Circulating fibrocytes traffic to the lungs in response to CXCL12 and mediate fibrosis

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Introduction

Idiopathic pulmonary fibrosis (IPF) is a chronic and often fatal pulmonary disorder with a survival rate as low as 30% 5 years after diagnosis (1, 2). The prevalence of this disease increases with age — with an incidence as high as 250 cases per 100,000 in the elderly (2, 3). There also appears to be an increase in the incidence of the disease in many Western countries, including the United Kingdom, New Zealand, and Germany (3). Conventional approaches to treating IPF have focused on immunosuppressive therapy, predominantly steroid usage, which has yielded poor results. The pathogenesis of pulmonary fibrosis is characterized by dysregulated and inappropriate tissue remodeling and enhanced vascular remodeling, fibroproliferation, and deposition of ECM (4). Without resolution, continued fibrosis leads to loss of lung function and ultimately death.

Fibrocytes present in the peripheral circulation were first identified a decade ago (5). These cells comprised a minor component of the circulating pool of leukocytes (less than 1%) and express a characteristic pattern of markers, including collagen I (Col I) and CD45 (5). When cultured in vitro, these cells became adherent and developed a spindle-shaped morphology (5). Subsequent studies have revealed that these circulating fibrocytes express chemokine receptors such as CXCR4 and CCR7 (6). Furthermore, circulating fibrocytes have been shown to traffic to wounds (6). They express ECM proteins, such as pro-Col I and pro–Coll III, which may be involved in connective tissue formation (6–8). No study has actually demonstrated that fibrocytes contribute to deposition of ECM and the promotion of fibrosis in vivo, however.

Tissue fibroblasts have classically been held to be responsible for repair and remodeling within the lung, although this view has been challenged since those fibroblasts observed in the fibrotic lung show phenotypic differences to their counterparts in the lungs of healthy individuals (9, 10). Thus, it is possible that, in common with their perceived role in fibrosis and wound repair, circulating fibrocytes may also be recruited to injured lungs as an integral component of the pathogenesis of pulmonary fibrosis.

Results

Human fibrocytes express CD45, Col I, and CXCR4 and migrate in response to CXCL12. To dissect the role of fibrocytes in pulmonary fibrosis, we initially isolated and characterized a population of fibrocytes from human peripheral blood (Figure 1 and Table 1). Although these cells are a minor component of the PBMC pool, by using a technique of immunodepletion and culturing cells in vitro, we were able to purify a population of long, spindle-shaped cells that morphologically resemble fibrocytes (6). We began our characterization of these purified fibrocytes by using flow cytometry to assess changes in the expression of the key fibrocyte markers CD45, CD34, Col I, and α-smooth muscle actin (α-SMA) as the cells matured in culture (Table 1). Our results show that these newly purified fibrocytes expressed high levels of CD45, CD34, and Col I, yet after 1 week in culture, expression of CD34 levels dropped dramatically and continued to decrease throughout the duration of the experiment (Table 1). CD45 expression began to fall at week 2, and by the third week (28–35 days after cell isolation) there was an approximate threefold reduction in the expression of CD45 by fibrocytes (Table 1). By contrast, Col I expression remained elevated at all time points examined (Table 1). Consistent with other published observations, there was no expression of α-SMA in freshly purified fibrocytes (6, 11), but after 3 weeks in culture almost one-third of fibrocytes were found to stain positive for α-SMA (Table 1). Furthermore, addition of TGF-β1 to cultured fibrocytes enhanced the kinetics of this differentiation (data not shown), which is again consistent with published observations (6, 11). Changes in the cell surface phenotype of cultured fibrocytes similar to those described above were also observed by immunostaining (Figure 1A). Here, purified fibrocytes...
cultured for 3 weeks (28–35 days after their initial isolation) were found to express intracellular Col I (Figure 1A), cell surface CXCR4 (Figure 1A), and α-SMA (Figure 1A).

Next, we triple-stained newly purified fibrocytes with Col I, CD45, and CXCR4 and analyzed them by flow cytometry (Figure 1B). Initially, we gated CD45+ fibrocytes and then examined the resulting population for dual expression of CXCR4 and Col I. In this way, we identified a major population of CD45+Col I+CXCR4+ human fibrocytes and a smaller population of CD45+Col I+CXCR4− fibrocytes (Figure 1B). To determine whether the expression of CXCR4 on the surface of purified fibrocytes was functional, we performed chemotaxis assays to look for specific migration in response to CXCL12. We observed significant chemotaxis at both 30 and 100 ng/ml CXCL12 (Figure 1C), indicating that CD45+Col I+CXCR4+ fibrocytes undergo directed movement in the presence of a chemotactic gradient in vitro.

Having established that cultured human fibrocytes can migrate in response to CXCL12 in vitro, we wanted to examine whether these fibrocytes could migrate in vivo and, in particular, whether they could specifically migrate into lungs undergoing a fibrotic response. To test this possibility we used a murine model of bleomycin-induced pulmonary fibrosis (12). Initially, cohorts of mice were treated with either bleomycin, saline, or nothing (naive mouse, day 0) and then sacrificed at day 1, 2, 4, 8, 16, and 20 to look for evidence of fibrosis. Exuberant collagen deposition within the lung is a key marker of fibrosis (9); on this basis we examined changes in pro-Col I and pro-Col III gene expression using real-time quantitative RT-PCR (Figure 2, A and B). Our data revealed that transcription of both pro–Col III and pro-Col I was dramatically upregulated in mice exposed to bleomycin (up to 30-fold higher) as compared with mice treated with saline alone (Figure 2, A and B). To confirm that collagen gene expression was translated into protein, we next measured total collagen deposition in the lung by the Sircol assay (13). We observed that the level of total collagen deposition was significantly greater in mice exposed to bleomycin compared with those lungs that received saline alone. These results suggest that a circulating pool of human fibrocytes exist that specifically migrate in response to CXCR4/CXCL12 and traffic to fibrotic lungs.

A kinetic analysis of the onset of fibrosis reveals dramatic collagen deposition beginning 4 days after bleomycin exposure. To more fully elucidate the role of fibrocytes in the development of fibroproliferative disorders of the lung, we focused our studies on the murine model of bleomycin-induced pulmonary fibrosis (12). Initially, cohorts of mice were treated with either bleomycin, saline, or nothing (naive mouse, day 0) and then sacrificed at day 1, 2, 4, 8, 16, and 20 to look for evidence of fibrosis. Exuberant collagen deposition within the lung is a key marker of fibrosis (9); on this basis we examined changes in pro-Col I and pro-Col III gene expression using real-time quantitative RT-PCR (Figure 2, A and B). Our data revealed that transcription of both pro–Col III and pro-Col I was dramatically upregulated in mice exposed to bleomycin (up to 30-fold higher) as compared with mice treated with saline alone (Figure 2, A and B). To confirm that collagen gene expression was translated into protein, we next measured total collagen deposition in the lung by the Sircol assay (13). We observed that the level of total collagen deposition was significantly greater in mice exposed to bleomycin compared with those lungs that received saline alone. These results suggest that a circulating pool of human fibrocytes exist that specifically migrate in response to CXCR4/CXCL12 and traffic to fibrotic lungs.

Table 1
Kinetics of percentage of positive Col I, CD45, CD34, and α-SMA expression in isolated and purified human fibrocytes as determined by FACS

<table>
<thead>
<tr>
<th>Marker</th>
<th>7–14 d after cell isolation immediately after purification</th>
<th>14–21 d after cell isolation 1 wk after purification</th>
<th>21–28 d after cell isolation 2 wk after purification</th>
<th>28–35 d after cell isolation 3 wk after purification</th>
</tr>
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<tr>
<td>Col I</td>
<td>90</td>
<td>90</td>
<td>88</td>
<td>91</td>
</tr>
<tr>
<td>CD45</td>
<td>90</td>
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<td>5</td>
</tr>
<tr>
<td>α-SMA</td>
<td>0</td>
<td>0</td>
<td>10</td>
<td>30</td>
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Figure 1
Characterization of human fibrocytes and their trafficking to bleomycin-induced lung fibrosis in SCID mice. (A) Human fibrocytes stained with control Ab, human Col I, human CXCR4, and human α-SMA 3 weeks after purification. Representative fields viewed at ×400 magnification. (B) Isolated human fibrocytes triple-stained for Col I, CD45, and CXCR4 were examined by FACS analysis. Data shows CD45+ fibrocytes that were examined for dual expression of CXCR4 and Col I. (C) Isolated human fibrocyte chemotaxis in response to 0, 30, and 100 ng/ml of CXCL12. Data are representative of three experiments. *P < 0.05. h.p.f., high-powered field. (D) Isolated human fibrocytes (106) were injected into the tail vein of SCID mice at day 4 after treatment with either intratracheal bleomycin (Bleo) or saline, followed by recovery of cells from the lungs at day 8. Cells were then stained for human CD45, Col I, and CXCR4 and analyzed by flow cytometry to determine trafficking of human fibrocytes to the lungs of SCID mice. *P < 0.05.
collagen protein increased dramatically in bleomycin-treated mice, especially between days 4 and 20 (Figure 2C). The levels of collagen found in saline-treated mice increased slightly in the first 2 days and then decreased to the levels observed in the naive mouse (day 0; Figure 2C). Thus, this data reveals that 4 days after mice have been exposed to bleomycin there is a significant increase in both collagen gene expression and protein deposition, which is indicative of the onset of fibrosis.

**CD45**+Col I+CXCR4+ fibrocytes accumulate in the lungs of bleomycin-treated animals. To characterize the role of murine CD45+Col I+CXCR4+ fibrocytes in our mouse model of bleomycin-induced pulmonary fibrosis, we isolated cells from the lungs and blood buffy coats of mice that had been exposed to bleomycin or saline for 1, 2, 4, 8, 16, and 20 days, triple-stained for CD45, Col I, and CXCR4, and examined them by flow cytometry (Figure 3A). Our results reveal a dramatic increase in the number of infiltrating CD45+Col I+CXCR4+ fibrocytes found in the lungs of mice exposed to bleomycin. Increased expression of CD45+Col I+CXCR4+ cells began to appear 2 days after bleomycin treatment, became maximal at day 8, and remained elevated at days 16 and 20. Expression of CD45+Col I+CXCR4+ cells in saline-treated mice also increased initially before returning to the levels observed in the naive mouse (day 0) by days 16 and 20 (Figure 3A). The most likely explanation for this latter phenomenon is that intratracheal instillation of saline itself is not a normal event and could have promoted an inflammatory response. In contrast to the time-dependent accumulation of murine fibrocytes in the lungs of bleomycin-exposed animals, the steady-state levels of circulating CD45+Col I+CXCR4+ fibrocytes in the blood remained similar in both groups (Figure 3A).

Previous studies have reported that fibrocytes can express CCR7 in addition to CXCR4 (6), and we have now identified a second fibrocyte population that is CD45+Col I+CCR7+ (Figure 3B and data not shown). These CCR7+ fibrocytes also traffic to the lungs of bleomycin-treated mice (Figure 3B); however, the absolute number of CCR7+ fibrocytes found in the fibrotic lung is twofold to threefold lower than the number of CXCR4+ fibrocytes present under similar conditions (i.e., compare A and B in Figure 3). Thus, these data suggest that although two chemotactic chemokine receptors are present on the surface of fibrocytes, CXCR4 appears predominant for the recruitment of fibrocytes to fibrotic lungs.

We have already shown that human CD45+Col I+CXCR4+ fibrocytes traffic through the bloodstream and are recruited into the lungs of bleomycin-treated animals (Figure 1D), and this suggests that murine CD45+Col I+CXCR4+ fibrocytes may be recruited in a similar manner. There appears to be insufficient peripheral blood fibrocytes present to drive the dramatic intrapulmonary recruitment of progenitor cells (Figure 3A), however. One explanation to account for this disparity is that the circulating pool of fibrocytes is continually replenished from the bone marrow. In support of this notion, recent work has suggested that mesenchymal stem cells originating from the bone marrow traffic to organizing alveolitis in pulmonary irradiated bone marrow chimeric mice (14). To formally test this hypothesis, therefore, bone marrow was recovered from mice treated with bleomycin or saline for 8 days, triple-stained for CD45, Col I, and CXCR4, and analyzed by flow cytometry (Figure 3C). Our results clearly indicate a dramatic increase in the number of CD45+Col I+CXCR4+ fibrocytes found in the bone marrow of bleomycin-challenged animals in comparison with the normal saline control mice. These data therefore suggest that bone marrow is at least one source of CD45+Col I+CXCR4+ circulating fibrocytes and that fibrocyte numbers in the bone marrow appear to be higher in mice treated with bleomycin.

For CD45+Col I+CXCR4+ fibrocytes to enter the lungs from the peripheral circulation, the cells must be actively recruited. One likely mechanism for achieving this is the chemokine receptor CXCR4 and its cognate ligand CXCL12, which has been shown to regulate cellular migration in other systems (6, 15, 16). To establish whether indeed a CXCL12 gradient exists between the lungs and the peripheral blood, lung homogenates were prepared and the plasma isolated from naive, saline-, or bleomycin-treated animals in each group. Data represent the mean ± SEM. *P < 0.05, significant differences between bleomycin and saline groups.
they decreased by day 20, again suggesting that the initial instillation of saline promoted an inflammatory response. Thus, these data support the notion that a CXCL12 gradient exists between the lungs and plasma of bleomycin-treated mice, which could promote the recruitment of CD45+Col I+CXCR4+ fibrocytes to the fibrotic lung. A similar chemokine gradient between the lungs and plasma of bleomycin-treated mice exists for CCL21 (6Ckine), but not for CCL19 (ELC), the putative ligands for CCR7. Here, peak levels of CXCL12 chemotactic axis reduces intrapulmonary recruitment of CX45+Col I+CXCR4+ fibrocytes by flow cytometry at day 16 (Figure 4A).

Our results show that significantly fewer CD45+Col I+CXCR4+ fibrocytes (P < 0.05) were observed in anti–CXCL12-treated bleomycin-exposed lungs than were observed in lungs exposed to bleomycin in the presence of control Ab’s (Figure 4A). Indeed, the number of infiltrating CD45+Col I+CXCR4+ fibrocytes observed in the anti–CXCL12-treated bleomycin-exposed mice was not significantly different from the number of CD45+Col I+CXCR4+ fibrocytes found in the lungs of mice exposed to saline alone. And in keeping with our previous observations (Figure 3A), there was a slight increase in the number of CD45+Col I+CXCR4+ fibrocytes in the lungs of saline-exposed mice in comparison with the naive control (Figure 4A).

Next we examined the effects of daily injections of anti-CXCL12 Ab’s or control Ab’s on collagen deposition in the lungs of mice exposed to bleomycin for 16 days (Figure 4B). Our results show that pulmonary collagen deposition was significantly reduced (P < 0.05) in anti-CXCL12–treated bleomycin-exposed mice in comparison with those mice receiving bleomycin and control Ab’s (Figure 4B).

The addition of neutralizing CXCL12 Ab’s to bleomycin-treated lungs did not, however, completely attenuate collagen deposition to the level of the saline control (Figure 4B). Taken together, therefore, these data indicate that inhibition of the CXCR4/CXCL12 chemotactic axis reduces intrapulmonary recruitment of CD45+Col I+CXCR4+ fibrocytes and significantly abrogates lung fibrosis in bleomycin-exposed mice.

To ascertain whether the reduction in collagen deposition was truly a reflection of fewer infiltrating CD45+Col I+CXCR4+ fibrocytes, rather than as a result of generalized migratory inhibition of other circulating CXCR4+ PBMCs, we also examined changes in the recruitment of a panel of CXCR4+ cells to bleomycin-exposed lungs treated with anti-CXCL12 Ab’s (Figure 4C). Thus, mice were treated with bleomycin and given daily injections of either anti-CXCL12 or control Ab’s for 8 days. Thereafter, cells were isolated from bleomycin-exposed lungs treated with either anti-CXCL12 or control Ab’s and stained with a panel of markers (CD3, CD4, CD8, NK1.1, Ly6, and Mac) prior to analysis by flow cytometry. Our results show that there was no statistical difference in the numbers of CD4+ (CD4) and CD8+ (CD8) T cells, NK cells (NK1.1), and CXCR4/CXCL12 chemotactic axis. To do this we used neutralizing anti-CXCL12 Ab’s that we have previously shown to specifically and selectively target only CXCL12 (16). Therefore, mice received intratracheal injections of bleomycin together with daily injections of either anti-CXCL12 or control Ab’s for 16 days. In addition, one cohort of mice received saline only, and a second cohort received nothing (naive control). Subsequently, all the mice were sacrificed and examined for infiltrating CD45+Col I+CXCR4+ fibrocytes by flow cytometry.
研究报告

Figure 4

中性化抗CXCL12 Ab抑制肺内招募的CD45⁺Col I⁺CXCR4⁺细胞和显著减少肺癌在肺部暴露于肺部暴露的小鼠中的纤维化。A. 单细胞悬浮液从肺部的无菌或含盐-或含肺部暴露的小鼠中隔离，其中含肺部暴露的小鼠也接受了每天注射不同抗-CXCL12 Ab或对照Ab 16天。这些细胞随后被三重染色为CD45，Col I，和CXCR4，并通过流式细胞术分析。

B. 总胶原存在于肺部组织中，来自无菌，含盐-或肺部暴露的小鼠，其中的含肺部暴露的小鼠也接受了每天注射不同抗-CXCL12 Ab或对照Ab 16天。这些细胞随后被三重染色为CD45，Col I，和CXCR4，并通过流式细胞术分析。n=5样本/每组。数据代表平均±SEM。*P<0.05。

C. 单细胞悬浮液被隔离于肺部组织的无菌，含盐-，或含肺部暴露的小鼠中，其中的含肺部暴露的小鼠也接受了每天注射不同抗-CXCL12或对照Ab 8天。这些细胞随后被三重染色为CD3，CD4，CD8，NK1.1，Ly6，和Mac519（Mac）并分析流式细胞术。n=3肺/每组。

discussion

此研究提供了，据我们所知，第一例证据，显示能够判断CD45⁺Col I⁺CXCR4⁺细胞对肺部纤维化的影响。

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 neutrophils (Ly6)，和单核细胞/巨噬细胞（Mac） trafficking to the lungs of bleomycin-exposed mice treated with either anti-CXCL12 or control Ab’s (Figure 4C). Although CXCL12 clearly does mediate recruitment of CXCR4⁺ cells to the lungs，the fact that neutralizing anti-CXCL12 Ab’s did not block migration here can be explained by the observation that these PBMC’s express chemokine receptors other than CXCR4 (17, 18)。It is possible, therefore，that inhibition of a single chemokine receptor/ligand combination does not prevent these cell types from intrapulmonary infiltration in response to chemokines other than CXCL12.

To further verify that systemic addition of anti-CXCL12 Ab’s selectively reduced intrapulmonary infiltration of CD45⁺Col I⁺CXCR4⁺ fibrocytes and attenuated bleomycin-induced pulmonary fibrosis，we performed histopathology on H&E-stained lung sections (Figure 5A)。In addition，we used the collagen-specific dye picrosirius red to stain lung sections and perform morphometric analysis (Figure 5B)。For the purposes of quantitation，fibrosis was defined as areas that had an Ashcroft grade of 7 or 8 (19)。These grades represent severe distortion of structure and large fibrous areas，including honeycomb lung or total fibrous obliteration of the field。By both H&E staining (Figure 5A) and morphometric analysis (Figure 5B)，anti-CXCL12 Ab’s significantly reduced bleomycin-induced pulmonary fibrosis.

In addition，we immunostained lung sections exposed to bleomycin for 16 days in the presence of daily injections of either control Ab’s (Figure 5C，left panel) or neutralizing CXCL12 Ab’s (Figure 5C，right panel) with specific anti-α-SMA Ab’s。Levels of α-SMA staining in those mice receiving neutralizing anti-CXCL12 was significantly lower than those mice receiving control Ab’s (Figure 5C)。Thus，by blocking the recruitment of CD45⁺Col I⁺CXCR4⁺ fibrocytes to bleomycin-exposed lungs，expression of α-SMA was also reduced。This suggests that CD45⁺Col I⁺CXCR4⁺ fibrocytes recruited from the peripheral circulation may ultimately develop an α-SMA⁺ phenotype in fibrotic lungs，which is compatible with our in vitro findings for human fibrocytes (Figure 1A and Table 1).
shown to be important in the regulated movement of fibrocytes (6). Indeed, flow-cytometric analysis of our purified human fibrocytes indicated that at least two phenotypically distinct fibrocyte populations (CD45+Col I/CXCR4+ and CD45−Col I/CXCR4−) were present. Thus it is possible that CD45+Col I/CXCR4+ fibrocytes actually express CCR7 and respond to its cognate ligands CCL19 and CCL21. To that end, we have identified a second population of fibrocytes in our mouse model that expresses CD45, Col I, and CCR7, and that also traffic to the lungs of bleomycin-exposed animals. Furthermore, it is possible that our neutralizing CXCL12 Ab was not completely effective in perturbing the putative CXCL12 chemotactic gradient that we believe develops between the lungs and blood plasma during bleomycin exposure and that recruits CD45+Col I/CXCR4+ fibrocytes to the lungs during the onset of pulmonary fibrosis. Thus, in future studies we hope to design appropriate strategies to more fully attenuate CXCR4 and CCR7 biology and subsequently to determine whether this has an additive effect on abrogating the fibrotic response in our mouse model of bleomycin-induced pulmonary fibrosis.

In addition, several different fibroblast phenotypes, including Thy-1+, Thy-1−, cyclooxygenase1+, and telomerase+, have been shown to be present at the site of evolving pulmonary fibrosis (10, 24–30). Although it would appear that these fibroblast phenotypes may contribute to the deposition of ECM components during lung fibrosis, it is not clear whether the origin of Thy-1+, Thy-1−, cyclooxygenase1+, and telomerase+ fibroblasts is from an intrapulmonary or extrapulmonary cellular source. If these fibroblast phenotypes originate from circulating CXCR4+ fibrocytes, it would be expected that neutralizing CXCL12 Ab’s would block both the recruitment of Thy-1+, Thy-1−, cyclooxygenase1+, and telomerase+ fibroblasts and attenuate their ability to secrete ECM components. Alternatively, if Thy-1+, Thy-1−, cyclooxygenase1+, and telomerase+ fibroblasts originate from a cellular source other than circulating CXCR4+ fibrocytes, then their role in mediating pulmonary fibrosis would not be impaired under conditions of blocking the CXCL12/CXCR4 biological axis. Thus, collagen deposition mediated by these additional fibroblast phenotypes could further explain why the reduction in collagen expression observed in bleomycin-exposed mice treated with anti-CXCL12 Ab’s failed to reach the levels of the saline or naive control. Other cell types, including pericytes, have also been postulated to play a role in the onset of fibrosis in the liver and dermis (31–33), although there is currently no evidence to suggest that these cells are an important source of collagen deposition in pulmonary fibrosis.

A key marker of IFP is the dramatic rise in both collagen gene expression and deposition of collagen protein within the lungs. In our murine model of bleomycin-induced pulmonary fibrosis, increased expression of both pro-Col I and pro–Col III was observed beginning 4 days after instillation of bleomycin. In addition, collagen deposition was clearly detectable in lungs within 4 days of bleomycin exposure. Morphometric analysis of picrosirius red staining of lung sections at day 16 after bleomycin exposure further verified the presence of pulmonary collagen deposition. This collagen deposition was strongly attenuated in bleomycin-exposed mice treated with daily injections of neutralizing anti-CXCL12 Ab’s. The magnitude of collagen reduction we observed here was similar to that observed in several other cytokine neutralization studies (34–39).

We measured total collagen deposition using the Sircol assay, which we felt was actually more reproducible and reliable than the hydroxyproline assay (our unpublished observations). The kinetics of collagen deposition we report here are in keeping with results found in other published work, where significant collagen deposition was detectable within 3 days of bleomycin treatment using the hydroxyproline assay (34). In addition, the concept that collagen is detectable very early in the inflammatory process is consistent with several studies that have shown markedly elevated levels of collagen during acute lung injury in patients (40–42). Indeed, increased collagen levels in the bronchoalveolar lavage (BAL) fluid of patients suffering from acute respiratory distress syndrome were detectable 3 days after the onset of the disease (40). Moreover, early detection of collagen in BAL fluid in these patients was strongly associated with an increased risk of mortality (40). This does not mean, however, that we will ultimately see fibrosis in these patients, especially in light of the magnitude of counterbalancing measures such as proteinases present at the same time, which would lead to collagen degradation, remodeling of collagen deposition, and resolution. This concept would be analogous to the saline control–treated lungs. In contrast, under conditions of bleomycin-induced pulmonary fibrosis, we speculate that the previously published find-
ings of elevated TGF-β in this model contributes to tipping the balance in favor of increased collagen production and at the same time decreased collagen degradation, which would result in more fibrosis. Therefore, although elevated numbers of fibrocytes may be present under conditions of saline exposure, as compared with naïve mice there is no marked accumulation of collagen.

We observed significant recruitment of CD45−Col I CXCR4+ fibrocytes into the fibrotic lung as early as 2 days after bleomycin exposure and 2 days before significant collagen deposition was detected, further suggesting that CD45+Col I CXCR4+ fibrocytes may be involved in the onset of fibrosis. Maximal recruitment of CD45+Col I CXCR4+ fibrocytes was observed 8 days after the instillation of bleomycin, and by days 16 and 20 the number of infiltrating CD45−Col I CXCR4+ fibrocytes appeared to decrease. The fate of these cells remains to be determined, however. Indeed, the apparent loss of CD45−Col I CXCR4+ fibrocytes from fibrotic lungs, may, in fact, reflect a change in the phenotype of these cells. Myofibroblasts (CD45−Col I α-SMA+), which have been found in pulmonary fibrotic lesions but not in normal lungs (9), is one potential differentiation phenotype for fibrocytes. Although it has been suggested that myofibroblasts arise from tissue fibroblasts within the lung itself (30, 43), the true origin of these cells has yet to be definitively established (9). In addition, myofibroblasts are known to be a major source of pro-Col I expression in pulmonary fibrotic lesions (9, 44), and we have clearly demonstrated that in the presence of anti-CXCL12 Ab’s collagen deposition is significantly reduced in bleomycin-exposed mice. Further support for the notion that circulating fibrocytes may ultimately differentiate into myofibroblasts within the lungs stems from our observations and other recent reports (6, 11) that cultured human fibrocytes eventually lose expression of CD34 and CD45 and upregulate α-SMA, which is consistent with the known myofibroblast phenotype (CD34−CD45−Col I α-SMA+) (9).

Another factor that may determine the fate of CD45−Col I CXCR4+ fibrocytes during pulmonary fibrosis is apoptosis. The inflammatory environment associated with the onset of fibrosis typically contributes extensively to the programmed cell death of injured cells, including those of the respiratory epithelium and infiltrating white blood cells. Thus, it is important for cells migrating into the lungs to have in place a mechanism to survive the apoptotic landscape. In this respect, CXCR4+ fibrocytes may be ideally situated to avoid programmed cell death since ligation of CXCR4 by CXCL12 has been shown to regulate the cell death machinery through downstream activation of the PI3K, AKT, and MAP kinase pathways (45–49).

In conclusion, we have shown, we believe for the first time, that CD45−Col I CXCR4+ circulating fibrocytes contribute to the pathogenesis of pulmonary fibrosis. These cells, therefore, offer a novel therapeutic target for developing more effective treatments against this devastating disease.

Methods
Animal model of pulmonary fibrosis. Female C57Bl/6 mice (6–8 weeks old) purchased from The Jackson Laboratory (Bar Harbor, Maine, USA) were anesthetized with ketamine and given intratracheal injections of either 0.025 U of bleomycin (0.15 U/kg Blenoxane; Sigma-Aldrich, St. Louis, Missouri, USA) or sterile saline. Mice were sacrificed for analysis 1, 2, 4, 8, 16, and 20 days after exposure to bleomycin. In separate experiments, mice treated with bleomycin were given daily injections (500 µl) of 10 mg/ml F(ab’); fragments of either neutralizing goat anti-CXCL12 Ab or control Ab’s for 8 or 16 days prior to sacrifice. Animal use for these studies was approved by the Department of Laboratory Animal Medicine, David Geffen School of Medicine at UCLA.

Lung tissue preparation for protein assay. Bleomycin- or saline control–treated lungs were homogenized and sonicated in complete buffer (Roche Diagnostics Corp., Indianapolis, Indiana, USA) as previously described (41). Homogenates were centrifuged at 900 g for 15 minutes, filtered through 1.2 µm Sterile Acrodiscs (Gelman Sciences Inc., Ann Arbor, Michigan, USA), and frozen at −70°C until required.

CXCL12 ELISA. Lung homogenate and plasma were assayed for CXCL12 levels as previously described (50).

Sircol assay. The Sircol collagen assay (Biocolor Ltd., Belfast, United Kingdom) was performed following the manufacturer’s instructions (13). Briefly, Sirius red reagent (50 µl) was added to each lung homogenate (50 µl) and mixed for 30 minutes. The collagen–dye complex was precipitated by centrifugation at 16,000 g for 5 minutes, washed with ethanol, and dissolved in 0.5 M NaOH. Finally, the samples were introduced into a microplate reader and the absorbance determined at 540 nm.

Total RNA isolation and real-time quantitative RT-PCR. Total RNA was prepared using TRIzol and by following the manufacturer’s instructions. Real-time quantitative RT-PCR was performed using specific TaqMan primers and probes, the ABI Prism 7700 sequence detector, and SDS analysis software (Applied Biosystems, Foster City, California, USA), as previously described (50).

Purification of circulating human fibrocytes. Human fibrocytes were isolated according to previously published methods (5, 6, 11). Briefly, PBMCs were isolated from leukopheresis packs (HemaCare Inc., Woodland Hills, California, USA) using Ficoll density centrifugation. Cells isolated in this way were then cultured in DMEM supplemented with 20% FCS, penicillin, streptomycin, and t-glutamine for 7–14 days. Typically, 2 x 10⁶ PBMCs were plated into each T150 tissue culture flask. After 7–14 days, the media was aspirated from the PBMC culture and the remaining adherent cells washed with sterile PBS. Subsequently, the adherent cells were harvested using Accutase (Innovative Cell Technologies Inc., San Diego, California, USA), which we have found to be more efficient than the standard trypsin protocol for removing adherent cells from plastic. Previous data (6) have indicated that the main contaminating cell types in these cultures are B lymphocytes (about 3%), T lymphocytes (about 13%), and monocytes/macrophages (up to 20% of the PBMC culture). Thus, we used immunomagnetic selection to deplete B lymphocytes (Dynabeads M-450 anti-CD19; Dynal Inc., Lake Success, New York, USA), T lymphocytes (Dynabeads M-450 anti-CD2), and monocytes/macrophages (Dynabeads M-450 anti-CD14) from our crude fibrocyte preparation. Purified fibrocytes prepared in this way were returned to culture (DMEM supplemented with 20% FCS, penicillin, streptomycin and t-glutamine) for an additional 5 days before analysis. Fibrocytes were routinely greater than 90% pure as determined by staining with CD45, Col I, and CXCR4. Typically, our isolated fibrocytes represented about 0.5% of the overall pool of peripheral white blood cells.

FACS analysis. Cells isolated from the bone marrow, lungs, and peripheral circulation of mice were initially stained with FITC-labeled CXCR4 or FITC-labeled CCR7 and peridinin chlorophyll-a protein–labeled (PerCP-labeled) CD45. Subsequently, the cells were permeabilized using cytofix/cytoperm (BD Biosciences — Pharmingen, San Diego, California, USA) to facilitate intracellular staining of Col I. Col I was stained initially with anti-human or anti-mouse Col I (Chemicon International, Temecula, California, USA) and then phycocerythrin-conjugated (PE-conjugated) goat anti-rabbit Ig (Molecular Probes Inc., Eugene, Oregon, USA). Three-color analysis of the stained cells was performed on a FACScan flow cytometer (BD Biosciences — Pharmingen) using Cellquest 3.2.1f software. In separate experiments cells isolated from the lungs were individually stained with PE-conjugated anti-CXCR4 and then phycoerythrin-conjugated (PE-conjugated) goat anti-rabbit Ig.
CD3, CD4, CD8, NK1.1, Ly6, and Mac519 (Serotec Inc., Raleigh, North Carolina, USA). Except where indicated, all mAb's were purchased from BD Biosciences — Pharmingen. To obtain sufficient cells from the blood for analysis by FACS, two buffy coats (500 μl per buffy coat) were combined from each treatment condition. Thus data from fibrocytes isolated from the peripheral circulation are presented as cells per milliliter of blood.

Purified human fibrocytes cultured for either 1, 2, or 3 weeks after their initial isolation were taken and stained with FITC-labeled CXCR4, APC-labeled CD34, and PerCP-labeled CD45. Next, the cells were permeabilized using cytofix/cytoperm to facilitate intracellular staining of Col I and α-SMA. Col I and α-SMA were then stained with unconjugated rabbit anti-human Col I and mouse anti-human α-SMA Ab’s followed by anti-rabbit FITC and anti-mouse FITC, respectively. Finally, single-color analysis of the stained cells was performed on a FACScan flow cytometer using Cellquest 3.2.1f1 software.

Preparation of F(ab′)2 fragments of goat anti-human CXCL12 whole Ig, F(ab′)2 fragments of goat anti-human CXCL12 (16) were prepared using the ImmunoPure F(ab′)2 kit (Pierce Chemical Co., Rockford, Illinois, USA). Briefly, immobilized pepsin was used to digest whole Ig into Fc and F(ab′)2 fractions. These fractions were then separated by protein A–affinity chromatography such that the Fc portion bound to the column, while the F(ab′)2 portion was eluted, collected, and further purified by dialyzing against PBS, pH 7.4, overnight.

Fibrocyte chemotaxis. Chemotaxis was performed using fibronectin-coated filters (5 μm pore) as previously described (16).

Histology. Lungs were perfused with saline and then inflated with 1 ml of 4% paraformaldehyde at a constant pressure of 30 cmH2O. Lungs were ligated at the trachea, removed en bloc, and immersed in 4% paraformaldehyde for 24 hours, at which time they were changed to 70% alcohol before paraffin embedding, followed by sectioning and H&E staining.

Morphometric analysis and picrosirius red staining. Paraffin-embedded sections (3 μm thick) of bleomycin-exposed lungs treated with either a control Ab or neutralizing anti-CXCL12 Ab were exposed to xylene (twice, 1 minute each) and then washed initially in 100% ethanol (twice, 1 minute), followed by 50% ethanol (twice, 1 minute), and finally rinsed in distilled water for 10 minutes. Subsequently, the sections were treated with 0.2% phosphomolybdic acid (2 minutes), then washed initially in 100% ethanol (twice, 1 minute), followed by 50% ethanol (twice, 1 minute), and finally rinsed in distilled water for 10 minutes. Subsequently, the sections were 0.05 or less.

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Immunostaining. Histopathologic sections of lung tissue were prepared from mice that had been exposed to bleomycin for 16 days in the presence of either control or neutralizing anti-CXCL12 Ab’s and assessed by immunostaining of α-SMA (R&D Systems Inc, Minneapolis, Minnesota, USA). Purified fibrocytes (104 cells per well) were cultured on TissueTechs (BD Biosciences — Pharmingen) for the times indicated and then immunostained. Briefly, the cells were fixed in 4% paraformaldehyde and then incubated with a 1:1 mixture of 3% hydrogen peroxide in methanol. Subsequently, the cells were exposed to Power Block (BioGenex Laboratories, San Ramon, California, USA) for 30 minutes and, after washing in PBS, stained with a primary Ab (human and murine Col I from Chemicon International; CXCR4 from R&D Systems Inc; α-SMA; or an isotype control) for a further 30 minutes at room temperature. After incubation with the primary Ab, the cells were washed with PBS and then exposed to a biotinylated anti-mouse secondary Ab for 30 minutes. Subsequently, the cells were washed in PBS and avidin-binding complex (Vector Laboratories, Burlingame, California, USA). Reagent was added, and the cells incubated again for 30 minutes. After an additional PBS wash, the cells were exposed to the chromogen 3,3′-diaminobenzidine tetrahydrochloride (Vector Laboratories), which turns the positively staining cells brown. Finally, the cells were counterstained with hematoxylin, neutralized in 10% ammonia, and left to dry overnight. After drying overnight, Permount was added, together with a coverslip, and the cells were then viewed by microscopy.

Statistical analysis. The animal studies involved 3–6 mice for each treatment group. Data were analyzed on a Dell PC (Dell, Round Rock, Texas, USA) computer using the Statview 5.0 statistical package (Abacus Concepts Inc., Berkeley, California, USA). Comparisons were evaluated by Student’s unpaired t test. Results were statistically significant if P values were 0.05 or less.

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