Deletion of c-FLIP from CD11b^{hi} Macrophages Prevents Development of Bleomycin-induced Lung Fibrosis

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Abstract

Idiopathic pulmonary fibrosis is a progressive lung disease with complex pathophysiology and fatal prognosis. Macrophages (M Φ) contribute to the development of lung fibrosis; however, the underlying mechanisms and specific M Φ subsets involved remain unclear. During lung injury, two subsets of lung $M\Phi$ coexist: Siglec- $F^{\rm hi}$ resident alveolar M Φ and a mixed population of CD11b^{hi} M Φ that primarily mature from immigrating monocytes. Using a novel inducible transgenic system driven by a fragment of the human CD68 promoter, we targeted deletion of the antiapoptotic protein cellular FADD-like IL-1β-converting enzyme-inhibitory protein (c-FLIP) to CD11b^{hi} MΦ. Upon loss of c-FLIP, CD11b^{hi} MΦ became susceptible to cell death. Using this system, we were able to show that eliminating CD11b^{hi} MΦ present 7–14 days after bleomycin injury was sufficient to protect mice from fibrosis. RNA-seq analysis of lung MΦ present during this time showed that CD11bhi MΦ, but not Siglec- F^{hi} M Φ , expressed high levels of profibrotic chemokines and growth factors. Human M Φ from patients with idiopathic pulmonary fibrosis expressed many of the same profibrotic

chemokines identified in murine CD11bhi M Φ . Elimination of monocyte-derived M Φ may help in the treatment of fibrosis. We identify c-FLIP and the associated extrinsic cell death program as a potential pathway through which these profibrotic M Φ may be pharmacologically targeted.

Keywords: macrophage; fibrosis; c-FLIP; RNA-seq; IPF

Clinical Relevance

Pulmonary fibrosis is a progressive, deadly disease of unclear etiology; understanding the underlying mechanisms is critical for developing effective treatments. We have identified a specific subpopulation of mainly monocyte-derived macrophages that drive the development of fibrosis and have described their unique profibrotic signature using RNA-seq. Removing this population abrogates the development of fibrosis.

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Macrophages (M Φ) are known to be involved in fibrogenesis. Lung M Φ numbers are elevated both in patients with idiopathic pulmonary fibrosis (IPF), a progressive lung disease with complex pathophysiology and fatal prognosis (1, 2), and in murine models of pulmonary fibrosis (3, 4). However, the mechanisms by which M Φ impact fibrosis are unclear. The objectives of this study were to determine the roles played by M Φ subsets in the development of lung fibrosis and to determine if fibrosis could be attenuated by promoting M Φ apoptosis.

During homeostasis, there are two main types of $M\Phi$ in the murine lung: resident Siglec-F^{hi}CD11b^{lo} MΦ that reside in the alveolar space, and CD11bhiSiglec-Flo $M\Phi$ that occupy the lung interstitium. Siglec-F^{hi} MΦ (often called resident alveolar MΦ and highly expressing CD11c) arise during embryogenesis, populate the airspaces immediately after birth, and self-renew throughout life without replacement from circulating monocytes (5–9). CD11 b^{hi} M Φ (often called interstitial M Φ [IMs]) also arise during embryogenesis, but are restricted to the lung interstitium and found in much lower numbers than Siglec-Fhi alveolar $M\Phi$ (5, 6).

After lung injury, $M\Phi$ numbers expand in both the alveolar and interstitial compartments. Studies using lung-shielded bone marrow chimeras, targeted $M\Phi$ or monocyte depletion, and CCR2^{-/-} mice all suggest that monocytes recruited from the circulation are the dominant source of new $M\Phi$ (often termed recruited $M\Phi$) after injury (10-12). Much like homeostatic resident IMs, newly recruited $M\Phi$ express high levels of CD11b and low levels of Siglec-F. Accordingly, during acute inflammation, three main M Φ populations exist in the lung. These include two types of resident MΦ (Siglec-F^{hi} CD11b^{lo} alveolar MΦ and CD11bhiSiglecFlo IMs), present before onset of inflammation and that remain throughout its course, and a third population of recruited $M\Phi$ in the alveolus and interstitium (also CD11bhiSiglecFlo) that derive from monocytes and are only present in the context of disease.

Studies that separate $M\Phi$ subsets are imperative for clarifying their roles in fibrogenesis (13). However, a lack of tools for distinguishing and manipulating $M\Phi$ subsets had previously hampered this

objective. The identification of reliable cell surface markers that distinguish resident alveolar M Φ from other subtypes constitutes a recent advance (7, 10, 14, 15). In addition, we have developed a new transgenic system to target M Φ subsets (16). Whereas traditional drivers of cre recombinase (cre) constitutively affect all lung M Φ starting early in development, we recently described a system in which CD11b^{hi} M Φ can be selectively targeted in a time-dependent fashion. Targeting is achieved using a fragment of the human CD68 promoter coupled with the reverse tetracycline-controlled transactivator (hCD68rtTA). Administration of doxycycline to these animals rapidly activates tetracycline-responsive elements in CD11b^{hi} IMs and recruited MΦ. Siglec-F^{hi} $M\Phi$ are not affected (16). As described subsequently here, we leveraged this new method to selectively manipulate the survival of CD11b^{hi} MΦ during bleomycin-induced lung injury and test their contribution to fibrogenesis.

Appropriately timed apoptosis of lung $M\Phi$ is beneficial for resolution of inflammation and effective tissue repair. $M\Phi$ resistance to apoptosis is implicated in the development of murine fibrosis and human IPF (17). We have previously demonstrated that CD11bhi MΦ are recruited to the alveolar space during LPS-induced lung injury, but disappear within 2 weeks as a consequence of cell surface Fas receptor (Fas)-FasL-mediated cell death (10). In comparison, $M\Phi$ numbers remain elevated for over 6 weeks after bleomycin-induced lung injury and fibrosis. Eventual decline in $M\Phi$ number correlates with resolution of fibrosis (4). Treatment of mice with TNF- α after bleomycin injury accelerates MΦ decline and hastens the resolution of fibrosis (4).

FasL and TNF- α both activate the extrinsic cell death pathway. To induce cell death, this pathway requires the simultaneous presence of a death-inducing ligand, expression of the corresponding death receptor, and a permissive intracellular environment. Ligation of FasL or TNF- α to their cognate surface receptors triggers the transproteolytic cleavage of pro–caspase-8 homodimers to active caspase-8. Whereas low levels of active caspase-8 activate NF- κ B and promote survival, higher levels trigger programmed cell death (18, 19). Cellular FADD-like IL-1 β -converting enzyme–inhibitory

protein (c-FLIP) is a critical death effector domain–containing protein that restricts caspase-8 activation (20–23). Accordingly, loss of c-FLIP sensitizes murine and human $M\Phi$ to cell death (24–26).

We hypothesized that deleting c-FLIP in CD11b^{hi} MΦ would promote their death and attenuate the development of fibrosis. We crossed hCD68rtTA mice with tet-On-Cre and cFLIP^{fl/fl} mice to create animals in which c-FLIP could be conditionally deleted in CD11 b^{hi} IMs and recruited M Φ upon administration of doxycycline. Deletion of c-FLIP sensitized M Φ to programmed cell death and specifically reduced CD11bhi MΦ numbers in the lungs. Moreover, deletion of c-FLIP in $CD11b^{hi}$ M Φ at both early (Day 1) and delayed (Day 7) time points significantly attenuated the development of fibrosis. Finally, using improved surface marker identification (7), we assessed the transcriptome of CD11bhi MΦ during bleomycin injury to identify potential mediators of their profibrotic function.

Methods

Animals

This study was approved and performed in accordance with the ethics guidelines of the Institutional Animal Care and Use Committee at National Jewish Health. cFLIP^{fl/fl} mice (27) were provided by Dr. You-Wen He (Duke University, Durham, NC). Details on other strains are provided in the data supplement.

Bleomycin Injury

Bleomycin (TEVA Pharmaceuticals) was administered in a dose of 3 U/kg, adjusted for the weight of the mouse, in 50 μ l of PBS. Bleomycin was instilled intratracheally using a modified gavage needle. Mice were sedated with isoflurane (Baxter) before intratracheal instillations.

Flow Cytometry

Cells were stained with surface antibody panels to identify myeloid populations and analyzed using an LSRII flow cytometer (BD) and FlowJo software (Tree Star) or sorted using Moflo XDP-100 with Summit v5.1 software (Beckman Coulter, Dako). Details can be found in the data supplement.

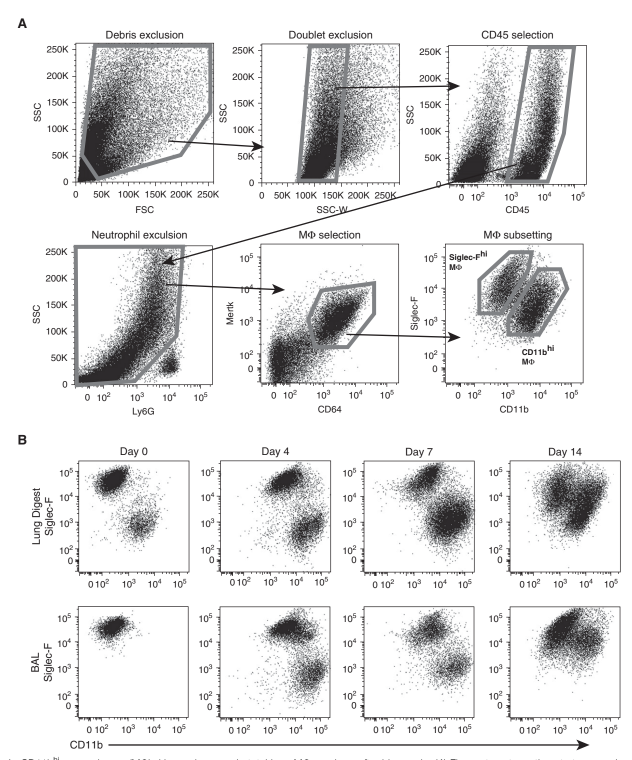


Figure 1. CD11b^{hi} macrophages (MΦ) drive an increase in total lung MΦ numbers after bleomycin. (A) Flow cytometry gating strategy used on lung digest and BAL to isolate Siglec-F^{hi} and CD11b^{hi} MΦ. Representative gating is shown on lung digest 14 days after bleomycin. Live cells are selected using size (forward scatter [FSC]) and granularity (side scatter [SSC]), doublets are excluded, and leukocytes are selected using CD45. Ly6G⁺ neutrophils are excluded, and CD64 and Mertk coexpression is used to select MΦ. (B) Siglec-F and CD11b are used to divide MΦ into two subpopulations. Representative dot plots from lung digest 4, 7, and 14 days after bleomycin are shown. (C and D) Total numbers of Siglec-F^{hi} and CD11b^{hi} MΦ were assessed in naive mice and mice 7 and 14 days after bleomycin. (C) MΦ in lung digest. (D) MΦ in BAL. Data from n = 3-5 mice/group shown as mean (±SEM). **P < 0.01, ***P < 0.001. IT = intratracheal.

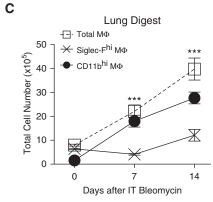


Figure 1. (Continued).

Measures of Caspase Activation

Cells recovered from BAL 7 days after bleomycin were adhesion purified to select for M Φ . Caspase-Glo (Promega, Madison, WI) assays were used per the manufacturer's instructions to measure caspase activity. Details can be found in the data supplement.

Assessing Fibrosis

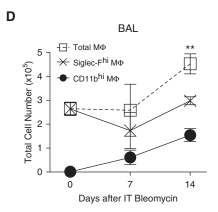
Immediately after death, static compliance, a measurement that reflects the static elastic recoil pressure of the lungs at a volume of 0.95 ml, was determined using flexiVent (SCIREQ). Lungs were perfused with PBS and the right lung was saved for hydroxyproline analysis (28), or lungs were inflated with low-melt agarose, fixed in 10% formalin overnight, then paraffin embedded; 10-µm slices were stained with Trichrome and imaged with an Aperio system (Leica Biosystems).

RNA Isolation and RT-PCR

RNA was isolated from M Φ using the RNeasy micro kit (Qiagen) and DNase treated as instructed by the manufacturer. RNA was then processed for RNA-seq or RNA was reverse transcribed to cDNA using Superscript IV VILO Master Mix (Invitrogen) per the manufacturer's instructions for RT-PCR. Primer sequences in can be found in the data supplement.

RNA-Seq

Isolated total RNA from sorted cells was processed for next-generation sequencing library construction as developed in the NJH Genomics Facility for analysis with a Life Technologies Ion Proton next-generation



sequencing platform. Details can be found in the data supplement.

Gene Expression Omnibus Data Sets

Microarray data from human M Φ (29) were obtained from the Gene Expression Omnibus (GEO; GEO accession no. GSE49072). Details can be found in the data supplement.

Statistical Analysis

Statistics for RNA-seq and GEO DataSet analyses were performed as described in the data supplement. For all other experiments, differences between two groups were examined using Student's unpaired t test. Differences among three or more groups were examined using ANOVA with Holm-Bonferroni's multiple comparison test *post hoc* analysis. A P value less than 0.05 was considered significant.

Results

CD11b^{hi} M Φ Account for the Increase in Total Lung M Φ after Bleomycin

MΦ contribute to lung fibrosis; however, the discrete contributions of MΦ subsets remain unclear. To develop a framework for studying the MΦ subsets, we first quantified MΦ during the initiation of fibrosis. We measured CD11 b^{hi} MΦ and Siglec- F^{hi} MΦ in BAL fluid and enzymatically digested lungs from bleomycin-treated mice, adapting previously published protocols (7, 10, 14, 15) (Figure 1A). For all subsets, MΦ were identified based on their coexpression of CD64 and MER proto-oncogene tyrosine kinase (Mertk), cell surface molecules that,

together, are specific for M Φ (30, 31). Notably, Ly6C hi "inflammatory" monocytes express low CD64 and negligible Mertk, and were thus excluded by our $M\Phi$ gating strategy (see Figures E1A and E1B in the data supplement). High expression of Siglec-F was recently identified as a marker of resident alveolar M Φ (7), and use of Siglec-F and CD11b was sufficient to distinguish M Φ subpopulations in the lung at all time points (Figure 1B). In contrast, use of CD11b in conjunction with other markers failed to clearly separate CD11bhi $M\Phi$ as inflammation progressed (Figures E1C and E1D). $M\Phi$ were examined in the first 2 weeks after injury, when fibrogenesis begins (4). Quantification of lung $M\Phi$ subsets revealed that CD11bhi $M\Phi$ were responsible for the increase in $M\Phi$ numbers in both the airspace and tissue compartments (Figures 1C and 1D).

CD11b^{hi} M⊕ Are Sensitized for Caspase-8–Dependent Cell Death after Deletion of c-FLIP

To specifically induce death of CD11 b^{hi} M Φ and block their accumulation, we conditionally deleted c-FLIP (gene cflar) using our newly described hCD68rtTA system (16). c-FLIP is a nonenzymatic caspase-8 homolog that dimerizes with caspase-8 and inhibits its proapoptotic activity. By crossing hCD68rtTA with tet-On-Cre and cFLIPfff mice, we were able to generate triple-transgenic mice in which administration of doxycycline leads to inducible deletion of cflar in CD11bhi IMs and recruited MΦ (but not Siglec-F^{hi} resident alveolar M Φ). For simplicity, we designated these animals as cFLIP $^{\Delta/\Delta}$. In comparison, control mice (designated cFLIP^{fl/fl}) were littermates possessing floxed cflar and tet-On-Cre, but lacking hCD68rtTA (Figure E2A). To validate this system, cFLIP $^{\Delta/\Delta}$ and cFLIP $^{fl/fl}$ mice were treated with bleomycin, but doxycycline was withheld. BAL M Φ were harvested 7 days later and doxycycline was added in vitro. Doxycycline led to the loss of cflar expression in $M\Phi$ from cFLIP^{Δ/Δ} mice; cflar was not deleted in $M\Phi$ from cFLIP $^{\rm fl/fl}$ animals (Figure E2B). c-FLIP deletion alone did not induce obvious spontaneous death of cultured MΦ.

We next sought, in ex vivo studies, to determine the effect of c-FLIP deletion on $M\Phi$ sensitivity to cell death by assessing quantitative changes in the activity of

caspase-8 (initiator caspase in extrinsic cell death) and caspase-3 (downstream terminal executioner caspase). We also examined phenotypic changes associated with cell death: loss of plasma membrane integrity and loss of cell adherence. In brief, $cFLIP^{\Delta/\Delta}$ mice and $cFLIP^{fl/fl}$ littermate controls were treated with bleomycin, and BAL M Φ were isolated 7 days later and placed in culture. Doxycycline was added to delete c-FLIP and an activating α-Fas antibody was added to trigger the extrinsic cell death pathway. We deliberately chose a dose of α-Fas that triggered minimal Fas activation. The caspase-8 inhibitor, Z-IETD-FMK, was added to some wells as a negative control. As shown in Figure 2A, when doxycycline or α -Fas alone was added to $cFLIP^{\Delta'/\Delta'}M\Phi$ there was no significant increase in caspase-8 activity. However, when α -Fas was added after doxycycline, caspase-8 activation increased. As anticipated, Z-IETD-FMK blocked this effect. Importantly, $M\Phi$ from littermate control mice (cFLIP^{fl/fl}) treated in parallel showed caspase-8 activity in response to α-Fas, but no additional effect of doxycycline (Figure 2B). Taken as a whole, these data demonstrate that death receptor ligation induces enhanced activation of caspase-8 when c-FLIP is deleted in lung $M\Phi$.

To determine whether c-FLIP deletion would lead to downstream activation of caspase-3, ex vivo studies were repeated, again using BAL MΦ from bleomycin-treated cFLIP $^{\Delta/\Delta}$ and cFLIP $^{fl/fl}$ mice. Caspase-3 activity showed the same pattern as caspase-8 activity. Cotreatment with doxycycline (to delete c-FLIP) and the α-Fas significantly enhanced caspase-3 activation (Figure 2C), whereas c-FLIP deletion alone did not. Moreover, blocking caspase-8 with the inhibitor, Z-IETD-FMK, prevented caspase-3 activation, confirming that caspase-3 activation was downstream of caspase-8 activity (Figure 2C). MΦ from cFLIP^{fl/fl} littermate control mice did not respond to doxycycline (Figure 2D).

Loss of plasma membrane integrity was observed in cFLIP $^{\Delta/\Delta}$ M Φ cotreated with doxycycline and α -Fas, but not in M Φ treated with α -Fas alone or in cFLIP $^{fl/fl}$ M Φ , providing additional evidence that c-FLIP deletion enables M Φ extrinsic cell death (Figure E2C). Leaving cFLIP $^{\Delta/\Delta}$ M Φ in culture for more than 12 hours after addition of α -Fas resulted in decreased cell

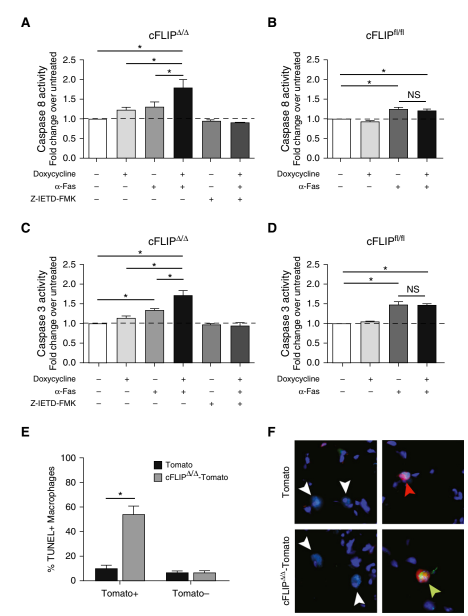


Figure 2. Cellular FADD-like IL-1 β -converting enzyme-inhibitory protein (cFLIP) $^{\Delta/\Delta}$ mice delete c-FLIP in CD11 b^{hi} M Φ and sensitize CD11 b^{hi} M Φ to caspase-8-dependent cell death. (A-D) Caspase activity was measured using Caspase-Glo luminescent assays. M Φ were collected by lavage 7 days after bleomycin and treated with doxycycline (20 ng/ml) in vitro for 36 hours to induce deletion of c-FLIP. Doxycycline was removed and cells were treated with α -Fas activating antibody (5 μ g/ml) for a further 8 hours (caspase-8) or 12 hours (caspase-3). Some wells were treated with the selective caspase-8 inhibitor, Z-IETD-FMK, at 10 µM for 30 minutes before α -Fas. (A and B) Caspase-8 activity of CD11b^{hi} M Φ from cFLIP^{Δ/Δ} (A) or cFLIP^{β/hi} mice (B) after doxycycline treatment and/or stimulation with activating α -Fas antibody. (C and D) Caspase-3 activity of CD11b^{hi} M Φ from cFLIP^{Δ/Δ} mice (C) or cFLIP^{fl/fl} mice (D) after doxycycline treatment and stimulation with activating α -Fas antibody. (E and F) Control Tomato-reporter mice and cFLIP $^{\Delta/\Delta}$ -Tomato mice were treated with bleomycin and then started on doxycycline at Day 7. At Day 10, lungs were collected, inflated, and frozen with optimal cutting temperature compound for histology. Double-stranded DNA breaks were labeled with terminal deoxynucleotidyl transferase dUTP nick end labeling (TUNEL; green), and tissue sections were costained with CD68 (teal) and DAPI (blue), along with endogenous Tomato reporter (red). Tomato^+ and $\mathsf{Tomato}^ \mathsf{M}\Phi$ were counted and assessed for TUNEL positivity. Representative images are shown, with white arrowheads pointing to Tomato viable MΦ, the red arrowhead pointing to a Tomato to viable M Φ , and the green arrowhead pointing to a TUNEL⁺ Tomato⁺ apoptotic M Φ . Data from n = 4-7 mice from three separate experiments shown as mean (\pm SEM). *P < 0.05. NS = not significant.

adherence and increased floating cells, also consistent with cell death (data not shown). Taken as a whole, these experiments demonstrate that c-FLIP deletion in lung $M\Phi$ amplifies caspase-8 activation in response to death receptor ligation, leading to downstream caspase-3 activation and cell death.

To confirm that loss of c-FLIP induced CD11bhi M Φ cell death *in vivo*, we crossed cFLIP $^{\Delta/\Delta}$ to Rosa-stopfl/f-Tomato reporter mice (cFLIP $^{\Delta/\Delta}$ -Tomato), such that administration of doxycycline simultaneously deletes c-FLIP in CD11bhi M Φ and marks these M Φ with Tomato. Control animals were CD68rtTA \times tetOncre \times Rosa-stopfl/f-Tomato reporter mice (control-Tomato) that marked CD11bhi M Φ with Tomato, but did not delete c-FLIP. cFLIP $^{\Delta/\Delta}$ -Tomato and control-Tomato mice were treated with bleomycin; 7 days later, mice were started on doxycycline to delete c-FLIP. Lung sections

were costained for CD68 to mark $M\Phi$ and terminal deoxynucleotidyl transferase dUTP nick end labeling (TUNEL) to mark dead cells (Figures 2E and 2F). In cFLIP $^{\Delta/\Delta}$ -Tomato mice, TUNEL $^+$ $M\Phi$ were increased in Tomato $^+$ (i.e., c-FLIP deficient) but not Tomato $^-$ (i.e., c-FLIP sufficient) $M\Phi$. In contrast, in Tomato-reporter mice, low numbers of TUNEL $^+$ cells were observed in both Tomato $^+$ and Tomato $^ M\Phi$. These data show that conditional deletion of c-FLIP *in vivo* results in $M\Phi$ death.

Deletion of c-FLIP from CD11b^{hi} MΦ Prevents Their Accumulation after Bleomycin

To assess the effect of c-FLIP deletion *in vivo*, we treated cFLIP $^{\Delta/\Delta}$ and littermate control mice (cFLIP $^{\rm fl/fl}$) with bleomycin; 1 day later, mice were started on doxycycline. Delaying doxycycline treatment allowed us

to avoid potential confounding effects of c-FLIP deletion on initial inflammation and injury. Previous hCD68rtTA reporter studies showed that maximum transgene expression takes at least 4 days, although expression began within 24 hours (16). The total number of lung M Φ in BAL and lung digest was enumerated on Days 0, 4, 7, and 14 (Figures 3A and 3B). Until Day 7, total MΦ numbers were similar between $cFLIP^{\Delta/\Delta}$ and $cFLIP^{fl/fl}$ mice. However, by Day 14, total numbers of $M\Phi$ were significantly lower in cFLIP $^{\Delta/\Delta}$ mice. This decrease in MD numbers was not due to a change in Siglec-F^{hi} MΦ, but rather to a loss of CD11 $\overset{\bullet}{b}{}^{hi}$ $M\Phi$ in both the BAL and lung digest (Figures 3C-3F). Importantly, early inflammation was similar between c-FLIP^{Δ/Δ} and cFLIP^{fl/fl} mice; neutrophil, eosinophil, and monocyte numbers were equivalent (Figures E3A-E3D). Dendritic cells, which are targeted by hCD68rtTA (16), also had similar numbers between

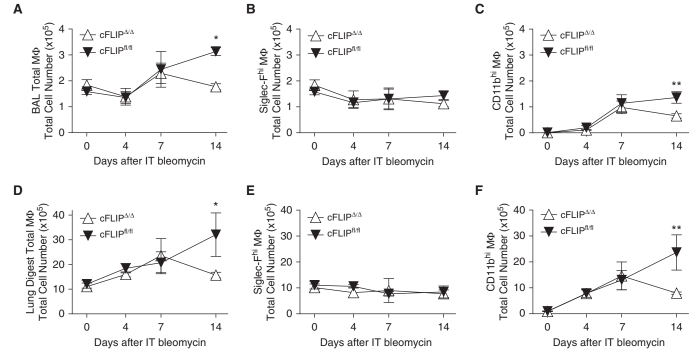


Figure 3. CD11b^{hi} MΦ, but not Siglec-F^{hi} MΦ, numbers are decreased in cFLIP^{Δ/Δ} mice after bleomycin. Flow cytometry was used to assess total MΦ numbers 0, 4, 7, and 14 days after bleomycin in lavage and lung digests. cFLIP^{Δ/Δ} and cFLIP^{β/fl} mice were started on doxycycline 1 day after bleomycin. (A–C) Total MΦ, Siglec-F^{hi} MΦ, and CD11b^{hi} MΦ numbers in BAL. (D–F) Total MΦ, SiglecF+ MΦ, and CD11b^{hi} MΦ numbers in lung digests. (G) Bleomycin-treated mice were given IT α -CD45 to label alveolar cells (IT CD45). BAL was performed 3 minutes later to sample alveolar cells and remove unbound antibody. Lung digestion was performed immediately after lavage. BAL and lung digest was processed for flow cytometry; Siglec-F^{hi} MΦ and CD11b^{hi} MΦ were separated as described in Figure 1A. IT CD45 staining is shown in BAL and lung digest. BAL staining acts as a positive control; lung digest from mice not given IT CD45 help set the negative gate. (H) CD11b^{hi} MΦ can be separated into high-and low-SSC populations using SSC versus CD11c. (H) Low-SSC CD11b^{hi} MΦ correlate with IT⁻ tissue MΦ, whereas high-SSC CD11b^{hi} MΦ correlate with IT⁺ alveolar MΦ. (H) Representative dot plots showing high- and low-SSC subpopulations in cFLIP^{fl/fl} and cFLIP^{Δ/Δ} mice 14 days after bleomycin. (H) Total numbers of high- and low-SSC CD11b^{hi} MΦ 14 days after bleomycin corresponding to alveolar and tissue CD11b^{hi} MΦ, respectively. Data from H = 3–5 mice/group shown as mean (±SEM). *H < 0.05, **H < 0.01.

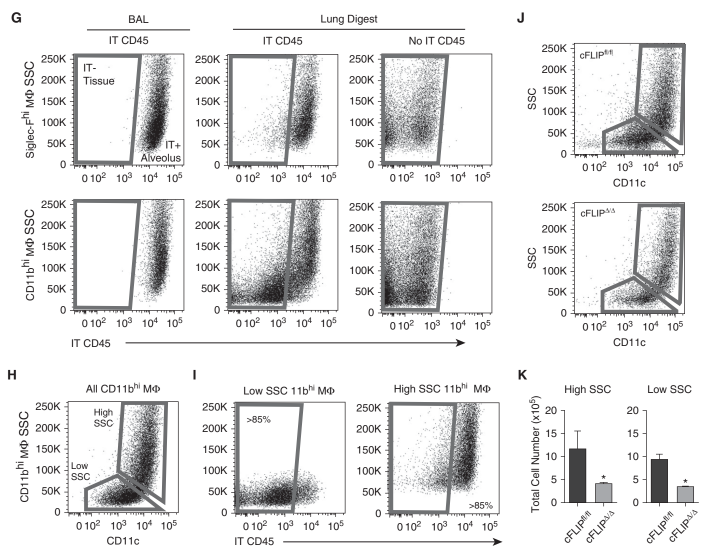


Figure 3. (Continued).

groups (Figures E3E–E3G). Furthermore, we have previously shown that no CD45-negative cells activate hCD68rtTA (16), but we verified that fibroblasts were not targeted in our system by staining lung sections from bleomycin-treated reporter mice with α -smooth muscle actin (Figure E4).

Because hCD68rtTA targets CD11bhi M Φ in the airspaces and tissue during inflammation (16), we separated CD11bhi M Φ by compartment to examine the effect of c-FLIP deletion. A fluorescently conjugated α -CD45 antibody was given intratracheally to bleomycin-treated mice immediately after being killed to selectively label M Φ in the airspaces (but not the tissues) (32). BAL was performed 5 minutes later, followed by lung digestion. As

expected, both Siglec-F^{hi} MΦ and CD11b^{hi} MΦ from BAL stained brightly with intratracheal α-CD45 (Figure 3G). Siglec-F^{hi} $M\Phi$ present in lung digests also stained brightly with intratracheal α-CD45, confirming that Siglec-F^{hi} MΦ existed exclusively in the alveolus. These data also demonstrate that lavage is inefficient in removing leukocytes from the alveolus, as previously shown (33). Notably, when CD11bhi MΦ from lung digests were examined for α-CD45 labeling, there were two populations. The first stained brightly with α -CD45, indicating that these cells were from the alveolar compartment and represented recruited CD11bhi alveolar MΦ. The second population was α-CD45 dim, indicating that these CD11bhi cells resided in the tissues and were IMs. Results were not compromised by any bleomycinor BAL-induced damage of the tissue; intravenous α -CD45 given before death was not found in BAL, demonstrating that the lung tissue is intact (Figure E3H).

Careful examination of the α -CD45 dim (i.e., interstitial) and bright (i.e., alveolar) CD11b^{hi} M Φ populations revealed that their side scatter (SSC) profiles were different (Figure 3G), as was expression of CD11c (data not shown). We therefore hypothesized that SSC and CD11c might serve as markers of tissue localization. Indeed, separating CD11b^{hi} M Φ from lung digest using SSC versus CD11c isolated two populations (Figure 3H) that correlated highly with alveolar and tissue M Φ (Figure 3I). Application of this technique to lung

digests from cFLIP $^{\Delta/\Delta}$ versus cFLIP $^{fl/fl}$ mice confirmed that deletion of c-FLIP resulted in decreased numbers of CD11 hi M Φ in both the interstitium and airspaces (Figures 3J and 3K). Taken as a whole, these results demonstrate that c-FLIP is required for maintenance of CD11 hi M Φ in both the alveolus and lung tissue, and that deletion of c-FLIP reduces CD11 hi M Φ numbers without impacting levels of other myeloid cell populations, including Siglec-F hi resident alveolar M Φ .

Deletion of c-FLIP from CD11b^{hi} $M\Phi$ and Loss of CD11b^{hi} $M\Phi$ Protected Mice from the Development of Lung Fibrosis

Because inducible deletion of c-FLIP in CD11b^{hi} MΦ significantly reduces their numbers in the lung, we used this system to test the hypothesis that elimination of CD11bhi MΦ would attenuate the development of fibrosis. As a first step, we treated $cFLIP^{\Delta/\Delta}$ and $cFLIP^{fl/fl}$ mice with bleomycin and began administration of doxycycline 1 day later. To provide a reference for noninjured lungs groups of $cFLIP^{\Delta/\Delta}$ and $cFLIP^{fl/fl}$, animals were given doxycycline without bleomycin. Fibrosis was assessed 14 days later using lung physiology, histology, and hydroxyproline measurements. As shown in Figure 4A, bleomycin-treated cFLIPfl/fl mice had a marked decrease in static lung compliance compared with healthy controls. Deletion of c-FLIP in CD11bhi MΦ reversed this defect; static compliance in bleomycintreated cFLIP $^{\Delta/\Delta}$ mice was similar to that in healthy controls. Similarly, lungs from bleomycin-treated cFLIP^{fl/fl} mice demonstrated a marked increase in hydroxyproline content, whereas cFLIP $^{\Delta/\Delta}$ mice were protected (Figure 4B). Trichrome staining of cFLIP $^{\Delta/\Delta}$ and cFLIP^{fl/fl} lungs showed patchy lung injury, typical of bleomycin. Larger areas of damage with inflammatory infiltrates and greater collagen deposition were observed in cFLIP^{fl/fl} mice compared with cFLIP $^{\Delta/\Delta}$ mice (Figures 4C and 4D). Taken together, these studies suggest that early deletion of c-FLIP from CD11bhi MΦ induces loss of CD11b $^{\rm hi}$ $M\Phi$ and attenuates fibrogenesis.

As shown in Figure 1, CD11 b^{hi} M Φ numbers in wild-type mice were increased 7 days after bleomycin; however, fibrosis in this system does not peak until Day 14 (4). Ergo, we hypothesized that the CD11 b^{hi}

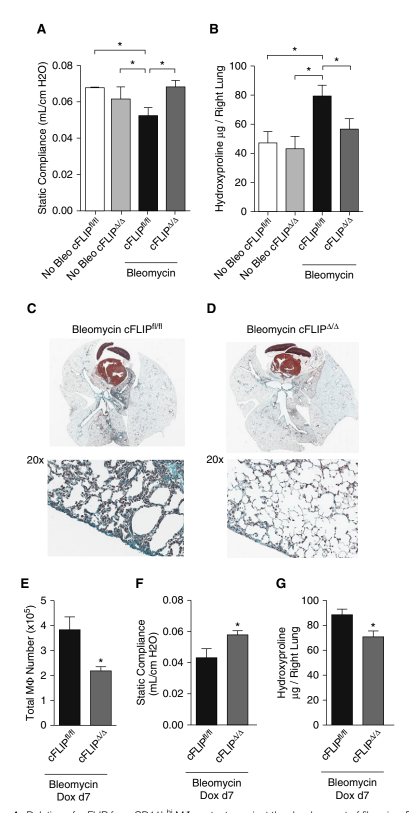


Figure 4. Deletion of c-FLIP from CD11b^{hi} M Φ protects against the development of fibrosis. cFLIP^{fl/fl} and cFLIP^{Δ/ Δ} mice were treated with bleomycin and started on doxycycline 1 day later; fibrosis was assessed at Day 14. Doxycycline-only controls comprised of both cFLIP^{fl/fl} and cFLIP^{Δ/ Δ} mice were given doxycycline for a matched 13 days before death. (A) Static compliance measured by flexiVent. (B) Hydroxyproline levels in the right lung. (C and D) Trichrome stain on paraffin-embedded lung

MΦ present 7–14 days after bleomycin were particularly important for the development of fibrosis. To test this, we treated mice with bleomycin, but waited 7 days before administering doxycycline. At Day 14, MΦ numbers were decreased in cFLIP $^{\Delta/\Delta}$ mice (Figure 4E), which were again protected from the development of fibrosis, as demonstrated by improved lung compliance and decreased hydroxyproline content (Figures 4F and 4G). These findings suggest that CD11b^{hi} MΦ present in the lungs 7–14 days after bleomycin promote the development of fibrosis.

CD11b^{hi} M⊕ Are Distinct from Siglec-F^{hi} M⊕ with High Expression of Profibrotic Chemokines after Bleomycin Injury

To investigate how CD11 b^{hi} M Φ might promote lung fibrosis, we performed RNAseq analysis of CD11b $^{\rm hi}$ M Φ and SiglecF $^{\rm hi}$ $M\Phi$ from wild-type mice after bleomycin (GEO accession no. GSE94699). Based on our data showing a profibrotic role for CD11bhi MΦ present 7–14 days after bleomycin, we chose to assess $M\Phi$ populations within this window, and isolated naive $\text{SiglecF}^{\text{hi}}$ $M\Phi$ and CD11b^{hi} MΦ and SiglecF^{hi} MΦ on Day 8. Our novel RNA-seq was merged with published RNAseq of resident IMs and SiglecF^{hi} MΦ from naive mice (GEO accession no. GSE94135) (31). Pearson correlation showed that CD11b^{hi} MΦ sorted 8 days after bleomycin were a distinct population from Siglec-Fhi $M\Phi$ and naive IMs (Figure 5A). In contrast, Siglec-F^{hi} MΦ were remarkably similar, whether collected from naive or bleomycintreated lungs. The most up- and downregulated genes are shown in the data supplement (Figure E5). To validate our sort strategy, we confirmed that all $M\Phi$ showed high expression of commonly accepted MΦ markers and low expression of genes associated with lymphocytes, dendritic cells, neutrophils, and eosinophils (Figure 5B).

Many researchers frame $M\Phi$ in terms of having an "M1" or "M2" polarization, where M2 is associated with prorepair or profibrotic $M\Phi$. However, we and others have questioned the ability of a binary

system to describe the complexities of $M\Phi$ polarization, particularly *in vivo* (34–36). RNA-seq showed that CD11bhi and Siglec-Fhi M Φ from bleomycin-treated mice expressed markers of both M1 and M2 polarization (Figure E6). Expression of these markers by the two M Φ subtypes was largely similar. In aggregate, the transcriptomal profiles of CD11bhi and Siglec-Fhi M Φ highlight the problem of ascribing a facile polarization state to M Φ populations *in vivo*. With this in mind, we sought to explore pathways that relate directly to fibrosis.

Closer inspection of the RNA-seq data showed that CD11b^{hi} MΦ had higher expression of epithelial and fibroblast proliferative factors (37), including hepatocyte growth factor, as well as higher expression of matrix-remodeling proteins (Figure E7A). More strikingly, CD11b^{hi} MΦ showed elevated expression of numerous chemokines compared with Siglec-F^{hi} MΦ (Figure 6A), many with established profibrotic functions, including *ccl2*, *ccl12*, and *ccl24* (38).

Human BAL M⊕ from Patients with IPF Show High Expression of Profibrotic Chemokines

To determine if human and murine $M\Phi$ from fibrotic lungs share common features that drive fibrosis, we used publically available gene-array data from the GEO to assess the overlap of our murine $\text{CD}11b^{\text{hi}}$ $M\Phi$ profibrotic $M\Phi$ with the signatures of BAL M Φ from patients with IPF (29) (GEO accession no. GSE49072). Most structural cell mitogens and matrixremodeling proteins, although upregulated in murine CD11bhi MΦ, were not upregulated in BAL M Φ from patients with IPF (Figure E7B). However, profibrotic chemokines were strongly upregulated in BAL M Φ from patients with IPF (Figure 6B); this overlapped with a group of murine homologs expressed by CD11b^{hi} MФ. Interestingly, this overlap comprised chemokines that recruit fibrocytes to the lung and promote fibroblast survival, including ccl2 and ccl24 (39-41). Although the human alveolar $M\Phi$ were obtained from

patients with established fibrosis (rather than the initiation phase), and were not divided into resident versus recruited subsets, these data suggest that one common mechanism by which murine and human $M\Phi$ contribute to the development of fibrosis may be elevated production of profibrotic chemokines that stimulate recruitment and survival of fibrocytes and fibroblasts.

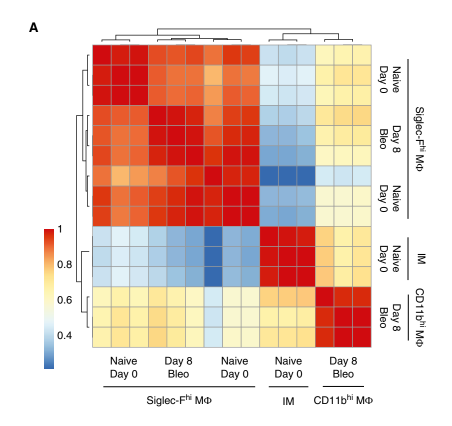
In summary, we show that murine CD11bhi M Φ promote the development of lung fibrosis after bleomycin. Using hCD68rtTA, we were able to delete c-FLIP in CD11bhi M Φ and induce CD11bhi M Φ death after bleomycin. Loss of CD11bhi M Φ was sufficient to prevent the development of fibrosis. RNA-seq suggests that this profibrotic function may be mediated through production of profibrotic factors, including chemokines specifically expressed by CD11bhi M Φ .

Discussion

 $M\Phi$ in the lungs can be divided into multiple subsets. Identifying their relative contributions to the development of fibrosis has become an important area of research. In this study, we provide a clear flow cytometry strategy to separate resident alveolar MΦ (Siglec-F^{hi}) from other MΦ present during fibrosis (CD11bhi), the majority of which are recruited M Φ that arise from immigrating monocytes. Using the novel hCD68rtTA system (16), we were able to discretely target CD11bhi MΦ without affecting Siglec-F^{hi} MΦ, and demonstrate the specific contribution of CD11b^{hi} MΦ to the development of bleomycin-induced fibrosis.

Using established M Φ markers, CD64 and Mertk (30, 31), we could clearly select M Φ within the BAL or lung digest. Recruited CD11b^{hi} M Φ in the alveolus adopt a resident-like phenotype over time, including down-regulation of CD11b, which can make their distinction difficult during the late phases of inflammation (42). However, we have identified high Siglec-F as a persistent marker of resident alveolar M Φ (7), and were able to leverage this

Figure 4. (Continued). sections from mice started on doxycycline 1 day after bleomycin and isolated at Day 14. Collagen stains blue–green. Zoom image of pleural surface highlights collagen deposition. (E–G) cFLIP $^{\text{II}/\text{II}}$ and cFLIP $^{\Delta/\Delta}$ mice were treated with bleomycin and started on doxycycline 7 days later. Fibrosis was assessed at Day 14. (E) M Φ numbers from lung digest of the left lung. (F) Static compliance measured by flexiVent. (G) Hydroxyproline content in the right lung. Data from n = 6–11 mice/group shown as mean (\pm SEM). *P< 0.05. Bleo = bleomycin; Dox = doxycycline.



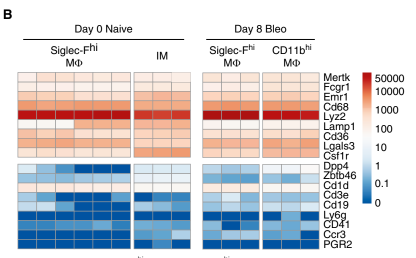


Figure 5. Global analysis of CD11b^{hi} MΦ and Siglec-F^{hi} MΦ transcriptomes shows that the two MΦ populations are distinct. Resident Siglec-F^{hi} alveolar MΦ or resident interstitial MΦ (lMs) from naive lungs (naive Day 0), and Siglec-F^{hi} and CD11b^{hi} MΦ from bleomycin-treated lungs after 8 days (Day 8 bleo) were sorted for RNA-seq analysis. (A) Individual replicates of Siglec-F^{hi} MΦ of IMs from naive lungs and Siglec-F^{hi} MΦ and CD11b^{hi} MΦ from bleomycin-treated lungs compared using pairwise Pearson correlation. (B) Heat map showing gene expression in transcripts per million (TPM) on a log scale of MΦ-lineage–associated genes and non–MΦ-lineage–associated genes.

observation to separate resident alveolar $M\Phi$ from other lung $M\Phi$ at all time points examined.

Importantly, we recently described a new inducible transgenic system,

hCD68rtTA, capable of targeting CD11b^{hi} IMs and recruited $M\Phi$ in the lung without affecting resident Siglec-F^{hi} $M\Phi$. Traditional cre drivers, such as lysozyme M and colony stimulating factor 1 receptor, do

not distinguish between $M\Phi$ subsets, and have considerable overlap with other myeloid cells. hCD68rtTA is the first transgenic system to discretely target CD11b^{hi} $M\Phi$ in the lung, presenting a unique opportunity to study their function. Using this system, we were able to manipulate the survival of CD11b^{hi} $M\Phi$ by deleting the antiapoptotic protein, c-FLIP, and thus study their function during fibrogenesis.

Previous approaches to target CD11bhi $M\Phi$ have relied on blocking their monocyte precursors (3, 11, 43). These techniques unavoidably impact all monocyte-derived cells, including dendritic cells and fibrocytes (41), as well as Ly6Chi inflammatory monocytes themselves, altering the early inflammatory response to bleomycin. With these confounding offtarget effects, it has been difficult to isolate the importance of CD11bhi MΦ. Our hCD68rtTA system improves on this in two ways: first, by directly targeting only 60% of monocytes compared with nearly 90% of CD11b^{hi} MΦ (16); second, c-FLIP expression is very low in monocytes, but strongly upregulated upon differentiation to M Φ (24). Accordingly, we found no evidence that monocytes were directly altered by loss of c-FLIP: early recruitment of CD11bhi M Φ was matched between cFLIP $^{\Delta/\Delta}$ and cFLIP $^{\rm fl/fl}$ mice, and monocyte recruitment to the alveolus was matched at all time points examined. Also importantly, fibroblasts are not targeted in our system; no α-smooth muscle actin⁺ or CD45⁻ cells had reporter expression. Moreover, we found no change in the numbers of any leukocyte population besides CD11bhi M Φ in cFLIP $^{\Delta/\Delta}$ mice. Therefore, cFLIP $^{\Delta/\Delta}$ mice discretely induce the death of CD11 b^{hi} M Φ .

In addition to superior targeting, deletion of c-FLIP was only induced upon administration of doxycycline. Taking advantage of this inducible system, we delayed doxycycline administration to exactly match initial injury between $cFLIP^{\Delta/\Delta}$ and $cFLIP^{fl/fl}$ mice, and define the window of CD11 b^{hi} M Φ action. For our initial studies, we began doxycycline 1 day after bleomycin; this substantially blocked the development of fibrosis. Waiting to administer doxycycline until 7 days after bleomycin afforded the same level of protection. Therefore, CD11bhi $M\Phi$ present in the lungs 7-14 days after bleomycin promote the development of fibrosis, whereas CD11bhi MΦ present between Days 1 and 7 do not. This model

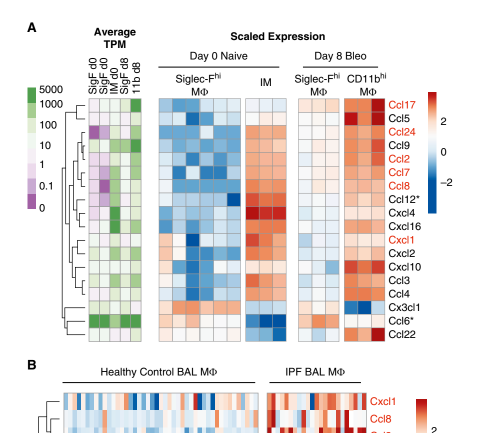


Figure 6. CD11b^{hi} MΦ express a profibrotic signature, including high expression of multiple profibrotic chemokines, that is validated in human MΦ from patients with idiopathic pulmonary fibrosis (IPF). (A) Heat maps showing expression of genes encoding chemokines. Unsupervised clustering was used to highlight associations between expression of these genes and MΦ subtypes. Raw expression is displayed on a log scale as average TPM per group (green–purple color scale) and scaled expression shows relative expression of each gene as the change from row mean scaled to a standard deviation of 1 (red–blue color scale). (B) Chemokines with significantly different expression in BAL MΦ from patients with IPF compared with normal volunteers were identified from publically available data in the Gene Expression Omnibus (accession no. GSE49072). Unsupervised clustering was used to highlight associations between expression of these genes and disease state. Scaled expression shows relative expression of each gene as the change from row mean scaled to a standard deviation of 1. Chemokines upregulated in both human IPF MΦ and murine CD11b^{hi} MΦ compared with control human MΦ or murine SiglecF^{hi} MΦ after bleomycin are highlighted in *red*. *Gene with no direct murine/human homolog.

may be adapted further, for example, by delaying doxycycline until after fibrosis has developed to test whether inducing CD11bhi M Φ death will accelerate the resolution of established fibrosis.

RNA-seq suggested multiple means by which CD11b $^{\rm hi}$ MP might exert a profibrotic function. We observed that

CD11b^{hi} M Φ are the predominant M Φ source of profibrotic chemokines, matrix remodeling proteins, and growth factors for epithelial cells and fibroblasts. Interestingly, we found that elevated expression of fibrogenic chemokines was echoed in human lavaged M Φ from patients with IPF. In particular, *ccl2* and *ccl24* were strongly upregulated in murine CD11b^{hi} M Φ

isolated from bleomycin-treated mice and in human $M\Phi$ from patients with IPF. In humans and mice, these chemokines can directly recruit fibrocytes to the lung. CCL24 also promotes the survival of fibroblasts, whereas CCL2 stimulates their proliferation and production of collagen (39, 40, 44, 45). CCL2 and CCL24 correlate with progression and severity of human IPF (46, 47). We speculate that production of CCL2 and CCL24 by CD11b^{hi} MΦ is one mechanism by which CD11bhi MΦ promote the development of fibrosis, which can be clarified in future studies. In addition, it is notable that the addition of exogenous apoptotic cells accelerates the resolution of fibrosis in mice (48). Accordingly, it is possible that inducing the death of CD11b hi M Φ in cFLIP $^{\Delta/\Delta}$ mice both removed a profibrotic signal and provided a resolving one.

Targeting profibrotic CD11bhi $M\Phi$ may be valuable in the treatment of fibrosis. We were able to induce the death of CD11b^{hi} M Φ by deleting c-FLIP, an apoptosis inhibitor downstream of death receptor ligation. This pathway is ripe for pharmacologic intervention. c-FLIP inhibitors, currently in development as antitumor drugs, may be used to block this key antiapoptotic regulator and sensitize profibrotic $M\Phi$ to death. Alternately, or even simultaneously, immunotherapies could focus on the exogenous addition of death ligands like FasL or TNF- α , increasing proapoptotic signals. Supporting this direction, administration of TNF-α to bleomycintreated mice was shown to accelerate the resolution of fibrosis (4). Profibrotic mediators produced by CD11bhi MΦ may also be fruitful targets. We speculate that CCL2 and CCL24 are key profibrotic chemokines. Although initial testing of α-CCL2 immunotherapy was halted due to complications (49), α-CCL24 immunotherapy is in development, and may be useful in the treatment of IPF.

Because much of the M Φ literature in lung fibrosis has focused on M Φ polarization, it is important to note that RNA-seq did not identify a clear M1 or M2 polarization in Siglec-F^{hi} or CD11b^{hi} M Φ . Although individual genes can be selected to argue for a specific polarization (i.e., Arg1), looking at the body of markers clearly demonstrates that both M1 and M2 pathways are

0

Ccl13*

Ccl18

Ccl17

activated in lung M Φ . Although this could reflect the presence of discrete M1 and M2 polarized subpopulations within M Φ subtypes (testable by single-cell analysis), our data best support the view that a binary system is not capable of categorizing the complex M Φ present in disease states (34, 36).

In contrast, separating subsets and studying functionality of $M\Phi$ provides important insight into disease pathology. We clearly show that CD11b^hi $M\Phi$ drive fibrogenesis, whereas resident Siglec-F^hi alveolar $M\Phi$ neither increase in number after bleomycin nor express high levels of profibrotic mediators. Nonetheless, our study is only a first step toward clarifying the complex $M\Phi$

subsets present in the lung during fibrosis. In the future, we anticipate that it will be possible to further divide CD11bhi MΦ into resident IMs and recruited alveolar and tissue $M\Phi$. It is intriguing that naive IMs express many chemokines and matrix-remodeling factors that can be considered profibrotic. Because cell origin and tissue environment shape MΦ function, phenotype, and transcriptome (7, 50, 51), whether alveolar and tissue CD11bhi MΦ are equally profibrotic is an interesting question for further study. Because hCD68rtTA targets both alveolar and tissue CD11bhi MΦ, our study was unable to address this question.

In summary, we have clearly identified CD11bhi M Φ as key participants in the development of fibrosis. Targeted induction of CD11bhi M Φ death prevents lung fibrosis. RNA-seq shows that CD11bhi M Φ are the M Φ source of profibrotic factors, particularly chemokines. Up-regulation of the same profibrotic chemokines is observed in human alveolar M Φ from patients with IPF. CD11bhi M Φ promote lung fibrogenesis, likely through the production of factors that support recruitment, growth, and differentiation of fibrocytes, fibroblasts, and myofibroblasts.

Author disclosures are available with the text of this article at www.atsjournals.org.

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