Platelets: Guardians of Tumor Vasculature

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Abstract

Solid tumors generate a prothrombotic environment capable of platelet activation. Recent findings indicate that the activated platelets are crucial regulators of tumor vascular homeostasis in that they prevent tumor hemorrhage. Surprisingly, this effect is independent of platelets’ capacity to form thrombi and instead relies on the secretion of their granule content. Thus, targeting platelet secretory activities may represent a new approach to specifically destabilize tumor vasculature. [Cancer Res 2009;69(14):5623–6]

Tumor-Platelet Cross-Talk

Since the first description by Trousseau (1), an association between cancer and thrombosis has been recognized. Tumor cells may express tissue factor, release procoagulant microparticles, and secrete cytokines rendering endothelium prothrombotic (2). Moreover, sluggish blood flow and contorted vessels are features of the tumor microcirculation that contribute to the tumor thrombogenic environment (3). Thus, tumors are capable of activating platelets, and it has been suggested that platelets in turn promote tumor growth (4, 5).

Clinical and experimental evidence indicates that platelets play a role in the spread of cancer. Both thrombocytopenia and antiplatelet treatments reduce the number of experimental metastases (6, 7). Coating circulating cancer cells with platelets protects them from the immune response (8) and enhances their extravasation by increasing their adhesion to the vessel wall (7). Whereas interactions of platelets with circulating cancer cells are well documented, the actual contribution of platelets to solid tumor homeostasis prompts further analysis.

Platelets are a rich source of pro- and antiangiogenic factors that can be differentially released upon platelet activation (9). Thus platelet activation could regulate angiogenesis, a process essential in cancer progression. Furthermore, platelets support the integrity of angiogenic (10) and inflamed microvessels (11) through the prevention of hemorrhage at sites of angiogenesis and inflammation. Because angiogenesis and leukocyte infiltration are both involved in the development of solid tumors, it is conceivable that platelets may have an impact on tumor vessel integrity and homeostasis. Indeed, recently platelets have been shown to continuously prevent tumor hemorrhage by releasing their granules (12). In this review, we summarize these findings and discuss their therapeutic potential.

Platelets and Tumor Vascular Homeostasis

The role of platelets in solid tumors was addressed by inducing acute thrombocytopenia with a platelet-depleting antibody in mice bearing subcutaneous tumors or established lung metastasis (12). Thrombocytopenia consistently resulted in severe bleeding that was specific to the tumor site and not seen elsewhere in the animal. The tumors from thrombocytopenic mice were characterized by massive hemorrhage at the tumor-stroma interface, reduced tumor cell proliferation, and increased apoptosis in the hemorrhagic areas. The profound effect of platelet depletion on tumors, independent of the tumor location and age, underlined the severity of the vessel damage. The vascular breaks occurred very rapidly as bleeding became noticeable as early as 30 minutes after induction of thrombocytopenia. The continuous need for platelets to prevent tumor bleeding and consequent cell death reveals the crucial contribution of platelets to tumor vascular homeostasis. Thus, apart from their supportive role in tumor metastasis, platelets also have to be considered as key regulators of tumor vessel stability.

Prevention of tumor bleeding by platelets displays surprising mechanistic features. In contrast to platelets’ role in primary hemostasis, firm platelet adhesion is not required for maintaining vascular integrity in tumors. Intravitral microscopy using fluorescently labeled platelets or immunostaining of platelets in tumor sections did not reveal the presence of thrombi in the tumor microcirculation (12). Previously, Manegold and colleagues reported platelet rolling but no adherence on tumor microvessels (13). The lack of adherent platelets led to the hypothesis that the mechanisms involved in the prevention of tumor hemorrhage by platelets differ from those required for platelet plug formation. This was confirmed genetically in experiments using mouse models of bleeding disorders due to defective platelet adhesion. These mice bleed profusely when injured but did not bleed at tumor sites. Thrombus formation relies on platelet adhesion through glycoprotein Ib alpha (GPIbα) to von Willebrand factor (VWF) and the activation of the integrin αIIbβ3 leading to platelet aggregation. VWF−/−, GPIbα inhibitor-treated mice, and mice with defective activation of integrins (CalDAG-GEFI−/−) showed no spontaneous intratumor hemorrhage (12). Moreover, transfusion of platelets mutant in GPIbα prevented thrombocytopenia-induced tumor bleeding. Interestingly, the ability of these platelets to prevent tumor bleeding was lost when platelet degranulation was induced before infusion. Thus, prevention of tumor hemorrhage by platelets most likely relies on the positive effects of the granular components. Activated platelets also shed several plasma membrane receptors and procoagulant microparticles that could contribute to the biological activities platelets exert on tumor vessels. Degranulated platelets characterized by low levels of ADP and serotonin have been found in patients with malignant solid tumors, even in the absence of active consumption coagulopathy, showing the capacity of the tumors to induce platelet secretion (14).
How Do Platelets Stabilize Tumor Vessels?

The platelet-derived factors involved in the prevention of tumor hemorrhage have not been identified. Additionally, the cell type that platelets influence most is not known and several types of cells could be affected. The soluble factors released by platelets may regulate the endothelial stability of the angiogenic tumor vessels and/or prevent vascular damages induced by the tumor cells or inflammatory cells associated with tumors.

Platelets have long been known to enhance endothelial barrier function and help preserve endothelium during organ reperfusion (15–17). Kitchens and Weis (18) reported thinning of endothelium in microvessels within 6 hours of platelet depletion in rabbits and an increase in vessel permeability. But platelet depletion was not sufficient to induce significant hemorrhage such as was observed in tumor vessels of the thrombocytopenic mice (12). Thus, tumor vessels have a greater need for the mediators released by platelets than the general microcirculation of the animal.

Several classes of molecules released by platelets were shown to reduce vascular permeability. The best known are angiopoietin-1 (ang-1) (refs. 12, 19), serotonin (5-HT) (ref. 20), and sphingosin-1-P (S1P) (ref. 21). Absence of the platelet-released factors was proposed to cause extravasation of red blood cells from postcapillary venules in thrombocytopenic patients (19, 22). In contrast to normal vessel wall, tumors abundantly express vascular endothelial growth factor (VEGF), also known as vascular permeability factor. As the name implies, it has a potent ability to increase vascular permeability. Therefore, in the tumors, platelets may have to counteract the effects of VEGF. Indeed, thrombocytopenia dramatically impairs the balance between pro- and antipermeability factors in tumor-bearing mice, in particular depleting blood of angiopoietin-1 and serotonin (12). Of note however, extravasation of red blood cells does not occur only through the endothelium but also through the basement membrane, which implies degradation or rupture of the basal lamina surrounding vasculature. It is likely that the massive extravasation of red blood cells observed in thrombocytopenic tumors involves factors capable of damaging basement membranes and/or abnormalities of the basement membrane in tumor vessels. Profound structural abnormalities of basement membrane in tumors, including broad extensions away from the vessel wall, were noted (23).

In addition to the beneficial effects platelets have on resting endothelium, platelets could diminish injuries produced by inflammation. Areas of inflammation are also unusually susceptible to hemorrhage in thrombocytopenia, both in the skin and in internal organs (11). Solid tumors show signs of chronic inflammation. These include the presence of infiltrating leukocytes, the expression of
cytokines such as tumor necrosis factor-α (TNF-α) or interleukin-1, chemokines, and active tissue remodeling with fibrin deposition, all potentially affecting vascular integrity (24). In addition, leukocytes secrete injurious products such as matrix metalloproteases (MMPs), serine proteases, and reactive oxygen species (ROS). Platelets could modulate the activity of leukocytes and also neutralize some of their harmful products through release of serpins, tissue inhibitors of metalloproteinases (TIMPs), and ROS scavengers from their granules (25–27). Therefore, the need for platelets to prevent tumor hemorrhage may involve not only their positive effects on endothelium (18) and role in angiogenesis (10) but also their ability to tighten endothelial junctions (15, 16, 19, 21) and to prevent potential vascular injuries by tumor-infiltrating leukocytes (Fig. 1) (ref. 11).

The fact that genetic inhibition of platelet adhesion does not affect their capacity to stabilize tumor vessels raises questions about how platelets deliver active compounds to tumor vessels. As previously hypothesized by Folkman and colleagues (5), platelet-tumor interactions might be facilitated by impaired blood flow and localized granular release by the procoagulant environment of the tumor. Platelets have two different types of granules, dense granules where serotonin is located and alpha granules containing protein components including angiogenic factors, growth factors, and immunomodulatory cytokines. The alpha granules further differ in their composition and can be differentially secreted by platelets depending on stimulus (9, 28) as is the case for endothelial storage granules (29). Thus, there could be a very fine regulation of expression of angiogenic, inflammatory, and vascular stability modulators by granular secretion within the tumor.

In parallel to studies reporting cancer-associated platelet activation, enrichment in intraplatelet angiogenic factors has also been described in cancer patients. Platelet lysate from breast cancer patients was noted to contain higher VEGF and angioptin-1 levels than from healthy subjects (30). Thus, it could be that solid tumors not only stimulate platelet granule secretion, but also load platelets with proangiogenic factors that stimulate tumor progression.

Therapeutic Implications for the Role of Platelets in Maintenance of Tumor Vessel Integrity

One important aspect of recent findings about platelet-tumor interactions is that platelet-mediated prevention of tumor vessel damage does not require platelet plug formation. The mechanistic dichotomy of platelet-dependent primary hemostasis and maintenance of tumor vessel integrity may allow specific pharmacological targeting of the two processes. Inhibitors of the platelet-derived soluble factors responsible for protection of tumor vessels may destabilize the tumor vasculature without affecting primary hemostasis and increasing the risk of bleeding outside the tumor. On the other hand, blockage of platelet activation that is involved in both integrin activation and degradation would probably alter both primary hemostasis and tumor vessel stability.

The identification of the specific vascular protective factor(s) delivered by platelets to solid tumors or discovery of inhibitors preventing their secretion could open new therapeutic perspectives. Targeting of these molecules or their secretion could selectively destabilize tumor vasculature and, thus, affect tumor cell viability. It is conceivable that destabilization of the tumor vessels might have additional beneficial consequences, such as promoting antitumor immunity through better exposure of tumor antigens to circulating immune cells, improving the delivery of chemotherapeutic agents, even larger nanoparticles, to solid tumors. Alternatively, mimicking the stabilizing effect of platelets on tumor vessels by providing the stabilizing factors might promote normalization of the tumor vasculature and its function. Normalizing tumor vessels is a strategy that was proposed to increase the efficiency of oxygen and drug delivery to solid tumors and that was shown to have a synergistic effect with cytotoxic therapy (31).

In conclusion, interference in platelet-tumor vessel cross-talk represents an interesting and challenging approach for manipulating tumor vasculature in order to improve anticancer therapies. New insights could be provided by the identification of the beneficial factors that platelets deliver to the tumor vasculature and of the cell type(s) platelets regulate to prevent tumor hemorrhage.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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