Priority Report

Recruitment of Myeloid but not Endothelial Precursor Cells Facilitates Tumor Regrowth after Local Irradiation

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Abstract

Tumor neovascularization and growth might be promoted by the recruitment of bone marrow–derived cells (BMDC), which include endothelial precursor cells and “vascular modulatory” myelomonocytic (CD11b+) cells. BMDCs may also drive tumor regrowth after certain chemotherapeutic and vascular disruption treatments. In this study, we evaluated the role of BMDC recruitment in breast and lung carcinoma xenograft models after local irradiation (LI). We depleted the bone marrow by including whole-body irradiation (WBI) of 6 Gy as part of a total tumor dose of 21 Gy, and compared the growth delay with the one achieved after LI of 21 Gy. In both models, the inclusion of WBI induced longer tumor growth delays. Moreover, WBI increased lung tumor control probability by LI. Exogenous delivery of BMDCs from radiation-naive donors partially abrogated the WBI effect. Myeloid BMDCs, primarily macrophages, rapidly accumulated in tumors after LI. Intratumoral expression of stromal-derived factor 1α (SDF-1α), a chemokine that promotes tissue retention of BMDCs, was noted 2 days after LI. Conversely, treatment with an inhibitor of SDF-1α receptor CXCR4 (AMD3100) with LI significantly delayed tumor regrowth. However, when administered starting from 5 days post-LI, AMD3100 treatment was ineffective. Lastly, with restorative bone marrow transplantation of Tie2-GFP–labeled BMDC population, we observed an increased number of monocytes but not endothelial precursor cells in tumors that recurred following LI. Our results suggest that an increase in intratumoral SDF-1α triggered by LI recruits myelomonocytes/macrophages which promotes tumor regrowth. Cancer Res; 70(14): 5679–85. ©2010 AACR.

Introduction

The recruitment of various blood-borne bone marrow–derived cells (BMDC) might be important for tumor neovascularization and growth, but their roles are often difficult to differentiate (1, 2). Endothelial precursor cells (EPC) may directly incorporate in 0% to 50% of newly formed tumor vessels, depending on the tumor type, organ site, and mouse strain (3). In addition, other BMDCs, referred to as “vascular modulatory/accessory” cells, might support angiogenesis in a paracrine manner. Among them, arguably the most important for angiogenesis are cells of myeloid/monocyte lineage (CD11b+), in particular, macrophages and Tie2-expressing monocytes (TEM; refs. 4, 5). The trafficking and tissue retention of BMDCs might depend, at least in part, on stromal-derived factor 1α (SDF-1α)–CXCR4 receptor pathway activation (6, 7).

EPCs might be increasingly attracted to tumor sites as a result of certain therapies and influence their outcome (e.g., vascular-disruptive and certain chemotherapeutic treatments; refs. 8, 9). But it remains unknown if EPCs contribute significantly after local irradiation (LI) to tumors, the neovascularization of which is presumed to be deficient (10, 11). Infiltration by other BMDCs—e.g., myelomonocytes/macrophages—has been previously documented in irradiated tumors (12–16). However, their role as modulators of tumor radiation response remains largely uncharacterized. Here, we evaluated the role of various BMDCs in tumor regrowth after LI in lung and breast tumor models.

Materials and Methods

Animals and tumors

54A human lung tumors were xenografted in male athymic NCr/Sed nude (nu/nu) mice, and MCA8 mouse mammary carcinomas were implanted in female syngeneic FVB mice (both subcutaneously in the hind limb; refs. 3, 11). To detect specific BMDC populations in tumors (i.e., Tie2+CD11b–EPCs and Tie2+CD11b+ TEMs), we irradiated wild-type FVB mice with 9 Gy of whole-body irradiation (WBI) followed by restorative bone marrow transplantation (BMT) from Tie2-GFP-FVB donors to create chimeric wild-type/Tie2-GFP-BMT mice (3). MCA8 tumors were implanted in these mice 6 weeks post-BMT.
Treatments and response evaluation

Tumor size was measured with a caliper at least thrice a week. When a tumor reached 5.5 mm in mean diameter, the mouse was randomized to a treatment group. Tumors were \( \gamma \)-irradiated using the same dose delivered either locally alone or locally plus a sublethal WBI dose of 6 Gy (for detailed treatment procedures, see Supplementary Material). Unsorted BMDCs or fluorescence-activated cell–sorted Sca1\(^{-} \)CD11b\(^{+} \) or Sca1\(^{+} \)CD11b\(^{+} \) BMDCs were i.v. infused in mice receiving LI plus WBI. CXCR4 inhibition, alone or after LI, was achieved using AMD3100 (5 mg/kg/d; Sigma-Aldrich) delivered by ALZET micro-osmotic pumps (DURECT Corporation) over 2 weeks. Therapeutic efficacy was measured as the time taken for tumors to grow to 7.5 mm in diameter (i.e., \(~2.5\)-fold increase compared with the pretreatment volume).

Immunohistochemistry and image analysis

Tumors were excised, cut in half, fixed for 2 hours in 4% formaldehyde in PBS, incubated in 30% sucrose in PBS overnight at 4°C and frozen in optimal cutting temperature compound (Tissue-Tek). Transverse tumors sections, 10 \( \mu \)m thick, were immunostained with antibodies to endothelial marker CD31, pan-myeloid marker CD11b or SDF-1\(^{\alpha} \), and counterstained by mounting with DAPI-containing medium (Vectashield, Vector Labs). Images were captured with a stack step of 5 \( \mu \)m using an Olympus confocal microscope, and the stained areas were quantified using in-house segmentation algorithms coded on a MATLAB platform (MathWorks). The antibodies used and details of image analysis are described in the Supplementary Material.

Flow cytometric analysis

Flow cytometry was performed on single-cell suspensions prepared from whole tumors after digestion with collagenase type II (Worthington Biochemical Corporation) and immunostaining for CD11b, Gr1, and F4/80 (using fluorescence-labeled monoclonal antibodies from BD PharMingen) as described (3), using an LSR-II flow cytometer (BD Biosciences).
Statistical analysis
Differences between mean values in each group were evaluated by $t$ test for independent samples and considered significant when $P < 0.05$. Data are presented as mean ± SEM. The comparison between collections of groups was carried out using $t$ tests for linear contrasts in one-way ANOVA. Tumor control probabilities were compared using $\chi^2$ test.

Results and Discussion

WBI delays tumor regrowth after irradiation
To diminish the potential involvement in tumors of host-derived cells following LI, we treated tumor-bearing mice with a sublethal dose of 6 Gy WBI. This dose is known to damage the bone marrow and deplete leukocytes temporarily (for 1.5–2 weeks) from the blood circulation (17, 18). Using this approach, we compared the efficacy of the same radiation dose (21 Gy) given to tumors either as two LI fractions or LI plus WBI; a flow chart of these treatments is presented in Fig. 1A, note the identical duration of irradiation. The growth of 54A tumors in nude mice was arrested in both groups, but tumor regrowth was significantly delayed when radiotherapy included WBI (Supplementary Fig. S1A). In the case of M5a8 carcinomas grown in immunocompetent FVB mice, tumors shrank by 1.5 to 2 mm posttreatment in both groups and WBI significantly delayed their regrowth (Supplementary Fig. S1B). In both tumors, the growth inhibition by 6 Gy of WBI alone was also longer than that by 6 Gy of LI. As summarized in Fig. 1B and C, WBI provided additional tumor growth delay in both models despite the difference in tumor postradiation dynamics and distinct immune profiles of the hosts. Finally, WBI significantly enhanced tumor curability following LI with higher doses compared with LI alone in 54A xenografts (Fig. 1D). Of note, WBI did not change the weight, appearance, or behavior of mice (data not shown).

BMDC infusion promotes tumor regrowth after LI plus WBI
The infusion of unsorted BMDCs in mice treated with 15 Gy of LI plus 6 Gy of WBI abrogated the significant regrowth delay achieved by WBI in both tumor models (Fig. 1B and C). The “tumor-rescuing” effect was greater for irradiated M5a8 tumors ($P < 0.05$, Fig. 1C). Moreover, a similar effect was seen after the infusion of myeloid progenitor BMDCs (Sca1+CD11b+) or radiotherapy included WBI (Supplementary Fig. S1A). In the case of M5a8 carcinomas grown in immunocompetent FVB mice, tumors shrank by 1.5 to 2 mm posttreatment in both groups and WBI significantly delayed their regrowth (Supplementary Fig. S1B). In both tumors, the growth inhibition by 6 Gy of WBI alone was also longer than that by 6 Gy of LI. As summarized in Fig. 1B and C, WBI provided additional tumor growth delay in both models despite the difference in tumor postradiation dynamics and distinct immune profiles of the hosts. Finally, WBI significantly enhanced tumor curability following LI with higher doses compared with LI alone in 54A xenografts (Fig. 1D). Of note, WBI did not change the weight, appearance, or behavior of mice (data not shown).

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more mature myeloid BMDCs (Sca1−CD11b+) in mice with 54A tumors treated with 15 Gy of LI plus 6 Gy of WBI (data not shown). These results indicate that the recruitment of radiation-naïve BMDCs could facilitate tumor regrowth after LI. As SDF-1α is a critical cytokine for BMDC recruitment (7), we then compared its intratumoral levels before and after LI.

LI upregulates SDF-1α expression in tumors
In both 54A and MCA8 models, 20 Gy of LI significantly increased SDF-1α protein expression in tumor tissues, as measured 2 days later (Fig. 2A). In addition, we found a trend for increased SDF-1α expression even after irradiation at a dose of 4 Gy (Supplementary Fig. S2). This rapid upregulation of
SDF-1α is likely induced directly by radiation, and is consistent with previous reports of SDF-1α upregulation shortly after irradiation of normal tissues or cancer cells in vitro (19, 20). SDF-1α might also be upregulated at later time points if tumors become hypoxic (16). As SDF-1α is thought to exert its effects on BMDCs via the CXCR4 receptor (6, 21), we next tested if CXCR4 blockade could delay tumor regrowth after LI.

**CXCR4 blockade delays tumor regrowth only when administered immediately after LI**

Inhibition of SDF-1α/CXCR4 signaling for 2 weeks using AMD3100-containing osmotic pumps did not affect tumor growth but significantly inhibited tumor regrowth of both 54A and MCa8 tumors when commenced immediately after 20 Gy of LI (Fig. 2B and C; Supplementary Fig. S3). This is consistent with recent data from glioma xenografts (16). However, this effect was abrogated when AMD3100 treatment was initiated 5 days after 20 Gy of LI. On the other hand, the combination of AMD3100 with 6 Gy of WBI immediately after 15 Gy of LI provided no additional growth delay in MCa8 tumors (Supplementary Fig. S4). This indicates that WBI and AMD3100 treatment have overlapping effects postirradiation. Collectively, these results suggest a critical role for the rapid, radiation-induced recruitment of BMDCs in tumors, mediated by SDF-1α-CXCR4 signaling. Therefore, we then studied the early BMDC infiltration in tumors after the treatment regimens shown in Fig. 1A.

**Rapid accumulation of myeloid BMDCs post-LI may facilitate tumor relapse**

We measured tumor accumulation of myeloid BMDCs 3 days after radiation treatments. Immunostaining for CD11b showed a significant increase in myeloid BMDC infiltration following LI compared to nonirradiated tumors (Fig. 4). This change in BMDC infiltration was observed in both 54A and MCa8 tumors.

| Figure 4. Analysis of BMDCs in tumors recurring after LI in wild-type/Tie2-GFP-BMT mice. A, representative confocal microscopy images of fluorescence immunohistochemistry in tumors recurring after 20 Gy of radiation versus nonirradiated size-matched (control) tumors. Note localization of GFP expression (green) in CD11b+ cells outside vessels (blue, arrows) but not in CD31+CD11b− vascular endothelial cells (red). B and C, quantification of BMDCs: overall myeloid cell infiltration (CD11b+) and CD31+ microvascular density were not significantly changed, but the total number of Tie2+ BMDCs increased in irradiated tumors (B). These represented mostly Tie2+CD11b+ TEMs, the majority of which were also CD31+ but had perivascular location; in contrast, the number of vessel-incorporated Tie2+CD31+CD11b− EPCs was negligible and not different in tumors growing after irradiation (C; *, P < 0.05). |
after LI alone in 54A tumors, whereas the regimens containing WBI abrogated this effect (Fig. 3A). Of interest, immunostaining for CD31 showed no significant change in tumor vessel density at day 3 in any group after radiation compared with nonirradiated tumors (Fig. 3A). Flow cytometric analysis of whole tumor lysates confirmed the significant increase in number of CD11b+ cells after LI and its reduction by WBI, both in 54A and MCa8 tumors (Fig. 3B and C). Further phenotypic analyses showed that the vast majority of these cells were F4/80+ macrophages. The number of F4/80+ macrophages was also significantly decreased in tumors by WBI (P < 0.05, using the linear contrast test for groups with versus without WBI). CD11b+Gr1+ immune-suppressive myeloid cells were elevated in tumors after 21 Gy of LI, but were not reduced by WBI. These results further support the importance of rapid myeloid cell accumulation for tumor resistance to LI. This LI-induced macrophage infiltration might also be associated with HIF-1α activation as well as inducible nitric oxide synthase and vascular endothelial growth factor overexpression and lead to better survival and further proliferation of irradiated endothelial cells (14).

TEM infiltration but not EPC vessel incorporation is increased in recurring tumors

Finally, we tested the contribution of various BMDCs to tumors recurring after LI. Although incompletely understood, tumor angiogenesis post-LI might be deficient compared with nonirradiated tumors (11, 15). In this context, vasculogenesis by nonirradiated EPCs could play a more substantial role than in the case of radiation-naïve tumor growth. To test this, we used MCa8 tumor growing in chimeric wild-type/Tie2-GFP-BMT mice, a tumor model in which EPC recruitment is negligible in the absence of treatment (3). We detected no significant effect of the prior BMT on the tumor growth rate and the growth delay after 20 to 25 Gy of LI, which suggested an efficient bone marrow reconstitution and a lack of apparent local “tumor bed effect” beyond 6 weeks after BMT in FVB mice. Because new vessel formation is mandatory only to support the increasing tumor mass postirradiation (10), we analyzed EPC incorporation in the vasculature of relapsed tumors (i.e., when they reached a diameter of 7.5 mm). Tumor tissue analysis by confocal fluorescence microscopy showed no difference in CD31+ microvascular density or total myeloid CD11b+ BMDC infiltration between LI-treated tumors and size-matched nonirradiated tumors (Fig. 4). However, the accumulation of Tie2-GFP+ BMDCs was significantly increased in tumors recurring after LI. The vast majority of Tie2-GFP+ BMDCs were CD11b+CD31+ myeloid cells localized in the tumor interstitium but not incorporated into the vessel wall. Thus, recurring tumors contained significantly more vascular-modulatory TEMs. In contrast, the number of EPCs incorporated in tumor vessels was substantially lower than TEM accumulation in tumor tissues and was not significantly different in recurring versus nonirradiated tumors. Collectively, these data support the potential role of TEM paracrine influence but not EPC-based vasculogenesis in recurring tumors after LI. The latter is consistent with the negligible incorporation of EPCs in the vasculature of tumors growing in preirradiated normal tissues (15).

Conclusions and implications

This study provides compelling evidence that host-derived BMDC infiltration in tumors is stimulated by LI and facilitates tumor recurrence through paracrine effects on irradiated tumor vasculature, inside and adjacent to the regressing/stabilized tumors. The rapid recruitment of nonirradiated myeloid BMDCs—primarily macrophages—is mediated at least in part by SDF-1α and may create a microenvironment that promotes the survival of tumors and endothelial cells and regrowth. Once tumors recur, an increased TEM, but not EPC infiltration, might promote tumor growth. These results suggest that targeting the SDF-1α-CXCR4 pathway in a specific “therapeutic window” or TEM accumulation may delay tumor recurrence after radiotherapy.

Disclosure of Potential Conflicts of Interest

R.K. Jain: commercial research grant, AstraZeneca and Dyax; honoraria, Genzyme and Ahlstrom; ownership interest, SynDevRx; consultant/advisory board, AstraZeneca, Dyax, Genzyme, Regeneron, and SynDevRx. The other authors disclosed no potential conflicts of interest.

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