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During tumor progression, immune system phagocytes continually clear apoptotic cancer cells in a process known as efferocytosis. However, the impact of efferocytosis in metastatic tumor growth is unknown. In this study, we observed that macrophage-driven efferocytosis of prostate cancer cells in vitro induced the expression of proinflammatory cytokines such as CXCL5 by activating Stat3 and NF-κB(p65) signaling. Administration of a dimerizer ligand (AP20187) triggered apoptosis in 2 in vivo syngeneic models of bone tumor growth in which apoptosis-inducible prostate cancer cells were either coimplanted with vertebral bodies, or inoculated in the tibiae of immunocompetent mice. Induction of 2 pulses of apoptosis correlated with increased infiltration of inflammatory cells and accelerated tumor growth in the bone. Apoptosis-induced tumors displayed elevated expression of the proinflammatory cytokine CXCL5. Likewise, CXCL5-deficient mice had reduced tumor progression. Peripheral blood monocytes isolated from patients with bone metastasis of prostate cancer were more efferocytic compared with normal controls, and CXCL5 serum levels were higher in metastatic prostate cancer patients relative to patients with localized prostate cancer or controls. Altogether, these findings suggest that the myeloid phagocytic clearance of apoptotic cancer cells accelerates CXCL5-mediated inflammation and tumor growth in bone, pointing to CXCL5 as a potential target for cancer therapeutics.

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Apoptosis-induced CXCL5 accelerates inflammation and growth of prostate tumor metastases in bone

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During tumor progression, immune system phagocytes continually clear apoptotic cancer cells in a process known as efferocytosis. However, the impact of efferocytosis in metastatic tumor growth is unknown. In this study, we observed that macrophage-driven efferocytosis of prostate cancer cells in vitro induced the expression of proinflammatory cytokines such as CXCL5 by activating Stat3 and NF-κB(p65) signaling. Administration of a dimerizer ligand (AP20187) triggered apoptosis in 2 in vivo syngeneic models of bone tumor growth in which apoptosis-inducible prostate cancer cells were either implanted with vertebral bodies, or inoculated in the tibiae of immunocompetent mice. Induction of 2 pulses of apoptosis correlated with increased infiltration of inflammatory cells and accelerated tumor growth in the bone. Apoptosis-induced tumors displayed elevated expression of the proinflammatory cytokine CXCL5. Likewise, CXCL5-deficient mice had reduced tumor progression. Peripheral blood monocytes isolated from patients with bone metastasis of prostate cancer were more efferocytic compared with normal controls, and CXCL5 serum levels were higher in metastatic prostate cancer patients relative to patients with localized prostate cancer or controls. Altogether, these findings suggest that the myeloid phagocytic clearance of apoptotic cancer cells accelerates CXCL5-mediated inflammation and tumor growth in bone, pointing to CXCL5 as a potential target for cancer therapeutics.

Introduction

The most common site of prostate cancer metastasis is bone, with an incidence of 65%–80% in patients with advanced disease (1). Once cancer cells spread to bone, they significantly alter normal bone remodeling, resulting in bone fractures, nerve compression, pain, and hypercalcemia (2). In bone, cancer cells find a protective and supportive microenvironment that promotes metastatic outgrowth. Prevention of bone metastasis is a major goal of treatment, and elucidation of the crucial factors contributing to the development of skeletal metastasis is key. Some mechanisms have been proposed, and monocytic myeloid cell populations attracted to the tumor microenvironment have been identified as critical mediators of inflammation (3, 4). One study found that CD11b+ myeloid cells expressing integrin αβ2 were mobilized to the skeletal tumor microenvironment via interaction with VCAM-1 expressed in breast tumor cells (5). Similarly, another breast cancer study showed that macrophages expressing αRC2 promote cancer cell survival and lung metastasis (6), suggesting that this interaction may be essential for both cancer colonization and osseous progression.

The bone microenvironment is rich in factors that promote the mobilization, proliferation, and differentiation of proinflammatory cells, which interact with disseminated tumor cells, promoting survival and colonization. This occurs through multiple inflammation-mediated molecular mechanisms that originate feed-forward amplification loops between tumor cells, inflammatory cells, osteoblasts, osteoclasts, and bone marrow stromal cells to perpetuate a chronic inflammatory state (7).

A successful antiinflammatory therapy that targets tumor metastasis has been associated with reduced infiltration of bone marrow–derived myeloid cells and prometastatic macrophages (8). Recently a crucial role of bone macrophages in the progression of prostate cancer skeletal metastasis was demonstrated in vivo using models of conditional macrophage depletion (9). Furthermore, special attention has been dedicated to macrophage-directed cancer immunotherapy based on the transformation of macrophages from M2 to M1 type to elicit antitumor responses (10). These approaches have the potential to inhibit tumor progression and metastasis.

Persistent inflammation also exacerbates cell stress and tissue damage, causing apoptotic/necrotic cell death. The clearance of dying cells occurs mainly through phagocytosis by macro-
Proinflammatory cytokines are induced in macrophages upon apoptotic cancer cell efferocytosis. Immunofluorescence and flow cytometric studies show that bone marrow–derived macrophages (MΦs) effectively efferocytose apoptotic cells (12, 14). We hypothesized that macrophages discriminate between different types of apoptotic cells and orchestrate a distinctive response accordingly. To investigate the production of proinflammatory cytokines, cocultures of macrophages and different highly apoptotic (HA) cells were analyzed (Supplemental Figure 1A; supplemental material available online with this article; https://doi.org/10.1172/JCI92466DS1). Two prostate cancer cell types (murine RM1 and human PC3) and 2 noncancer cell types (murine osteoblastic MC4 and murine primary bone marrow stromal cells [BMSCs]) were used. RM1 cells, derived from C57BL/6J mice, represent a valuable phages and other cells (11). Phagocytosis of apoptotic cancer cells (termed efferocytosis) is a process often underestimated during tumor progression due to rapid clearance by phagocytes (11). Tumor cell death, significantly amplified by chemotherapies or other targeted therapies, triggers extensive efferocytosis, which has been suggested to accelerate further tumor growth at least in part by inducing M2-like macrophage polarization and resultant protumoral functions (12, 13).

Using in vivo models of skeletal tumor growth and apoptosis-inducible prostate cancer cells, the role of apoptotic cancer cell clearance in skeletal tumor progression was investigated. A critical role of efferocytosis-induced inflammation mediated by macrophage-derived CXCL5 was discovered as a novel mechanism underlying skeletal metastasis and a viable target for cancer therapeutics.

**Results**

Proinflammatory cytokines are induced in macrophages upon apoptotic cancer cell efferocytosis. Immunofluorescence and flow cytometric studies show that bone marrow–derived macrophages (MΦs) effectively efferocytose apoptotic cells (12, 14). We hypothesized that macrophages discriminate between different types of apoptotic cells and orchestrate a distinctive response accordingly. To investigate the production of proinflammatory cytokines, cocultures of macrophages and different highly apoptotic (HA) cells were analyzed (Supplemental Figure 1A; supplemental material available online with this article; https://doi.org/10.1172/JCI92466DS1). Two prostate cancer cell types (murine RM1 and human PC3) and 2 noncancer cell types (murine osteoblastic MC4 and murine primary bone marrow stromal cells [BMSCs]) were used. RM1 cells, derived from C57BL/6J mice, represent a valuable...
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ed in macrophages interacting with apoptotic MC4 cells (Supplemental Figure 1B). Other cytokines, like CCL2 and MIP-1α, were increased in cocultures with PC3(HA) but not in cocultures with RM1(HA) cells, and a small increase in the proinflammatory cytokines sTNFRI and sTNFRII (soluble TNF receptors I and II) was observed for both RM1 and PC3 (Supplemental Figure 1B). Apoptotic cells alone showed reduced levels of cytokines compared with macrophages alone (Supplemental Figure 1C). Intriguingly, the proinflammatory factors CCL5, CXCL1, CXCL5, and IL-6 have been shown to have tumor-promoting functions in different contexts (18, 19). To determine the transcriptional regulation in macrophages, quantitative PCR mRNA expression analyses were performed for CXCL1, CXCL5, and IL-6 to compare macrophage responses to the different apoptotic cells. Figure 1C indicates that these cytokines were transcriptionally upregulated in macrophages interacting with apoptotic cancer cells in contrast with noncancer cells, which correlates with the cytokine array results. Since pros-

model for investigations of prostate cancer interactions with bone marrow stroma as they promote osteolytic lesions accompanied by perosteal bone deposition in immunocompetent C57BL/6J mice (9, 15). Human PC3 cancer cells, originally isolated from prostate cancer bone metastasis (16), have been used extensively for their ability to metastasize to the bone when injected in immunocompromised mice via intracardiac inoculation (17). Secreted proteins from cocultures of macrophages and apoptotic cells were analyzed using inflammation arrays (Figure 1A), and results were quantified and normalized relative to macrophage-alone controls (Figure 1B). The cytokines expressed upon efferocytosis of apoptotic cancer cells differed from noncancer cells, suggesting that different pathways were activated in macrophages. Cytokine activation of CCL5, CXCL1, CXCL5, IL-6, and IL-12(p40/p70) was observed in macrophages cocultured with apoptotic cancer cells, but not apoptotic noncancer cells. Upon noncancer apoptotic cell engulfment, CCL2 was the only cytokine significantly upregulat-

Figure 2. Efferocytosis of apoptosis-inducible prostate cancer RM1-iC9 cells. (A) Apoptosis-inducible RM1-iC9 cells were treated with VEH or AP and analyzed by Western blot for the activation of inducible caspase-9 (IC9) and apoptosis-associated caspase-3. AP treatment increased processed caspase-9 and cleaved caspase-9, reflecting apoptosis (see complete unedited blots in the supplemental material). (B) Bone marrow MΦs were cocultured with RM1-iC9 cells stained with Cell Trace–CFSE (Invitrogen), and treated with VEH or AP for 18 hours. The cocultures were stained with F4/80-APC antibody and analyzed by flow cytometry. Representative FACS images from ImageStream showing MΦs alone (APC+, red gate), RM1-iC9 alone (CFSE+, green gate), and 3 different examples of MΦs engulfing RM1-iC9 (APC+CFSE+, yellow gate). (C) Bar graphs indicating the percentage of efferocytic MΦs (APC+CFSE+ cells), MΦs with high internalization of RM1-iC9, and engulfed RM1-iC9 (APC+CFSE+) relative to total cancer cells (CFSE+ plus APC+CFSE+) after treatment with VEH or AP. (D) mRNAs isolated from APC+CFSE+ gated cells (efferocytic MΦs) of MΦ+RM1-iC9(AP) and APC+ gated cells (nonefferocytic MΦs) of MΦ+RM1-iC9 (VEH) samples were analyzed by qPCR for selected genes. Fold changes were expressed relative to the nonefferocytic MΦ values (VEH). Data in C and D are mean ± SEM, n = 3 per group; **P < 0.01, *P < 0.001 (2-tailed Student’s t test).
The cytokine array results and aligned with the mRNA analyses. Altogether, these results suggest that the clearance of apoptotic cells by macrophages induced a selective response dependent on the cell type engulfed and a common inflammatory expression pattern in macrophages efferocytosing apoptotic cancer cells.

To investigate the transcription factor activity in macrophages in response to the apoptotic cells, we used TRACER (transcriptional activity cell array) technology (Figure 1F and ref. 20). The activity of 13 transcription factors was investigated in cocultures with apoptotic RM1 or MC4, 2 cell lines that induced a different response in macrophages. Macrophages were transduced with a reporter luciferase construct containing the DNA binding site for each transcription factor or a control vector and the luciferase cancer cells are of epithelial origin, we investigated whether the inflammatory responses induced in macrophages were a result of their interaction with apoptotic epithelial cells. Similarly to the experiments above (Supplemental Figure 1A), we used apoptotic mouse primary prostate epithelial cells (MPECs), apoptotic total bone marrow cells (BMCs) (isolated from mouse tibiae), or apoptotic RM1 cells (all originated from C57BL/6J mice) in cocultures with bone marrow macrophages to analyze changes in gene expression of the different inflammatory cytokines (CCL5, CXCL1, CXCL5, and IL-6). Only apoptotic RM1 cancer cells were capable of inducing an inflammatory response in bone marrow macrophages (Figure 1D). ELISA evaluation for CXCL1 and CXCL5 proteins in the coculture media for apoptotic cancer cells (Figure 1E) confirmed the cytokine array results and aligned with the mRNA analyses.

Figure 3. Efferocytosis-induced activation of Stat3 and NF-κB signaling mediates inflammatory response in macrophages. (A) MΦs were cocultured 20 hours alone or with apoptotic RM1(HA) cells. Protein from cocultures was analyzed by Western blot for activation of NF-κB and Stat3 signaling using consecutively the following antibodies (panels from top): phospho–NF-κB(p65) [p–NF-κB(p65)], p-Stat3, and p100/p52. Bottom panel shows total protein for each sample. (B) Protein bands in the blots were quantified and normalized to total protein for each sample. Graphs depict the fold changes for p-p65, p-Stat3, p52/p100 ratio, and p52 relative to MΦ signal, respectively. (C) Statistic, a selective inhibitor of Stat3 activation, was incubated with MΦs (12.5 μM) for 1 hour, then removed before coculture with RM1(HA) for 5 hours. Protein was analyzed by Western blot using the p-Stat3 antibody. (D) Bay11-7082, an inhibitor of NF-κB signaling, was preincubated with MΦs (20 μM) for 1 hour before coculture with RM1(HA) for 5 hours. Protein was analyzed by Western blot using the p-p65 antibody. Graphs in C and D depict the quantification of p-Stat3 and p-p65 signals normalized to total protein (lower panel) relative to the average MΦ control (VEH), respectively. See complete unedited blots in the supplemental material. (E) mRNAs were isolated from cocultures at 5 and 20 hours of incubation as described in the experiments in C and D and analyzed by qPCR for indicated genes. Graphs show the fold change relative to MΦ control for each group. Data in B are mean ± SEM, n = 4 per group (2-tailed Student’s t test), or n = 3 per group in C–E (1-way ANOVA); **P < 0.01, *P < 0.001, †P < 0.0001.
cells, RM1-iC9, from murine RM1 cells using the viral construct for inducible caspase-9 (iC9) (23). The induction of apoptosis and resultant caspase-3 activation were validated by treatment with the dimerizer drug AP20187 (AP) or control vehicle (VEH) followed by Western blot analysis of cell extracts (Figure 2A). Formation of processed caspase-9 and corresponding cleaved caspase-3 confirmed apoptosis activation in AP-treated cells. To verify that the inducible apoptotic RM1-iC9 cells were able to be efferocytosed, cells were prelabeled with CFSE dye, cocultured with macrophages, and treated with VEH or AP. After 16–18 hours the cells were collected, labeled with F4/80-APC antibody, and analyzed using the ImageStream flow cytometer (Abcam), which provides microscopic event images. Double-positive APC+CFSE+ cells indicate efferocytic macrophages (macrophages engulfing apoptotic RM1-iC9 cells) as depicted in Figure 2B (yellow gate).

Efferocytosis induces an inflammatory response via activation of Stat3 and NF-kB signaling. To better understand the role of efferocytosis, we generated apoptosis-inducible prostate cancer cells, RM1-iC9, from murine RM1 cells using the viral construct for inducible caspase-9 (iC9) (23). The induction of apoptosis and resultant caspase-3 activation were validated by treatment with the dimerizer drug AP20187 (AP) or control vehicle (VEH) followed by Western blot analysis of cell extracts (Figure 2A). Formation of processed caspase-9 and corresponding cleaved caspase-3 confirmed apoptosis activation in AP-treated cells. To verify that the inducible apoptotic RM1-iC9 cells were able to be efferocytosed, cells were prelabeled with CFSE dye, cocultured with macrophages, and treated with VEH or AP. After 16–18 hours the cells were collected, labeled with F4/80-APC antibody, and analyzed using the ImageStream flow cytometer (Abcam), which provides microscopic event images. Double-positive APC+CFSE+ cells indicate efferocytic macrophages (macrophages engulfing apoptotic RM1-iC9 cells) as depicted in Figure 2B (yellow gate). The APC+CFSE+ gate exhibited images with green apoptotic cancer cells engulfed by red F4/80+ macrophages (Figure 2B) with...
high internalization, indicating efferocytosis. In the APC–CFSE− gate, efferocytosis was observed at different stages of digestion correlating to the position of the cell in the plot. The brightest cells in the CFSE axis showed less digested cancer cells inside macrophages (Figure 2B). As expected, the percentage of cells (gated from single cells in focus) with high internalization was strikingly higher in the samples treated with AP relative to VEH (Figure 2C). Furthermore, the percentages of highly internalized cells were similar to the percentages of cells gated as APC–CFSE+, validating that this gate demonstrates that efferocytosis increased with the induction of apoptosis in cancer cells. When apoptosis was induced with AP, the proportion of engulfed RM1-iC9 cells relative to total cancer cells was approximately 30%, compared with less than 5% when the cells were treated with VEH (Figure 2C). Macrophages from the APC–CFSE− gate, which had been cocultured with apoptosis-induced RM1-iC9 (AP-treated) cells, were sorted by flow cytometry and compared with APC−-gated macrophages sorted from VEH samples. Quantitative PCR analyses revealed higher (5- to 10-fold) mRNA expression of the proinflammatory cytokines CXCL5, CCL5, and IL-6 in efferocytic macrophages (AP–CFSE−-sorted) as compared with nonefferocytic (AP−-sorted) macrophages (Figure 2D).

Simultaneous activation of Stat3 and NF-κB signaling in the tumor microenvironment has been identified as a critical element of inflammation-associated tumor progression (24, 25), and here, NF-κB was identified as a critical transcription factor induced in efferocytic macrophages (Figure 1F). Thus the increase in phosphorylation/activation of NF-κB(p65) and Stat3 occurring in macrophages via efferocytosis of apoptotic RM1 cells was further investigated. Protein lysates isolated from cocultures of macrophages and RM1(HA) cells were analyzed by Western blot and quantified. Figure 3A depicts consecutive blotting of the membrane with antibodies for phosphorylated (p) p65, p-Stat3, and NF-κB2(p100/ p52), respectively. Quantification of Western bands normalized to total protein in each lane (bottom panel in Figure 3A) showed activation (phosphorylation) of both canonical NF-κB(p65) (p-p65) and Stat3 (p-Stat3) signaling in efferocytic macrophages (Figure 3B). No significant changes in the processing of NF-κB2(p100) precursor to produce p52 (noncanonical NF-κB signaling) were induced by efferocytosis (Figure 3B), which suggests the predom-
inant role of canonical signaling in the efferocytic inflammatory response. The apoptotic RM1-only lysates demonstrated insignifi-
cant signals for p-Stat3 or p-p65, which is consistent with the apop-
totic stage of these cells.

To evaluate the dependence of Stat3 and NF-κB in inflamma-
tory cytokine production, cocultures of macrophages and apop-
totic RM1 cells were treated with the small-molecule inhibitors
Stattic (Stat3 phosphorylation inhibitor; ref. 26) and Bay11-7082
(IκBα phosphorylation inhibitor; refs. 27, 28). Figure 3, C and D,
demonstrates the inhibition of p-Stat3 and p-p65 after 5 hours of
coculture pretreated with Stattic or Bay11-7082, respectively. Nei-
ther Stattic nor Bay11-7082 inhibited efferocytosis (Supplemen-
tal Figure 2A); however, both inhibitors significantly mitigated
proinflammatory cytokine expression in macrophages at 5 and 20
hours of coculture (Figure 3E).

Treatment with emetine, an IκBα phosphorylation inhibi-
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hours of coculture (Figure 3E).

Treatment with emetine, an IκBα phosphorylation inhibi-
tor (29), reduced efferocytosis in cocultures of macrophages with apoptosis-induced RM1-iC9 (Supplemental Figure 2B).
This correlates with Supplemental Figure 2, C and D, which shows that emetine blocked the AP-induced increase of both
p-NF-kB(p65) and p-Stat3 in the F4/80+ macrophage popu-
lation. Furthermore, Ly6B (Ly6-B.2 or antigen 7/4), a bone
marrow–derived inflammatory macrophage antigen (30),
was increased when efferocytosis was activated with the AP
dimerizer (Supplemental Figure 2E), while inhibition of effe-
rocytosis with emetine reduced Ly6B, which is consistent with
the inflammatory response of macrophages upon efferocyto-
sis of apoptotic cancer cells. However, emetine inhibited not
only p-NF-kB(p65) but also p-Stat3 and efferocytosis, differ-
ning from the Bay11-7082 inhibitor results. Therefore, the com-
bined effect of Stattic and Bay11-7082 on efferocytosis was
determined. Pretreatment of macrophages (1 hour) with the
combination of these inhibitors (same concentrations used in
Supplemental Figure 2A) resulted in a complete abrogation of
efferocytosis (Supplemental Figure 2F). These findings sug-
gest that the effect of emetine on efferocytosis may be due do a
simultaneous blocking of Stat3 and NF-κB activation.

Altogether these findings correlate efferocytosis with per-
sistent inflammation within the tumor microenvironment via acti-
vation of Stat3 and NF-κB signaling in macrophages.
cell numbers in the AP-treated tumors relative to VEH, while no significant changes were observed in mitotic cell numbers between groups (Figure 4E). Considering that mice were sacrificed 2 days after the last AP treatment, it was expected that a major proportion of apoptotic cells would be cleared. Nevertheless, this result validates the effectiveness of AP treatment for the induction of cancer cell apoptosis in vivo.

As described above (Figure 1), in vitro results showed that CXCL5 was highly induced in macrophages upon efferocytosis of cancer cells. CXCL5 expression has been shown to correlate with prostate cancer progression and metastasis and is associated with stromal inflammation (32). To investigate whether expression of proinflammatory CXCL5 correlates with the induction of apoptosis in AP-treated mice, total protein from tumor vossicles was collected and analyzed by ELISA. As observed in Figure 4F, a significant increase in CXCL5 was detected in the AP-treated group, aligning with induction of apoptosis and consequent efferocytosis. A significant correlation between tumor weight and CXCL5 concentrations was found for both VEH- and AP-treated tumor vossicle groups (Figure 4G).

The immune cell composition of tumor vossicles was analyzed via flow cytometry. The AP-treated group (induced efferocytosis) showed significantly increased F4/80+CD206+ myeloid cells (Figure 5A), which characterizes M2-like macrophages (33). Other myeloid cells also showed increased infiltration in the AP-treated tumor vossicles, including: CD11b+Gr-1+ (monocytic/granulocytic cells associated with tumor progression and antitumor immunity) (34, 35) (Figure 5B), total CD206+ and Gr-1+ (characterizes inflammatory monocytes/granulocytes; ref. 36) (Figure 5, C and D), and CD68+ within the CD11b+ population (marker of phagocytic myeloid cells; aligns with higher efferocytosis induced by apoptosis) (Figure 5E). Other flow results are shown in Supplemental Figure 3A. Furthermore, whole blood count of VEH- and AP-treated mice revealed no significant differences (Supplemental Figure 3B), suggesting that the results observed in the tumor microenvironment are local and not related to general changes in the circulating monocytes/neutrophils.

CXCL5, a crucial inflammatory-microenvironment cytokine that accelerates tumor progression. As CXCL5 was found to increase with cancer cell death and efferocytosis in the tumor...
microenvironment, the specific role of CXCL5 was addressed using CXCL5−/− mice in the vossicle implant model. To avoid any possible compensation with CXCL5 produced by immune cells, vossicles from CXCL5−/− mice were inoculated with the RM1-iC9 cells and implanted in CXCL5−/− mice. Similarly, WT (CXCL5+/+) vossicles were implanted in WT mice (Figure 6A). To induce apoptosis and efferocytosis and hence increase the expression of CXCL5 (in accordance with results in Figure 4), both groups were injected with AP at days 7 and 11 as described. Figure 6A and Supplemental Figure 4A demonstrate tumor growth deceleration in the CXCL5−/− mice compared with WT, and CXCL5−/− mouse tumors had significantly reduced weight at study end (Figure 6B).

Proinflammatory cytokine profiling in tumors from WT and CXCL5−/− mice revealed that CXCL5 was the only cytokine that was significantly different with distinctively lower levels in the CXCL5−/− mice (Figure 6C). These results were confirmed by ELISA (Figure 6D) and demonstrate that the major contribution of CXCL5 in the tumor microenvironment originated from the host and not from the tumor. However, since the cytokine array has limited detection, a more sensitive ELISA revealed a significant decrease in CXCL1 levels (Figure 6E). This may be explained by reduced inflammation in CXCL5−/− mice as a consequence of strikingly lower CXCL5 levels. These findings suggest a crucial role of host-derived CXCL5 in prostate cancer tumor progression in the bone microenvironment.

Apoptotic cells were counted inside the non-necrotic tumor sections, and no differences were found (Supplemental Figure 4B), which confirmed that AP treatment induced apoptosis in WT and CXCL5−/− vossicles similarly. There were no differences in mitotic cell numbers even though tumor growth was reduced. This could be due to a temporal effect, since the analysis of the tumors was performed at the experimental endpoint (2 days after AP treatment), which does not account for earlier differential effects in tumor growth.
Fluorescence IHC of sections inside the tumor vossicles was performed. While areas inside the tumor vossicles were highly infiltrated by inflammatory F4/80+ macrophages (red) and Ly6B+ cells (green) (Figure 7A), quantitative analysis revealed that CXCL5−/− mice exhibited fewer F4/80+ macrophages and Ly6B+ inflammatory cells (Figure 7B). Figure 7C depicts the engulfing morphology observed in some macrophages when interacting with large nuclei, typical of tumor cells in comparison with the smaller nuclei of macrophages (37).

Cell analysis of tumor vossicles via flow cytometry revealed significantly reduced F4/80+ macrophages, F4/80+CD206+ M2-like macrophages, and CD11b+Gr-1+ myeloid cells in CXCL5−/− mice (Figure 8, A–C), as well as reduced CD11b+Ly6B+ and total Ly6B+ inflammatory cells (Figure 8, D and E). These results align with the IHC findings (Figure 7) and the in vitro data (Supplemental Figure 2E), in which Ly6B was increased via efferocytosis and in correlation with inflammation. Furthermore, a decrease in total CD68+ phagocytic macrophages (the majority also F4/80+) was observed (Figure 8F). In contrast, an increase in cells expressing the T lymphocyte-activating antigen CD86+ was detected in the CXCL5−/− tumor vossicles (Figure 8G). Additional flow results are shown in Supplemental Figure 4C. Interestingly, the blood count of mice bearing tumor vossicles revealed significantly lower concentrations of leukocytes, lymphocytes, and monocytes in WT versus CXCL5−/− mice, where CXCL5−/− mice demonstrated values within the normal range (Supplemental Figure 4D). These findings suggest that increased recruitment of inflammatory cells into the tumors of WT mice may be responsible for a reduction of these cells in circulation, an effect that was not observed in WT mice or alleviated in CXCL5−/− mice.

For more in-depth analyses, flow cytometric results were compared between tumor vossicles and bone marrow myeloid cells from tumor-free WT and CXCL5−/− mice. In the bone marrow of the CXCL5−/− mice the percentages of Gr-1, Ly6B, Ly6C, Ly6G, and CD11b myeloid cells were higher than in WT mice.
inducer of inflammation-accelerating tumor growth in the bone microenvironment.

Figure 10. Cancer cell death induces accelerated tumor growth and bone osteolysis in the intratibial tumor model. (A) Experimental schematic. Mice injected with RM1-iC9 cells were randomized before VEH or AP treatment. (B) Representative μCT images showing trabecular bone in VEH and AP cancer-inoculated tibiae and the corresponding contralateral controls (nontumor) (scale bars: 400 μm). (C) Bone parameters quantified by μCT: trabecular bone volume relative to total volume (Tb.BV/TV), trabecular number (Tb.N), trabecular thickness (Tb.Th), trabecular spacing (Tb.Sp), cortical thickness (Cortical Th), and total bone volume relative to total volume (BV/TV); VEH (n = 11) and AP (n = 12). (D) Representative images of H&E sections of tumor-inoculated VEH and AP tibiae. Tumor areas are highlighted in yellow and necrosis in green (scale bars: 400 μm). (E) Quantification of tumor area relative to total bone area (Tm.Ar/T.Ar). (F) Necrotic area relative to tumor area inside the bone. (G) Images inside the tumor areas of sections stained using F4/80 and Ly6B antibodies for VEH and AP (×50) (scale bars: 200 μm). (H) Quantification of F4/80+ (n = 11 per group) and Ly6B+ VEH (n = 11) and AP (n = 12) staining (3 fields inside tumor area at ×20 per sample). Data are mean ± SEM; *P < 0.05, **P < 0.01, †P < 0.0001 (2-tailed Student’s t test).

(Supplemental Figure 5A). These findings agree with the previously reported increase in Gr-1+ cells in CXCL5−/− mice and the suggested role of this cytokine in bone marrow neutrophil homeostasis (38). Increased numbers of neutrophils were also found in the blood of CXCL5−/− mice relative to WT, but no changes in other leukocytes were observed (Supplemental Figure 5B). In contrast, the bone marrow analysis of tumor-free mice showed a significantly lower percentage of CD86+ and no changes in total F4/80+ cells (Supplemental Figure 5A). Since the intratumoral changes in myeloid cell populations are skewed in the opposite direction relative to tumor-free bone marrow, the differences observed in the tumor infiltration are not likely related to overall changes in the normal bone marrow of these mice; instead these changes highlight tumor environment specificity.

Altogether these findings demonstrate the critical role of the CXCL5 produced by the efferocytic macrophage as an inducer of inflammation-accelerating tumor growth in the bone microenvironment.

Cancer cell death induces accelerated tumor growth and bone destruction, while CXCL5 deficiency hinders tumor progression. Since the accelerated growth of cancer RM1-iC9 cells in the osseous vossicle model resulted in a large proportion of cells growing outside the bone, the effect of cancer cell death in tumor progression was further analyzed in an intratibial inoculation model. Engulfment of cancer cells after apoptosis induction mediated by AP treatment was evaluated. RM1-iC9 cells were GFP-labeled upon transfection with a lentiviral reporter construct prior to their inoculation into the left tibia of mice. Mice were randomized into groups treated with AP or VEH for 5 hours at day 5 after cancer inoculation (Figure 9A). Following this treatment, the bone marrow cells of the left tibia (tumor-inoculated) and the contralateral right tibia (control) were iso-
To further understand the effects of cancer cell apoptosis and subsequent efferocytosis on tumor progression and bone remodeling, the RM1-iC9 cells were inoculated via intratibial injection (day 0) and treated with AP, similarly to Figure 10A. Representative μCT images showing trabecular bone in the tumor tibiae for WT and CXCL5−/− mice (scale bars: 400 μm). Bone parameters were quantified by μCT in tumor-injected tibiae for WT and CXCL5−/−, similarly to Figure 10C. Representative images of H&E sections for WT and CXCL5−/− tibiae. Tumors are highlighted in yellow (scale bars: 400 μm). Quantification of tumor area relative to total bone area (Tm.Ar/T .Ar). Data are mean ± SEM; *P < 0.05, **P < 0.01, *P < 0.001 (2-tailed Student’s t test).

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To further understand the effects of cancer cell apoptosis and subsequent efferocytosis on tumor progression and bone remodeling, the RM1-iC9 cells were inoculated via intratibial injection (day 0) and treated with VEH or AP at days 6 and 8 (Figure 10A). Tibiae (tumor-inoculated and control) were collected at day 9 and analyzed by micro–computed tomography (μCT). The analysis of AP-Tumor tibiae showed significant reduction in trabecular and total bone volume and reduced trabecular number and thickness with a corresponding increase in trabecular spacing relative to VEH-Tumor (Figure 10, B and C). No changes were observed in the cortical bone (Figure 10C). These results suggest that escalated bone osteolysis was induced by increased tumor growth via amplified efferocytosis and the predominant osteolytic nature of RM1 cells. However, to exclude the possibility that differences in bone volume could be due to an effect of AP in nontumor cells, μCT analysis revealed no differences in bone between VEH-Control and AP-Control contralateral tibiae (Supplemental Figure 7), confirming that the differences observed between AP and VEH in the tumor are a result of the tumor-induced osteolysis mediated by cancer cell apoptosis. Furthermore, osteoclast tartrate-resistant acid phosphatase–positive (TRAP+) staining measured in bone inside the tumors (regions of interest [ROIs] depicted in Supplemental Figure 8A) demonstrated no differences in either osteoclast surface per bone surface or osteoclast number per bone surface (Supplemental Figure 8B), suggesting that the changes in trabecular and total bone volume are concomitant with osteolytic activity but not with increased osteoclast numbers.

The percentage of nonengulfed GFP+ RM1-iC9 cells was calculated relative to total cells. While higher numbers of GFP+ RM1-iC9 cells were observed in the bone marrow of tumor tibiae relative to control, AP treatment significantly increased this population when compared with VEH, indicating rapid growth of cancer cells after AP treatment (Figure 9, D and G).
Changes in the bones of AP-Tumor tibiae relative to the VEH-Tumor were correlated with the increased tumor progression seen in the AP-treated vossicle model. Tumor areas normalized to total bone area demonstrated that tumors were larger in AP-treated mice compared with VEH (Figure 10, D and E). The percentage of necrosis calculated as the ratio of necrotic areas divided by the total tumor area inside the bone demonstrated that increased tumor necrosis is induced by AP (Figure 10, D and F). These results suggest that the induction of apoptosis, resulting in larger tumors, also induces necrosis, perhaps enhanced by the limited capacity of macrophages to effectively clear the large proportion of apoptotic cells at an advanced stage of tumor growth. Consistent with the induction of apoptosis by AP, IHC inside the tumor revealed increased numbers of F4/80+ and Ly6B+ cells suggesting increased inflammation in tumors where cell death was stimulated (Figure 10, G and H).

Altogether these experiments demonstrated that induction of apoptosis with AP increased efferocytosis, rapidly stimulated cancer growth in the bone, and increased bone osteolysis.

Next, the role of CXCL5 in inducing tumor acceleration was investigated. An experiment comparing tumor growth in the bones of WT (CXCL5+/+) and CXCL5−/− mice was performed. Using the same approach described above (Figure 10A), RM1-iC9 cells were inoculated via intratibial injection in 6-week-old mice. To maximize proinflammatory CXCL5 expression, WT and CXCL5−/− mice were treated with AP to induce cancer cell apoptosis. At endpoint, μCT analysis was performed on tumoral tibiae. CXCL5-deficient mice had significantly higher trabecular bone volumes relative to WT, increased trabecular numbers, and reduced trabecular spacing (Figure 11, A and B).

H&E-stained tumor sections showed reduced tumor areas in CXCL5−/− mice relative to WT (Figure 11, A and C), confirming the proinflammatory role of CXCL5 induced by efferocytosis of apoptotic cancer cells.

Nonclassical (CD16+) phagocytic peripheral blood mononuclear cells and CXCL5 serum levels are associated with human prostate cancer skeletal metastasis. Mononuclear cells were isolated from whole peripheral blood of noncancer (Normal, n = 21) and prostate cancer bone-metastatic patients (Bone met., n = 47). Monocyte populations were assessed via flow cytometry. (A) CD68+ monocytes were assessed, and representative FACS images are displayed at right. (B) Subpopulations of CD14−/−CD16+ (blue boxes), CD14−/−CD16− (red boxes), and CD14−/−CD16+ (yellow boxes) were gated from total cell populations and quantified for Normal and Bone met. samples. Representative FACS plots are shown. (C) Freshly isolated monocytes (CD14+) from normal (n = 8) and prostate cancer bone-metastatic (n = 14) patients were cultured with phosphatidylserine-coated fluorescently labeled apoptotic-mimicry beads (3:1) and efferocytosis assessed by flow cytometry for CD14+ (APC) cells with ingested beads (representative FACS plots are shown). (D–F) Human serum isolated from normal (n = 20), localized (high-risk) prostate cancer (Localized, n = 40), and bone-metastatic prostate cancer (Bone met., n = 22) patients was analyzed by ELISA for CXCL5 (D), CXCL6 (E), and IL-8 (F). CXCL6 and IL-8 analysis included n = 38 for Localized. Data are mean ± SEM; **P < 0.01, *P < 0.001, †P < 0.0001 (Wilcoxon 2-sample test and Kruskal-Wallis test with Bonferroni’s correction).
metastatic to bone with multiple lesions (≥3 lesions), with no prior chemotherapy for metastatic disease and no signs of ongoing infection or another malignant disease, was analyzed (Supplemental Table 2). These samples were compared with cohorts of patients with either localized high-risk prostate cancer or non-cancer patients. The age distribution from each group is shown in Supplemental Figure 10A. All samples were analyzed by ELISA to measure 3 closely related cytokines: CXCL5, CXCL6 (a human cytokine homologous to mouse CXCL5), and IL-8 (CXCL8). Human IL-8 is a proinflammatory cytokine with no direct murine homolog, suggested to be functionally homologous with murine CXCL1, CXCL5, and CXCL6 (43). Intriguingly, normal samples displayed significantly higher serum CXCL5 relative to localized cancer samples (Figure 12D). Importantly, considerably higher levels of CXCL5 were detected in the serum of bone-metastatic patients compared with other groups, while no significant differences were observed for CXCL6 (Figure 12E). Contrary to the CXCL5 results, serum samples from bone-metastatic patients showed significantly reduced IL-8 concentrations relative to normal and localized samples (Figure 12F). Furthermore, no significant correlation was found between the CXCL5 or IL-8 serum concentrations and the age of patients for each group, respectively (Supplemental Figure 10B). Altogether these findings suggest the predominant role of proinflammatory CXCL5 in prostate cancer bone metastasis in concert with the increased phagocytic circulating CD68+CD14+CD16+ monocytic cell population.

Discussion
Persistent inflammation in the tumor microenvironment is a common denominator of cancer progression characterized by the infiltration of myeloid cells that facilitate tumor expansion and metastasis (44). Chronic inflammation increases cell stress and tissue damage, which induces apoptotic/necrotic cell death. Consistent with these results, we hypothesized that phagocytic monocytes/macrophages would increase serum levels of the inflammatory CXCL5 cytokine in bone metastasis patients because of tumor cell clearance both in circulation and in the bone marrow compartment. A cohort of serum samples from patients with progressive advanced castration-resistant prostate cancer metastatic to bone with multiple lesions (≥3 lesions), with no prior chemotherapy for metastatic disease and no signs of ongoing infection or another malignant disease, was analyzed (Supplemental Table 2). These samples were compared with cohorts of patients with either localized high-risk prostate cancer or non-cancer patients. The age distribution from each group is shown in Supplemental Figure 10A. All samples were analyzed by ELISA to measure 3 closely related cytokines: CXCL5, CXCL6 (a human cytokine homologous to mouse CXCL5), and IL-8 (CXCL8). Human IL-8 is a proinflammatory cytokine with no direct murine homolog, suggested to be functionally homologous with murine CXCL1, CXCL5, and CXCL6 (43). Intriguingly, normal samples displayed significantly higher serum CXCL5 relative to localized cancer samples (Figure 12D). Importantly, considerably higher levels of CXCL5 were detected in the serum of bone-metastatic patients compared with other groups, while no significant differences were observed for CXCL6 (Figure 12E). Contrary to the CXCL5 results, serum samples from bone-metastatic patients showed significantly reduced IL-8 concentrations relative to normal and localized samples (Figure 12F). Furthermore, no significant correlation was found between the CXCL5 or IL-8 serum concentrations and the age of patients for each group, respectively (Supplemental Figure 10B). Altogether these findings suggest the predominant role of proinflammatory CXCL5 in prostate cancer bone metastasis in concert with the increased phagocytic circulating CD68+CD14+CD16+ monocytic cell population.

**Discussion**
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quently, several aggressive tumors are characterized by high apoptotic rates, which are often underestimated as a result of efficiency of clearance (efferocytosis) (11). Efferocytosis is a fundamental function of professional phagocytes (mainly macrophages) and an essential mechanism of normal tissue homeostasis that also prevents inflammation. However, here we present new evidence suggesting that the response of bone marrow macrophages when engaged in efferocytosis of prostate cancer cells differs from the response when engaged with cells that bone marrow macrophages would normally encounter, such as osteoblasts and stromal cells.

Cancer cell efferocytosis was found to induce an inflammatory response in macrophages characterized by the expression of CCL5, CXCL1, CXCL5, and IL-6, which have been correlated with tumor inflammation mechanisms and metastasis in several cancer types (45, 46). Accordingly, these findings suggest that the induction of efferocytosis leads to activation of NF-κB(p65) and Stat3 signaling and proinflammatory cytokine expression. Blocking NF-κB(p65) or Stat3 signaling mitigated the expression of inflammatory cytokines in macrophages, while simultaneous inactivation of these pathways blocked the efferocytic function of macrophages. Since neither Static nor Bay11-7082 inhibited efferocytosis when used alone, these findings further suggest that these two pathways are not only activated by efferocytosis, but are critical components of this mechanism, where at least one has to be fully active for this essential function of macrophages. In accordance, NF-κB(p65) and Stat3 are steadily activated in immune cells, where they selectively regulate each other to trigger amplification loops that stimulate the production of protumor inflammatory factors (7, 24, 25). It has been suggested that continuous activation of Stat3 is a prerequisite for these tumor-promoting immune responses (47). Furthermore, activation of Stat3 by efferocytosis mediates the M2-like polarization of bone marrow macrophages (12). In vivo studies in breast and colon cancer showed that efferocytosis induced M2 polarization and metastasis, while increased cell death correlated with poorer outcomes (13, 48). Moreover, the proinflammatory cytokine IL-6 induced M2-like macrophage polarization in human macrophages (49). Our findings suggest that efferocytosis induces tumor-promoting inflammation mediated by the persistent activation of Stat3 and NF-κB(p65) in macrophages.

The bone marrow is a myeloid-rich microenvironment for seeding and colonization of metastatic cells. Using apoptosis-inducible murine RM1 prostate cancer cells and bone vossicles as a syngeneic model of osseous environmental tumor growth, it was demonstrated that induced cancer death accelerated tumor progression. RM1 cells were chosen in this study as these cells are derived from prostate epithelium of C57BL/6J mice and overexpress the Ras and Myc oncogenes, which resembles the oncogene-specific gene expression signatures of prostate cancer patient samples and is associated with prostate cancer progression (50, 51). The vossicle model provides an osseous scaffold for tumor growth where cancer cells are implanted directly in the interacting bone niche. This interaction between tumor and bone is critical, especially at the early stages of tumor growth. The importance of bone in the vossicle serving as scaffold was demonstrated in previous experiments in which tumor growth was accelerated in the vossicles isolated from a Gas6–/– mouse relative to controls after implantation in the same host. In these experiments tumor growth was assessed in vossicles where tumors overgrew the bone scaffolds (31). Another example of the vossicle model highlighting the bone microenvironment in tumor progression revealed that ablation of the osteoblast niche hindered prostate cancer growth in tumor vossicles (17). Furthermore, it was recently reported that macrophage ablation decelerated tumor growth using the RM1/vossicle model (9).

Elevated numbers of apoptotic cells were observed in the tumor vossicles where apoptosis was induced, and cell death induced a higher percentage of F4/80+CD206+ (M2-like) macrophages in tumors, which correlates with increased polarization induced by efferocytosis (12). Furthermore, CD11b+Gr-1+ cells, expressing markers of myeloid-derived suppressor cells (MDSCs), were also increased in tumors where cell death was induced. Although overlapping values of different immune cell populations were detected by flow cytometry between VEH- and AP-treated groups, this could be explained by the significant level of apoptosis found in RM1-iC9 control tumors and/or the possibility that AP treatment did not equally reach all cancer cells because of the complexity of large tumors. For similar reasons, a high variability in the CXCL5 concentrations was detected in the tumors linked to the various levels of apoptosis, which would overlap in some cases with controls. While we cannot exclude the important role of other proinflammatory cytokines, the critical role of CXCL5 is further supported by the significant correlation found between CXCL5 levels and tumor weight.

Other studies have identified CXCL5 as a predictor for bone-metastatic prostate cancer (52), and in a breast cancer metastatic model it was shown that this cytokine is a factor secreted by tumor-associated osteoblasts to function as an inducer of epithelial-to-mesenchymal transition and metastasis (53). Furthermore, in a model of rhabdomyosarcoma, it was demonstrated that antibody-mediated neutralization of CXCR2 (a receptor for CXCL5) enhanced anti-PD1 effectiveness by preventing the tumor infiltration of the CD11b+Ly6G+ suppressor cells (54). However, none of these works identified the proinflammatory efferocytic macrophage as a trigger of tumor growth in the bone microenvironment.

Using the CXCL5+ mice in a syngeneic tumor vossicle model delineated the crucial role of host-derived CXCL5 in the mechanism of cell death–induced tumor progression. CXCL5 deficiency significantly inhibited tumor progression in bone vossicles. Although the cytokine array might have missed other cytokines represented at lower concentrations, CXCL5 was expressed at considerable levels, and the major contribution was from the host microenvironment and not the cancer cells. Intriguingly, ELISA analysis showed reduced levels of CXCL1 in CXCL5–/– vossicles versus WT. Although CXCL1, a cytokine that interacts with the same receptor (CXCR2) as CXCL5, was expressed at lower levels in the same receptor (CXCR2) as CXCL5, these results are consistent with reduced cell death–induced inflammation mediated by CXCL5 deficiency in the vossicles.

Analysis of CXCL5–/– tumor vossicles revealed reduced percentages of M2-like (F4/80+CD206+) and phagocytic (CD68+F4/80+) macrophages as well as inflammatory Ly6B+ cells and granulocytic CD11b+Gr-1+ cells that characterize MDSCs, all of which aligns with reduced inflammation and efferocytosis and the role of CXCL5 as a crucial chemoattractant of MDSCs. Importantly, and in contrast, increased influx of CD86+ cells was identified within CXCL5–/– vossicles.
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RESEARCH ARTICLE

The accelerated tumor progression in the bone reported here resulted in increased osteolysis, necrotic cell death, and inflammation as detected by increased F4/80+ and Ly6B+ cell infiltration. Previous in vivo studies in prostate cancer models suggest that increased macrophage infiltration into the tumors results in CXCL5 and other proinflammatory cytokines that facilitate recruitment and activation of cancer therapies, the identification of CXCL5 as a crucial factor in this mechanism provides an important clue for designing additional successful therapies.
Methods

Cell lines and animals. Murine prostate cancer RM1 cells were originally a gift from Timothy C. Thompson (Baylor College of Medicine, Houston, Texas, USA) (63, 64). Apoptosis-inducible RM1-iC9 cells were developed by transduction of murine RM1 with the viral construct for inducible caspase-9 (iC9) as previously described (23). The newly constructed RM1-iC9 cells were characterized by an independent laboratory (IDEXX BioResearch), which confirmed origin from the C57BL/6 mouse strain, the same profile as the parent RM1, and no contamination was detected. PC3 cells (also tested by IDEXX BioResearch) were obtained from the American Type Culture Collection (ATCC). MC3T3-E1 subclone 4 (MC4) cells (ATCC CRL2593) were provided by Renny T. Franceschi (University of Michigan). C57BL/6J mouse primary prostate epithelial cells (MPECs) were obtained from Cell Biologics (C57-6038).

All animal experiments were performed with approval from the University of Michigan Institutional Animal Care and Use Committee. Control WT mice (C57BL/6J) were purchased from the Jackson Laboratory. CXCL5−/− knockout mice on a C57BL/6J background were donated by George Scott Worthen and Junjie Mei (Division of Nephrology, Children’s Hospital of Philadelphia, University of Pennsylvania, Philadelphia, Pennsylvania, USA) (38).

Vossicle (vertebral body) subcutaneous RM1-iC9 tumors. The vossicle model, which provides an osseous scaffold for tumor growth in a bone microenvironment, has been previously described (9, 31). Detailed description is provided in Supplemental Methods.

Intratibial injection model. Apoptosis-inducible RM1-iC9 murine prostate cancer cells (2 × 103) or RM1-iC9-GFP (RM1-iC9 cells transduced with GFP-reporter lentiviral construct; Lenti-GFP, University of Michigan Vector Core) were injected in the left tibiae of C57BL/6J mice under anesthesia as previously described (9). Detailed description is provided in Supplemental Methods.

Human mononuclear cell isolation. Human peripheral blood mononuclear cells were isolated from 7.5 ml of venous blood collected in EDTA-coated vacutainer tubes (Becton Dickinson). Samples were collected at the University of Michigan Comprehensive Cancer Center from patients with prostate cancer bone metastases as well as from healthy donors without any cancer history. Complete description is provided in Supplemental Methods.

Statistics. Statistical analyses were performed in GraphPad Prism 6 (GraphPad Software Inc.) using unpaired 2-tailed Student’s t test to compare differences between 2 groups with significance of P less than 0.05. One-way ANOVA with multiple-comparisons tests was used to compare 3 or more groups with significance of P less than 0.05. The in vivo tumor progression was analyzed using 2-way ANOVA with multiple comparisons. Human peripheral blood mononuclear cell populations and human ELISA data were analyzed using nonparametric rank tests; 2-group comparisons were tested with a Wilcoxon rank test, and when more than 2 groups were tested the Kruskal-Wallis rank test was used at significance of P less than 0.0083 (using the Bonferroni multiple-comparisons adjustment).

TRACER data from multiple experimental repeats were combined. Measurements were log2-transformed and normalized to the average intensity of the control reporter, which contained only the minimal TATA-box promoter. The data were further normalized to background, defined as cells transfected with reporter virus but not administered apoptotic cells. Finally, data were normalized to the initial reporter measurement for each treatment condition at 0 hours. Statistical significance among background and experimental groups was determined using the R package limma (65). To generate heatmaps, the log, fold change for each condition and time point was averaged among replicates. Reporters were grouped into 4 clusters using k-means clustering.

Study approval. All participating patients were enrolled in our prospective study (HUM0052405), which was approved by the University of Michigan Medical School Institutional Review Board (Ann Arbor, Michigan, USA). All patients provided their written informed consent prior to their participation in the study.

Additional methods and extended descriptions are provided in Supplemental Methods.

Author contributions

HR, JDJ, and LKM conceived and designed the study. YW, KJP, TMM, and ETK provided human samples, and JEN provided the apoptosis-inducible construct and expertise. HR, MCP, JDJ, SW, AJK, and RK performed experiments. HR, JDJ, MCP, AJK, RK, LDD, and KLM contributed to experimental design and data analysis. JEW contributed to tissue data analysis, images and expertise. SDN performed statistical analysis. MCP, SW, and HR created figures. HR, LKM, AJK, JDJ, MCP, SW, RK, and LDD contributed to manuscript writing. All authors revised and edited the manuscript.

Acknowledgments

We are indebted to George Scott Worthen and Junjie Mei for providing CXCL5−/− mice. We thank Fabiana Soki, Megan Michalski, and Benjamin Sinder for technical assistance, and thoughtful discussions. We also thank Sasha Meshinchi for confocal microscopy assistance (University of Michigan Microscopy Facility), David Adams for assistance with Anims Imaging flow cytometer, Michelle Lynch for μCT assistance, and Chris Strayhorn and Theresa Cody for histology sectioning. We are very grateful to Victoria Zakrzewski, Kenneth Rieger, and Serk In Park for artistic contributions. This work was supported by NIH awards P01-CA093900 to LKM, KJP, and ETK, R01-CA173745 to LDS, R01-DK053904 to LKM, R01-DE21139 and R01-DE23220 to JEN, and R01GM093183 to LDS; and Department of Defense awards W81XWH-14-1-0408 to JDJ, W81XWH-14-1-0287 to TMM, and Prostate Cancer Foundation (PCF) Young Investigator Award to TMM.

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