Visualization of Proliferative Vascular Endothelial Cells in Tumors in Vivo by Imaging Their Partner of Sld5-1 Promoter Activity

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Vascular endothelial cells (ECs) isolated from tumors characteristically express certain genes. It has recently been suggested that tumor vessel normalization facilitates effective drug delivery into tumors; however, how tumor vessel normalization can be recognized on the basis of the molecules expressed by tumor ECs is not clearly defined. The degree of cell proliferation is an important indicator to characterize the condition of the ECs. Herein, we generated transgenic mice expressing enhanced green fluorescent protein (EGFP) under the transcriptional control of the DNA replication factor partner of Sld5-1 (PSF1; official name GINS1) promoter to assess whether active ECs can be distinguished from dormant ECs. Predictably, ECs in the adult skin exhibited no EGFP signals. However, after s.c. injection of tumor cells, some ECs shifted to EGFP positivity, enabling distinction of EGFP-positive from EGFP-negative cells. We found that only a fraction of the EGFP-negative ECs strongly expressed the glycosylphosphatidylinositol-anchor protein CD109 associated with the phosphatidylinositol 3-kinase pathway. Taken together, these data indicate that areas of vascular normalization in tumors can be detected by CD109 expression, and this provides a window of opportunity for timing chemotherapy. (Am J Pathol 2018, 188: 1300–1314; https://doi.org/10.1016/j.ajpath.2018.01.015)

For suppression of cancer cell proliferation and metastasis, the focus has been on tumor angiogenesis as a therapeutic target. A conventional strategy is to inhibit angiogenesis by blocking angiogenic factors, to reduce the supply of nutrients and oxygen, and to prevent cancer cell invasion through blood vessels.1 On the other hand, vascular normalization is proposed as a new therapeutic concept for anticancer therapy, which improves drug delivery into tumor tissues, resulting in tumor regression.2 Accordingly, it is necessary to establish endothelial cell characteristics in the tumor to monitor the state of the tumor vasculature precisely.

Endothelial cells (ECs) with at least three different phenotypes can be identified during angiogenesis (namely, so-called tip, stalk, and phalanx cells).3 Tip cells possessing filopodia are responsible for determining the direction of migration of newly developing blood vessels. Stalk cells are located behind tip cells and are actively proliferative.4 Phalanx cells emerge in the final step during angiogenesis. These latter cells are covered with mural cells, such as pericytes and smooth muscle cells, and contribute to smooth vessel lumen formation by strong cell-cell adhesion via vascular endothelial—cadherin. In addition, vascular ECs exhibit tissue-specific diversity and structural differences (ie, capillary, artery, and vein).5,6 Therefore, gene expression in ECs may be modified by the microenvironment in each organ.

A previous report indicated that blood vessel ECs in normal organs and tumor tissues differ in their gene

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D.Y. and W.J. contributed equally to this work.

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expression profiles. Particularly, differences in the expression of genes for the cell surface molecules CD276, CD137 (soluble CD137), minK-related peptide 2 (Mrp2), Doppel, protein tyrosine phosphatase receptor type N (Ptprn), CD109, and Aneulysis, the secreted factors Apelin, placenta growth factor (Pigf), and collagen type VIII (Coll VIII) are found, and the intracellular molecules vascular SH2-containing protein (VSCP), phosphatidylinositol-4-phosphate 5-kinase and related FYVE finger-containing proteins signal transduction (Etsv4), and ubiquitin D have been noted. On the other hand, it has been suggested that blood vessels in the tumor microenvironment are more unstable than in normal organs. However, whether there are gene expression patterns characteristic of such unstable blood vessels in tumors has not been established. Therefore, it would be useful to be able to identify newly developing unstable blood vessels in tumors by gene expressing profiling of ECs.

We previously reported that partner of Sld5-1 (PSF1; official name GINS1), a member of the GINS complex composed of PSF1, PSF2, PSF3, and SLDS, is essential for rapid proliferation of cells, such as epiblasts, with stem cell properties in embryos and for hematopoietic stem cell proliferation during recovery from bone marrow ablation by the anticancer drug 5-fluorouracil (5-FU). PSF1 regulates DNA replication as a member of the GINS complex and functions to regulate chromosomal segregation as a single molecule. On the other hand, cells that are not proliferating express little PSF1 protein, and PSF1 promoter activity is silenced in G0/G1. Therefore, in the present study, we asked whether quiescent or proliferating cells can be distinguished by any differences in PSF1 promoter activity and specific gene expression patterns in the tumor vasculature.

Materials and Methods

Mice

C57BL/6 mice and KSN nude mice (7 to 8 weeks of age) were purchased from Japan SLC (Shizuoka, Japan). To generate PSF1 promoter–enhanced green fluorescent protein (EGFP) mice, previously cloned PSF1 promoter was used. pBlue-script KS + PSF1 promoter-EGFP-neo was cleaved by Apal and EcoRI, and the purified product was injected into fertilized eggs to generate transgenic mice. Genotyping to confirm the expression of the transgene was performed by PCR using the following primers: forward: 5'-CACATGGAAGCAGCAGCATGACTTTTCG-3'; reverse: 5'-TGCTCAGTTGATGGTGTGCG-3'. All experiments were performed in accordance with the guidelines of Osaka University Committee for animal and recombinant DNA experiments.

Antibodies

Anti-GFP antibody (Ab) (MBL, Nagoya, Japan; BioLegend, San Diego, CA), anti–Ki-67 Ab (DakoCytomation, Glostrup, Denmark; eBioscience, San Diego, CA), anti–chondroitin sulfate proteoglycan 4 (NG2) Ab (Millipore, Temecula, CA), anti–α-smooth muscle actin–Cy3 Ab (Sigma, St. Louis, MO), anti–F4/80 Ab (AbD Serotec, Kidlington, UK), anti–cytokeratin 5 Ab (Abcam, Cambridge, UK), and anti–CD31, anti–CD45, and anti–Ter119 Abs (BD Pharmingen, Franklin Lakes, NJ) were used as the primary Abs in immunofluorescence. Anti-GFP and anti–α-smooth muscle actin–Cy3 Abs were diluted 1:500. Other Abs were diluted 1:200. Alexa Fluor 488 or Alexa Fluor 546–labeled goat anti-rabbit and anti-rat IgGs (Molecular Probes, Eugene, OR) and Alexa Fluor 647–labeled goat anti-Armenian hamster IgGs (Jackson ImmunoResearch, West Grove, PA) were used as the secondary Abs (dilution, 1:200). Nuclei were stained by Hoechst or TO-PRO-3 (Molecular Probes) (dilution, 1:1000).


Cell Culture

Mouse pancreatic endothelial cell line (MS1; ATCC, Manassas, VA), mouse embryonic fibroblast cell line (NIH3T3), mouse melanoma cell line (B16), and mouse Lewis lung carcinoma cell line (LLC; RIKEN Bioresource Centre, Tsukuba, Japan) were grown in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum and 1% penicillin/streptomycin (Life Technologies, Carlsbad, CA). The LLC-derived highly metastatic line (Ex-3LL; JCRB Cell Bank, Tokyo, Japan) was grown in Hams F10 and L15 (mixed 3:7) medium (Thermo Fisher Scientific, Waltham, MA) supplemented with 10% fetal bovine serum. Human colorectal adenocarcinoma cell line HT29 (ATCC) was cultured in RPMI 1640 medium with 10% fetal bovine serum and 1% penicillin/streptomycin. MS1 cells were used at 1 × 10^3 cells (sparse seeding) or 5 × 10^5 cells (for confluence) in 12-well culture plates. After day 2, MS1 cells were used for analysis. Phosphatidylinositol 3-kinase (PI3K) inhibitor LY294002 (Merck, Kenilworth, NJ) was used at 50 μmol/L concentration and cultured for 24 hours with the cells. LLC-Mock and LLC–mouse angiopoietin-1 were established by overexpressing pEGFP-N1 (Clontech, Mountain View, CA) or mouse angiopoietin-1 plasmids. Stable cells were selected using 1 mg/mL geneticin (Gibco, San Diego, CA).
Aorta Ring Assay

Aortas were isolated from 6- to 8-week-old C57BL/6 mice and cut into 0.5- to 1-mm slices for making aorta rings. Aorta rings were implanted into collagen type I gels (Nitta-gelatin, Osaka, Japan) with 50 ng/mL human VEGF (PeproTech, Rocky Hill, NJ) and cultured for 1 week in HuMedia-EG2 (Kurabo, Osaka, Japan) with 50 ng/mL human VEGF.11

Matrigel Plug Assay

Eight-week-old mice were inoculated with 500 µL growth factor-reduced Matrigel (BD Biosciences, San Jose, CA) with 150 ng/mL human VEGF and 30 U/mL heparin (Sigma-Aldrich, St. Louis, MO). After 1 week, gels were dissected with skin and fixed with 4% paraformaldehyde/phosphate-buffered saline for 4 hours at room temperature. After fixation, gels were substituted by 20% sucrose and mounted with OCT compound (Sakura Finetek Japan, Tokyo, Japan).

Tumor Implantation

LLC, B16, Ex-3LL, LLC-Mock, or LLC—mouse angiopoietin-1 tumor cells (1 × 106 per mouse in 0.1 mL phosphate-buffered saline) were inoculated subcutaneously into wild-type C57BL/6 or PSF1 promoter—EGFP mice (7 to 8 weeks of age). For tumor growth inhibition, on day 7 after s.c. inoculation of B16 cells, mice were treated with i.p. injection of saline or 60 mg/kg body weight 5-FU (Kyowa Hakko Kirin, Tokyo, Japan) every other day. For the detection of the normalization window, a single dose of 5 mg/kg bevacinuzumab was injected into KSN nude mice bearing HT29 tumor once the tumor volume had reached 45 to 55 mm3.

Confocal Laser Scanning Microscopy

Immunostaining was performed as described previously.12 The slides were observed under a Leica TCS SP5 Ver1.6 (Leica Microsystems, Wetzlar, Germany) using HC PL APO CS 20 × 0.7 DRY. Images were processed using Adobe Photoshop CS5 Extended software (Adobe Systems, San Jose, CA).

Flow Cytometric Analysis and Cell Sorting

Cultured cells were detached from the culture dish by Versene (Gibco). Mouse tissues were separated by dispase II, collagenase type I, and collagenase type II, as described previously.13 Red blood cells were removed from mouse tissues using red cell lysis buffer, and cells were reacted with anti-CD16/32 antibody for blocking. Pyronin Y/Hoechst staining was performed as described previously.13 Flow cytometric analysis was performed by FACSCalibur (BD Biosciences), and a FACSARia (BD Biosciences) was used for cell sorting. Data analysis was performed by FlowJo7.6 (FlowJo, LLC, San Carlos, CA).

Quantitative Real-Time PCR

RNA was extracted from cultured cells or sorted cells by using QIAshredder and RNeasy Plus Mini Kit (Qiagen, Hilden, Germany). To generate cDNA, the PrimeScript RT reagent Kit (TaKaRa, Shiga, Japan) was used. Quantitative real-time PCR was performed using the MX3000P Real-Time PCR System (Stratagene, San Diego, CA). Specific primers are used per previous reported sequences or new sequences in this study (Table 1).10,13—24

Statistical Analysis

All experiments were performed in triplicate and repeated at least three times (n = 3). All data are displayed as the means ± SD and were analyzed by repeated-measures two-way analysis of variance or t-tests. P < 0.05 was considered statistically significant.

Results

Generation of Mice Expressing EGFP under the Transcriptional Control of the PSF1 Promoter (PSF1 Promoter—EGFP Mice), and Assessment of EGFP Expression in Embryonic Tissues

Mice were generated in which PSF1 promoter activity can be visualized by EGFP expression, and it was tested whether EGFP can mark proliferating cells reflecting stem or progenitor properties. A total of 5.5 kb of the PSF1 5′-flanking fragment, including the first intron, was used to generate these transgenic mice (Figure 1A).10 EGFP transgene expression was confirmed using specific primers (Figure 1B).

The appearance of embryos under stereoscopic fluorescence microscopy revealed stronger EGFP signals at embryonic day 13.5 in the head of transgenic mice than in wild-type mice (Figure 1C). EGFP expression was observed in the ventricular zone of the brain in sections from these transgenic mice (Figure 1D). A previous report had shown that neural progenitor cells in the ventricular zone during embryogenesis undergo expansion. This study had used a fluorescent ubiquitination-based cell cycle indicator (Fucci) system able to visualize the cell cycle without immunostaining.25 Therefore, coimmunostaining was performed with the cell proliferation marker Ki-67 and EGFP. Most EGFP signals in cells merged with Ki-67 (Figure 1E). Taken together, these results suggest that PSF1 promoter activity can mark proliferating cells in vivo.

PSF1 Promoter—EGFP Positivity Marks Proliferating Cells in Adult Tissues

Next, it was assessed whether PSF1 promoter—EGFP—positive cells are present in adult tissues. Intestine and skin was selected because terminally differentiated cells are constantly being
Visualization of Proliferative ECs

Table 1 Mouse Primer List

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*This work.
F, forward; R, reverse.

The level of PSF1 protein expression and PSF1 promoter activity is correlated with cell growth in stem or progenitor cells. In addition, the immortalized fibroblast cell line NIH3T3 also expressed PSF1 protein.
Therefore, it was assessed whether PSF1 mRNA expression is induced when cells proliferate. Cultured NIH3T3 cells and mouse EC MS1 cells both expressed little PSF1 mRNA at confluence, where the cell cycle is arrested, but they up-regulated PSF1 in sparsely seeded cell cultures (Figure 3A). Using Pyronin Y and Hoechst double staining as a method to analyze the cell cycle (where Pyronin Y binds to RNA and thus signals RNA synthesis), in sparse cultures, the MS1 line contained both Pyronin Y-positive and Pyronin Y-negative cells (Figure 3B). Similar to the expression of PSF1 as influenced by cell density, Pyronin Y-positive cells expressed higher levels of PSF1 mRNA than Pyronin Y-negative cells (Figure 3C). Culture to confluence increased the number of Pyronin Y-negative cells, but some Pyronin Y-positive cells were still present (Figure 3B). In confluent cultures, Pyronin Y-positive cells also expressed higher levels of PSF1 mRNA than their negative counterparts (Figure 3C).

Next, the PSF1 promoter activity of vascular ECs was studied in an *ex vivo* model of angiogenesis using the aorta ring assay. On collagen type I gels, ECs in neovessels developing from the aorta ring of PSF1 promoter–EGFP mice expressed EGFP (Figure 3D). Matrigel plug assay was also used; when Matrigels containing VEGF and heparin were injected into PSF1 promoter–EGFP mice subcutaneously, CD31-positive neovessels derived from the skin vasculature were EGFP positive. This was seen despite the fact that CD31-negative EGFP-positive cells, possibly fibroblasts, also expressed EGFP (Figure 3E).
addition, it was investigated whether proliferating EGFP\(^+\) ECs can be detected \textit{in vivo} using these PSF1 promoter–EGFP mice. First, brain tissues were stained on embryonic day 13.5. Some of the CD31\(^+\) vessels were stained for EGFP (Supplemental Figure S2A). Moreover, EGFP\(^+\) ECs were detected in neonatal retina (Supplemental Figure S2, B and C). Taken together, PSF1 promoter activity is up-regulated in ECs when angiogenesis is taking place, whereas dormant ECs in preexisting blood vessels express little or no PSF1.
Visualization of Endothelial PSF1 Promoter Activity in Tumor Angiogenesis

During tumor growth, neovascularization is induced by the sprouting and extension of ECs from the preexisting vasculature to supply nutrients and oxygen through new vessels. As previously described, preexisting blood vessels express little PSF1 promoter–driven EGFP (Figure 2, E and F). Proliferating ECs in the tumor derived from preexisting skin vasculature were visualized. B16 mouse melanoma cells were inoculated subcutaneously into mice, and after 3 days, tumor tissue was harvested together with skin. Interestingly, part of the skin vasculature near the tumor had become EGFP positive (Figure 4A). Moreover, EGFP-positive ECs extending from the skin vasculature toward the tumor cells were detected (Figure 4B). At a little far from tumor area, there are some EGFP-positive cells that seem to be hematopoietic cells and ECs. For the initiation of sprouting angiogenesis, many
monocyte lineage cells and hematopoietic progenitor cells are recruited from preexisting blood vessels to the tumor; these cells have the ability to proliferate. Therefore, one possibility is that EGFP-positive and PSF1-activated hematopoietic cells localize at a short distance from the tumor. Moreover, hematopoietic cells secrete angiogenic cytokines, such as VEGF and platelet-derived growth factor, which would affect endothelial cell growth in the locality.

Figure 4 Initiation of tumor angiogenesis up-regulates PSF1 promoter activity in endothelial cells (ECs). A: Skin vasculature 3 days after B16 melanoma cell injection into the mouse. White dashed line indicates localization of tumor. Arrows indicate EGFP-positive ECs around the tumor. Two pieces of images were put together and edited to one piece. B: Sprouting new capillary from preexisting blood vessels around the tumor. White dashed line indicates localization of tumor. Arrows indicate budding ECs that express EGFP and CD31 (red). C: Flow cytometric analysis of endothelial PSF1 promoter activity in skin tissues around tumors 3 days after tumor cell inoculation. PSF1 promoter activity in ECs is increased in the skin vasculature, with B16 tumor inoculation (orange) relative to skin from wild type (WT; red) or transgenic (Tg; blue) mice in the steady state. D: PSF1 promoter activity of tumor vasculature 7 days after tumor cell inoculation. Section was stained with anti-CD31 antibody (red). E: Immunostaining of hematopoietic (leukocyte, CD45⁺; macrophage, F4/80⁺; erythrocyte, Ter119⁺) and α-smooth muscle actin (α-SMA)⁺ smooth muscle cells (red) in the skin of Tg mice around the tumor tissue 3 days after tumor cell inoculation. Scale bars: 100 μm (A); 75 μm (B); 250 μm (D); 50 μm (E). Max, maximum.
Figure 5  Proliferative capacity and cell cycle analysis of tumor endothelial cells (ECs) monitored by PSF1 promoter activity. A: Growth inhibition of EGFP<sup>high</sup> ECs by 5-fluorouracil (5-FU) analyzed by flow cytometry. B: Flow cytometric analysis of Ki-67 positivity in EGFP<sup>high</sup> and EGFP<sup>−</sup> ECs from tumors generated by s.c. inoculation of B16 melanoma cells. C: Detection of extracellular signal-regulated kinase (ERK) phosphorylation in tumor ECs from transgenic (Tg) mice. EGFP<sup>high</sup> ECs showed higher ERK phosphorylation (pERK) levels compared with EGFP<sup>−</sup> ECs. D: Profiling of cell cycle arrest-related gene expression in EGFP<sup>high</sup> ECs from B16 tumors. The level of mRNA expression in EGFP<sup>−</sup> ECs from tumor was set as unity. E: Hoechst and Pyronin Y staining of ECs from the skin or B16 tumor of wild-type (WT) mice. Black boxed areas indicate Pyronin Y<sup>−</sup> or Pyronin Y<sup>+</sup> ECs within Hoechst-low cells, respectively. F: Forward and side scatter (FSC and SSC, respectively) plots of B16 tumor EC fraction divided by Pyronin positivity, as indicated in E. G: Forward and side scatter plots of EGFP-negative or highly positive B16 tumor EC fractions in Tg mice. EGFP positivity was determined as indicated by red or blue bars in the left panel. Middle and right panels indicate EGFP negative or high positive ECs. Data are expressed as means ± SD (D), n = 3 (D). *P < 0.05, **P < 0.01 EGFP negative versus high positive ECs.
To confirm that cells double labelled by CD31 and EGFP were ECs, flow cytometry was used to show that CD31⁺ CD45⁻ gated ECs in tumors express a higher level of EGFP than those in the skin in the steady state (Figure 4C). Seven days after tumor cell inoculation, most CD31⁺ ECs in the tumor expressed EGFP (Figure 4D). Part of the round cell population expressing the leukocyte marker CD45 or the monocyte marker F4/80 expressed EGFP (Figure 4E). However, Ter119-positive erythrocytes and α-smooth muscle actin—positive mesenchymal cells expressed little EGFP (Figure 4E). These results suggest that CD31⁺ ECs and some hematopoietic cells expressed EGFP in the skin near the tumor. Similar results were also obtained using the mouse Lewis lung carcinoma cell line LLC as the s.c. tumor instead of B16 cells (Supplemental Figure S3, A and B). One difference relative to the B16 tumor model was that LLC tumors contained abundant EGFP-positive hematopoietic cells (Supplemental Figure S3, C–E), suggesting that visualization of EGFP⁺ ECs in the tumor microenvironment is more readily achieved in the B16 model.
PSF1 Promoter—Active ECs in Tumors Have High Proliferative Activity

PSF1 promoter activity in ECs is up-regulated in the tumor. The correlation of PSF1 promoter activity with cell growth in tumor ECs was studied next. First, the DNA synthesis inhibitor 5-FU was used during tumor development. As expected, 5-FU treatment inhibited the growth of ECs with high PSF1 promoter activity (EGFPhighECs) (Figure 5A). The relevance of expression of the cell growth marker Ki-67 and the status of extracellular signal–regulated kinase phosphorylation in such EGFPhighECs was also studied. EGFP- or EGFPhighECs were isolated from the tumor, and it was found that the latter expressed higher levels of Ki-67 and contained more phosphorylated extracellular signal–regulated kinase than EGFP-ECs (Figure 5, B and C). These data strongly suggest that PSF1 promoter activity correlates well with cell growth.

Next, the relationship between the expression of cell cycle arrest genes (p15, p16, p18, p19, p21, p27, p53, and...
p57) and PSF1 promoter activity was examined. This analysis revealed that most cell cycle arrest gene expression was lower in EGFP<sup>high+</sup> ECs than in EGFP<sup>−</sup> ECs (Figure 5D), suggesting that EGFP<sup>high+</sup> ECs enter cell cycle progression. It is well known that migrating cells and dividing cells have a specific different morphology. Most obviously, proliferating cells contain many intracellular compartments necessary for the active synthesis of RNA and DNA. Therefore, EC morphology was characterized in cells with or without PSF1 promoter activity. Using both the Pyronin Y and Hoechst cell staining methods, it was found that ECs in normal skin synthesized little RNA, whereas there is vigorous RNA synthesis in tumor ECs (Figure 5E). Next, the morphological complexity of the intracellular compartment was studied by flow cytometry. Interestingly, those ECs with higher capacity RNA synthesis were larger (as determined by forward scatter) and had greater intracellular complexity (as determined by side scatter) than cells with a low RNA synthesis capacity (Figure 5F). Similarly, EGFP<sup>high+</sup> ECs in the tumor were larger and had greater intracellular complexity than EGFP<sup>−</sup> ECs (Figure 5G). These data indicate that PSF1 promoter activity in tumor ECs correlates with cell growth, suggesting that PSF1 promoter—EGFP mice can be used to visualize proliferating ECs by their EGFP expression.

Classification of Quiescent or Proliferating ECs in the Tumor as Monitored by PSF1 Promoter Activity

PSF1 promoter activity can be used to visualize proliferating ECs in tumors in vivo. To determine specific gene expression in cells differing by the intensity of their PSF1 promoter activity, tumor ECs were sorted and the expression of genes related to vascular formation was assessed using real-time activity. Tumor ECs were sorted and the expression of genes in cells differing by the intensity of their PSF1 promoter activity was assessed using real-time activity. VEGFR1 signaling induces vascular stabilization. In this case, PSF1 promoter activity in tumor ECs correlates with cell growth, suggesting that PSF1 promoter—EGFP mice can be used to visualize proliferating ECs by their EGFP expression.

Vascular Stabilization Induces Endothelial CD109 Expression

Nonproliferative ECs (EGFP<sup>−</sup> ECs) strongly expressed VEGFR1 and CD109 (Figure 6). We hypothesized that VEGFR1 and CD109 expression was induced by vascular stabilization in the tumor microenvironment. To test this, two models of vascular stabilization using 5-FU or the anti-VEGF monoclonal antibody bevacizumab were used. 5-FU was used in the same way as in Figure 5A. As the number of EGFP<sup>−</sup> ECs increased, the fraction of CD109-positive cells also significantly increased (Figure 7A). Although bevacizumab inhibits VEGF-dependent endothelial cell growth and migration, it can also induce vessel maturation, depending on time of exposure. To evaluate whether increased CD109 expression occurred in the so-called normalization window, a previously validated method was used. Bevacizumab was injected once into human colorectal adenocarcinoma HT29-bearing mice. With time, vessel density decreased, but on day 5 after bevacizumab injection, pericyte-covered mature vessels were detected. At this time, functional vessels increased and hypoxic regions significantly decreased. However, on day 8 after bevacizumab injection, hypoxic regions increased again. Because of this, we had previously concluded that normalization of tumor vessels was induced from days 3 to 5 after bevacizumab injection in our model. Accordingly, as
expected, enhanced CD109 expression was detected on days 3 and 5 after bevacizumab injection (Figure 7, B and C).

Next, in vitro culture experiments were performed because it is known that cell density affects cell proliferation. Sparse cell seeding induces proliferation and migration, whereas cell-cell contact in confluent cultures suppresses proliferation and migration and induces cell cycle arrest. Interestingly, MS1 cells express little CD109 in sparsely seeded cultures, but it was induced at confluence (Figure 7D).

Previously, we have shown that angiopoietin-1 (Ang1)/Tie2 signaling enhances EC-EC cell adhesion, resulting in structurally stable blood vessels in which mural cells cover the ECs. In addition, we previously reported that overexpression of Ang1 in tumor cells induced intratumoral blood vessel maturation, whereby mural cells covered ECs and inhibited tumor growth more than in tumors derived by inoculation with parental tumor cells. Herein, the same Ang1-overexpressing LLC tumor model was used. In these tumors, the number of CD109-positive ECs was approximately twice that seen in parental LLC tumors (Figure 7E). PI3K-Akt signaling is a major vascular stabilization pathway under Tie2 activation with Ang1. Inclusion of the PI3K inhibitor LY294002 in MS1 cultures suppressed CD109 induction, even in confluent cells (Figure 7F). These results suggest that CD109 expression in tumor ECs was induced in parallel with vascular stabilization via the PI3K-Akt pathway.

It has been suggested that pericycle coverage is involved in the process of vascular maturation and that ECs are rendered dormant after pericycle coverage. Therefore, finally, the relationship between pericycle coverage and CD109 expression and PSF1 promoter activity was analyzed. Endothelial CD109 expression was tested by immunostaining, but was found to be technically challenging. Instead, the relationship between PSF1 promoter activity and pericycle coverage was assessed. The results showed that some EGFP-negative/low vessels are present in pericycle-covered vessels but others are in vessels that are not covered (Supplemental Figure S6). Moreover, some highly EGFP-positive vessels were covered with pericytes (Supplemental Figure S6). Therefore, these findings suggest that pericycle coverage is, in fact, not correlated with tumor vasculature dormancy.

Discussion

Normalization of the tumor vasculature is an effective method for improving drug delivery into tumors. However, a major clinical problem with this treatment is how to recognize the vascular normalization window for optimal timing of drug delivery. To approach this issue, the identification of specific markers would be extremely useful. In the present study, we distinguish between proangiogenic proliferating ECs and quiescent ECs in tumors on the basis of their PSF1 promoter activity. CD109 was highly expressed in dormant ECs and could be used to detect normalized blood vessels, thus allowing identification of the window of opportunity for optimal delivery of chemotherapeutics.

Several methods for detecting proliferating cells can be applied, such as bromodeoxyuridine incorporation, proliferating cell nuclear antigen staining, Ki-67 staining, or Pyronin Y/Hoechst staining. These methods take time to detect proliferating cells, and only bromodeoxyuridine and Pyronin Y/Hoechst staining can be used to detect cell cycling in living cells. Moreover, it is unclear whether any of these methods reflects the genuine in vivo condition of living cells. To overcome this potential difficulty, the Fucci system has been developed. However, the Fucci system can identify G1, S, G2, and M phases but cannot distinguish between G1 and G0.

Our system monitoring PSF1 promoter activity resolves this problem. PSF1 promoter–EGFP ECs had lower levels of Ki-67 positivity than PSF1 promoter–EGFP ECs and were dormant in the tumor (Figure 5). Therefore, PSF1 promoter activity is especially suitable for detection of cell cycle status in living dormant ECs.

VEGFR1 is known as a VEGF decoy receptor and is involved in antiangiogenesis. In addition, sVEGFR1 regulates sprouting at the angiogenic front in a concentration-dependent manner. The consensus is that VEGFR1 functions as a decoy receptor by trapping VEGF and, thus, regulates angiogenesis. Consistent with this, inhibition of PlGF/VEGFR1 signaling suppressed tumor growth without affecting healthy vessels. Interestingly, in our model, VEGFR1 expression is down-regulated in PSF1 promoter–EGFP ECs. These data suggest that PSF1 promoter–EGFP ECs may be highly sensitive to VEGF stimulation.

It is widely accepted that ECs proliferating during angiogenesis are stalk cells expressing specific markers. Previous reports have shown that the transforming growth factor family cell surface receptor CD105 (endoglin) and the transcription factor SRY-related HMG-box transcription factor (Sox)17 are proangiogenic endothelial markers. It has been reported that more ECs with high levels of Sox17 and VEGFR2 are in S/G2/M phase in tumors, relative to Sox17-low ECs. In our experiments, PSF1 promoter activity did not correlate with the amount of VEGFR2 expressed by the cells. This suggests that Sox17 is expressed by ECs at the onset of sprouting angiogenesis and does not distinguish between stalk and phalanx cells.

Interactions between tip and stalk cells during sprouting angiogenesis have been well studied at the molecular level, showing that Notch1-Dll4 and Jagged1 expression and regulation is involved in this process. It has been reported that disruption of Notch signaling induces excessive generation of tip cells. However, in this study using PSF1 promoter activity, the level of expression of these genes was only marginally changed, suggesting that the
EGFP-negative EC population contains not only phalanx cells but also nonproliferative stalk cells. However, further analysis is required to determine how to distinguish between tip, stalk, and phalanx cells, according to their proliferative status.

Seaman et al. have reported that tumor ECs express 13 specific genes, of which CD109, CD137, and CD276 were tested at the protein level and only CD109 was identified as different, depending on PSF1 promoter activity in tumor ECs. Tumor-specific gene expression was expected in proliferating ECs, but CD109 was also present on non-proliferating ECs.

Previous reports have shown that the ectodomain of CD109 is cleaved, and that soluble CD109 inhibits transforming growth factor-β signaling through direct binding to transforming growth factor-β. It is possible that inhibition of transforming growth factor-β signaling promotes tumor vascular formation because vascular maturation is perturbed. This suggests that intact CD109 expression on the cell surface reflects endothelial maturation and dormancy.

Ang1 overexpression in tumors induced CD109 and the PI3K signaling pathway is involved in this process, strongly suggesting that CD109 is a vascular stabilization marker. Although Nolan et al. proposed that ECs are heterogeneous, depending on the organ of origin, it is likely that CD109 expression is a common phenotype, not depending on physiological or pathologic conditions.

New therapeutic strategies to improve drug delivery for more effective antitumor therapy are highly desirable. As previously alluded to, one approach to this is the induction of vascular stabilization in the tumor. This would reduce hypoxia in the tumor and inhibit progression of tumor malignancy. Therefore, monitoring CD109 expression in tumor vessels may be an effective screening method to recognize the normalization window and determine optimal timing for treatment with anticancer drugs.

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Supplemental Data

Supplemental material for this article can be found at https://doi.org/10.1016/j.ajpath.2018.01.015.

References

22. Franco CA, Mersckay M, Parla...


