Angiogenic factors FGF2 and PDGF-BB synergistically promote murine tumor neovascularization and metastasis

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Tumors produce multiple growth factors, but little is known about the interplay between various angiogenic factors in promoting tumor angiogenesis, growth, and metastasis. Here we show that 2 angiogenic factors frequently upregulated in tumors, PDGF-BB and FGF2, synergistically promote tumor angiogenesis and pulmonary metastasis. Simultaneous overexpression of PDGF-BB and FGF2 in murine fibrosarcomas led to the formation of high-density primitive vascular plexuses, which were poorly coated with pericytes and VSMCs. Surprisingly, overexpression of PDGF-BB alone in tumor cells resulted in dissociation of VSMCs from tumor vessels and decreased recruitment of pericytes. In the absence of FGF2, capillary ECs lacked response to PDGF-BB. However, FGF2 triggers PDGFR-α and -β expression at the transcriptional level in ECs, which acquire hyperresponsiveness to PDGF-BB. Similarly, PDGF-BB–treated VSMCs become responsive to FGF2 stimulation via upregulation of FGF receptor 1 (FGFR1) promoter activity. These findings demonstrate that PDGF-BB and FGF2 reciprocally increase their EC and mural cell responses, leading to disorganized neovascularization and metastasis. Our data suggest that intervention of this non-VEGF reciprocal interaction loop for the tumor vasculature could be an important therapeutic target for the treatment of cancer and metastasis.

Introduction

Similar to growing healthy tissues, expansion of malignant tissues and tumor metastasis are dependent on neovascularization, which is accomplished by processes of angiogenesis, vasculogenesis, and vascular remodeling (1–13). The tumor vasculature usually consists of disorganized, leaky, premature, tortuous, and hemorrhagic blood vessels that provide a structural basis for cancer cell invasion and spread (1, 3, 9, 14). These unusual features of tumor blood vessels represent the consequence of an imbalanced production of various angiogenic factors and the hypoxic environment within the tumor tissue. For example, VEGF is usually expressed at high levels in most tumors and has become an obvious therapeutic target for cancer therapy (1, 3, 6, 14). Indeed, current antiangiogenic strategies for cancer therapy are based on blocking VEGF functions and anti-VEGF agents have successfully been used for the treatment of certain types of human cancers (15–17). However, tumors also produce multiple non-VEGF angiogenic factors, and anti-VEGF monotherapy could potentially encounter drug resistance, suggesting that tumors could use non-VEGF angiogenic factors to grow blood vessels (18, 19).

The tumor tissue consists of heterogeneous and genetically unstable malignant cells and a diversity of various other cell types including inflammatory cells, stromal cells, blood vessel ECs, lymphatic ECs, and VSMCs and/or pericytes, which are constantly exposed to hypoxic and stressful environments (18, 19). Both genetic instability of tumor cells and diversity of cell types determine expression of multiple angiogenic factors in the tumor tissue (20). Both PDGF-BB and FGF2 are frequently expressed at high levels in various tumor tissues (21, 22). While PDGF-BB displays potent biological activity on PDGFR-expressing VSMCs, it usually lacks biological effects on ECs that do not express detectable levels of PDGFRs (21, 23, 24). Thus PDGF-BB is considered as a mitogenic and chemotactic factor for VSMCs/pericytes, but not for ECs. Indeed, deletion of PDGF-B or its prominent receptor, PDGFR-β in mice leads to embryonic lethality, manifesting leaky and hemorrhagic phenotypes due to lack of pericytes and/or VSMCs in blood vessels (23, 24). In contrast to PDGF-BB, FGF2 is a potent angiogenic factor directly stimulating EC proliferation, although it also acts on VSMCs in vitro (25). However, delivery of FGF2 in vivo mainly induces angiogenesis without significantly increasing recruitment of VSMCs (26). Although the roles of individual angiogenic factors in promoting tumor angiogenesis are relatively well studied, little is known about the interplay between various angiogenic factors and their combined effects in tumor neovascularization, growth, and metastasis. The tumor vasculature is constantly exposed to multiple growth factors, and the complex interactions between various factors determine the ultimate outcome of tumor vessel growth, which might involve activation of MAPK and other signaling components in ECs and other vascular cells (27).

In this study, we provide compelling evidence that FGF2 acts as a sensitizer for ECs to respond to PDGF-BB, which feeds back to VSMCs to enhance their responses to FGF2 stimulation. The underlying mechanisms of this reciprocal interaction involve upregulation of PDGFR expression in ECs by FGF2 and of FGFR1 expression in VSMCs by PDGF-BB. The biological consequence of such a reciprocal interaction in tumors is manifested by hyperneovascularization and high degree of disorganized primitive tumor vasculatures, which are poorly coated with pericytes and VSMCs. These altera-
tions of tumor blood vessels lead to an accelerated tumor growth rate and metastasis. Thus our results provide what we believe to be new mechanistic insights on the cooperative role of various angiogenic factors in promoting tumor growth and metastasis.

Results

Prerequisite role of FGF2 in facilitating PDGF-BB–induced capillary EC migration. To study EC responses to FGF2 and PDGF-BB stimulation, bovine capillary endothelial (BCE) cells were used for in vitro studies. Previous studies have demonstrated that these capillary ECs were highly relevant for in vivo angiogenesis and were sensitive to various angiogenic stimuli and inhibitors (28, 29). As expected, FGF2 at a low concentration (10 ng/ml) significantly stimulated BCE cell proliferation (Figure 1A). In contrast, PDGF-BB exhibited a minor proliferative effect on these cells at a high concentration (100 ng/ml), but pretreatment of BCE with FGF2 did not significantly potentiate the effect of PDGF-BB on cell proliferation. Similarly, pretreatment of BCE cells with PDGF-BB did not increase BCE responses to FGF2 stimulation. Interestingly, FGF2 did not significantly induce BCE cell migration, suggesting that FGF2 acts as a mitogenic factor but not a chemotactic factor for ECs (Figure 1B). These results are consistent with previous findings that FGF2 is mainly a proliferative growth factor for capillary ECs (30). In the absence of FGF2, PDGF-BB did not significantly induce BCE cell motility. However, PDGF-BB remarkably stimulated the migration of BCE cells preexposed to FGF2, and this migratory effect could be inhibited by an anti-PDGFR agent, STI571, in a dose-dependent manner (Figure 1B). These results show that pretreatment of capillary ECs with FGF2 is a prerequisite for PDGF-BB–induced EC migration.

FGF2 and PDGF-BB synergistically induced angiogenesis. Reciprocal interactions between FGF2 and PDGF-BB on ECs and VSMCs suggested that these 2 factors cooperatively regulated vessel growth in vivo. To study this possibility, FGF2 and PDGF-BB were coimplanted into the mouse cornea. Indeed, FGF2 plus PDGF-BB synergistically induced corneal angiogenesis (Figure 2). Confocal analysis of whole-mount CD31-stained corneal tissues showed that at day 5 after implantation, FGF2 plus PDGF-BB–induced vessels consisted of a relatively disorganized vasculature as compared with single factor–induced vessels (Figure 2, E–G and I–K). We should emphasize that the disorganized vasculatures could only be detected at the early phase of the vessel formation, and they were remodeled into a well-defined tree-like vascular network after long-term exposure to FGF2 and PDGF-BB (31).
To validate these in vitro findings, various angiogenic factor–implanted corneal blood vessels were detected in situ for mRNA expression of PDGFRs. Consistent with the in vitro EC data, extremely high levels of both PDGFR-α and PDGFR-β were found in the FGF2-induced blood vessels (Figure 4, C–J). ECs in the FGF2-induced new blood vessels expressed high levels of both PDGFR-α and PDGFR-β. Uptregulation of PDGFR-α and PDGFR-β by FGF2 in BCE cells was further validated by RT-PCR analysis (Figure 4A). These data demonstrate that the 5′-end region, the GC-rich box, and the CAAT box are all crucial for FGF2-triggered upregulation of PDGFR expression.

To validate these in vitro findings, various angiogenic factor–induced corneal blood vessels were detected in situ for mRNA expression of PDGFRs. Consistent with the in vitro EC data, extremely high levels of both PDGFR-α and PDGFR-β were found in the FGF2-induced blood vessels (Figure 4, C–J). ECs in the FGF2-induced new blood vessels expressed high levels of both PDGFR-α and PDGFR-β. In contrast, PDGF-BB–induced vessels only expressed a moderate level of PDGFRs. These in vivo findings provide further evidence that FGF2 at the transcriptional level triggers PDGFR-β–induced EC and blood vessel growth.

Elevated protein levels of PDGFR-β in FGF2-treated EC. To investigate if the elevated mRNA levels of PDGFRs in FGF2-treated BCE cells also led to increased protein expression, cell lysates treated with different concentrations of FGF2 were immunoblotted with an anti–PDGFR-β–specific antibody. At the concentrations of 5 and 10 ng/ml, FGF2 remarkably increased PDGFR-β expression in BCE cells (Figure 4B). In addition to elevation of total receptor molecules, the phosphorylated PDGFR-β molecules were also proportionally increased, suggesting that high levels of PDGFR-β led to activation of these receptors.

Activation of intracellular signaling pathways. Treatment of BCE cells with FGF2 led to an elevated level of phosphorylated Erk (p-Erk), which could be further increased following PDGF-BB treatment. High levels of p-Erk persisted for more than 6 hours before returning to the basal level (Figure 4K). A modest and transient effect of PDGF-BB alone on elevation of p-Erk was also detected (data not shown). Interestingly, after 60 minutes of exposure to PDGF-BB, the FGF2-pre-treated cells began to show an elevated level of phospholipase C-γ (PLC-γ), and high levels of PLC-γ persisted for the entire time course of experiments. Levels of phosphorylated Src and Akt were not altered by FGF2 pretreatment. These findings validate that PDGF-BB directly induces EC activity in FGF2-pre-treated cells.

PFDF-BB potentiated FGF2–induced VSMC proliferation. To study the role of PDGF-BB in modulating biological effects of FGF2 on VSMCs, we isolated rat aorta VSMCs, which expressed α-SMA but lacked an ability of internalizing acetylated LDL (Ac-LDL) as BCE cells (Figure 1, C–F). As expected, both FGF2 and PDGF-BB significantly stimulated VSMC proliferation and migration (Figure 1, C–F). Pretreatment of FGF2 did not alter PDGF-BB–induced VSMC proliferation and migration responses as compared with the FGF2-treated group. Remarkably, pretreatment of VSMCs with PDGF-BB significantly potentiated FGF2-induced cell proliferation but not cell migration (Figure 1, G and H). These findings demonstrate that PDGF-BB modulates FGF2-induced cell proliferation activity on VSMCs.

PDGF-BB induced FGFR1 promoter activity in VSMCs. To study if FGF2 and PDGF-BB reciprocally communicate with each other at the receptor signaling level, we studied the regulation of FGFR expression by PDGF-BB in both ECs and VSMCs. The promoter region of FGFR1 was fused with the chloramphenicol acetyltransferase (CAT) reporter gene, and the fusion construct was used to transfect both ECs and VSMCs. Interestingly, PDGF-BB induced about a 3-fold increase of reporter gene activity in VSMCs (Figure 3C). The PDGF-BB–induced FGFR1 activity seemed to be restricted to VSMCs, and stimulation of FGFR1/CAT–transfected BCE cells with PDGF-BB did not result in any increase of promoter activity (data not shown). In contrast to FGFR1, PDGF-BB did not induce promoter activity of FGFR3 and FGFR4 when assayed by promoter luciferase activity (data not shown).

To define the critical regions of the PDGFR promoter responsible for FGF2 activation in ECs, serial deletions or mutations were introduced into different regions of the PDGFR-β promoter (Figure 3, D–G). Deletion or mutation of the 5′-end region, the GC-rich box, or the CAAT box resulted in loss of the promoter activity in FGF2–treated BCE cells (Figure 3, D–G). Uptregulation of PDGFR-α and PDGFR-β by FGF2 in BCE cells was further validated by RT-PCR analysis (Figure 4A). These data demonstrate that the 5′-end region, the GC-rich box, and the CAAT box are all crucial for FGF2-triggered upregulation of PDGFR expression.
not shown). Thus our data indicate that PDGF-BB may upregulate FGFR1 expression in VSMCs but not in ECs.

**FGF2 and PDGF-BB synergistically stimulated tumor neovascularization.** To study the reciprocally interactive role of FGF2 and PDGF-BB in promoting tumor angiogenesis, secreted forms of FGF2 and PDGF-BB were overexpressed in a murine fibrosarcoma. As expected, implantation of PDGF-BB tumors in syngeneic mice resulted in an accelerated tumor growth rate as compared with control tumors (Figure 5A). Implantation of both PDGF-BB and FGF2 tumors in mice led to a further increase of the tumor growth rate. Immunohistochemical analysis of tumor tissues revealed an increased vascular density in FGF2- and PDGF-BB–overexpressing tumors (Figure 5, B and C). Strikingly, in the FGF2- and PDGF-BB–coexpressing tumors, not only was the vascular density remarkably increased, but the tumor vascular structure also underwent remarkable changes (Figure 5B). The FGF2/PDGF-BB–induced tumor blood vessels appeared to be primitive vascular plexuses, which exhibited disorganized large pseudo-vessels. Intriguingly, high numbers of capillary sprouts or tip cells existed in these primitive vascular plexuses, suggesting coordinated effects of these 2 factors in promoting tumor vessel growth. These findings demonstrate that FGF2 and PDGF-BB together not only synergistically stimulate tumor neovascularization but also change the architecture of the tumor vasculature.

**Uncoupling of tumor microvessel growth and VSMC coating.** PDGF-BB has previously been reported to be a potent mitogen for VSMCs (21). However, overexpression of PDGF-BB alone in tumors did not significantly increase the number of α-SMA–positive structures (Figure 6, A and B). Surprisingly, PDGF-BB promoted disassociation of VSMCs from tumor microvessels, which exhibited random distribution of VSMCs throughout the tumor tissue (Figure 6A). Although FGF2 and PDGF-BB together slightly increased the percentage of association between CD31-positive structures and VSMCs, the total number of tumor vessels coated with VSMCs was significantly lower than that of controls (Figure 6A). In contrast, a nearly completely overlapping pattern of CD31-positive and α-SMA–positive structures was seen in vector plus FGF2 tumors (Figure 6, A and C). Similarly, virtually all microvessels of vector and FGF2 tumors were coated with VSMCs. These data demonstrate surprising findings that expression of PDGF-BB in tumor cells results in disassociation of VSMCs from the tumor vasculature.

**Inhibition of pericyte recruitment by tumor-produced PDGF-BB.** We then analyzed pericyte distribution within the tumor tissue. In the vector-transfected tumors, a significant number of neuro/glial cell 2 chondroitin sulfate proteoglycan–positive (Ng2-positive) pericytes was found in the tumor tissue, and virtually all of them remained in association with tumor microvessels (Figure 7, A–C). FGF2 significantly increased both the total number of pericytes in the tumor tissue and their recruitment onto tumor microvessels. In contrast, very few Ng2-positive pericytes were present in PDGF-BB tumors, but almost all of them remained in association with tumor vessels (Figure 7, A–C). Similarly, only very sparsely distributed pericytes were found in the PDGF-BB and FGF2–coexpressing tumor tissue, and they mainly remained in association with relatively large tumor vessels. Almost all microvessels in PDGF-BB and PDGF-BB plus FGF2 tumors lacked pericyte coating. These unexpected findings demonstrate that PDGF-BB inhibits pericyte recruitment in the tumor environment.

**Reciprocal vascular interactions between FGF2 and PDGF-BB promoted metastasis.** To study the functional consequence of FGF2 plus PDGF-BB on tumor vasculature, we performed experiments of spontaneous metastasis in mice in which subcutaneous primary tumors were surgically removed at the size of 1.5 cm³. FGF2 tumor cells were labeled with luciferase, and PDGF-BB tumor cells were labeled with GFP for monitoring tumor spreads. Approximately 4 weeks after removal of primary tumors, about 30% of animals in the FGF2 plus PDGF-BB group developed pulmonary luciferase-positive metastatic nodules as detected by bioluminescence analysis (Figure 8, B and E). In contrast, FGF2 tumor–bearing mice or vector plus FGF2 tumor–bearing mice did not show any signs of lung metastasis (Figure 8A). These findings demonstrate that PDGF-BB could facilitate metastasis of FGF2 tumor cells despite the fact that PDGF-BB displayed no effects on tumor cells in vitro. Necropsy analysis of tumor-bearing...
mice revealed that about 30% of FGF2 plus PDGF-BB tumor-bearing mice had visible surface lung metastases (Figure 8, D and F). In contrast, none of the vector plus FGF2 tumor-bearing mice developed visible pulmonary lung metastasis (Figure 8, C and F). Histological analysis of lung tissues confirmed that FGF2 plus PDGF-BB stimulated lung metastases (Figure 8, H and I). Because vector plus PDGF-BB tumor cells also expressed GFP, we detected GFP-positive tumor cells in the lung tissue. Interestingly, about 44% of mice showed GFP-positive pulmonary micrometastases, and none of the vector plus PDGF-BB tumor-bearing mice had GFP-positive cells in their lungs (Figure 8, G, J, and K). These findings show that reciprocal interplay between FGF2 and PDGF-BB in the tumor environment promotes pulmonary tumor metastasis.

Discussion

Both genetic and epigenetic factors contribute to the switch of an angiogenic phenotype in tumors, which produce multiple angiogenic factors. Although angiogenic activity of individual factors is relatively well studied, little is known about the interplay between various tumor-produced angiogenic factors and their cooperative efforts in promoting tumor neovascularization. Here we report on the impact of the interplay between FGF2 and PDGF-BB on vas-
cular cells in promoting tumor neovascularization, vessel maturation, and metastasis. FGF2 triggers PDGF-BB responses in ECs and enhances PDGF-BB–induced tumor angiogenesis. Conversely, PDGF-BB augments the effect of FGF2 on VSMCs. The underlying mechanisms of this reciprocal interaction involve transcriptional upregulation of their receptors on ECs and VSMCs. Activation of this interactive loop in a tumor environment leads to accelerated but uncoupled growth of ECs and VSMCs, leading to the formation of primitive tumor vasculature, which promotes metastasis.

PDGF-BB is a well-characterized growth factor displaying potent biological effects on mural cells including pericytes and VSMCs, but not on ECs (21, 23, 24). Inactivation of both PDGF-B and PDGFR-β genes results in lethal embryonic phenotypes of improper development of the vasculature due to lack of pericytes and VSMCs (23, 24). Although a recent report demonstrates that PDGFR-α but not -β is expressed in isolated ECs, it is not known how PDGFR-α expression is regulated or what its role is in mediating PDGF-BB–induced angiogenic responses (32). All members of the PDGF family display potent angiogenic activity in vivo (33). For example, PDGF-AA, PDGF-AB, PDGF-BB, and PDGF-CC potently induce neovascularization in mouse corneas. If VSMCs are the primary vascular target cells for PDGFs, how could they induce the growth of angiogenic vessels consisting mainly of ECs? These and other in vivo studies suggest that the endothelial effects of PDGFs are probably modulated by other factors. In the present study, we provide compelling in vitro and in vivo evidence that FGF2 induces promoter activities of both PDGFR-α and PDGFR-β in ECs and the increased transcripts lead to high levels of PDGF-α and PDGFR-β in ECs. When high levels of PDGFR-α and PDGFR-β in ECs become activated by available ligands, the PDGF-induced angiogenic response is overwhelming.

Figure 5
Tumor growth rates and vasculature. (A) Growth factor– or vector-transfected tumor cells were subcutaneously implanted in SCID mice, and tumor growth was measured daily. Green line, FGF2 + PDGF-BB; red line, FGF2; blue line, vector + FGF2; purple line, vector and black PDGF-BB. (B) At day 13 after tumor cell implantation, tumors were removed and stained with an anti-CD31 antibody, and tumor blood vessels were analyzed by confocal microscopy using 3D projections. GFP-expressing tumor cells are green in color, and tumor blood vessels are presented in red. Arrows point to tip-cell sprouts from the vascular plexuses induced by FGF2 and PDGF-BB. (C) Quantification of CD31-positive tumor vessels from 8–12 randomized cryosectioned fields. The data represent means of average determinants ± SEM. *P < 0.05; **P < 0.001. Scale bar: 100 μm (B, upper and lower panels); 50 μm (B, middle panels). V, vector.
FGF2 potently stimulates EC proliferation but has almost no effect on chemotaxis (30). Additionally, our present study shows that PDGF-BB is able to induce EC migration but not proliferation. Angiogenesis requires both EC proliferation and migration, and FGF2 triggered PDGF response in ECs, probably to compensate for its defective effect on cell migration. When both systems became activated, coordinated EC proliferation and migration contributed to guided vessel growth. Increased levels of PDGF-BB positively fed back to the FGF system in VSMCs by transcriptional upregulation of FGFR1 expression (Figure 9). This result is consistent with previous findings that PDGF-BB upregulates FGFR1 expression in VSMCs (34). PDGF-BB only facilitates FGF2-induced VSMC proliferation but not migration. In a similar scenario as for ECs, PDGF-BB cross-communicates with FGF2 to keep a balance between VSMC proliferation and migration. However, the overall coordinated control mechanism of EC and VSMC growth and association in newly formed blood vessels is not understood. Our findings uncover the underlying molecular mechanisms by which FGF2 and PDGF-BB synergistically and coordinately stimulate neovascularization. The angiogenic synergism is particularly limited to the interactive loop between FGF and PDGF systems because other potent angiogenic factors such as VEGF-A are unable to induce PDGFR expression and promote synergistic angiogenesis in vitro and in vivo (31). There seems to be a functional difference of FGF2 plus PDGF-BB–induced vasculatures in the tumor versus healthy tissue environments. In the ischemic muscle tissue, delivery of dual factors induced functional and relatively mature blood vessels, which improved functional outcomes of ischemic tissues. However, in the tumor environment, FGF2 plus PDGF-BB promoted relatively premature vasculatures that mediated metastasis. Although it is unclear why these 2 factors induced premature blood vessels in tumors, it is possible that other factors such as VEGF-A could also play a role in a further complex interplay between different factors. For example, it is known that FGF2 and VEGF-A could also synergistically induce angiogenesis (35). The other possibility is that tumors constitutively produce these factors at high levels, whereas a slow-release system is used in the ischemic settings. The third possibility is that PDGF-BB released by a slow-release scaffold could build up a growth factor–release gradient and attract vessel growth and mural cell recruitments toward the gradient. In contrast, the tumor-released PDGF-BB is evenly distributed within the tumor environment.
Figure 7

Interaction between pericytes and ECs in tumors. At day 10 after implantation, tumor tissues were double-stained with an anti-CD31 antibody and an anti-NG2 antibody. (A) The CD31- (red) and NG2-positive (blue) signals were revealed by Alexa Fluor 555- and Cy5-labeled antibodies, respectively, using single-layer projections in a confocal microscope. Tumor cells were GFP positive (green). T, intratumoral area; PT, peritumoral area. (B) Total numbers of NG2 positive vessels were randomly counted from 9 fields/group, and (C) percentages of NG2-positive vessels relative to total CD31-positive vessels were calculated. The data represent means of average determinants ± SEM. *P < 0.05; **P < 0.01; ***P < 0.001. Scale bar: 50 μm.

One of the most intriguing findings in our present study is that the tumor-produced PDGF-BB promotes disassociation of VSMCs from the tumor vasculature. This finding contradicts the known effect of PDGF-BB on recruitment of VSMCs into the newly formed blood vessels. Why does PDGF-BB repel VSMCs from tumor vessels? What is so special about the tumor environment? Although these questions may be involved in complex genetic and epigenetic mechanisms of many molecular players that regulate tumor growth and vessel formation, a simple and straightforward answer could be implicated in a spatiotemporal relationship between the site of PDGF-BB production and tumor vessel development. In most other studies, PDGF-BB is produced by growing ECs, which play guiding and recruiting roles in the growing cone of vessel tips for attracting pericytes and VSMCs. However, when PDGF-BB is produced by tumor cells, VSMCs and pericytes might become “confused” about their migration and lose their guidance movement due to lack of chemoattractant gradient. Thus they could move away from blood vessels and remain in close contact with tumor cells. In addition to VSMCs, ECs might also become confused about their migration, as their tip cells are randomly distributed in the vascular plexuses (Figure 5B). The underlying mechanism of the inhibitory effect of PDGF-BB on recruitment of pericytes into the tumor vasculature remains unknown. These findings are crucially important for understanding the role of PDGFs in tumor vessel maturation, patterning, and drug target definition.

Bloodstream metastasis is dependent not only on vessel density but also on quality of vascular structure. Primitive tumor vessels are vulnerable for tumor cell invasion and provide a structural basis...
Stimulation of metastasis by coexpression of FGF2 and PDGF-BB in tumors. Subcutaneous tumors were removed when they reached 1.5 cm³. (A and B) Mice were followed for 4 weeks, and pulmonary metastases of vector/FGF2 tumor–bearing (A) and PDGF-BB/FGF2 tumor–bearing (B) mice were examined by bioluminescence. Arrows point to luciferase–positive lung metastases. (E) Numbers of pulmonary luciferase–positive animals relative to total numbers of animals are presented as percentages. (C and D) Examples of lung morphology of vector/FGF2 tumor–bearing (C) and PDGF-BB/FGF2 tumor–bearing (D) mice and arrows in D panel point to visible lung metastases in the PDGF-BB/FGF2 group. (F) Numbers of lungs with visible pulmonary metastases versus total numbers of lungs are presented as percentages. (H and I) Lung tissues were stained with H&E and metastatic nodules were validated in the PDGF-BB/FGF2 group (I). (H) No metastasis was visible in the vector/FGF2 tumor-bearing control group. (J and K) GFP-positive metastases were revealed by analysis of lung sections under a fluorescent microscope of vector/FGF2- (J) and PDGF-BB/FGF2-tumor bearing (K) mice. (G) Numbers of lungs with GFP-positive metastases relative to total numbers of lungs are presented as percentages. Met, metastasis; HL, healthy lung tissue. Scale bars: 100 μm.

expect a similar drug-resistant mechanism to also exist in cancer patients. PDGFs are attractive targets for cancer therapy, and anti–PDGF agents are effective for the treatment of experimental and human tumors (37). Paradoxically, PDGF and probably other vascular remodeling factors could play an important role in normalization of tumor vasculature, which might increase chemotherapeutic drug delivery (14, 38). This paradoxical issue needs to be further clarified in future studies. Although PDGF-BB and FGF2 may be expressed at low levels in some tumors, their coordinated activity in the promotion of synergistic angiogenesis and uncoupling of vascular remodeling in a tumor environment should not be underestimated for tumor growth and metastasis.

Taken together, our work provides what we believe to be a novel mechanistic insight on transcriptional regulation of angiogenic responses and vascular remodeling induced by FGF2 and PDGF-BB. To our knowledge, this is the first report describing how the interplay between different angiogenic factors can promote tumor neovascularization and metastasis. As tumor blood vessels are constantly exposed to various angiogenic factors, understanding molecular mechanisms of interplays between various angiogenic factors is crucial for the development of effective therapeutic agents. Our findings provide one such example of reciprocal interactions and a potentially novel therapeutic option.

Methods

Reagents and animals. The antibodies used in our studies include a rat anti-mouse CD31 monoclonal antibody (BD — Pharmingen), a mouse anti-human α-SMA (Dako), and a rabbit anti-mouse NG2 antibody (Chemicon International). Immunocompetent 6- to 8-week-old female SCID mice were acclimated and caged in groups of 6 or less. Animals were followed up to 3–6 weeks and anesthetized by an injection of Hypnorm (fentanyl citrate, fluanisone; VetaPharma) and Midazolam (dormicium; Roche) (1:1) before all procedures and sacrificed by a lethal dose of Hypnorm; about 10–50 μl/mouse. All animal studies were reviewed and approved by the animal care and use committee of the North Stockholm Animal Board (Stockholm, Sweden).

EC proliferation assay. A 72-hour BCE cell proliferation assay was performed as previously described (39). Cells growing in gelatinized 6-well plates were
dispersed in 0.25% trypsin solution and resuspended with DMEM containing 0.5% FCS. We added 1 x 10^5 cells to each gelatinized well of 24-well plates and incubated them at 37°C for 1 hour, followed by addition of various concentrations of FGF2 and PDGF-BB in quadruplicates to each well and further incubation for 12 hours. The old medium was replaced with fresh medium in the presence or absence of growth factors (10 ng/ml FGF2 or 100 ng/ml PDGF-BB) and incubated for 60 hours. After a total of 72 hours of incubation, cells were trypsinized, resuspended in Isoton II solution (Beckman Coulter), and counted with a Coulter counter (Beckman Coulter).

**VSMC proliferation.** VSMCs were isolated from the aortic media of male F344 rats by collagenase digestion and cultured in F-12 Ham medium (GIBCO) supplemented with 50 μg/ml L-ascorbic acid, 50 μg/ml streptomycin, 50 IU/ml penicillin, and 20% FCS (GIBCO) as previously described (40). These cells were kindly provided by Eric Wahlberg’s laboratory at the Karolinska Hospital, Stockholm, Sweden. Cells growing in 6-well plates were dispersed in a trypsin solution and resuspended in Isoton II solution (Beckman Coulter), and counted with a Coulter counter (Beckman Coulter).

**EC and VSMC chemotaxis assay.** The motility responses of BCE or VSMC cells to FGF2 and PDGF-BB were assayed by using the modified Boyden chamber technique described in ref. 13. Briefly, the ability of BCE or VSMC cells to migrate through a micropore nitrocellulose filter (8 μm pore size) was measured as a criterion for chemotactic stimuli. Cells were either pretreated or not pre-treated with 10 ng/ml FGF2 or 100 ng/ml PDGF-BB for 24 hours, and cells were trypsinized and resuspended in fresh medium supplemented with 0.5% FCS. Approximately 5 x 10^4 cells were added into each of the upper wells of the Boyden chamber (48-well chamber), which contained 10 ng/ml of FGF2 or 100 ng/ml of PDGF-BB in the lower chamber. The cells were incubated for 4 hours at 37°C, after which the Boyden chamber was disassembled and cells attached to the filter were fixed in methanol and stained with a Giemsa solution. Four replicate samples were used in each experiment, and all experiments were performed at least 4 times. Cells that had migrated through the filter were counted using a light microscope and plotted as numbers of migrating cells per optic field.

**Ac-LDL and α-SMA staining of VSMCs and ECs.** VSMCs and ECs were grown on coverslips in 6-well culture dishes under growth conditions described above. Cells in approximately 80% confluency were fixed in acetone for 20 minutes and vigorously washed with PBS prior to incubation with an α-SMA antibody (1:1000 dilution). An anti-mouse IgG-FITC–labeled antibody was used as the secondary antibody (Vector Laboratories). Ac-LDL labeled with the fluorescent dye Dil was diluted to 10 μg/ml in DMEM supplemented with 10% FCS. Medium was then added to 6-well plates, where VSMCs or ECs were growing on coverslips as described above and incubated for 4 hours at 37°C. Media was removed from the culture, and cells were washed several times with probe-free media and examined under a fluorescent microscope.

**Transient transfection and reporter gene assay.** PDGF-Rα and PDGF-Rβ promoter–luciferase fusion gene constructs were generated as previously described (41–43). BCE–human telomerase reverse transcriptase–positive (BCE-hTERT+) cells were maintained in DMEM containing 10% FBS, 100 μg/ml penicillin, and 100 μg/ml streptomycin. About 1 x 10^5 cells were seeded in each well of 12-well plates for 16 hours prior to transfection, yielding approximately 95% of confluence. Before transfection, the medium was changed into serum-free and streptomycin/penicillin-free DMEM. The cells were then transfected with FuGENE6 (Roche) reagent according to the manufacturer’s protocol. Briefly, 1.2 μl FuGENE6 reagent was diluted in 100 μl serum-free and penicillin/streptomycin-free DMEM, and then 350 ng PDGFR-α and PDGFR-β promoter–reporter gene constructs and 50 ng CMV–β-galactosidase expression vector were vector. These optimal values were determined experimentally several times. The resulting complex was added to BCE or VSMC cells, and 10 ng/ml FGF2 or PDGF-BB were added. Cells were incubated for 40 hours, and luciferase activity was determined at the end of incubation according to the protocol described in the Bright-Glo luciferase kit (Promega). Measurements were done in quadruplicate using a Sirius luminometer (Berthold).

The FGFR1-CAT was kindly provided by J. DiMario, Rosalind Franklin University, Chicago, Illinois, USA. The promoter-reporter fusion gene constructs were used for transfection of BCE and VSMCs according the procedures described above. The transfected cells were treated with FGF2 or PDGF-BB as described above. For measurement of CAT activity, a standard protocol provided by the manufacturer was used (Roche).

**RT-PCR.** A quantitative RT-PCR method was performed to amplify PDGFR-α and PDGFR-β using cDNAs prepared from BCE cells as templates. The following primers were used: a 27-mer 5′-end nucleotide sequence for PDGFR-α (5′-GGCCGACCTTGTGGAGGTTTGTGACG-3′) and PDGFR-β (5′-AACTTCGTTAGAACAGGAGGGAGAGA-3′).
3′), a 26-mer 3′-end nucleotide sequence for PDGFR-α (5′-CGTTGC-
GAGCCGGAGTCTCGGATG-3′), a 21-mer 5′-end nucleotide sequence for
PDGFR-β 5′-GCAGACTGTTGCCAGCAAGG-3′, and a 24-mer 3′-end nucleotide sequence for PDGFR-β 5′-CGACCCGGTAGCATTTT-
GATAGCC-3′. As a loading control bovine GAPDH cDNA was used, and
paired primers of a 29-mer 5′-end nucleotide sequence for bovine GAPDH
(5′-CGAGATGAGAGAGGCCAGGGGCG-3′) and a 30-mer 3′-
end nucleotide sequence for bovine GAPDH (5′-CCAGGCCTCTCCATT-
GTTAGTAGGCACCC-3′) were used. The PCR-amplification program
consisted of 35 cycles of the following: 94°C for 2 minutes, 50°C for 2
minutes, and 72°C for 2 minutes.

Mouse corneal micropocket assay. The mouse corneal assay was performed
as previously described (28). Micropockets containing 40 ng of FGF2 (Phar-
macia & Upjohn), 160 ng of PDGF-BB (Peprotech Inc.), or 40 ng of FGF2
plus 160 ng of PDGF-BB were implanted into mouse corneal micropock-
ets. Circumferential neovascularization and vascularization areas were
measured at various time points.

In situ hybridization. We used 2 probes complementary to PDGFR-α
(nucleotides 423–470 and 3083–3130) and 2 probes complementary to
PDGFR-β (nucleotides 946–996 and 2610–2657). All probes were used sep-
arately and did not match any known sequence in GenBank except those
of the intended genes. Corneal histologic sections were hybridized with the
32P-labeled probes. Slides were rinsed, dehydrated, and emulsion dipped.
After 5 weeks of exposure, slides were developed and counterstained with
cresyl violet. Specific labeling was confirmed by similar expression patterns
revealed by 2 probes each (complementary to different parts of the mRNA)
for PDGFR-α and PDGFR-β. See Supplemental Methods for details and
quantification of autoradiographic signals (supplemental material avail-
able online with this article; doi:10.1172/JCI32479DS1).

Immunoblot analysis. Monolayers of BCE cells were stimulated with
10 ng/ml FGF2 or 100 ng/ml PDGF-BB for various time points. After exten-
sive washing with PBS, cells were lysed with a lysis buffer of 20 mM Hepes,
pH 7.1, 150 mM NaCl, 50 mM NaF, 1% Triton X-100, 30 mM Na2PO4,
5 mM ZnCl2, 20 mM β-glycerophosphate, 10 mM 4-nitrophenyl phosphate,
1 mM DTT, 100 μM Na3VO4, and 0.5 mM PMSF. Cell lysates were centrifuged in
a tabletop centrifuge at the maximum speed (18,000 g) for 20 minutes,
and the supernatants were collected for protein determination by using a
modified Lowry method (Bio-Rad) with BSA as a standard. Equal amounts
of each sample were loaded onto 4%–20% acrylamide SDS/PAGE gradi-
et gels (NOVEX). Proteins were transferred onto nitrocellulose mem-
branes (Schleicher & Schuell), which were subsequently blocked overnight
at room temperature with 2.5% BSA-PBST (140 mM NaCl, 2.7 mM KCl,
8.1 mM Na2HPO4, pH 7.3, and 0.3% Triton X-100), followed by incubation
at 4°C overnight with primary antibodies against PLCγ-1 (kindly provided
by Carl-Henrik Heldin, Uppsala, Sweden; ref. 44), including phosphospe-
cific Src (PY418; Biosource), phosphospecific-Akt (Ser473; Cell Signaling
Technology), phosphospecific-p44/p42 MAPK (Thr202/Tyr204; Cell Sig-
naling Technology), phosphospecific-p38 MAPK (Thr180/Tyr182; Cell Sign-
naling Technology), and phosphospecific-SAPK/JNK (Thr183/Tyr185;
Cell Signaling Technology) antibodies. After extensive washing with PBST,
membranes were incubated for about 1 hour at room temperature with
peroxidase-coupled secondary antibodies. After further washing, immuno-
reactive signals were revealed by chemiluminescence. We used an antibody
specific for β-actin (monoclonal anti–β-actin, clone AC-15 [mouse ascites
fluid, Mouse IgG1]; Sigma-Aldrich) as a loading control.

Tumor growth assay. A murine fibrosarcoma T241 cell line was used for
the generation of transfectant cell lines overexpressing enhanced GFP and
hPDGF-BB, or the empty vector as previously described (45). The fibro-
sarcoma cells expressing a secreted form of hGF2 were established as
previously described (46). In some experiments, vector and FGF2 tumor
cells were further stably transduced with the luciferase gene as previously
reported (47). Approximately 0.5 × 106 vector-, FGF2-, and PDGF-BB-
transduced tumor cells; 0.25 × 106 vector-transduced plus 0.25 × 106 FGF2-
transduced tumor cells; or 0.25 × 106 PDGF-BB–transduced plus 0.25 × 106
FGF2-transduced tumor cells were subcutaneously implanted on the back of
each 6- to 8-week-old female SCID mouse, and tumor volumes were
measured as previously reported (39, 48).

Metastasis assay. When primary tumors reached a volume of 1.5 cm3 (the
ethical limit), primary tumors were surgically removed under anesthetic
conditions (n = 10/group). The open wounds were sutured, and mice were
observed for 4 weeks. Mice were i.p. injected with 1.5 mg/200 μl of L-Diuc-
ferin (Xenogen) and scanned for luciferase-positive metastases by biolumi-
nescence imaging analysis (IVIS 100; Xenogen). The mice were sacrificed at
end of week 4 after removal of the primary tumors. After opening the chest
and abdomen, the exposed organs and tissues were further scanned for
luciferase-positive metastases. Lungs were resected, and surface metastases
were examined visually under a light microscope. Several organs including
lung, liver, spleen, kidney, ovary, and brain were sectioned and analyzed for
GFP-positive tumor metastases under a fluorescent microscope.

Histology and whole-mount staining. Malignant and nonmalignant paraf-
fin-embedded tissues were sectioned at 5 μm thickness and stained with
H&E according to our previously described methods (13). Whole-mount
staining was performed according to previously published methods (45).
Briefly, small pieces of tissues were cut into thin slices and fixed in 3%
PFA overnight, followed by treatment with proteinase K (20 μg/ml). Rat
anti-mouse CD31 antibodies were used as primary antibodies and goat
anti-rat Alexa Fluor 555–conjugated antibodies (Invitrogen) were used as
secondary antibodies. Additionally, a mouse anti-human a-SMA primary
antibody (Dako) and a rabbit anti-mouse secondary antibody Alexa Fluor
647–conjugated antibody were used (Invitrogen). An anti-NG2 antibody
(Chemicon) was used for detection of pericytes. Sections were examined
under a confocal microscope (Zeiss Confoil LSM510 microscope). By
scanning 10 thin sections of each sample (4–5 μm long), 3D images of
each tissue sample were assembled. Quantitative analysis from at least
5 different tissue sections was performed using the color range tool of
Adobe Photoshop CS software program. Alternatively, slides were exam-
ined using a custom-built dual-mode Ultraview LCI confocal system
(PerkinElmer) at the Karolinska Institute visualization core facility. 3D
reconstruction was carried out using Imaje, and images were further
processed in Adobe Photoshop CS.

Immunofluorescent staining. Cryostat tissue sections 20 μm thick were
incubated with a specific antibody against CD31 according to standard
immunochemistry procedures. Briefly, after 3 washes in PBS, speci-
mens were incubated for 30 minutes in a blocking solution containing 4%
nonimmune goat serum (Vector Laboratories) in PBS, followed by incu-
bation with primary antibodies for 2 hours at room temperature. After
3 rigorous washes with PBS, tissues were incubated for 1 hour at room
temperature with secondary antibodies labeled with Alexa Fluor 555–
coujugated red (1:500). Sections were examined using a confocal
microscope (Zeiss LSM710 microscope). Positive signals were photo-
graphed under a confocal microscope (IVIS 100; Xenogen). The mice were sacrificed at
end of week 4 after removal of the primary tumors. After opening the chest
and abdomen, the exposed organs and tissues were further scanned for
luciferase-positive metastases. Lungs were resected, and surface metastases
were examined visually under a light microscope. Several organs including
lung, liver, spleen, kidney, ovary, and brain were sectioned and analyzed for
GFP-positive tumor metastases under a fluorescent microscope.

Statistics. Statistical analyses of in vitro and in vivo results were performed
using the standard 2-tailed Students t test on Microsoft Excel. Some exper-
imental data were analyzed using the 2-way ANOVA method in Micro-
soft Excel, followed by a Bonferroni post hoc test using the GraphPad
QuickCalcs online Web site (http://www.graphpad.com/quickcalcs/post-
test1.cfm). P < 0.05, P < 0.01, and P < 0.001 were deemed as significant,
highly significant, and extremely significant, respectively.

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