced host immune suppression that may lead to improved prevention and treatment of primary or secondary infections in the lungs.

Materials and Methods

\textbf{Influenza viruses}

We used the St. Jude Children’s Research Hospital strain of mouse-adapted influenza virus A/Puerto Rico/8/34 (H1N1), referred to as “PR8,” as well as the human clinical isolate of the pandemic influenza virus A/California 04/09 (H1N1), referred to as “pdm H1N1.” All viruses were passaged once through Madin-Darby canine kidney cells, stocks were grown by a single passage through eggs, and allantoic fluid was stored at ~80°C. The viral titers of the stocks were characterized via median tissue culture-infective dose (TCID\textsubscript{50}) assay in Madin-Darby canine kidney cells.

\textbf{Bacterial strains}

\textit{S. pneumoniae} A66.1, a type 3 encapsulated strain, was engineered to express luciferase (Kevin Francis and Jun Yu, Xenogen Corporation, Alameda, CA). Pneumococci were grown in Todd Hewitt broth (Difco Laboratories, Detroit, MI) to an OD\textsubscript{600} of ~0.4 and then frozen at ~80°C mixed 2:1 with 5% sterile glycerol. The titers of the frozen stocks were quantitated on tryptic soy agar (Difco Laboratories, Detroit, MI) supplemented with 3% v/v sheep erythrocytes (blood agar). In all instances, the infectious dose administered was confirmed by serial dilution and plating of the bacterial suspension on blood agar plates.

\textbf{Mice}

Six- to 8-wk-old female BALB/c mice (Jackson Laboratory, Bar Harbor, ME) were maintained in a Biosafety Level 2 facility in the Animal Resource Center at St. Jude Children’s Research Hospital. Animals were given general anesthesia that consisted of 2.5% inhaled isoflurane (Baxter Healthcare Corporation, Deerfield, IL) before all interventions, and all studies were approved by the Animal Care and Use Committee at St. Jude Children’s Research Hospital.

\textbf{Infectious model}

Infectious agents were diluted in sterile PBS and administered intranasally in a volume of 100 \(\mu\)l (50 \(\mu\)l nostril) to anesthetized mice held in an upright position. In all experiments of influenza infection, PR8 influenza virus was given at a dose of 25 doses infectious for 50% of tissue culture wells (TCID\textsubscript{50}) per 100 \(\mu\)l per mouse, which caused ~10% weight loss and no mortality when given alone. In coinfection experiments, PR8 infection was followed at the specified time point by pneumococcal challenge with 200 CFU/mouse. Infected mice were weighed and assessed daily for illness and mortality for 7 d after pneumococcal challenge. The virus infectious dose in the pdm H1N1 experiment was 600 TCID\textsubscript{50} per mouse, which caused morbidity and weight loss comparable with PR8 infection (data not shown).

\textbf{In vivo labeling of lung-resident macrophages}

We intranasally administered 100 \(\mu\)l of 10 mM PKH26-phagocytic cell labeling dye (Sigma-Aldrich, St. Louis, MO) into anesthetized mice 5 d before influenza or mock infection, as previously described (21).

\textbf{Imaging of live mice}

Mice were then imaged for 60 s using an IVIS CCD camera (Caliper Life Sciences, Alameda, CA) daily after pneumococcal challenge to monitor in vivo pneumococcal pneumonia development. Total photon emission from selected and defined areas within the images of each mouse was quantitated using LivingImage software (Caliper Life Sciences) as described previously (22) and expressed as the flux of relative light units per minute. Pneumonia was defined as visible bioluminescence within the thorax and detection of a flux of >11,000 relative light units per minute.

\textbf{Flow cytometric analysis of immune cells in bronchoalveolar lavage fluid, postlavage lungs, and mediastinal lymph nodes}

After euthanasia by CO\textsubscript{2} inhalation, the trachea was exposed and cannulated with a 24-gauge plastic catheter (Becton Dickinson Infusion Therapy Systems, Sandy, UT). Lungs were lavaged four times with 1 ml cold sterile HBBS supplemented with 0.1 mM EDTA. The whole lungs after lavage or mediastinal lymph nodes were harvested and physically homogenized by syringe plunger against a 40-\(\mu\)m cell strainer and washed in FACS buffer consisting of HBBS, 0.1 mM EDTA, and 1% heat-inactivated FBS. Cell suspension of bronchoalveolar lavage fluid (BALF), postlavage lung homogenate, or mediastinal lymph nodes homogenate was centrifuged at 4°C, 350 \(\times\) g for 7 min, and the BALF supernatant was stored at ~80°C. Flow cytometry (LSRII and LSRII Fortessa; Becton Dickinson, San Jose, CA) was performed on the cell pellets after incubation with 75 \(\mu\)l of 1:200 dilution of Fc block (anti-mouse CD16/CD32; BD Bioscience, San Jose, CA) on ice for 10 min, followed by surface marker staining with anti-mouse Abs: CD11c (eFluor 450), F4/80 (FITC), Ly6G (PerCp-Cy5.5), Ly6C (allophycocyanin), and CD11b (allophtycocyanin-eFluor 780; eBioscience, San Diego, CA). The data were analyzed using FlowJo 8.8.6 (Tree Star, Ashland, OR) where viable cells were gated from a forward scatter/side scatter dot plot. First, we gated out neutrophils (CD11b\textsuperscript{hi} Ly6G\textsuperscript{hi}), then subgated macrophages (CD11c\textsuperscript{lo} F4/80\textsuperscript{lo}) based on CD11b surface expression into AMs (CD11b\textsuperscript{lo}), IMs (CD11b\textsuperscript{hi} F4/80\textsuperscript{lo}), and recruited exudate macrophages (CD11b\textsuperscript{hi}). Viable and nonviable cells were counted before surface marker staining, and the percentage viability was counted via the trypan blue exclusion method using a Cell Countess System (Invitrogen, Grand Island, NY). The absolute numbers of different cell types were calculated based on the proportion of viable events analyzed by flow cytometry as related to the total number of viable cells per sample. Live/dead aqua dye was added during flow cytometric surface staining to determine the total numbers of dead AMs and IMs. This dye binds to free amines after penetrating the impaired cell membrane of dead cells. First, AMs and IMs were gated as mentioned earlier but without prior gating for viable events based on a forward scatter/side scatter dot plot. Live/dead aqua positively stained AMs and IMs were gated as dead cells, and their numbers were calculated based on the proportion of dead events from the total events, analyzed by flow cytometry. Average mean fluorescence intensity of PKH26-PCL was measured for different subsets of macrophages in the PE channel.

\textbf{Confocal laser scanning microscopy of PKH26-labeled lungs}

Five days after in vivo PKH26-PCL labeling of lung macrophages, euthanasia by CO\textsubscript{2} inhalation was performed, and the trachea was exposed and cannulated with a 24-gauge plastic catheter (Becton Dickinson Infusion Therapy Systems). Lungs were harvested after instillation of 1.2 ml of 4% freshly prepared formaldehyde with PBS. Harvested lungs were fixed in 4% formaldehyde with PBS at room temperature for 1 h. Then every lung was washed with PBS and cut into four pieces before mounting into PBS in four wells of Nunc Lab-Tek chambered cover glass (Thermo Scientific, Ashville, NC) onto glass slides, and stained with Diff-Quik (Quik-Dip stain; Mercedes Medical, Sarasota, FL). Neutrophils, monocytes/macrophages, and lymphocytes were identified by morphometry, and images were taken from different representative fields of stained macrophages under high-power fields.
Early pneumococcal clearance determination

Early pneumococcal clearance was determined by measuring the pneumococcal count remaining within alveolar airspace. In brief, mice were euthanized by CO$_2$ inhalation 3 h after pneumococcal inoculation. BALF was harvested using sterile HBSS supplemented with 0.1 mM EDTA (lavage by 1 ml twice); then half the BALF was spread on blood agar plates supplemented with 0.4 mg kanamycin/ml blood agar to select for kanamycin-resistant luciferase-expressing pneumococci (22) and incubated at 37°C overnight.

Local GM-CSF treatment regimen

We treated anesthetized mice intranasally with 25 µg recombinant mouse GM-CSF (Invitrogen, Grand Island, NY) in 100 µl sterile PBS on days −1 and +1 before and after PR8 infection. We treated control mice with vehicle (PBS) only.

Statistical analysis

Comparison of survival and pneumonia development between groups of mice was done with the log-rank $\chi^2$ test on the Kaplan–Meier survival data. Bacterial titers, cell counts, and mean fluorescence intensity of PKH26-PCL in BALF and postlavage lung homogenates were compared between groups using ANOVA. Comparison of weight loss between groups of mice was done using Student $t$ test for pairwise comparisons. A $p$ value $<0.05$ was considered significant for these comparisons. Prism 4 for Windows (GraphPad Software, V 4.03) was used for all statistical analyses.

Results

In vivo labeling of lung-resident macrophages can distinguish AMs from IMs

Before examining any alterations in the lung-resident innate immune cells in influenza-infected hosts, we determined critical techniques for the differential analysis of the heterogeneous population of macrophages in both alveolar airspace and postlavage lung tissue. First, we refined a flow cytometry technique for gating resident macrophages so that clear differentiation between different cells types could be accomplished. We used the common markers CD11c$^+$ and F4/80$^+$, and added gating by another marker, CD11b ($\beta_2$ integrin). CD11b has been shown to be weakly expressed on AM cell surface (23–25) whereas highly expressed on granulocytes, exudate macrophages, monocytes, and some dendritic cells (26–29). Therefore, AMs are better gated as CD11c$^{hi}$ F4/80$^{hi}$ CD11b$^{dim}$ in BALF as shown recently (25, 30). This allowed differentiation in our flow cytometric analysis of the two major subsets of lung-resident macrophages in mock-infected mice. Thus, AMs could be gated as CD11c$^{hi}$ F4/80$^{hi}$ CD11b$^{ve}$ and IMs as CD11c$^{ch}$ F4/80$^{hi}$ CD11b$^{lo-int}$ in both BALF and postlavage lung homogenate (Fig. 1A).

To confirm our gating strategy for both subsets of lung-resident macrophages and to differentiate them from recruited macrophages, we did in vivo labeling of lung-resident macrophages before influenza infection using intranasally administered PKH26-PCL dye. Both subsets of lung-resident macrophages were intensely labeled by the dye, showing high mean fluorescence intensities of PKH26-PCL dye (MFI-PKH26). Nonetheless, significant differences in MFI-PKH26 were observed between AMs and IMs in both BALF and postlavage lung homogenates, with IMs showing significantly lower MFI-PKH26 than AMs (Fig. 1B). Selective labeling of lung-resident macrophages was also confirmed by confocal microscopy of fixed naive PKH26-labeled lungs showing the same pattern of difference in PKH26-MFI (Supplemental Fig. 1). These differences reflect the spatial and functional differences between the two major subsets of lung-resident macrophages, with IMs having less accessibility to the intranasally administered dye and lower phagocytic capacity than AMs (14, 24, 31). Thus, in vivo labeling method confirmed the validity of our flow cytometric gating strategy for the lung-resident macrophage subsets.

Influenza virus depletes AMs

To determine whether influenza infection causes any alteration in the numbers of lung-resident macrophages, we determined the percentages of lung-resident macrophages in flow cytometric dot
plots and calculated their absolute numbers. We infected BALB/c female mice intranasally with the mouse-adapted H1N1 influenza virus strain A/Puerto Rico8/34 (PR8) using a sublethal dose (25 doses infectious for 50% of tissue culture wells [TCID_{50}] per 100 μL). Seven days after influenza virus infection, >90% of the AM pool was depleted (Fig. 2A). To exclude the possibility that the observed depletion of AMs is unique to the use of a mouse-adapted influenza virus strain (PR8), we examined changes in the AM pool size using a human clinical influenza isolate from the 2009 influenza pandemic, pdm H1N1. We found that AMs were significantly depleted within 7 d after pdm H1N1 infection to levels comparable with PR8 infection (Fig. 2B).

**Tracking of AM and IM dynamic changes during influenza infection**

To track depletion kinetics of AMs and changes of the IM pool size during influenza infection, we again did in vivo labeling of lung-resident macrophages followed by sublethal PR8 influenza infection and studied different time points in the first 2 wk after influenza virus infection. We found that AMs were significantly depleted in the alveolar airspace shortly after PR8 infection starting on the first day postinfection (p.i.; Fig. 3A). Considering that some AMs may not be completely harvested in BALF and still adhere to the respiratory tract lining, we also analyzed postlavage lungs and found that higher numbers of AMs remained in the postlavage lung homogenate. These remaining AMs were significantly depleted starting 3 d p.i. compared with those in mock-infected mice (Fig. 3B). Consequently, 3 d after PR8 influenza infection was identified as the earliest time point of significant whole-lung AM depletion. The AM pool was partially replenished 9 d p.i. (in BALF only), whereas complete replenishment appeared 11 d p.i. (Fig. 3A, 3B). Conversely, the IM pool was not significantly depleted during the course of influenza infection; indeed, it was expanded at later time points (Fig. 3A, 3B).

To confirm the validity of the tracking process of lung-resident macrophages during the course of influenza infection, we measured dynamic changes in their MFI-PKH26. At any time point after influenza infection, AMs showed the highest MFI-PKH26 compared with IMs or recruited exudates macrophages (Supplemental Fig. 2), confirming our gating strategy that differentiates between AM and IM subsets. The MFI-PKH26 of AMs was significantly decreased shortly after influenza infection, however, no significant changes in their absolute numbers were observed in postlavage lungs in the first week of infection (Fig. 3B, 3D). In the second week, the IM pool significantly expanded with a significant increase in absolute numbers harvested from BALF (Fig. 3A).

To test whether these observed changes in PKH26-MFI of lung-resident macrophages were due to changes in their absolute numbers or due to instability of the dye, we measured the stability of PKH26-PCL dye in vivo in mock-infected lungs. PKH26-PCL dye demonstrated efficient labeling and stability inside the lung-resident macrophages for at least 14 d after mock infection (i.e., 19 d after labeling), thereby covering the tracking period during which we did the kinetics analyses (Supplemental Fig. 3).

We did further characterization of the different macrophage subsets in the lungs during influenza infection based on their surface expression levels of Ly6C Ag that can be used to differentiate between resident and recruited inflammatory macrophage subsets, in addition to their maturation stages (27, 28). We found that resident AMs did not express surface Ly6C, whereas resident lung IMs express intermediate levels, reflecting their intermediate stage of maturation and supporting the previous studies suggesting them as precursors for AMs (24, 32). In contrast, the recruited exudative macrophages with a CD11b^{high} phenotype showed significantly high expression levels of Ly6C (Supplemental Fig. 4).

**Influenza infection induces cell death of AMs**

To determine whether AM depletion is due to a cell death process, we measured the total numbers of dead AM cells during PR8 infection using a cell viability dye (live/dead aqua). The total numbers of dead AMs were significantly higher in both BALF and postlavage lungs of influenza-infected mice than in mock-infected mice (Fig. 4A). In contrast, there was no significant difference between the total numbers of dead IMs in influenza-infected and mock-infected mice (Fig. 4B). To determine the type of AM cell death process, we examined alveolar airspace macrophages for any morphologic changes associated with influenza infection. Diff-Quik–stained cytospin slides of BALF cells harvested from influenza-infected mice 3 d p.i. showed many macrophages with cellular damage manifestations. They were characterized by distorted nuclei and more vacuoles in cytoplasm than in mock-infected mice (Fig. 4C). This suggests that cell death was due to a secondary necrotic process.

To determine whether AMs were lost due to recruitment to the lung-draining lymph nodes, we examined the mediastinal lymph node 7 d after PR8 influenza infection and did not observe significant recruitment of PKH26-PCL–labeled AMs to it (data not shown).

**Influenza infection enhances susceptibility to secondary pneumococcal pneumonia**

Influenza-mediated death of AMs is likely to have significant effects on primary and secondary immunity. To explore one potential defect, we studied the permissiveness of influenza-infected hosts to secondary respiratory bacterial infections. We first infected BALB/c female mice intranasally with PR8 influenza virus using a sublethal dose (25 TCID_{50}). Influenza-infected mice showed mild morbidity manifested as loss of ∼10% of their original body weight within 7 d after influenza infection (Fig. 5A). At that time point, we induced secondary bacterial infection via intranasal administration of a small inoculum (200 CFU) of the serotype 3 clinical isolate of *S. pneumoniae*, A66.1. Influenza-infected mice showed high susceptibility to secondary pneumococcal infection with a significant increase in morbidity and continuous body weight loss (Fig. 5A).
All coinfected mice died within 3–5 d after pneumococcal inoculation (Fig. 5B). In contrast, all single influenza- or single pneumococcus-infected mice recovered quickly with no mortality (Fig. 5A, 5B). To determine whether the coinfected mice died because of pneumococcal pneumonia, we monitored pneumococcal growth in vivo through bioluminescent imaging of the lungs (22). Only coinfected mice experienced development of serious pneumococcal infections; pneumonia occurred within 48 h after pneumococcal inoculation (Fig. 5C).

**AM depletion during influenza infection impairs early pneumococcal clearance**

To determine whether the observed depletion of the AM pool contributes to enhanced susceptibility to secondary pneumococcal infection in influenza-infected hosts, AM phagocytic function was assessed in vivo by measuring early pneumococcal clearance with or without influenza infection. First, we determined the earliest time point at which a small pneumococcal inoculum (200 CFU) could be efficiently cleared within the alveolar airspaces of naive mice. Pneumococcal clearance was tested at different time points (1–4 h) after bacterial inoculation. Complete pneumococcal clearance was observed within 48 h after pneumococcal inoculation (Fig. 5C).

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higher doses of pneumococcus, whereas influenza virus-infected mice were unable to clear the basal inoculum (Fig. 5D).

**Influenza-infected mice demonstrate increased susceptibility to secondary pneumococcal pneumonia during the AM depletion phase**

To determine whether AM depletion during influenza infection correlates with susceptibility to secondary pneumococcal infection, we examined the early pneumococcal clearance at different time points after influenza infection. Early pneumococcal clearance was significantly impaired during PR8 infection from 3 until 9 d p.i. (Fig. 6A). This period of early pneumococcal clearance impairment closely mirrors the phase of the AM pool depletion observed during PR8 infection (Fig. 3A, 3B). Later, the impaired early pneumococcal clearance was restored (Fig. 6A).

To determine whether the impaired early pneumococcal clearance during influenza infection enhances susceptibility to secondary pneumococcal pneumonia development, we monitored mouse lungs via bioluminescence imaging 24 h after bacterial inoculation. Among the influenza-infected mice, 100% experienced development of secondary pneumococcal pneumonia when they were secondarily infected by pneumococcus 3, 5, or 7 d after influenza infection. In contrast, all influenza-infected mice that were secondarily infected either 1 or 14 d p.i. cleared the bacterial dose efficiently and did not have secondary pneumococcal pneumonia (Fig. 6B). Secondary pneumococcal pneumonia development was less frequent in influenza-infected mice that were secondarily infected by pneumococcus on 9 or 11 d p.i., with only 60 or 40% of pneumonic mice, respectively (Fig. 6B). As expected in this model, all coinfected mice that experienced development of secondary pneumococcal pneumonia died within a few days of bacterial pneumonia (Fig. 6C).

**Local GM-CSF treatment decreases secondary pneumococcal pneumonia development in influenza-infected mice**

To restore the early bacterial clearance efficiency that was impaired during the AM depletion phase, we tested local recombinant GM-CSF treatment as a means to accelerate replenishment of the depleted AM pool in influenza-infected mice. rGM-CSF was intranasally administered in two doses on days −1 and +1 before and after PR8 infection (Fig. 7A). First, we analyzed the effect of this GM-CSF treatment regimen on the size of the AM and IM pools. GM-CSF treatment in influenza-infected mice resulted in a significant expansion of the IM pool with partial replenishment of the AM pool (Fig. 7B).

To evaluate the efficacy of local GM-CSF treatment in the co-infection model, we administered pneumococcus 3 d after PR8 infection into GM-CSF–treated and mock-treated mice groups (Fig. 7A). Local GM-CSF treatment led to better early pneumococcal clearance in influenza-infected mice than in mock-treated, mock-infected mice. Conversely, early pneumococcal clearance remained impaired in mock-treated, influenza-infected mice (Fig. 7B). Interestingly, some of the GM-CSF–treated influenza-infected mice could efficiently clear the bacterial inoculum, whereas others could not. Next, we tested the ability of local GM-CSF treatment to prevent secondary pneumococcal pneumonia development after...
bacterial inoculation 3 d p.i. Local GM-CSF treatment protected 50% of coinfected mice against secondary pneumococcal pneumonia. Meanwhile, all mock-treated coinfected mice experienced development of pneumococcal pneumonia (Fig. 7D).

Discussion
Influenza is well-known to increase susceptibility to secondary bacterial infections, such as secondary pneumococcal pneumonia. The increased permissiveness of influenza-infected lungs to pneumococcal outgrowth suggests a defect in the host innate immune defenses that establishes a niche for bacterial infections. In this study, we examined a novel mechanism of influenza-mediated immune suppression: depletion of AMs. The resulting immune defect is likely to have pleiotropic effects on primary and secondary immunity. We examined one potential effect: bacterial escape from early innate immunity contributing to enhanced vulnerability to respiratory bacterial superinfections. Resident AMs are essential for early bacterial clearance and protection against bacterial infections (13, 21). We found that influenza infection induces depletion of AM pool for certain period during which early clearance of small pneumococcal inocula is significantly impaired. This influenza-associated damaging effect on AMs is correlated with enhanced vulnerability to secondary pneumococcal pneumonia. It is likely that other effects on immunity may be seen based on the lack of AMs at this crucial period of viral infection and clearance.

One observation stemming from this study is that examination of lung-resident macrophages in the setting of influenza infection should be cautiously performed. During influenza infection, diverse chemokines are upregulated in lungs, activating an influx of heterogeneous populations of innate immune cells, such as monocytes, inflammatory macrophages, and monocyte-derived dendritic cells. The dynamic changes in the phagocyte populations in influenza-infected lungs require critical methods to distinguish resident macrophages from recruited ones. Based on the surface phenotype of lung-resident macrophages in naive mice, we used an extensive gating strategy to analyze and differentiate between resident and recruited macrophages. Thus, AMs were gated as CD11ch F4/80hi CD11b−ve, whereas IMs and recruited macrophages were gated as CD11ch F4/80hi CD11blo-int and CD11ch F4/80hi CD11bhigh, respectively. Further characterization showed the ability of Ly6C marker to differentiate between resident and recruited macrophages. Thus, AMs were gated as CD11cch F4/80hi CD11b−ve, whereas IMs and recruited macrophages were gated as CD11cch F4/80hi CD11blo-int and CD11cch F4/80hi CD11bhigh, respectively. Further characterization showed the ability of Ly6C Ag is expressed by circulating blood monocytes and macrophages that can be recruited to tissues under inflammation conditions (27). However, Ly6C expression is downregulated during differentiation of blood monocytes or macrophages into tissue-resident macrophages after migration to tissue. Therefore, it can be added as a suitable marker to differentiate macrophage subsets and their stages of maturation into tissue-resident phenotype (28). Moreover, in vivo labeling of lung-resident macrophages was done to

FIGURE 5. Murine influenza–pneumococcal coinfection model. Body weight loss (A) and survival rate (B) of single pneumococcus-infected (PBS and S. pneumoniae, n = 6), single influenza virus-infected (influenza and PBS, n = 8), and coinfected (influenza and S. pneumoniae, n = 9) mice with 7-d interval between two doses in each group. Data represent the average ± SD. ***p < 0.001, Student t test at each time point compared with single influenza-infected mice group (A); ***p < 0.001, log-rank test on the Kaplan–Meier survival data (B). (C) Thorax bioluminescence of luciferase-expressing A66.1 pneumococcus from three representative coinfected mice showing development of pneumococcal pneumonia. Images were taken 48 h after secondary bacterial challenge using 200 CFU pneumococcus on day 7 after influenza infection. (D) Pneumococcal titers harvested 3 h after bacterial inoculation (inoculum of ≥200 CFUs) from alveolar airspace of mock- and influenza-infected mice 7 d p.i. are shown as percentage of inoculum. ***p < 0.001 compared with mock-infected (naive) mice.
Kaplan–Meier survival data (0.001 compared with day 1 or day 14 coinfection groups, log-rank test on the pneumonia development (200 CFUs) from the alveolar airspace of mock-infected and influenza-P. pneumonia harvested 3 h after bacterial inoculation (inoculum of 200 CFUs) from the alveolar airspace of mock-infected and influenza-infected mice 1, 3, 5, 7, 9, 11, and 14 d p.i. Secondary pneumococcal pneumonia development (B) and mortality (C) are manifested in influenza-infected mice that are secondarily infected by pneumococcus during the AM depletion phase (D.1 group: n = 5, D.3 group: n = 6, D.5 group: n = 4, D.7 group: n = 10, D.9 group: n = 5, D.11 group: n = 5, and D.14 group: n = 5). Other studies were not able to identify depletion of resident AMs because of some limitations in their analyses. Mostly they used gating strategies not sufficient to distinguish between resident AMs and recruited macrophages. The PKH26–PCL dye used to label macrophages forms fluorescent microparticles, which can be taken up by resident phagocytes in lungs and remain stable for >21 d (21), and emits high-fluorescence intensity for at least 19 d (21). Based on their analysis, they gated AMs as CD11c$^{+ve}$ Mac-1$^{-ve}$, decreased on day 3 p.i (35), which is consistent with our findings. In contrast, they did not analyze absolute numbers of cells.

In most studies of primary or secondary pneumococcal pneumonia in animal models, AM-mediated protection has not been rigorously studied (21). This appears mainly to be because of the use of high doses of pneumococcus (≥1 × 10^6 CFU) to induce a robust and reproducible pneumonia, doses that overwhelm the phagocytic capacity of resident AMs. Instead, an influx of neutrophils in the setting of development of type-specific Ab, the second line of innate immune defense coupled with the early adaptive response, was found in these models to be important for host protection against primary pneumococcal pneumonia (36). However, in the perhaps more physiologically relevant setting of a relatively small dose of pneumococcus (200 CFU) after sublethal mild influenza infection, we can demonstrate a strong effect mediated by AMs. This low bacterial inoculum successfully caused lethal secondary pneumococcal pneumonia within 48–72 h after bacterial inoculation, pneumonia that was dependent on an absence of AMs.

In vivo labeling of lung-resident macrophages in this study was a novel approach that allowed successful differentiation of AMs from recruited macrophages. Furthermore, monitoring changes of MFI-PKH26 for lung-resident macrophages reflected the dynamic changes in their absolute numbers during influenza infection. For instance, expansion of the AM pool during the recovery phase was accompanied by significant decrease in their MFI-PKH26. Nevertheless, the AM population showed the highest MFI-PKH26 of AMs, even during their replenishment phase, implies that the AM pool was replenished mainly via differentiation of the PKH26-labeled IMs into an AM surface phenotype rather than maturation of PKH26-unlabeled recruited blood macrophages. These results support the findings that the origin of the AM pool is mainly through differentiation of intermediate lung macrophages rather than self-proliferation (24). In contrast, the quick drop in MFI-PKH26 of IMs in the first week after influenza infection, without apparent changes in their absolute cell numbers, suggests that influenza infection may induce partial depletion of the IM pool. However, this partial depletion can be quickly recompensated by the proliferative capacity of IMs, maturation of the recruited macrophages, or both (14, 24).

Other studies were not able to identify depletion of resident AMs because of some limitations in their analyses. Mostly they used gating strategies not sufficient to distinguish between resident AMs, IMs, and recruited exudate macrophages. For example, gating for a single marker like F4/80 Ag or macrophage-specific esterase staining of macrophages isolated from BALF during influenza infection is targeting mixed populations of resident and recruited macrophages whose total cell numbers increase during infection (37). In addition, although the same in vivo labeling method was used before, AM depletion could not be identified during influenza (21). Based on their analysis, they gated AMs as CD11c$^{+ve}$ PKH26$^{+ve}$ that may be mixed with IMs. They analyzed AM numbers on day 9 after influenza infection in BALF only. Based on our model, we showed a certain period for AM depletion. Therefore, analyzing AMs at a single time point (day 9 p.i.) is not conclusive regarding AM depletion event during influenza infection. Furthermore, analyzing cells in BALF only may dismiss significant numbers of AMs remaining adherent in the lungs after lavage, leading to underestimation of resident AM pool size during comparison between influenza- and mock-infected mice. Likewise,
in another study, they did not show AM depletion, because of missing CD11b marker in their AM gating, as well as harvesting only BALF for kinetics analysis (17). Taken together, missing an important marker that distinguishes between different lung-resident macrophage subsets and recruited monocytic cells or using BALF only for analyzing resident AMs may lead to erroneous conclusions regarding their kinetics during infections.

GM-CSF is a cytokine with diverse functions critical for effective innate immunity in lungs. Among these activities, it regulates AM differentiation and activation (23, 38), enhances proliferation of resident pulmonary macrophages (28), and expands the pool of resident AMs (39). It also has an important role in pulmonary surfactant homeostasis (40). Recent studies showed that GM-CSF overexpression in lungs has prophylactic activity against lethal influenza and pneumococcal pneumonias (39, 41, 42). We found that local rGM-CSF treatment in influenza-infected mice induced significant expansion of the IM pool. In addition, absolute numbers of AMs increased under this treatment regimen in influenza-infected mice. This increase was not to the degree as that of IMs, possibly because they were analyzed at an early time point (3 d p.i.). Thus, our data support the previous studies showing GM-CSF enhances the proliferation capacity of lung-resident macrophages and maturation of AMs. Considering the higher proliferative capacity of IMs over AMs, our short treatment regimen could significantly expand the pool of IMs more than AMs. As a result of partial replenishment of the AM pool with pulmonary GM-CSF treatment, influenza-infected mice had partial restoration of efficient early pneumococcal clearance. Furthermore, they manifested improved protection against secondary pneumococcal pneumonia. Although pulmonary GM-CSF treatment has some drawbacks such as inflammatory activity (43) that can render its use in humans problematic by exacerbating inflammatory lung injury, the results of our study are promising. They suggest that strategies seeking to balance the protective AM replenishment effects and the adverse effect of exuberant inflammation induction via combining an adjunctive anti-inflammatory therapy with GM-CSF treatment might be possible.

Overall, this study suggests a novel mechanism of influenza-mediated immune suppression that resulted in increased vulnerability of influenza-infected hosts to bacterial superinfections. Resident AM depletion during influenza infection establishes a niche for secondary pneumococcal infection by altering early cellular innate immunity in the lungs, thereby allowing pneumococcal outgrowth causing lethal pneumococcal pneumonia. The precise functional characterization of this novel finding can change the way researchers look at the alteration of pulmonary cellular

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**FIGURE 7.** Local rGM-CSF treatment improved secondary pneumococcal pneumonia development in influenza-infected mice. (A) Intranasal administration of rGM-CSF into PR8-infected mice on days −1 and +1 before infection and p.i., followed by 200 CFU pneumococcus on day 3 improved early pneumococcal clearance from the alveolar airspace (C) and secondary pneumococcal pneumonia development (n = 9) (D) compared with mock-treated coinfected mice (n = 4). (B) The absolute numbers of AMs and IMs increased in GM-CSF-treated influenza-infected mice analyzed 3 d p.i. (n ≥ 4). *p < 0.05, **p < 0.01, ***p < 0.001, Dunnett’s multiple-comparison test (ANOVA), compared with mock-treated mock-infected mice (B, C). Bar graphs show the average ± SD. *p < 0.05 compared with day 1 or day 14 coinfection groups, log-rank test on the Kaplan–Meier survival data (D).
immunity during sublethal influenza infections. There are likely to be important effects on immunity beyond the bacterial escape studied in this article. Furthermore, these findings should open avenues for novel immunomodulating therapeutic interventions to prevent respiratory bacterial superinfections by quick replenishment of the critical innate immune effectors during both pandemic and seasonal influenza.

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Disclosures

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References


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Supplemental Fig. 1: Fluorescence microscopic examination of PKH26-labeled naive lungs confirms selective in vivo labeling of lung-resident macrophages.

(A) Photomicrographs show that both AMs (arrow) and IMs (asterisk) are selectively labeled by PKH26-PCL dye and emit red PKH26 fluorescence. Green autofluorescence can be detected from AMs, IMs, and elastin and collagen fibers. (B) AMs have significantly higher PKH26-MFI and autofluorescence compared with that of IMs measured from 3 different fields. Images were obtained by a Nikon Eclipse TE2000-E microscope (n=3).

**P < 0.01 Mann Whitney Student t-test. The bar graphs show the average ± SD.
Supplemental Fig. 2: Differential MFI of PKH26-PCL dye of different pulmonary macrophage subsets during influenza infection in BALF (A) and post-lavage lungs (B) are compared during the course of influenza infection (n ≥ 4). *P < 0.05, **P < 0.01 Dunnett's multiple comparison test (ANOVA) compared with AMs at each time point. The bar graphs show the average ± SD.
Supplemental Fig. 3: Fluorescence intensity of PKH26-PCL dye is stable in the resident pulmonary macrophages for 2 weeks after mock infection. % PKH26-MFI values are calculated as percentages of the average PKH26-MFI of AMs on day 3 after PBS administration (n ≥ 4). *P < 0.05 Dunnett's multiple comparison test (ANOVA), compared with % PKH26-MFI of AMs on day 3 after PBS. The bar graphs show the average ± SD.
Supplemental Fig. 4: Differential surface expression of Ly6C marker on different pulmonary macrophage subsets. Ly6C-MFI are compared at different timepoints after influenza infection. **P < 0.01, ***P < 0.001 Mann Whitney Student t-test, where AMs are compared with IMs, and IMs are compared with recruited Mφ at each time point (n ≥ 4). The bar graphs show the average ± SD.