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Activated Alveolar Epithelial Cells Initiate Fibrosis through Secretion of Mesenchymal Proteins

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Address correspondence to Kevin K. Kim, M.D., 109 Zina Pitcher Place, BSRB 4061, Ann Arbor, MI 48109. E-mail: kevkim@med.umich.edu. Fibrosis is characterized by accumulation of activated fibroblasts and pathological deposition of fibrillar collagens. Activated fibroblasts overexpress matrix proteins and release factors that promote further recruitment of activated fibroblasts, leading to progressive fibrosis. The contribution of epithelial cells to this process remains unknown. Epithelium-directed injury may lead to activation of epithelial cells with phenotypes and functions similar to activated fibroblasts. Prior reports that used a reporter gene fate-mapping strategy are limited in their ability to investigate the functional significance of epithelial cell-derived mesenchymal proteins during fibrogenesis. We found that lung epithelial cell-derived collagen I activates fibroblast collagen receptor discoidin domain receptor-2, contributes significantly to fibrogenesis, and promotes resolution of lung inflammation. Alveolar epithelial cells undergoing transforming growth factor- β -mediated mesenchymal transition express several other secreted profibrotic factors and are capable of activating lung fibroblasts. These studies provide direct evidence that activated epithelial cells produce mesenchymal proteins that initiate a cycle of fibrogenic effector cell activation, leading to progressive fibrosis. Therapy targeted at epithelial cell production of type I collagen offers a novel pathway for abrogating this progressive cycle and for limiting tissue fibrosis but may lead to sustained lung injury/inflammation. (Am J Pathol 2013, 183: 1559-1570; http:// dx.doi.org/10.1016/j.ajpath.2013.07.016)

Progressive fibrosis can occur as a serious complication of lung injury, as a sequel of many inflammatory chronic diseases, or as a primary disease, such as idiopathic pulmonary fibrosis (IPF).¹ Progressive fibrosis often has a devastating clinical course without good therapeutic options. Tissue fibrosis is characterized by accumulation of activated fibroblasts and extensive matrix remodeling. Primary functions of activated fibroblasts include deposition of fibrotic matrix proteins and secretion of profibrotic factors.¹⁻⁴ The cellular source of activated fibroblasts remains unknown and controversial.⁵ The traditional model has been that resident fibroblasts respond to injury by proliferating and acquiring a profibrotic, activated phenotype accounting for all of the deposition of fibrotic matrix proteins. A newer hypothesis is that structural cells with normal physiological functions can respond to injury by down-regulating some of their physiological proteins and activities in favor of a profibrotic phenotype that overlaps with activated fibroblast activities. Proposed cells that can respond in this way include epithelial cells, endothelial cells, and pericytes. A third possible source of fibrillar collagens are from circulating bone marrow-derived fibrocytes that can be rapidly recruited to sites of injury. These different possibilities are not mutually exclusive and may potentially have nonredundant functions during fibrogenesis.^{5–9} In addition to the secretion of fibrotic matrix proteins, activated fibroblasts have an important function in recruiting more activated fibroblasts through secretion of specific profibrotic factors in response to transforming growth factor (TGF)- β -mediated activation.^{2,5} Thus, a proposed model is that injury leads to activated not a profibrotic phenotype similar to activated

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fibroblasts, leading to early fibrotic matrix deposition and activation of other profibrotic effector cells. Understanding these events may lead to new targets for inhibiting progressive fibrogenesis.

Activation of lung epithelial cells during fibrogenesis is an attractive hypothesis, given the likely propensity for environmental lung epithelial injury and the extensive animal model and human data suggesting that epithelial cell dysregulation is an important contributor of lung fibrogenesis.¹⁰⁻¹³ We and others have identified expression of mesenchymal genes, such as type I collagen, within epithelial cells of human fibrotic lung tissue, suggesting partial mesenchymal transition.^{14–16} A number of different lung epithelial cells, including alveolar and airway epithelial cells, can undergo epithelial-mesenchymal transitions (EMTs) in vitro in response to cytokines known to be highly expressed in the fibrotic lung.^{17–20} Fate-mapping studies that use an epithelial cell-specific Cre transgene and a lox-stoplox reporter transgene to permanently and specifically label epithelial cells were proposed as a definitive method for proving or disproving EMTs during animal models of fibrogenesis.²¹ However, these studies have severe limitations leading to continued controversy.²² For example, in the lung at least three separate groups have identified EMTs during fibrogenesis by this technique, although a recent report failed to find evidence of EMTs.²³⁻²⁶ A similar controversy exists in fibrogenesis of the kidney and liver.^{6,21,22,27–29} One of the points of controversy has been the potential for artifact from overlapping cells, leading to the appearance of a cell costaining the reporter protein and a mesenchymal protein. The costaining approach lends itself to cytoskeletal or cell surface protein such as vimentin, α -smooth muscle actin (SMA), fibroblast-specific protein, or N-cadherin, but these proteins are of uncertain functional significance to fibrogenesis. Most importantly, gene expression studies in human samples and the murine fate-mapping reporter gene costaining approaches are both ultimately descriptive without addressing these more important questions: to what extent and in what way do different cell types contribute to fibrosis? Several recent reports have found that epithelial cell-specific deletion of EMT transcription factors and cell surface receptors can dramatically attenuate fibrosis well beyond the fractional contribution of epithelial-derived cells to the pool of activated fibroblasts, suggesting that activated epithelial cells may have a nonredundant function during fibrogenesis.^{30–33}

The present study offers a new and more functional approach to understanding the contribution of epithelial cells. We have developed a novel mouse model with conditional deletion of type I collagen that enables lung epithelial cell-specific deletion of this gene to determine the ultimate consequences on fibrogenesis. In addition to type I collagen, we found that activated lung epithelial cells express a number of profibrotic proteins and can activate lung fibroblasts. These studies confirm an early and important epithelial cell expression of type I collagen and shift the focus away from arbitrary definitions of EMTs to better understanding the function of epithelial cells during fibrogenesis.

Materials and Methods

Mice

The floxed collagen, type I, alpha 1 (Collal) targeting vector was generated with standard cloning and recombineering techniques to incorporate loxP sites within introns 2 and 5 and a FRT [flippase (Flp) recombination target]flanked neomycin resistance gene within intron 5 (Supplemental Figure S1). Embryonic stem (ES) cell electroporation, ES cell culture, and blastocyst injection to generate chimera were performed by inGenious Targeting Laboratory (Ronkonkoma, NY). ES clones were screened by PCR and Southern blot analysis. Southern blot analysis was performed as previously described.³⁴ ES cell DNA was digested with SphI and 5' probe that was generated with the following primers: forward, 5'-AATAGTGTTATGCTCT-GGTTTC-3' and reverse, 5'-CTGCAGTGGCTAGAAAA-GTCA-3'. The 5' probe recognizes a 23-kb DNA fragment for wild-type (WT) ES cells and a 15-kb fragment for correctly targeted floxed Col1a1 ES cells. ES cell DNA also was digested with HindIII and 3' probe generated with the following primers: forward, 5'-CTGGTCGCCCCGGTGA-3' and reverse, 5'-GACTTGGGTGTGACTATCACATA-AAAAGACC-3'. The 3' probe recognizes a 10-kb DNA fragment for WT ES cells and a 5-kb DNA fragment for correctly targeted floxed Col1a1 ES cells. For PCR analysis of floxed Collal, forward primer (F1), 5'-ATCCATCAT-GGCTGATGCAATGCG-3', located with neomycin resistance and reverse primer (R1), 5'-TGACTTACGGGTTCT-CCTTTGGCA-3', located 7 kb from the start site confirm integration of targeting vector to the Collal gene. Forward primer (F2), 5'-GGTAGCTCTGGCATGCATAAC-3', overlaps with 5' loxP site and reverse (R2), 5'-AGCTAGCTT-GGCTGGACGTAAACT-3', within neomycin resistance confirm integration of the 5' loxP site and neomycin gene on the same allele. Forward primer (F3), 5'-TGGTACAGCA-CTTTACAGCGCACA-3', located upstream of 5' loxP insertion site and reverse prime (R3), 5'-TTACTCGGCCTGG-GTCACTTCTTT-3', are located downstream of 5' loxP insertion site; WT allele yields a 138-bp PCR product, floxed Collal with insertion of 40-bp loxP, and SphI site yields a 178-bp band. Removal of neomycin gene confirmed by PCR using forward primer (F4), 5'-ACTGTCCTCCAATA-AACTGCAGTTTTCTTT-3', located upstream of neomycin insertion site and reverse primer (R4), 5'-TAGCAGTAAT-GGGACAAACGGATGTAG-3', located downstream of neomycin insertion site. Mice with constitutive expression of flpE (The Jackson Laboratory, Bar Harbor, ME) were used to delete the FRT-flanked neomycin resistance gene. Floxed Col1a1 mice were backcrossed with C57bl/6 mice for at least eight generations.

To delete floxed *Colla1* in lung epithelial cells mice were crossed with mice expressing the surfactant proteins-C promoter-reverse tetracycline transactivator (SPC-rtTA) and tetO-CMV (cytomegalovirus) promoter-Cre recombinase (tetO-Cre) transgenes as previously described.²⁴ Triple transgenic SPC-rtTA/tetO-Cre/Col1a1^{f/f} (homozygous floxed Col1a1) mice are abbreviated SCcol. Smad3-null mice were purchased from The Jackson Laboratory (Bar Harbor, ME). Mice were genotyped by PCR. Six- to 8-week-old mice were given 50-µL intratracheal injections of saline or saline with dissolved bleomycin (1.3 U/kg) via surgical tracheotomy. At various time points after injection, mice were euthanized, and bronchoalveolar lavage (BAL) and lung samples were collected for analysis. For BAL, the lungs were lavaged with 1 mL of PBS per mouse. Samples were centrifuged, and the supernatants were used for total protein count and immunoblot analysis. The cell pellets were resuspended for total cell counting with a hemocytometer under light microscopy. Lung physiological measurements were obtained from 6- to 8-weekold anesthetized SCcol mice and littermate controls as previously described.32,35

All mice were bred and maintained in a specific pathogenfree environment, and all animal experiments were approved by the University Animal Care and Use Committee at the University of Michigan.

Reagents

Plasma fibronectin (FN), bleomycin, and phospho-Smad2 antibody were purchased from Millipore (Billerica, MA); collagen I, Matrigel (MG), biotin-conjugated rat anti-mouse CD16/32 and CD45 antibodies, and 70-µm and 40-µm nylon filters from BD Biosciences (San Jose, CA); purified human keratinocyte growth factor from PeproTech (Rocky Hill, NJ); purified human TGF-β1 from R&D Systems (Minneapolis, MN); small airway basal media and small airway growth media (SAGM) from Lonza Inc. (Allendale, NJ); horseradish peroxidase-conjugated secondary antibodies from Santa Cruz Biotechnology, Inc. (Santa Cruz, CA); immunofluorescentconjugated secondary antibodies, streptavidin-coated magnetic beads, and magnetic particle separator from Invitrogen (Carlsbad, CA); collagen I and discoidin domain receptor-2 (DDR2) antibodies from Abcam (Cambridge, MA); horseradish peroxidase-conjugated phospho-tyrosine and glyceraldehyde phosphate dehydrogenase (GAPDH) antibodies from Cell Signaling Technology Inc. (Danvers, MA); adenoviruses expressing Cre (AdCre) or green fluorescent protein (AdGFP) were obtained from the University of Iowa Gene Transfer Vector Core Facility; and dispase from Roche (Indianapolis, IN). All other reagents were purchased from Sigma-Aldrich (St. Louis, MO).

Mouse Fibroblast and Type II Cell Isolation and Culture

Cells were cultured in a 37° C, 5% CO₂ incubator. Murine primary lung fibroblasts were harvested from 6- to 10-week-old

mice as described previously.³⁶ Cells were used at passages two to four. In some experiments cells were treated with three daily doses of AdGFP or ADCre (50 pfu/cell) and then cultured for an additional 7 days before analyzing the cells. Adult murine fibroblast cell line, MLg (CCL-206), was purchased from the American Type Culture Collection (Manassas, VA) and also maintained in Dulbecco's modified Eagle's medium (DMEM) with 10% fetal bovine serum, penicillin, and streptomycin. Murine type II alveolar epithelial cells (AECs) were isolated as previously described with minor modifications.^{24,37} Briefly, after mice were sacrificed, lungs were exposed, perfused, and lavaged. Lungs were filled with dispase, followed by low-melt agarose. Lungs were digested at room temperature for 45 minutes then dissected. Dissected lungs were then treated with DNase for 10 to 15 minutes, then serially filtered through 70-µm, 40-µm, and 20-µm filters. The crude suspensions were then labeled for CD16/32 and CD45 with the use of biotin-conjugated antibodies, streptavidinconjugated magnetic beads, and a magnetic separator. AECs were further negatively selected by incubating for 1 to 2 hours on a tissue-culture-treated plastic plate.

H&E Staining and Masson's Trichrome Assay

Lungs from sacrificed mice were inflated to a pressure of 25 cm H_2O and fixed with formaldehyde. Lungs were embedded in paraffin, sectioned, and stained with H&E and Masson's trichrome by the McClinchey Histology Laboratory (Stockbridge, MI).

Hydroxyproline Assay

Hydroxyproline was measured by methods previously described.³⁶ Briefly, homogenates from the entire lungs were baked in 12N HCl overnight at 120°C. Aliquots of the samples were added to citrate buffer and chloramine T solution and incubated at room temperature for 20 minutes. Erlich's solution was added, and the samples were incubated at 65°C for 15 minutes. Absorbance at 540 nm was measured. Hydroxyproline content was quantified by comparison against a hydroxyproline standard curve.

Immunofluorescence Staining

Lungs were perfused with PBS, inflated with intratracheal OCT, removed, and immediately frozen in a dry-ice alcohol bath. Lungs were stored at -80° C. Seven-micron lung sections were stained as previously described.²⁴ Stained sections were visualized on an Olympus BX-51 fluorescence microscope (Olympus, Tokyo, Japan), and images were captured with an Olympus DP-70 camera and analyzed with DP controller software version 3.1.1.267.

Collagen I Deletion in Fibroblasts

Col1a1^{f/f} lung fibroblasts were treated with three daily doses of AdGFP or AdCre (50 pfu/cell). After 1 week equal numbers of

cells were plated on tissue culture plastic or collagen I-coated plates. Cells were maintained in DMEM and cultured on collagen I-coated plates, which were treated with an additional 10 μ g/mL collagen I. After 24 hours, RNA was collected for gene expression analysis. In some experiments, AdCre-treated Col1a1^{*f/f*} fibroblasts were seeded on six-well tissue culture plates and then stimulated with 1 mL of DMEM supplemented with 0.5 mL of BAL fluid from control and SCcol mice collected 1 week after saline or bleomycin injury or treated with 10 μ g/mL purified collagen I. After 15 minutes cells were lyzed and analyzed for DDR2 phosphorylation.

CM Stimulation

AECs were cultured on either MG- or FN-coated six-well plates. Some wells were treated with 10 μ mol/L TGF- β receptor inhibitor SB431642, 5 pfu per cell lentivirus-expressing siRNA, or 50 pfu/cell AdGFP or AdCre. Conditioned media (CM) was generated by changing the AEC media to serum-free SAGM plus 0.1% bovine serum albumin 24 to 48 hours before collection (free of inhibitors or lentivirus). Harvested CM was filtered, then either stored at 80°C or immediately added to cultured lung fibroblasts. After 48 hours, fibroblasts were lyzed for various assays.

Gene Expression Analysis

RNA was isolated from cells and tissue with TRIzol Reagent (Invitrogen) according to the manufacturer. Reverse transcription was performed with the SuperScript III First-strand synthesis kit (Invitrogen), and RT-PCR was performed with POWER SYBR Green PCR MasterMix Kit (Applied Biosystems, Foster City, CA) and Applied Biosystems 7000 sequence detection system. The relative expression levels of gene were calculated against β -actin or GAPDH. Eighty-four genes related to mouse fibrosis plus five housekeeping genes were evaluated by RT-PCR (RT^2) Profiler PCR Arrays PAMM-120A; Qiagen, Valencia, CA) according to the manufacturer. The gene expression level was normalized to the average of five housekeeping genes and was calculated as $2^{-\Delta\Delta Ct}$. PCR primers for floxed Collal DNA recombination were as follows: forward, 5'-GGTAGCTCTGGCATGCATAAC-3', and reverse, 5'-AG-CTAGCTTGGCTGGACGTAAACT-3'. PCR of cDNA isolated from whole lungs was used to detect the truncated Collal with forward primer 5'-ATGGCCAAGAAGA-CAAACTTT-3' (overlaps adjacent exon 1 and 6) and reverse primer 5'-GGCCTTGGAAACCTTGTGGAC-3'. RT-PCR was performed with the Power SYBR Green PCR MasterMix Kit (Applied Biosystems). Relative expression levels of genes in fold change were calculated against β-actin or GAPDH. The following primers were used: collagen 1a1, forward, 5'-TGACTGGAAGAGCGGAGAG-TACT-3', and reverse, 5'-GGTCTGACCTGTCTCCATG-TTG-3'; connective tissue growth factor (CTGF), forward, 5'-CACTCCGGGAAATGCTGCAAGGAG-3' and reverse,

5'-GTTGGGTCTGGGCCAAATGT-3'; collagen, type III, alpha 1 (Col3a1), forward, 5'-CAAGGTCTTCCTGGT-CAGCCT-3' and reverse, 5'-TGCCACCAGGAGATCCA-TCTC-3'; α -SMA, forward, 5'-AGAGACTCTCTTCCAG-CCATC-3' and reverse, 5'-GACGTTGTTAGCATAGA-GATC-3'; β -actin, forward, 5'-CCGTGAAAAGATGACC-CAGATC-3' and reverse, 5'-CACAGCCTGGATGGCTA-CGT-3'; GAPDH, forward, 5'-AACTTTGGCATTGTGGA-AGG-3' and reverse, 5'-ACACATTGGGGGTAGGAACA-3'. Other primers used have been previously reported.³⁸⁻⁴¹

Immunoblot Analysis

Immunoblot analysis and immunoprecipitation of tissue and cells were performed as previously described.²⁴ Scanned immunoblots are representative of at least three separate experiments, and densitometry was quantified by ImageJ software version 1.47 (NIH, Bethesda, MD).

Statistical Analysis

Data were expressed as means \pm SEM. For evaluation of group differences, the two-tailed Student's *t*-test was used assuming equal variance. A *P* value of <0.05 was accepted as significant.

Results

Generation and Validation of Mice with Lung Epithelial Cell-Specific Deletion of Col1a1

Floxed Col1a1 mice were generated by integrating loxP sites within introns 1 and 5 of the Collal gene which enabled Cremediated deletion of exons 2 to 5, resulting in frameshift and early translational termination of the truncated mRNA (Figure 1A and Supplemental Figure S1). Conditional deletion of collagen I expression was confirmed with the use of murine embryonic fibroblast (MEF) cells from Col1a1^{f/f} embryos. Col1a1^{f/f} MEF cells treated with an AdCre had near complete loss of collagen I compared with cells treated with a control AdGFP or control MEFs (Col1a1^{w/w}) treated with either AdCre or AdGFP (Figure 1B and Supplemental Figure S2A). Lung epithelial cell-specific recombination was achieved by crossing with mice carrying the human SPC-rtTA and tetO-CMV-Cre transgenes (Figure 1C).²⁴ Recombination within lung epithelial cells was verified with PCR primers encompassing the floxed region. DNA from lungs of SCcol mice showed a 2.2-kb PCR product that corresponded to nonrecombined flox Col1a1 in nonepithelial cells and a 0.5-kb PCR product consistent with removal of the 1.7-kb floxed region within epithelial cells. PCR of AECs from SCcol mice yielded only the 0.5-kb recombined band, whereas PCR from lung fibroblasts yielded only the 2.2-kb non-recombined band. DNA from littermate control mice, lacking one of the transgenes, only produced the 2.2-kb PCR product. We and others have previously reported that primary AECs cultured on provisional matrix proteins such as FN undergo EMTs



Generation and validation of mice with lung epithelial-specific deletion of Col1a1. A: Schematic of floxed Col1a1 (Col1a1^{f/f}) mice generated by Figure 1 inserting loxP sites flanking exons 2 to 5 in the Col1a1 gene. Cre-mediated recombination of the floxed Col1a1 leads to removal of exons 2 to 5, shift in the translational reading frame in subsequent exons, and permanent inactivation of collagen I expression. B: MEFs isolated from Col1a1^{f/f} mice have deletion of collagen I expression after AdCre, whereas cells treated with AdGFP or WT Col1a1 MEFs treated with AdGFP or AdCre have normal expression of collagen I by immunoblot analysis. C: Lung epithelial cell-specific and permanent deletion of Col1a1 is achieved with the use of transgenic mice that carry the SPC-rtTA and tetO-Cre. Triple transgenic SPC-rtTA/tetO-Cre/Col1a1^{f/f} mice are abbreviated SCcol. The SPC promoter yields rtTA expression specifically within lung epithelial cells, in the presence of doxycycline, rtTA activates the tetO-CMV promoter, leading to expression of cre recombinase and permanent deletion of DNA flanked by loxP sites. Deletion within lung epithelial cells is confirmed by PCR with the use of primers that encompass the floxed region. Intact floxed Col1a1 yields a 2.2-kb PCR product, and removal of the floxed Col1a1 yields a 0.5-kb PCR product. D and E: Uninjured lungs from SCcol (E) mice have normal histology compared with littermate genotype control mice (D), by H&E staining. F: AECs from SCcol mice have diminished EMT-induced expression of collagen I by immunoblot analysis compared with AECs from littermate control mice lacking one of the three transgenes. Primary lung fibroblasts from SCcol mice have similar collagen I expression compared with control lung fibroblasts, verifying robust lung epithelial-specific deletion of Col1a1. Original magnification: ×200 (D and E). AdCre, adenoviral-mediated expression of Cre; AdGFP, adenovirus expressing green fluorescent protein; AEC, Alveolar epithelial cell; Col1a1, collagen, type I, alpha 1; Ctrl, control; EMT, epithelial-mesenchymal transition; Fib, fibroblast; MEF, murine embryonic fibroblast; SPC-rtTA, surfactant proteins-C promoter-reverse tetracycline transactivator; tetO-Cre, tetO-cytomegalovirus promoter-Cre recombinase; WT, wild-type.

through production and activation of endogenous TGF-β1.^{24,42} AECs from SCcol mice expressed significantly reduced levels of collagen I after FN-mediated EMTs compared with littermate control AECs (Figure 1F and Supplemental Figure S2B). Lung fibroblasts from SCcol mice showed similar levels of collagen I as control lung fibroblasts. Thus, SCcol mice have robust lung epithelial cell-specific loss of Col1a1. As expected SCcol mice have normal histology (Figure 1, D and E) and physiology (Table 1) because lung epithelial cells express little to no collagen I under physiological conditions.^{14,24}

Mice with Lung Epithelial Cell Deletion of Col1a1 Have Abnormal Response to Injury

Groups of SCcol mice and littermate control mice lacking at least one of the transgenes were given intratracheal saline or bleomycin. Genotype control mice developed robust fibrosis after bleomycin, visualized by trichrome and H&E staining. In contrast, SCcol lungs appeared injured but had less fibrosis (Figure 2A and Supplemental Figure S3, A and B), resulting in similar physiological impairment (Supplemental Table S1). Immunoblot analysis of lung lysates indicated that control mice have increased levels of collagen I in response to bleomycin, whereas collagen I up-regulation was attenuated in SCcol mice (Figure 2B and Supplemental Figure S4). Hydroxyproline analysis also showed a robust increase in collagen deposition in bleomycin-injured control mice and an attenuated response in SCcol mice (Figure 2C).

Surprisingly, a trend was observed for more death in the SCcol mice after bleomycin treatment than in the genotype control mice. No deaths were found in either saline treatment group (Figure 3A). To determine whether loss of collagen deposition in SCcol mice led to increased or

	Control	SCcol	P value
IC (mL)	$\textbf{0.473} \pm \textbf{0.027}$	$\textbf{0.467} \pm \textbf{0.017}$	0.83
VC (mL)	$\textbf{0.562} \pm \textbf{0.034}$	$\textbf{0.559} \pm \textbf{0.019}$	0.93
ERV (mL)	$\textbf{0.089} \pm \textbf{0.009}$	0.092 ± 0.003	0.73
FRC (mL)	$\textbf{0.586} \pm \textbf{0.017}$	$\textbf{0.575} \pm \textbf{0.012}$	0.64
TLC (mL)	$\textbf{1.059} \pm \textbf{0.038}$	$\textbf{1.042} \pm \textbf{0.029}$	0.71
RV (mL)	0.497 ± 0.016	$\textbf{0.483} \pm \textbf{0.013}$	0.53
Cch (mL/cm H_2 0)	0.034 ± 0.002	0.034 ± 0.002	0.82

 Table 1
 Baseline Physiology of Littermate Control and SCcol

 mice

Values indicate means \pm SEM. n = 9 to 10 per group.

Cch, chord compliance; ERV, expiratory reserve capacity; FRC, functional reserve capacity; IC, inspiratory capacity; RV, residual volume; SCcol, triple transgenic surfactant proteins-C promoter-reverse tetracycline transactivator/ tet0-cytomegalovirus promoter-Cre recombinase/homozygous floxed collagen, type I, alpha 1; TLC, total lung capacity; VC, vital capacity.

sustained lung injury/inflammation we measured total protein and cell count in the BAL fluid. Genotype control mice and SCcol mice have a similar approximate fivefold increase in BAL protein and an approximate threefold increase in BAL cell count 1 week after bleomycin. However, 2 weeks after bleomycin, control mice have a reduction in BAL protein and cell count, whereas persistent elevation was observed in SCcol mice (Figure 3, B and C).

Because the trend toward increased death in SCcol mice may affect the differences observed in the hydroxyproline and collagen I immunoblot analysis, we wanted to assess for differences in collagen I production at a time point before bleomycin-induced death. One week after bleomycin, mRNA from whole lungs of genotype control mice and SCcol mice had a similar increase in Colla1 mRNA compared with saline-treated mice (Figure 4A). However, mRNA from AECs isolated 1 week after bleomycin indicated an increased expression of Col1a1 within control AECs and a significant attenuation within isolated SCcol AECs (Figure 4B). Deletion of the floxed exons 2 to 5 resulted in frameshift and early termination of translation; however, transcription and generation of truncated Colla1 mRNA lacking exons 2 to 5 (Col1a1 Δ ex2-5) may still occur within lung epithelial-derived cells. PCR primers were generated to detect the presence of a truncated Col1a1 Aex2-5 mRNA in which exon 1 is directly adjacent to exon 6. Indeed, mRNA from lungs of bleomycin-treated SCcol mice showed the presence of Col1a1 Aex2-5 mRNA, indicating transcription of Col1a1 specifically within lung epithelial-derived cells (Figure 4C). Collectively, these studies indicated bleomycininduced expression of Col1a1 within lung epithelial cells. Furthermore, genotype control mice had small foci of collagen I deposition 1 week after bleomycin, and this was largely absent in SCcol mice (Supplemental Figure S5, A and B). Several studies have shown soluble procollagens in the BAL fluid of patients with fibrotic lung disease.^{43–45} BAL samples from SCcol and littermate control mice were collected 1 week after bleomycin or saline treatment and were analyzed by immunoblot for collagen I (Figure 4D and

Supplemental Figure S6A). A dramatic increase was observed in collagen I in the BAL fluid of bleomycin-injured control mice and attenuation in the SCcol mice treated with bleomycin, indicating that accumulation of collagen I in the BAL after bleomycin is dependent on the presence of an intact *Col1a1* gene within lung epithelial cells. Essentially, no detectable collagen I was observed in the saline-treated mice. We next wanted to determine whether deficient collagen accumulation could lead to attenuated collagen signaling. DDR2 has been identified as a novel receptor tyrosine kinase that uses type I collagen as its major ligand.⁴⁶ Whole lung lysates were analyzed by immunoprecipitation against DDR2, followed by immunoblot analysis for phospho-tyrosine. As expected, genotype control mice had increased phosphorylation of DDR2 in response to



Figure 2 Lung epithelial cell-derived type I collagen contributes to fibrogenesis. **A**: Lung sections from littermate control and SCcol mice 3 weeks after bleomycin injection, stained with H&E and trichrome. Original magnification, ×100. Genotype control mice have robust fibrosis compared with SCcol mice. **B**: Whole lung lysate from mice 3 weeks after treatment with saline or bleomycin was analyzed for collagen I by immunoblot analysis. Control mice have robust induction of collagen I after bleomycin compared with SCcol mice. **C**: Hydroxyproline assay from entire lungs 3 weeks after saline or bleomycin in SCcol (black bars) or littermate control (white bars) mice. SCcol mice have less fibrosis after bleomycin; Sal, saline; SCcol, triple transgenic surfactant proteins-C promoter-reverse tetracycline transactivator/tet0-cytomegalovirus promoter-Cre recombinase/homozygous floxed collagen, type I, alpha 1.



Figure 3 Lung epithelial cell-derived collagen I inhibits sustained lung inflammation. **A**: Survival of littermate control (white circles; n = 25) and SCcol (black circles; n = 18) mice after bleomycin. SCcol mice have a trend toward less survival. **B** and **C**: BAL from control (white bars) and SCcol (black bars) mice at 0, 7, and 14 days after bleomycin injury was analyzed for protein (**B**), and cell count (**C**) shows that SCcol and control mice have similar increases in proteins and cell count 1 week after bleomycin, whereas SCcol mice have greater persistence of protein, n = 5 to 7 per group. *P < 0.05 versus control bleomycin. BAL, bronchoalveolar lavage; SCcol, triple transgenic surfactant proteins-C promoter-reverse tetracycline transactivator/tet0-cytomegalovirus promoter-Cre recombinase/homozygous floxed collagen, type I, alpha 1.

bleomycin, whereas SCcol mice had attenuated DDR2 phosphorylation, indicating deficient collagen signaling in these mice (Figure 4E and Supplemental Figure S6, B and C). Collectively, these results support an important contribution of type I collagen by lung epithelial cells after lung injury.

AEC Activation Promotes Fibroblast Activation

An initial collagen deposition by epithelial cells may promote activation and collagen production by lung fibroblasts. Prior

studies of collagen signaling within cultured fibroblasts are limited because of significant expression of collagen I in the confined space of the culture dish. To address this problem we isolated lung fibroblasts from Col1a1^{f/f} mice. AdCre-treated Col1a1^{f/f} lung fibroblasts were stimulated with BAL from control and SCcol mice collected 1 week after saline or bleomycin injury. Fifteen minutes after BAL stimulation cells were lysed and analyzed for phospho-DDR2. BAL from control mice injured with bleomycin induced DDR2 phosphorylation, indicating the presence of bioactive collagen. DDR2 phosphorylation was not induced by BAL from SCcol mice (Figure 5A and Supplemental Figure S7A). To determine the influence of type I collagen signaling on fibroblast extracellular matrix (ECM) production, Col1a1^{f/f} lung fibroblasts were treated with AdCre to delete Collal or AdGFP as a control. Lung fibroblasts with Cre-mediated deletion of Collal had decreased expression of Colla1, as expected, and some suppression of Colla2, Col3a1, and FN expression. Culturing Cre-treated Col1a1^{f/f} lung fibroblasts in the presence of collagen I reversed the effects of AdCre treatment, suggesting that the presence of collagen I modestly enhances transcription of these genes within lung fibroblasts (Figure 5B). Treating WT or AdGFPtreated Col1a1^{f/f} cells with exogenous type I collagen had little effect on the expression of these genes. Given the unexpected sustained inflammation in SCcol mice after bleomycin, AdGFP- and AdCre-treated Col1a1^{f/f} lung fibroblasts were also analyzed for expression of several inflammatory cytokines. Surprisingly, deletion of Col1a1 led to increased expression of IL-1 α , IL-1 β , and tumor necrosis factor α (Supplemental Figure S8A).

Activated AECs may secrete other factors that more profoundly promote fibroblast activation. We and others have previously reported that primary AECs cultured on laminin or MG maintain an AEC phenotype, whereas cells cultured on provisional matrix proteins such as fibrin and FN undergo TGF-β-mediated EMTs.^{24,42} Murine lung fibroblasts (MLg) were treated with CM from primary AECs cultured on MG (AEC CM) or FN (EMT CM). EMT CM promoted fibroblast expression of collagen I, SMA, and FN compared with fibroblasts treated with AEC CM (Supplemental Figure S8B). Enhanced expression of α -SMA and collagen I by fibroblasts stimulated with EMT CM was confirmed by immunoblot analysis (Figure 5C and Supplemental Figure S7B). Lung fibroblasts treated with EMT CM had levels of collagen I and SMA similar to TGF-β1-stimulated lung fibroblasts, whereas fibroblasts treated with AEC CM or plain media had lower levels of collagen I and SMA. EMT CM did not promote Smad2 phosphorylation consistent with our prior observations that TGF- β production and activation in this in vitro system is required for AEC EMTs, but the active TGF- β remains attached to the ECM rather than being secreted into the CM. We next wanted to confirm that production of the EMT-derived fibroblast activating factor(s) required TGF-\beta-mediated AEC EMTs. TGF-\beta signaling was inhibited in primary AECs undergoing EMTs with the



Figure 4 Lung epithelial cell-deletion of collagen I attenuates collagen I accumulation acutely after bleomycin treatment. A: Whole lung mRNA was analyzed for Col1a1 mRNA 1 week after bleomycin or saline, n = 4 to 9. *P <0.05 versus control saline. Difference between control (white bars) and SCcol (black bars) bleo is not statistically significant. P = 0.3. **B**: mRNA from AECs isolated 1 week after bleomycin or saline, n = 4. $^{\dagger}P < 0.01$ versus control saline; ${}^{\ddagger}P < 0.01$ versus control bleomycin. **C**: Whole lung mRNA was analyzed with primers specific for a truncated Col1a1 mRNA missing exons 2 to 5 (Col1a1 Δ ex2-5), forward primer bridged the adjacent exons 1 and 6, and reverse primer was within exon 8. Col1a1 Δ ex2-5 was detected in SCcol mice 1 week after bleomycin, indicating native promoter activation and transcription of the recombined Col1a1 gene within lung epithelial cells after bleomycin. D: Immunoblot of 50 μ L of a 1-mL BAL from littermate control and SCcol mice 1 week after saline or bleomycin. Bleomycin induces collagen I accumulation in BAL of control mice. BAL of SCcol mice have less collagen I accumulation after bleomycin. E: One week after bleomycin, SCcol mice have attenuated activation of the collagen receptor DDR2 observed by immunoprecipitation of whole lung lysate for DDR2 followed by immunoblot analysis for phosphotyrosine. Immunoblot of 1% of the whole lung lysate indicates increase in DDR2 expression after bleomycin in SCcol and control mice. AEC, alveolar epithelial cell; BAL, bronchoalveolar lavage; Bleo, bleomycin; Ctrl, control; DDR2, discoidin domain receptor 2; Sal, saline; SCcol, triple transgenic surfactant proteins-C promoter-reverse tetracycline transactivator/tetO-cytomegalovirus promoter-Cre recombinase/homozygous floxed collagen, type I, alpha 1.

use of 10 μ mol/L TGF- β receptor (TGF- β R) kinase inhibitor (SB431642) and by using Smad3-deficient AECs. CM from AECs treated with SB431542 and from Smad3-null AECs had a diminished capacity to activate lung fibroblasts

(Figure 5D and Supplemental Figure S7C). Collectively, these results suggest that TGF- β signaling is required for AECs to produce/secrete the fibroblast stimulating factor(s) but that the secreted factor is not TGF- β itself.

To identify candidate TGF-\beta-induced, AEC-derived fibroblast-activating factors we used a mouse fibroblast pathway-specific reverse transcription-PCR array. mRNA from AECs cultured on MG, FN, or FN plus TGF-BR inhibition were compared. As expected, AECs cultured on FN had marked up-regulation of collagen I, collagen III, and SMA, and expression of these genes was suppressed by TGF- β R inhibition. The highest expressed genes by AECs on FN included several secreted profibrotic proteins: CTGF, thrombospondin, collagen III, and collagen I (Figure 5E and Supplemental Table S2). Expression of these genes was suppressed by inhibition of TGF- β signaling. Several of these factors have been used as markers of fibrotic fibroblasts and have been shown to induce fibroblast expression of type I collagen, suggesting that fibroblast-activating factors secreted by activated lung epithelial cells may not be limited to type I collagen.

Discussion

These findings support a model in which lung epithelial cells can acquire functions and protein expression that overlap with activated fibroblasts in response to injury. In this model, injury leads to activation of the epithelial cells which initiates secretion of matrix proteins early in response to injury, and these proteins promote activation of fibroblasts, leading to progressive fibrosis. We found that activated lung epithelial cells produce collagen I and that deletion of type I collagen within lung epithelial cells limits fibrosis. Several prior reports have suggested the ability of lung epithelial cells to express type I collagen in vitro when stimulated with TGF-B.^{24,47} Recent reports indicate that deletion of the TGF- β receptor in either lung epithelial cells or resident fibroblasts leads to a similar attenuation of fibrosis in mouse models.^{33,48} These findings are in agreement with at least one point from a recent report from Rock et al,²⁶ in which type II AECs were found to have an approximately twofold induction in type I collagen 10 days after bleomycin. Rock et al²⁶ did not find other evidence for EMT, in contrast to several other reports that favored EMT during lung fibrosis. The differences among these reports are likely because of differences in the fate-mapping strategy. Studies favoring EMT labeled lung epithelial cells broadly, whereas Rock et al²⁶ initiated fate-mapping within a subtotal population of type II AECs. Indeed in vitro evidence is strong that a number of different lung epithelial cell types can express type I collagen when activated.^{17,20,24,47,49} Further, the fraction of labeled type II AECs was dramatically reduced after bleomycin injury in Rock et al,²⁶ suggesting that the dTomato reporter used in that study might be toxic to injured cells as previously



Figure 5 Alveolar epithelial cell-derived factors promote fibroblast activation. A: DDR2 phosphorylation of collagen I-deleted fibroblasts by BAL from control and SCcol mice collected 1 week after saline or bleomycin injury. Cells treated with 10 µg/mL exogenous type I collagen and cells treated with BAL from bleomycin-injured control mice have induction of DDR2 phosphorylation. **B**: Reverse transcription-PCR of primary lung fibroblasts from Col1a1^{f/f} mice treated with AdGFP, AdCre, or AdCre plus exogenously added type I collagen (collagen I-coated plus 10 µg/mL added to media). Loss of collagen I expression led to reduction in expression of several fibroblast activation markers; the expression of these genes was restored by exogenous collagen. C: Immunoblot of lung fibroblasts stimulated with plain media, CM from AECs cultured on Matrigel (AEC CM), CM from AECs cultured on FN (EMT CM), and media supplemented with 4 ng/mL TGF- β . TGF- β and EMT CM activate fibroblast expression of α -SMA and collagen I. TGF- β promotes smad2 phosphorylation but EMT CM does not. D: Immunoblot of lung fibroblasts stimulated with plain media, CM from WT AECs cultured on FN (WT CM), smad3-null AECs (Sm3-CM) cultured on FN, and WT AECs cultured on FN with addition of TGFβ receptor inhibitor SB431542 (SB), 10 μ mol/L (WT + SB CM). E: Mouse fibrosis reverse transcription-PCR array of AECs cultured on Matrigel, FN, or FN with addition of SB. Expression of genes on FN versus Matrigel (black) and FN with SB versus Matrigel (red) are shown. Blue lines indicate fivefold differential expression. Several genes of interest are indicated. AdCre, adenovirus expressing Cre; AdGFP adenovirus expressing green fluorescent protein; AEC, alveolar epithelial cell; BAL, bronchoalveolar lavage; Bleo, bleomycin; CM, conditioned media; Col I, type I collagen; Col1a1^{f/f}, homozygous floxed Col1a1; Col1a2, collagen, type I, alpha 2; Col3a1, collagen, type III, alpha 1; ctgf, connective tissue growth factor; Ctrl, control; DDR2, discoidin domain receptor-2; EMT, epithelial-mesenchymal transition; FN, fibronectin; GAPDH, glyceraldehyde phosphate dehydrogenase; lox, lysyl oxidase; Sal, saline; SB, transforming growth factor β receptor inhibitor SB431542; SCcol, triple transgenic surfactant proteins-C promoter-reverse tetracycline transactivator/tet0cytomegalovirus promoter-Cre recombinase/homozygous floxed collagen, type I, alpha 1; SMA, smooth muscle actin; sma, smooth muscle actin; TGF- β , transforming growth factor β ; tsp, thrombospondin; WT, wild-type.

suggested⁵⁰ or that a plastic stem cell-like population capable of regenerating the epithelium and up-regulating mesenchymal proteins during injury was not labeled. A consensus definition of EMT has been elusive, and demonstrating an important contribution of proteins typically associated with activated fibroblasts may not be sufficient to prove EMTs. However, this approach may lead to a better understanding of the functions of different cell types during fibrogenesis, which may ultimately move the field beyond this controversy.

One critical limitation of a fate-mapping/costaining approach is the descriptive nature of the findings as opposed to a functional analysis. We found that epithelial cell-derived type I collagen affects both the overall amount of fibrosis and may have an effect on survival through regulating the lung injury response. Some of this effect may be through collagen-mediated cell signaling. We found that type I collagen is produced quickly in response to injury and leads to activation of the collagen signaling acutely after bleomycin injury. Indeed, there is mounting evidence that collagen signaling has important functions in inflammation and fibrogenesis.^{51,52} We used phosphorylation of the fibroblast collagen receptor, DDR2, as an indicator of epithelial cellderived collagen signaling on neighboring fibroblasts. In vitro, we found collagen-dependent activation of fibroblasts (Figure 5). Deleting endogenous collagen expression resulted in attenuated expression of several ECM genes, and addition of exogenous collagen restored expression of these genes (Figure 5B). These studies suggest the presence of a collagen signaling pathway, leading to augmented fibroblast production of ECM. It is unclear whether the cellular source of collagen I is important in determining this response in vivo, but differences in the timing and localization of collagen production after injury are potentially important. We are currently investigating the role of specific collagen receptors in regulating this response. Several studies support an influence of DDR2 activation on fibroblast function and that collagen I can signal through other cell surface receptors.⁴⁶ If type I collagen indeed promotes further production of type I collagen, as our data suggest, then epithelial cells may represent an important early source of collagen I necessary for progressive fibrosis. Thus, the contribution of type I collagen from different cell types is unlikely to be a simple sum calculation.

Deletion of type I collagen within lung epithelial cells surprisingly promotes sustained lung inflammation after bleomycin injury. These findings suggest that removing a significant source of type I collagen impairs the lungs ability to replace the damaged normal matrix with a fibrotic matrix, leading to prolonged inflammation. The bleomycin model is often criticized as a poor model of IPF in part because of the robust inflammation that occurs after bleomycin contrasted with the paucity of inflammation in IPF.^{53,54} However, fibrogenesis often occurs in the context of acute and chronic inflammation, and factors produced during the injury/inflammation phase are thought to facilitate the fibrogenic phase by promoting fibroblast activation.^{1,55} Our data suggest that factors produced during the fibrogenic phase, particularly collagen I, may regulate termination of the injury/inflammatory phase. As mentioned,

some evidence supports an anti-inflammatory role for type I collagen. Indeed we found that deleting type I collagen within primary lung fibroblasts augments expression of several inflammatory cytokines, suggesting autocrine and paracrine inhibition of inflammatory cytokine expression through type I collagen signaling. Our results may thus translate better to inflammatory fibrotic disease rather than IPF and suggest that targeting collagen production after robust lung injury must be taken with caution.

This study used type I collagen as a prototypical secreted mesenchymal protein that is critical for fibrogenesis, but activated lung epithelial cells likely produce and secrete other mesenchymal proteins that may promote fibroblast activation. We found that AECs have high expression of a number of mesenchymal genes, including CTGF, fibrillar collagens, and thrombospondin. CTGF expression is increased within fibroblasts in patients with IPF.⁵⁶ CTGF expression is induced by TGF- β in a number of fibroblast cell lines and studies that used a CTGF-promoter GFP transgenic reporter mouse reported CTGF promoter activity exclusively within activated fibroblasts⁵⁷ in a model of skin fibrosis. CTGF has thus been used as an effective marker of an activated, fibrotic fibroblast.⁵⁸ CTGF expression has been found within lung epithelial cells and endothelial cells during fibrogenesis and has been used as a marker of EMTs, but the functional importance of epithelial cell-derived CTGF has not been reported.^{59,60} Inhibition of CTGF has been shown to attenuate fibrosis in animal models.^{61,62} Thrombospondin expression has also been used as a marker of fibrogenic cells, and thrombospondin is thought to enable activation of latent TGF- β , potentially promoting fibroblast activation. A recent report however found that thrombospondin-null mice are not protected from bleomycin-induced fibrosis.⁶³ In addition to type I collagen, type III collagen is often used as a marker of activated fibroblasts, and mice with deletion of type III collagen have impaired type I collagen fibrillogenesis.⁶⁴

Future studies that use this more functional approach may help resolve controversies from descriptive fate-mapping studies. The contribution of collagen I from other cell types can be pursued in future studies, and epithelial cell-specific deletion of other genes typically associated with activated fibroblasts (collagen III, fibroblast-specific protein, SMA, CTGF, etc) could be applied to better understand the contribution of epithelial cell activation to fibrogenesis. We used PCR primers to specifically detect the presence of a truncated Col1a1 mRNA (with exon 1 directly adjacent to exon 6), indicating lung epithelial cell-specific origin of this important fibrotic protein. We cannot exclude the possibility that translation of this modified mRNA results in a truncated polypeptide affecting epithelial cell phenotype, but this limitation does not affect the overall conclusion that activated epithelial cells are actively expressing Colla1. This PCR approach does not enable histological localization of the epithelial-derived cells with activation of the Colla1. This approach importantly uses the intact native promoter rather than reporter transgenes with a fragment of the promoter. This is important because different cell types often activate transcription of the same genes by using different promoter regions, some of which are distant from the transcriptional start site.^{65,66} Expression of mesen-chymal proteins from nontraditional cell types may use previously uncharacterized promoter regions. Future studies will investigate differences between epithelial cells and fibroblasts in promoter activation sites. A more complete understanding of cellular contributions to progressive fibrogenesis may lead to new targets for potential therapeutic intervention.

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Supplemental Data

Supplemental material for this article can be found at *http://dx.doi.org/10.1016/j.ajpath.2013.07.016*.

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