Tumorigenesis and Neoplastic Progression

MCP1 Directs Trafficking of Hematopoietic Stem Cell-Derived Fibroblast Precursors in Solid Tumor

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Our previous studies have demonstrated that hematopoietic stem cells (HSCs) are a novel source of carcinoma-associated fibroblasts. However, the mechanisms regulating recruitment and homing of HSC-derived carcinoma-associated fibroblasts or their precursors to the tumor microenvironment are unknown. Herein, we demonstrate using a single cell transplantation model that circulating fibroblast precursors (CFPs) are of HSC origin. This population increased with tumor burden in vivo and functional in vitro studies showed that CFPs preferentially migrated and differentiated into fibroblasts in response to tumor, suggesting that HSC-derived CFPs serve as an intermediate between the bone marrow and tumor. Based on this chemotactic ability and our demonstration of a monocyte lineage origin for CFPs, we investigated the role of monocyte chemoattractant protein (MCP1) in mediating CFP recruitment/homing. Blocking tumor-produced MCP1 inhibited in vitro migration of CFPs in response to multiple tumor types, indicating broad biological significance for this CFP/chemokine interaction. In vivo, CCR2-expressing CFPs increased in circulation during the period of active tumor growth and stromal development. Inhibition of MCP1 during tumor development resulted in decreased tumor volume in tumor-bearing mice. Together these findings confirm an HSC origin for CFPs, demonstrate a role for MCP1 in regulating their contribution to the tumor microenvironment, and suggest a potential therapeutic target. One of the most prevalent stromal components of solid tumors is the fibroblast, termed carcinoma-associated fibroblasts (CAFs). Only recently have CAFs been appreciated for their potential ability to promote tumor progression,1,2 by stimulating growth of the cancer cell population, sustaining vascularization, inhibiting the immune response, and promoting invasion and metastasis.3–7 While the importance of the CAFs is becoming evident, the source of these cells is less clear. It was generally thought that all stromal fibroblasts were recruited from resident tissue fibroblasts. Recently, alternative sources for CAFs have been proposed. Several studies have suggested that fibroblasts/myofibroblasts in fibrotic lesions and tumors may arise from transdifferentiation of epithelial cells to myofibroblasts, a process termed epithelial-to-mesenchymal transdifferentiation.8–10 While a rare event, because epithelial-to-mesenchymal transdifferentiation–derived CAFs are thought to retain genetic and/or epigenetic insults of the tumor cell progenitor, it is thought that these cells have a distinct behavioral and molecular signature from CAFs of other origins.10 Recent studies have also suggested that CAFs may have a bone marrow (BM) origin.9–11

Adult bone marrow is thought to contain at least two types of stem cells, hematopoietic stem cells (HSCs) and mesenchymal stem cells (MSCs). HSCs are defined by their ability to repopulate the bone marrow and blood lineages of lethally irradiated mice. MSCs are a population of bone marrow cells that were defined in vitro based on their adherent properties12,13 and later by their ability to give rise to a variety of mesenchymal cell types including adipocytes,14,15 chondrocytes,14,16,17 osteoblasts/osteocytes,14,18 endothelial cells,19 and fibroblasts.20 Based on their adherent properties and their ability to give rise to mesenchymal cell types, it is widely held that the bone marrow source of fibroblasts is the MSCs. However, recent studies by our group have raised the intriguing

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Recent studies have shown that the tumor stromal microenvironment co-evolves with tumor and suggest that the tumor–stroma interaction holds exciting potential as a therapeutic target.
possibility that HSCs also give rise to fibroblasts. Our studies using clonally engrafted animals in which the bone marrow is reconstituted by a clonal population of cells derived from a single enhanced green fluorescent protein-positive (EGFP\(^+\)) HSC demonstrated an HSC origin for two types of fibroblasts precursors, bone marrow colony forming unit fibroblasts and circulating fibrocytes.\(^{21}\) Using these clonally engrafted mice, we have also established an HSC origin for multiple tissue fibroblasts (reviewed in\(^{22}\)), including those found in the tumor stroma.\(^{23}\)

Based on our previous studies demonstrating an HSC origin for at least a portion of CAFs,\(^{23}\) herein, we tested the hypothesis that an HSC-derived circulating fibroblast precursor (CFP) is recruited from the bone marrow to the tumor microenvironment. Using a single cell transplantation model in conjunction with the Lewis lung carcinoma model (LLC), our findings demonstrate that CFPs are of HSC origin. We also show that CFPs increase in the peripheral blood with tumor burden and contribute to tumor stroma formation \textit{in vivo}. Functional \textit{in vitro} studies show that CFPs preferentially migrate and differentiate to fibroblasts in response to tumor. To elucidate the mechanisms by which these cells are recruited to tumor microenvironment, we examined the role of chemokines with known roles in HSC and/or tumor biology including CXCL12 (stromal-derived factor-1), CCL2 (secondary lymphoid chemokine) and MCP1/CCL2 (monocyte chemotactic protein; MCP1). The stromal-derived factor-1 pathway has been shown to be a major mediator of cell trafficking in both HSC and tumor biology\(^{25}\) and fibrocytes have been shown to express the stromal-derived factor-1 receptor, CXCR4.\(^{26,27}\)

C57Bl/6-CD45.2 background) were provided by Dr. Masaru Okabe (Osaka University, Japan).\(^{31}\) Mice were bred and maintained at the Animal Research Facility of the Veterans Affairs Medical Center (Charleston, SC). Research was conducted in accordance with guidelines set by the PHS Policy on Human Care and Use of Laboratory Animals and the Institutional Animal Care and Use Committee, Department of Veterans Affairs Medical Center.

### Antibodies and Reagents

The following were purchased from BD Pharmingen (San Jose, CA): phycoerythrin (PE)-conjugated anti-Scal (anti-Ly-6A/E[D7]); allophycocyanin (APC)-conjugated anti-c-kit (2B8); biotinylated and fluorescein isothiocyanate-conjugated anti-CD34 (RAM34); PE-conjugated, biotinylated, or purified anti-Gr-1 (anti-Ly-6G[RB6-8C5]); PE-conjugated or biotinylated anti-CD45R/B220 (RA3-6B2); PE-conjugated anti-Thy-1.2 (30-H12); biotinylated or purified anti-TER-119; biotinylated anti-CD3e, PE and APC-cy7-conjugated anti-CD45 (Leukocyte Common Antigen, Ly-5.5;30-F11); PE and fluorescein isothiocyanate-conjugated anti-Mac-1 (CD11b; M1/70); PE-conjugated and biotin-conjugated anti-CD8a (53-6.7); PE-conjugated and biotin-conjugated anti-CD4 (GK1.5); and PE-conjugated anti-CD45.1 (A20). PE-conjugated anti-F4/80 (BM8), and fluorescein isothiocyanate-, APC-, and biotin-conjugated anti-CD34 (RAM34) were obtained from eBioscience (San Diego, CA). Discoidin domain receptor 2 (DDR2) goat polyclonal antibodies were purchased from Santa Cruz Biotechnology (Santa Cruz, CA). Biotin-conjugated purified anti-collagen type 1 (rabbit) antibody (Col-1; BAC) and biotin-conjugated Rabbit IgG whole molecule was purchased from Rockland Immunodiagnostic Systems (Gilbertsville, PA). Purified rabbit monoclonal CCR2 antibody (CCR2; E68) and antibodies to vimentin were purchased from Abcam (Cambridge, MA). Antibodies to \(\alpha\)-smooth muscle actin (\(\alpha\)-SMA) were obtained from Sigma (St. Louis, MO) and Abcam. MCP1/CCL2 neutralizing antibody was purchased from Rockland Immunodiagnostic Systems (Gaithersburg, MD). PE-conjugated, APC-conjugated, Cy5- and Cy3-conjugated secondary antibodies were purchased from Jackson ImmunoResearch (West Grove, PA).

### Transplantation

Ten- to fourteen-week-old male or female EGFP mice were used as donors. Cional cell transplantation was performed using previously described methods with minor modifications.\(^{21,23}\) Bone marrow cells were flushed from femurs and tibiae, pooled, and washed in PBS containing 0.1% bovine serum albumin (BSA). Mononuclear cells were isolated by gradient separation using Lympholyte M (Cedarlane Laboratories Limited, Ontario) and lineage-negative (Lin\(^-\)) cells were prepared by negative selection using biotin-conjugated antibodies (anti-CD20, -Gr-1, -CD4, -CD8, -TER-119) and MACS anti-biotin microbeads (Miltenyi Biotec Inc, Auburn, CA). Lin\(^-\) cells were stained with PE-conjugated anti-Scal, APC-conjugated anti-c-kit, and biotinylated anti-CD34 followed by streptavidin-conjugated APC-Cy7 and resuspended (1 \times 10^6 cells/ml) in Ca^{2+}-, Mg^{2+}-free Hanks balanced salt solution (Invitrogen, Gaithersburg, MD) with 2% fetal bovine serum (FBS; Atlanta Biologicals, Lawrenceville, GA), 10 mmol/L HEPES, and 1% penicillin/streptomycin. Cells were then incubated with Hoechst 33342 (Sigma; 5 mg/
ml) at 37°C for 60 minutes. After washing, propidium iodide was added and cell sorting was performed using a MoFlo Fluorescence Activated cell sorter (Dako Cytomation, Fort Collins, CO). The gate used for side population cells corresponded to R3 and R4 subfractions described by Goodell et al.32 and R1 and R2 described by Matsuzaki et al.33 Single Lin− Sca-1+ ckit− CD34+ side population cells were deposited into wells of Corning U-bottom 96-well culture plates (Corning, NY) using the CyCLONE cell deposition system. Eighteen hours after single-cell deposition, wells containing single cells were identified and the incubation continued for a total of 7 days at 5% CO2, 37°C in media containing α modification of Eagle’s medium (ICN Biomedicals, Aurora, OH), 20% FBS, 1% deionized fraction V BSA, 1 × 10–6 mol/L 2-mercaptoethanol (Sigma), 100 ng/ml SCF, and 10 ng/ml G-CSF. This combination of cytokines stimulates proliferation of primitive hematopoietic progenitors.34 Because the majority of HSCs are dormant in cell cycle and begin cell division a few days after initiation of cell culture, we selected clones consisting of no more than 20 cells after incubation. This method significantly enhanced the efficiency of generating mice with high-level multilineage engraftment.21,23 Approximately one-fourth of all mice transplanted using these methods resulted in high-level multilineage engraftment above 50%.

Recipient C57Bl/6-CD45.1 mice (10- to 14-week-old) were given a single 950-cGy dose of total-body irradiation using a 4 × 10–6 V linear accelerator. Contents of wells (≤20 clonal cells) were injected via tail vein into lethally irradiated mice with 500 CD45.1 (EGFP−) bone marrow Lin− ckit− Sca-1+ CD34+ radioprotective cells. Osawa et al35 showed that these CD34+ cells are effective radioprotective cells and contain almost no HSCs. For generation of mice transplanted with unmanipulated bone marrow cells, bone marrow was flushed, washed, and prepared from EGFP mice as described above. One million white blood cells were delivered to lethally irradiated C57Bl/6-CD45.1 mice as described.

For hematopoietic engraftment analysis, peripheral blood was obtained from the retro-orbital plexus of anesthetized mice and red blood cells were lysed (1× PharM Lyse; BD Pharmingen). Donor-derived cells (EGFP+) in T cell, B cell, granulocyte, and monocyte/macrophage lineages were analyzed by staining with PE-conjugated anti-Thy-1.2, anti-CD45R/B220, anti-Gr-1, and anti-Mac-1, respectively. Mice with total engraftment levels below 50% were not used for studies.

CFP Sorting

Peripheral blood (1.5 to 2 ml) was obtained via cardiac puncture from isoflurane-anesthetized mice using a 28½ gauge syringe resulting in euthanasia by exsanguination. Samples were then lysed (BD PharmLyse) for 15 minutes, washed in PBS containing DNase (Invitrogen, Carlsbad, CA) and blocked with normal goat serum (Jackson ImmunoResearch) for 20 minutes at 4°C. Cells were stained using optimum concentrations of primary antibodies for 20 minutes at 4°C, washed and incubated with appropriate secondary antibodies for 20 minutes at 4°C. Samples were analyzed and/or sorted using a MoFlo high-performance cell sorter.

Cell Lines

Murine Lewis lung carcinoma (LLC 1), mammary carcinoma (EMT6), osteosarcoma (K7M2) and mouse lung epithelial (MLE-12) cell lines were obtained from American Type Culture Collection (Manassas, VA) and propagated and harvested using standard cell culture methods. All cells were grown to 95% confluency in 5% CO2, 37°C. LLC and K7M2 cells were cultured in Dulbecco’s Modified Eagle medium (DMEM; Invitrogen) containing 10% FBS. EMT6 cells were grown in Waymouth’s MB 752/1 Media with 2 mmol/L L-Glucose (Invitrogen) containing 15% FBS. MLE-12 cells were cultured in 50:50 DMEM: Ham’s F12 Media (Sigma) containing 2% FBS and insulin-transferrin-sodium selenite media supplement (Sigma).

For injection into mice, recovered LLC cells were resuspended in serum-free medium at a concentration of 1 × 106 cells/0.2 ml and injected subcutaneously into anesthetized engrafted mice. Tumor length and width was measured daily to calculate tumor size (L × W).

Migration Assays

Migration assays were performed using 8.0 μm pore size Costar Transwell inserts (Corning) according to the manufacturer’s protocol. Peripheral blood, sorted CFPs or control MLE-12 cells were resuspended at 1 × 107 cells/ml in DMEM containing 10% FBS. Experimental wells received 600 μl of 24-hour conditioned tumor cell media (LLC-CM, EMT6-CM, or K7M2-CM), or CM with the addition of 4.5 μg neutralizing antibodies to MCP1 and control wells received 600 μl DMEM containing 10% FBS. Transwell inserts were placed and peripheral blood cells, sorted CFPs or control MLE-12 cells were added in 200 μl. After a 3-hour incubation at 37°C in 5% CO2, migrated cells were recovered from the lower well using Trypsin-EDTA 0.25% and counted. Experiments were conducted with four to six replicates using four mice per replicate.

Culture of Sorted CFPs

Sorted CFPs were grown in human fibronectin coated 4- or 8-well culture slides (BD Biosciences, San Jose, CA) at 37°C in 5% CO2 for up to 14 days. Experimental wells received 24-hour tumor-CM, or tumor-CM containing 4.5 μg of neutralizing MCP1 antibody. Control wells received 10% FBS/DMEM (control), 10% FBS/DMEM containing 10% mouse serum (Equitech-Bio, Kerrville, TX), or tumor-CM alone (without sorted cells added to well). In some experiments, macrophage-colony stimulating factor (M-CSF; R&D Systems) was added to CFP cultures at 10 μg/ml.

Enzyme-Linked Immunosorbent Assay

Enzyme-linked immunosorbent assay (ELISA) was performed on 24-hour conditioned cell culture supernatants.
using R&D Systems Quantikine ELISA kits (R&D Systems) according to the manufacturer’s protocol and analyzed with a SPECTRAmax M2 microplate reader running SoftMax Pro software (Molecular Devices, Sunnyvale, CA).

**Immunohistochemistry and Microscopy**

Cultured cells were fixed in 4% paraformaldehyde for 20 minutes and tumor tissues were fixed in 4% paraformaldehyde for 6 hours. Tumors from engrafted mice were processed for paraffin sectioning as previously described using protocols that allow for preservation of fluorescence intensity. For staining, samples were permeabilized in 0.01% Triton X-100/PBS (15 minutes), blocked in 3% BSA/5% normal donkey serum/PBS (2 hours), and incubated with appropriate primary antibodies diluted in 3% BSA/5% normal donkey serum/PBS (1 hour). Samples were washed in PBS and incubated for 1 hour with fluorochrome-conjugated secondary antibodies diluted in 3% BSA/5% normal donkey serum/PBS. The samples were again washed in PBS, incubated with Hoechst 33342 nuclear dye (diluted 1:20,000 in PBS; 8 minutes), washed in PBS and coverslipped. Epifluorescence and differential interference contrast (DIC) microscopy was conducted using a Leica DMR microscope with a narrow band-pass green fluorescent protein excitation cube. Images were processed using Adobe Photoshop CS2 (Adobe Systems Inc., San Jose, CA).

**Quantitative Real-Time PCR Analysis**

Cells were sorted using the MoFlo cell sorter. Total RNA was extracted from sorted cells using TRIzol (Invitrogen) following the manufacturer’s instructions. Two μg of total RNA was converted to cDNA by the High Capacity cDNA Reverse Transcription Kit (Applied Biosystems, Foster City, CA) following company recommended protocol. Quantitative real-time PCR analysis (qPCR) was performed for different genes in separate wells (singleplex assay) of 96-well plate in a reaction volume of 20 μl. Predesigned unlabeled gene-specific PCR primers and TaqMan dye-labeled probes (Assays-on-Demand Gene Expression mix) were purchased from Applied Biosystems. Glyceraldehyde-3-phosphate dehydrogenase served as the endogenous control for all experiments. Three replicates of each sample were amplified to give one measurement for a sample. Primers were tested for equal efficiency over a range of target gene concentration before use. All primers were designed to cross exon—exon boundaries. The PCR reaction mixture (including 2× TaqMan Gene Expression Master Mix, 20× Assays-on-Demand Gene Expression Assay Mix, 100 ng of cDNA) was run in Applied Biosystems ABI Prism 7500 Sequence Detection System instrument using universal thermal cycling parameters. The relative quantification of target gene expression in a test sample to a control calibrator sample (ΔΔCt Method) was used for data analysis.

**Inhibition of CFP Trafficking in Vivo**

C57Bl/6-CD45.1 mice were subcutaneously injected with LLC cells as described above. Half of the mice received MCP1 neutralizing antibody (66 μg/mouse; stock 1 μg/ml) via tail vein injection on days 2, 4, 6, 8, 10, 12, 14, and 16 after tumor cell delivery. This dose was based on ELISA studies of MCP1 production by LLC cells. Mice were euthanized on day 18 of tumor growth. Tumor volume was calculated using the ellipsoid volume formula ($\pi/6 \times L \times W \times H$). Tumors were then divided into two samples, one half was fixed and processed for paraffin sections and the other half was digested overnight with trypsin-EDTA 0.25%, then stained and analyzed using the MoFlo cell sorter.

**Figure 1. Multilineage engraftment.** Panel A shows forward and side scatter gate (R1), B shows isotype control for CD45.1 staining, and C and D show positive control for CD45.1 staining and EGFP expression, respectively. A representative analysis of peripheral blood cells from a clonally engrafted mouse 2 months after transplantation shows that EGFP+ donor cells represented 93.58% of total nucleated cells (E). EGFP+ donor derived cells also represented 99.75% of granulocytes-macrophages (F), 98.49% of B-cells (G), and 62.43% of T-cells (H).
peripheral blood of this clonally engrafted mouse was 96.3%. Isotype controls are shown in Figure 2, A–C and a population of CAFs in both melanoma and LLC tumor models.23 mice used in these tracing studies were transplanted with un-manipulated bone marrow cells from EGFP+ mice. This allowed for optimization of the number of HSC-derived cells for identification in tumor microenvironment. Resulting tumors were harvested at day 13 of growth (~1 cm in diameter) and either digested for flow cytometric analysis or processed for paraffin sections. When digested tumor was examined for the presence of EGFP+ CFPs, the EGFP+CD45−DDR2+ was found to compose 4.39% of the total tumor population.

Paraffin sections were then immunostained and imaged for the presence of EGFP+ cells that expressed DDR2 (Figure 3, A–K). Analysis of high magnification images showed numerous EGFP+DDDR2+ cells within the tumor microenvironment (Figure 3, B, and G–J). Several EGFP+DDR2+ cells had an elongated, spindle-shape characteristic of fibroblasts (Figure 3, A, B, H, and J, arrowheads). EGFP+DDR2+ cells with a rounded morphology, a shape consistent with that of a cell either in or just extravasated from circulation were also seen in the tumor stroma (Figure 3, A–D, G, I–K, arrows). EGFP+DDR2+ cells with a rounded morphology were also found among red blood cells (asterisk) within the lumen of tumor vasculature (Figure 3, C, D, and K). To further characterize these HSC-derived cells, tumor sections were examined for the presence of EGFP+ cells that expressed α-smooth muscle actin (αSMA, a protein whose expression is associated with myofibroblasts and CAFs) within the tumor microenvironment (Figure 4, A–F). High magnification images show that a subpopulation of EGFP+ cells within the tumor microenvironment expressed αSMA (Figure 4, arrows). These findings are supported by our previous studies demonstrating that HSCs give rise to αSMA+, collagen-expressing CAFs and perivascular cells.23 Together, these data suggest that HSC-derived CFPs arrive at the tumor via the circulation where they transition from a rounded circulating phenotype to more mature elongated fibroblast phenotype within the tumor stroma.

Results

CFPs Are of HSC Origin

Our previous in vitro studies identified a population of fibroblast precursors from the bone marrow and peripheral blood that were of HSC origin.21 When cultured, these precursors showed a decrease CD45 expression and an increase in expression of collagen I (Col I) and Discoidin Domain Receptor 2 (DDR2; a collagen receptor), demonstrating that differentiation of precursors to fibroblasts is accompanied by reciprocal expression of HSC and fibroblast markers.21 Our recent studies examining differentiation of human peripheral blood-derived HSC and fibroblast markers.21 Our previous studies identified a population of fibroblast precursors from the bone marrow and peripheral blood that were of HSC origin.21 When cultured, these precursors showed a decrease CD45 expression and an increase in expression of collagen I (Col I) and Discoidin Domain Receptor 2 (DDR2; a collagen receptor), demonstrating that differentiation of precursors to fibroblasts is accompanied by reciprocal expression of HSC and fibroblast markers.21 Our present study, we first confirmed the HSC origin of fibroblast precursors in circulation using clonally engrafted mice. CFPs were defined as CD45+DDR2+ cells.21 Mice engrafted with clonal populations derived from a single EGFP+ HSC were generated as described previously.21,23,36,39 Briefly, single Lin−Sca-1+ckit+CD34− side population cells were deposited individually and cultured for one week in the presence of SCF and G-CSF. Clones consisting of 20 or fewer cells were transplanted into lethally irradiated recipients. Two months after transplantation, multilineage engraftment was confirmed by flow cytometry (Figure 1, A–H). Mice exhibiting high levels of multilineage engraftment from a single HSC were then examined for the presence of HSC-derived EGFP+ CFPs. Flow cytometric analysis of peripheral blood demonstrated that CFPs represented ~20% of the total population (Figure 2A). Further analysis showed that 99.18% of the CD45−DDR2+ cells were EGFP+ (Figure 2B), demonstrating that they were derived from the single-sorted EGFP+ HSC.

CFPs Are Found in Vivo in the Tumor Stroma

Our next studies showed that the HSC-derived CFPs observed in the peripheral blood incorporated into the LLC tumor microenvironment in vivo. For these studies, mice were engrafted with EGFP+ whole bone marrow and LLC cells were delivered subcutaneously. Because our studies using clonally engrafted mice established the HSC origin of both the CFP (Figure 2, A–C) and a population of CAFs in both melanoma and LLC tumor models.23 mice used in these tracing studies were transplanted with un-manipulated bone marrow cells from EGFP+ mice. This allowed for optimization of the number of HSC-derived cells for identification in tumor microenvironment. Resulting tumors were harvested at day 13 of growth (~1 cm in diameter) and either digested for flow cytometric analysis or processed for paraffin sections. When digested tumor was examined for the presence of EGFP+ CFPs, the EGFP+CD45−DDR2+ was found to compose 4.39% of the total tumor population.

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CFPs Increase with Tumor Development

We next examined the peripheral blood from normal and tumor bearing mice for the presence of CFPs. Normal mice were injected with LLC cells and tumors were allowed to progress to various time points (d3, d10, and d17). Flow cytometric analysis of peripheral blood from LLC-bearing mice showed that the number of circulating CD45\(^{-}/\)DDR2\(^{+}\) cells increases with tumor development (Figure 5). To correlate the increase in CFPs seen in the peripheral blood with CFP participation in the tumor microenvironment, we analyzed cells from digested LLC tumors. In tumors that had been allowed to progress to maximum size (1.5 cm, day 21 post delivery), 8.15 ± 2.35% (n = 4) of the total tumor population was made up of CD45\(^{-}/\)DDR2\(^{+}\) cells. This was almost double that seen at day 13 (4.39%, as described above). These findings demonstrate that the number of CFPs in circulation increase with tumor burden and confirm that CFPs incorporate into developing tumors.

CFPs Cultured in Presence of Tumor-Conditioned Media Give Rise to Fibroblasts

To demonstrate that CFPs were able to differentiate into fibroblasts in vitro in response to tumor, CD45\(^{-}/\)DDR2\(^{+}\) cells were sorted from the peripheral blood of normal mice and cultured in the presence of conditioned medium from Lewis lung carcinoma cells (LLC-CM) to mimic tumor microenvironment (Figure 6). At the end of the 7-day culture period, cells were fixed and imaged using DIC microscopy to visualize the morphology of cells. Morphological analysis showed that differentiation into...
fibroblasts, ie, elongated, spindle-shaped cells, is hastened by LLC-CM (Figure 6, C and D) relative to control media (Figure 6A) or media containing mouse serum (Figure 6B), which was shown to support fibroblast growth from stem/progenitor cells in 14 to 28 days. To demonstrate that cells with morphological characteristics of fibroblasts expressed proteins associated with fibroblasts, cultured cells were stained with antibodies to collagen I (Col I, a matrix molecule that is produced by activated fibroblasts), αSMA, and vimentin (an intermediate filament protein associated with fibroblasts) (Figure 7, A–F). These findings demonstrate that tumor-derived factors promote the differentiation of CD45⁺DDR2⁺ CFPs to fibroblasts.

**CFPs Preferentially Migrate in Response to Tumor-Conditioned Media**

To begin to determine the mechanism by which HSC-derived CFPs traffic to tumor, migration assays were conducted. Transwell assays were first conducted using freshly isolated nucleated peripheral blood cells from normal mice. Quantitative analysis showed that a population of cells preferentially migrated in response to LLC-CM (LLC-CM, 16.10 ± 6.49% cells; control, 4.67 ± 2.87% cells; P ≤ 0.05, n = 3 replicates, 4 mice per experiment). To determine whether CFPs were the cells responsible for migration seen in response to LLC-CM, CD45⁺DDR2⁺ CFPs were sorted from peripheral blood of normal mice (Figure 8, A and B) and analyzed in transwell assays. The percentage of migrating cells responding to LLC-CM (43.87 ± 13.3% cells; n = 11) was higher than the percentage of cells responding to control medium (26.6 ± 9.6% cells; n = 6), demonstrating that sorted CFPs preferentially migrate in response to LLC-CM (Figure 8C, P = 0.02). Control MLE-12 cells showed no preferential migration in response to LLC-CM (P = 0.39; n = 4).

**CFPs Are of the Monocyte Lineage**

Based on our data demonstrating that CFPs were able to respond to chemotactic signals, a characteristic associated with the monocyte, and data from other groups suggesting a close relationship between the fibroblast and monocyte lineages, we next sought to define the lineage origin of the HSC-derived CFP. For these studies, the CD45⁺DDR2⁺ population was analyzed for expression of markers associated with the monocyte lineage using multiparameter flow cytometry. These studies showed that 71.22% to 79.92% of CD45⁺DDR2⁺ cells expressed Mac1 (CD11b). Further analysis showed that...
this population was heterogeneous in its expression of Mac1 and F4/80, a marker that increases as monocytes mature to macrophages (Figure 9, A–D).

To test the ability of these subpopulations to give rise to fibroblasts in vitro, CD45^+DDR2^+ Mac1^hi^F4/80^+ (9.22% of CD45^+DDR2^+ cells), CD45^+DDR2^+ Mac1^lo^F4/80^− (38.39% of CD45^+DDR2^+ cells), CD45^+DDR2^+ Mac1^lo^F4/80^− (3.09% of CD45^+DDR2^+ cells), and CD45^+DDR2^+ Mac1^lo^F4/80^+ (0.26% of CD45^+DDR2^+ cells) cells were sorted (Figure 9B) and cultured in the presence of LLC-CM. The numbers of sorted cells in each subpopulation that gave rise to fibroblasts was calculated based on morphology daily over 14 days (Figure 9E). Mac1^hi^ CFPs had significantly higher fibroblast growth than Mac-1^lo^, with ~80% of the cells differentiating into fibroblasts within two weeks in LLC-CM. Morphological analysis at day 14 of culture showed that cells with high expression of Mac-1 had the greatest fibroblast growth per well and that expression of F4/80 was not a significant determinant of fibroblast potential (Figure 9, F–H). CD45^+DDR2^+ Mac1^lo^F4/80^− cells, which represented 0.26% of the CD45^+DDR2^+ population, showed no cells by the end of the culture period and were not used for subsequent quantitative or immunohistochemical studies. Staining with antibodies to Col I and α-SMA demonstrated that cells in each sorted population were able to express Col I to some degree, with cells from Mac^lo^ populations showing greatest Col I production. However, the numbers of cells with a mature phenotype, indicated by an elongated morphology and expression of Col I and α-SMA, were greater in the cultures of sorted CFPs based on high expression of Mac-1 (Figure 9, I–P).

To examine the potential of CD45^+DDR2^+ cells based on Mac1 expression at the level of the single cell, single cells from each population were sorted into individual wells of a 96-well culture plate and cultured in the presence of LLC-CM for 7 days. Calculation of the number of fibroblasts based on morphology showed that 84% of CD45^+DDR2^+ Mac1^hi^ cells (42/50 wells) were able to give rise to fibroblasts within 7 days. Cultured CD45^+DDR2^+ Mac1^lo^ cells resulted in few fibroblasts (8%; 4/50 wells). Together these studies suggest that the CD45^+DDR2^+Mac1^hi^ CFP population represents an immature circulating fibroblast precursor that is derived from the monocyte and can give rise to fibroblasts in the presence of tumor-CM.

To further confirm that CFPs were of monocyte origin, sorted CD45^+DDR2^+ cells were cultured for 11 days in the presence of exogenous M-CSF, a cytokine that is known to regulate the maturation of the monocyte lineage and whose increased levels has been associated with chronic diseases and cancer (reviewed in42). Based on morphology and immunostaining with antibodies to Col I and α-SMA, M-CSF was shown to promote differentiation of sorted CFPs when compared with 10%FBS/DMEM (control) or 10% FBS/10% mouse serum/DMEM (Figure 10, A–J). M-CSF-induced differentiation was even more robust than that seen when cells were cultured in
LLC-CM (Figure 10, compare panels E, F and G, H). ELISA analyses also showed that LLC-CM contained M-CSF (1 x 10^5 LLC cells produced 95.13 pg/ml M-CSF), suggesting that M-CSF may play a role in the differentiation of CFPs to mature carcinoma-associated fibroblasts.

**CFPs Express CCR2, Migrate in Response to Tumor-CM and Migration is MCP1-Dependent**

We next investigated the mechanism by which CFPs from the peripheral blood traffic to tumor. Given the shared lineage between CFPs and monocytes, we first examined LLC cell production of monocyte-associated chemokines including CXCL12 (stromal-derived factor-1), CCL21 (secondary lymphoid chemokine), and MCP1/CCL2 (monocyte chemoattractant protein 1; MCP1). ELISA for chemokine production by LLC cells showed that LLC-CM contains MCP1 in a cell concentration-dependent manner (1 x 10^5 LLC cells produced 2.21 ng/ml MCP1), but not stromal-derived factor-1α or secondary lymphoid chemokine.

Flow cytometric analysis using a MoFlo cell sorter for low incidence populations showed that ~0.5 to 1% (mean 0.74% ± 0.21%) of CFPs express CCR2, the MCP1 chemokine receptor (Figure 11, A, B). To confirm that the CD45^-DDR2^-CCR2^- cells sorted were not an artifact of flow cytometry, the CD45^-DDR2^-CCR2^- and CD45^-DDR2^-CCR2^+ populations were sorted (see gates, Figure 11C) and analyzed either by flow cytometry for purity or by qPCR. Flow cytometric analysis showed that 99.86% of the sorted CD45^-DDR2^-CCR2^- cells were negative for CCR2 expression (Figure 11D). Analysis by qPCR showed that the CCR2 message was significantly higher in the...
CD45^+ DDR2^+ CCR2^+ population (mean 1.58 ± 0.18) as compared with the CD45^+ DDR2^+ CCR2^- population (0.88 ± 0.13; P = 0.03). To determine whether CCR2-expressing CFPs could migrate in response to LLC-CM, peripheral blood CD45^+ DDR2^+ CCR2^- cells were placed into transwells. Results showed that CD45^+ DDR2^+ CCR2^- cells migrated in response to LLC-CM (85% cells) as compared with control media (30% cells). Sorted CD45^+ DDR2^+ CCR2^- cells were also shown to give rise to fibroblasts, based on morphology, in response to LLC-CM (data not shown).

Figure 10. M-CSF promotes CFP differentiation. Sorted CFPs (CD45^+ DDR2^+) cultured in the presence of 10 μg/ml exogenous M-CSF (G, H) gave rise to fibroblasts earlier than those cultured in control media (A, B), media with mouse serum (C, D) or LLC-CM (E, F). Analysis of differentiation was based on morphology (DIC images, A, C, E, G) and expression of Col 1 (green) and αSMA (red). Panels B, D, F, H, and J show superimposition of images taken in the green (Col I), red (αSMA) and blue (Hoechst) channels. Yellow indicates expression of both Col1 and αSMA. In some cases, superimposition of αSMA expression and Hoechst images resulted in nuclei that appeared pink. Panels I and J show DIC and fluorescent images of tissue stained only with secondary antibodies and Hoechst (blue). Scale bar = 25 μm.

Figure 11. CFP migration to tumor is dependent on MCP1. A representative image from flow cytometric analysis of peripheral blood showing that the percentage of circulating CD45^+ DDR2^+ cells (R1, A) expressing CCR2 (R2, B) is 0.83% (R3, C). Panel C also shows the gates used to isolate CD45^+ DDR2^+ CCR2^- (R3) and CD45^+ DDR2^+ CCR2^- (R4) populations for flow cytometry or qPCR. Reanalysis of CCR2 expression in the CD45^+ DDR2^+ CCR2^- cells showed that this population was 99.86% pure (D). Isotype controls are shown in E and F. Sorted CD45^+ DDR2^+ CFPs (gray bars) or MLE-12 control cells (black bars) were placed into transwell assays in the presence of control media (DMEM 10% FBS), or tumor-CM (ie, LLC-CM, EMT6-CM, or K7M2-CM) with or without the addition of neutralizing anti-MCP1 antibody (G). Increased migration of sorted CFPs was seen in response to LLC-CM (* P = 0.02), EMT6-CM († P = 0.00003), and K7M2-CM (‡ P = 0.02) as compared with control media. Addition of neutralizing anti-MCP1 antibody to tumor-CM significantly reduced migration as compared with tumor-CM-induced migration of CFPs (LLC-CM/anti-MCP1, § P = 0.03; EMT6-CM/anti-MCP1, ¶ P = 0.02; K7M2-CM/anti-MCP1, ‡‡ P = 0.04). CFP migration in the presence of tumor-CM and neutralizing anti-MCP1 antibody was at (‡‡ for LLC; ‡‡ for K7M2) or below (†† for EMT6) control. No significant migration was seen with MLE-12 cells under any condition (P ≥ 0.05).
The effects of inhibiting MCP1 produced by tumor on CFP migration were then examined in vitro. CFPs were placed into transwell culture plates with control media, tumor-CM, or tumor-CM with the addition of MCP1 neutralizing antibody. Tumor-CM was harvested from the LLC, EMT-6 (murine mammary carcinoma) and K7M2 (murine osteosarcoma) tumor cell lines, all of which have been shown to produce MCP1 (data above and references 43, 44). Analysis showed that exposure to conditioned media from each tumor line resulted in preferential migration of CFPs when compared with response of CFPs to control media (Figure 11G, \( P \leq 0.05 \)). The addition of MCP1 neutralizing antibody to LLC-CM, EMT-6-CM, or K7M2-CM resulted in significant decrease in migration of sorted CFPs, when compared with tumor-CM alone (Figure 11G, \( P \leq 0.05 \)). Inhibition of MCP1 reduced migration of CFPs to percentages at (Figure 11G, \( P \geq 0.05 \)) or below (Figure 11G, \( P \leq 0.05 \)) control levels. Together, these findings suggest that tumor production of MCP1 directs migration of HSC-derived CFPs in multiple tumor types.

**Discussion**

Multiple origins for CAFs have been suggested, including resident tissue fibroblasts, epithelial-to-mesenchymal transdifferentiation, and the bone marrow. Our previous studies, based on single HSC transplantation, demonstrated that HSCs give rise to a population of collagen-producing CAFs and perivascular cells in tumor stroma. In the present study, we examined the mechanisms by which these HSC-derived cells contribute to the tumor stroma. Using transplantation of a clonal population of cells derived from a single HSC, we establish that CFPs are of HSC origin. We found these HSC-derived cells both incorporated into the tumor stroma and circulating in the tumor vasculature in vivo, indicating that CFPs contribute to tumor via the circulation. We also showed that a population of these HSC-derived cells expressed αSMA, a marker of CAFs. In addition, the CFP population increased in the peripheral blood and tumor stroma with tumor burden. In vitro studies confirmed that sorted CFPs were able to give rise to mature fibroblasts/myofibroblasts that expressed collagen, vimentin, and αSMA in response to tumor conditioned media. These findings are supported by our previous works demonstrating that differentiation of both mouse and human fibroblast precursors is accompanied by an increase in fibroblastic markers and a concomitant decrease in hematopoietic markers. Together, these findings demonstrate that HSC-derived CFPs are, at least, a contributing source of CAFs during tumor development.

We also sought to determine the lineage origin of HSC-derived CFPs. Based on their HSC origin, ability to preferentially migrate and previous studies suggesting similarities between the fibroblast precursor, fibroblasts, and the monocyte, it was our hypothesis that the CFP represents a transitional phenotype from the monocytic lineage to the fibroblastic lineage with the ability to respond to and participate in solid tumor stromal formation. Our data show that the CFP represents a subpopulation of
circulating monocytes that resides in the Mac1\textsuperscript{+} subset of peripheral blood cells. These findings also show that while 80% to 84% of sorted Mac1\textsuperscript{+} CFPs are able to give rise to fibroblasts in response to tumor. Mac1\textsuperscript{+} CFPs did not possess this potential to the same extent, suggesting that the CFP is derived from the monocyte rather than the monocyte precursor. The CFP was also shown to differentiate in the presence of M-CSF, a cytokine known to regulate the monocyte lineage, whose increased presence is associated with several chronic diseases and cancer (reviewed in\textsuperscript{42}). Our finding of a monocyte lineage origin for HSC-derived CFPs is in agreement with others demonstrating that fibrocytes, which are a mesenchymal cell type in peripheral blood thought to be a source of fibroblasts,\textsuperscript{45} may differentiate from monocytes\textsuperscript{26,40,41} (reviewed in\textsuperscript{46,47}).

Based on this demonstrated relationship between CFPs and monocytes and their ability to preferentially migrate in response to tumor, we examined the potential role of chemokines in the recruitment/homing of HSC-derived CFPs to tumor. A subset of CFPs was shown to express functional CCR2, the chemokine receptor for MCP1, and inhibition of MCP1 produced by multiple tumor types (ie, LLC, EMT6, and K7M2) was shown to reduce CFP migration. We also found that the number of CFPs expressing CCR2 increased in the peripheral blood as tumor size increased. Based on tumor size, this period of active CFP recruitment and homing corresponded to the period of tumor growth in which the stromal component, including the vasculature was established. Together with our previous studies demonstrating an HSC origin for CAFs, myofibroblasts and perivascular cells,\textsuperscript{23} the present studies suggest that at least a portion of HSC-derived CFPs use the CCR2 chemokine receptor to home to tumor microenvironment to contribute to these populations. Finally, we demonstrate that disruption of normal MCP1/CCR2 signaling results in reduced tumor burden. While it is likely that systemic inhibition of MCP1 affected both the CFP and monocyte/macrophage populations, given our findings of shared lineage origin between CFPs and monocytes/macrophages, these findings suggest the therapeutic potential of targeting the MCP1/CCR2 axis during tumor development to alter tumor growth rate and/or limit tumor progression and metastasis.

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