Thrombospondin 1 and Vasoactive Agents Indirectly Alter Tumor Blood Flow\textsuperscript{1,2}

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Abstract

Nitric oxide (NO) plays important physiological roles in the vasculature to regulate angiogenesis, blood flow, and hemostasis. In solid tumors, NO is generally acknowledged to mediate angiogenic responses to several growth factors. This contrasts with conflicting evidence that NO can acutely increase tumor perfusion through local vasodilation or diminish perfusion by preferential relaxation of peripheral vascular beds outside the tumor. Because thrombospondin 1 (TSP1) is an important physiological antagonist of NO in vascular cells, we examined whether, in addition to inhibiting tumor angiogenesis, TSP1 can acutely regulate tumor blood flow. We assessed this activity of TSP1 in the context of perfusion responses to NO as a vasodilator and epinephrine as a vasoconstrictor. Nitric oxide treatment of wild type and TSP1 null mice decreased perfusion of a syngeneic melanoma, whereas epinephrine transiently increased tumor perfusion. Acute vasoactive responses were also independent of the level of tumor-expressed TSP1 in a melanoma xenograft, but recovery of basal perfusion was modulated by TSP1 expression. In contrast, overexpression of truncated TSP1 lacking part of its CD47 binding domain lacked this modulating activity. These data indicate that TSP1 primarily regulates long-term vascular responses in tumors, in part, because the tumor vasculature has a limited capacity to acutely respond to vasoactive agents.


Introduction

Solid tumor growth requires the generation of new blood vessels from an established vascular network, a process termed angiogenesis. The newly formed vascular network provides necessary nutrients and oxygen to meet the increased metabolic demand of the growing tumor. Tumor angiogenesis is an intensely pursued therapeutic target for treating cancers. Although several antiangiogenic drugs have attained US Food and Drug Administration approval and significantly delay tumor progression, hypertension is a frequent adverse effect of these agents [1]. If arteries supplying the tumor are selectively constricted, this adverse effect could contribute to the efficacy of antiangiogenic drugs. If instead these drugs induce systemic hypertension, the resulting increased tumor perfusion could limit their efficacy [2,3].

Although long-term control of angiogenesis can clearly limit tumor growth, acute regulation of tumor blood flow and dynamics may be more difficult. The intratumor vasculature is poorly organized, and it displays variability in flow and tissue perfusion. Intratumoral vessels typically lack anatomic elements required for normal vasoactive responses including autonomic neural elements [4]. Additionally, some

Abbreviations: BAEC, bovine aortic endothelial cell; BOLD, blood oxygen level--dependent; DEA/NO, diethylamine NONOate; DETA/NO, diethyltriamine NONOate; MSME, multislice multiecho; sGC, soluble guanylyl cyclase; TSP1, thrombospondin 1; VSMC, vascular smooth muscle cell

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tumors lack lymphatic vessels, which combined with increased vascular permeability increases interstitial pressure and decreases tumor perfusion [5]. Increased tumor interstitial pressure encourages vascular and lymphatic collapse, stagnation, and reversal of arterial flow [5,6]. Consequently, tumor blood flow is very heterogeneous, with areas of significant under perfusion and hypoxia [7]. Additionally, tumor vessels display dynamic changes that are specific for the growth stage of the tumor [8]. Such irregularity in flow confounds effective delivery of therapeutic agents to tumor tissue [9,10]. In contrast, adjacent blood vessels that feed tumor vessels do contain normal vascular anatomy and should respond to vasoactive challenges in a predictable fashion. Limiting tumor perfusion with agents that can predictably enhance or retard tumor blood flow represents a potential therapeutic avenue [11,12]. An alternative strategy to acutely control flow using agents that cause rapid tumor vascular thrombosis and obliteration has also shown promise [13].

Nitric oxide (NO) is an important signaling molecule for acute regulation of blood flow and for long-term angiogenic responses. It is constantly produced by vascular cells, and it relaxes the arterial component of blood vessels [14]. The direct effect of vessel relaxation is to increase blood flow and tissue perfusion. Chronic exposure to NO drives the formation of new blood vessels by stimulating both endothelial and vascular smooth muscle cell (VSMC) proliferation and migration as part of a physiologic response to normal wound healing and during tumor growth and metastasis [15,16]. Furthermore, endogenous NO can be limiting for tumor blood flow [17]. Several studies have reported that inhibition of NO synthase using either nitro-L-arginine methyl ester or N^e-nitro-L-arginine results in decreased total tumor blood flow in animal models [18–22] or has inhibitory effects on tumor growth [23]. These reports are supported by studies demonstrating that tumor blood flow is increased by exogenous NO donors [24] (and reviewed in Refs. [25–27]). Other studies, however, found that NO can decrease tumor blood flow in animal models, which has been attributed to a steal effect [28].

Consistent with these conflicting studies in rodents, attempts to demonstrate therapeutic responses through manipulation of tumor blood flow have been modest in people [23,29–31]. Nonspecific inhibition of endogenous NO production in a recent phase I trial in patients with several tumor types documented immediate decreases in tumor blood volume using contrast-enhanced computed tomography [30]. In some cases, tumor blood volume remained decreased for 24 hours. In contrast, treatment of patients with hepatic tumors using a continuous infusion of epinephrine increased tumor blood flow in 66% of cases [32].

The matricellular protein thrombospondin 1 (TSP1) was the first identified endogenous angiogenesis inhibitor [33,34]. Its expression is frequently lost in some types of human tumors. Conversely, overexpression of TSP1 in experimental tumors dramatically decreases tumor growth rate, neovascularization, and metastasis [35]. Thrombospondin 1 inhibits NO signaling in both endothelial and VSMC by inhibiting activation of soluble guanylyl cyclase (sGC) [15,36]. In VSMC and in vivo, TSP1 limits vasorelaxation by NO [37]. Under an ischemic stress, blocking TSP1 acutely increases tissue blood flow and perfusion [37]. The ability of TSP1 to abrogate NO-stimulated angiogenesis and blood flow changes requires the cell surface receptor CD47 [36]. Targeting this receptor mimics the inhibitory actions of TSP1 on NO-driven vascular responses [38].

Given the pivotal role that TSP1 plays in controlling NO-driven vascular responses and blood flow in nontumor vascular networks, we hypothesized that, in addition to its antiangiogenic activity, TSP1 may acutely regulate flow in the peritumoral or tumor vasculature. A recent histochemical analysis of pancreatic tumors in mice treated with recombinant TSP1 type 1 repeats inferred that this treatment decreased tumor blood flow [39], but a direct assessment of flow was not performed. Here, we approach this question by assessing the role of host and tumor TSP1 expression in regulating acute tumor blood flow responses. We present evidence that vasoactive agents such as NO and epinephrine primarily alter tumor perfusion in an indirect manner. Thrombospondin 1 expressed in the tumor vasculature does not alter such blood flow responses to vasoactive stimuli. In contrast, overexpression of TSP1 by tumor cells moderately decreases tumor blood flow responses to NO and epinephrine. These results provide important insights for evaluating efforts to directly alter tumor blood flow using vasoactive agents and for maximizing the efficacy of angiogenesis inhibitors.

### Materials and Methods

#### Animals

Wild type and TSP1 null C57BL/6 mice were housed in a pathogen-free environment and had ad libitum access to standard rat chow and water. Cr(Cnr) athymic nude Fbf female mice were obtained from Charles River (Tacomin, MD). Care and handling of animals comply with standards established by the Animal Care and Use Committee of the National Cancer Institute.

#### Reagents and Cells

The murine B16F10 melanoma cell line was kindly provided by Dr. Lyuba Varticovski. Cells were expanded in standard growth medium (RPMI with 10% FCS; Gibco, Grand Island, NY) and frozen at uniform passage for injection in mice. The human melanoma cell line, MDA-MB-435, was previously transfected with a CMV-THBS1 expression plasmid, and TSP1 expression was quantified as described [40]. The parental cell line, a TSP1-overexpressing clone (TH26) and a clone expressing a C-terminal truncation of TSP1 (TH50) were used. Cells were MAP tested and expanded in complete DMEM containing 10% fetal bovine serum and 750 µg/ml G418 (Gibco-BRL, Gaithersburg, MD) and frozen for future use. Porcine femoral artery VSMC were harvested from Yucatan white hairless miniature pigs and cultured in SM-GM +2.5% FCS (Lonza, Belgium). Bovine aortic endothelial cells (BAECs) were prepared from fresh aortic segments and grown in DMEM containing 10% fetal bovine serum, penicillin G (100 units/ml), and streptomycin (100 µg/ml).

#### Blood Oxygen Level–Dependent Magnetic Resonance Imaging

Magnetic resonance imaging (MRI) scans were acquired using a 4.7-T scanner (Bruker BioSpin, Karlsruhe, Germany) and isoflurane anesthesia. Muscle and tumor tissue was scanned at rest, so alterations in oxygenation reflected changes in perfusion rather than in oxygen consumption. Magnetic resonance measurements were started after the mouse’s body temperature reached 37°C. Before the experiments, a multislice multiecho (MSME) sequence was used to determine the target slice location. A series of T2*-weighted gradient echo blood oxygenation level–dependent (BOLD) image data sets transverse to the midpoint of the femur were repeatedly acquired.
Figure 1. NO-stimulated angiogenesis in B16F10 melanomas is inhibited by exogenous TSP1. B16F10 tumor biopsies (1 mm³) excised from sex- and age-matched wild type and TSP1 null mice were embedded in three-dimensional collagen matrix and incubated in growth medium ± DETA/NO (10 μM) (A, left panel) or ± TSP1 (2.2 nM; A, right panel). Sex- and age-matched athymic nude mice were injected with equal numbers of control MDA-MB-435, clone TH26, or clone TH50 tumor cells to the lateral thigh. Tumor size was determined with calipers by the same examiner at the indicated time intervals, and volumes were calculated (B). Sex- and age-matched athymic nude mice were injected with equal numbers of MDA-MB-435, Neo vector-transfected, clone TH26, or clone TH50 cells to the lateral thigh. Equal sized tumors were excised, and 1-mm³ biopsies were embedded in collagen matrix with growth medium ± NO (DETA/NO 10 μM) (C). After 7 days of incubation, vascular cell invasion was quantified as the distance of farthest cell invasion from the biopsy border in each of four quadrants. Sex- and age-matched athymic nude mice were injected with equal numbers of MDA-MB-435, Neo vector-transfected, clone TH26, or clone TH50 cells to the lateral thigh. Equal sized tumors were excised, and 1-mm³ biopsies were embedded in collagen matrix with growth medium. After 7 days of incubation, cytospin immunocytochemistry of matrix-invading cells was performed. CD31-positive staining cells were counted in 10 random high-power fields from each tumor explant and presented as mean ± SD (D). Bovine aortic endothelial cells were cultured under empty Transwell chambers or chambers that contained parental MDA-MB-435 or clone TH26 cells, and the BAECs were imaged (E). Results are representative of three independent experiments.
for 30 minutes to monitor temporal changes in blood oxygenation and blood flow. Diethylamine NONOate (DEA/NO; 100 nmol/g body weight) was injected with normal saline through rectal cannula 5.0 minutes after starting the scan. Imaging parameters used were as follows: \( T_R \), 450 milliseconds; flip angle, 45°; number of excitations, 2; slice thickness, 2 mm; matrix size, \( 64 \times 64 \); number of scans, 30; total imaging time for the series, 29 minutes.

**Laser Doppler Analysis of Tumor and Hindlimb Perfusion**

Cr(Cnr)athymic<nu>IBf nude female mice 8 weeks in age were injected subcutaneously with \( 10^6 \) tumor cells to the lateral thigh. Doppler analysis of tumor perfusion was performed when tumors reached 1 cm\(^3\) in size. Core temperature was monitored through rectal probe and maintained precisely at 35°C by a heated stage and heat lamp. Anesthesia was obtained with 1.5% isoflurane. Under loop magnification, the cutaneous envelope over each tumor was carefully excised, protecting the vascular network decorating the tumor mass. A MoorLD1-2k scanner (Moor Instruments, Devon, UK) was used with the following parameters: scan area, \( 1.6 \times 2.5 \) cm; scan speed, 4 msec/pixel; scan time, 1 minutes 54 seconds; override distance, 20 cm. After obtaining baseline tumor perfusion data, animals received either 0.05 \( \mu \)g of epinephrine i.p. in sterile warmed saline or 1 \( \mu \)l/g body weight of a 100-mM stock solution of DEA/NO. Tumor perfusion was then measured. Perfusion was also determined in normal hind limbs in sex-, age-, and weight-matched C57BL/6 or athymic mice after DEA/NO administration (1 \( \mu \)l/g weight of a 100-mM stock) using the same anesthetic/physiologic parameters and imaging conditions.

**Intracellular cGMP Assay**

Cells were plated at a density \( 1 \times 10^6 \) cells/well in 12-well culture plates (Nunc) in standard medium and weaned over 48 hours of serum and treated in basal medium + 0.1% BSA with the indicated agents. Total cGMP was determined through immunoassay (Amersham, GE Healthcare, UK). In some experiments, equal numbers of MDA-MB-435, clone TH26, and clone TH50 cells were plated in six-well plates in standard growth medium and were allowed to adhere for 6 hours; medium was replaced with serum-free RPMI with 0.1% BSA. Cells were incubated for 12 hours overnight; conditioned medium was collected, diluted at the indicated concentrations in fresh serum-free RPMI with 0.1% BSA, applied ± DEA/NO to porcine VSMC; and cGMP was determined.

**Tumor Growth Studies**

To confirm growth rates of the cancer cell lines, separate cohorts of 8-week-old female Cr(Cnr)athymic<nu>IBf mice were injected subcutaneously with \( 10^6 \) tumor cells to the lateral thigh. Tumor growth was measured by the same investigator at twice weekly intervals.

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**Figure 2.** Differential BOLD response and effect of TSP1 in tumor and normal leg. Anatomic and BOLD MR scans for (A) WT and (B) TSP1 null mice bearing B16F10 melanoma tumors 1000 mm\(^2\) in size were obtained from \( T_2^* \)-weighted MSME sequences. DEA/NO (100 nmol/g body weight) was injected with saline through an intrarectal cannula 5 minutes after starting the scan. Green and red colors show positive and negative BOLD MRI signals, respectively, at the indicated times after NO administration. The BOLD images were superimposed with the corresponding anatomic images to determine exact locations in the lateral thigh sections. Averaged BOLD MRI signal changes (C) in tumor and normal leg as a function of time after NO challenge. The green and red plots show increased and decreased BOLD MRI signals, respectively. Values are presented as mean ± SE from five and six experiments in WT and TSP1 null mice, respectively. Age- and sex-matched C57BL/6 wild type and TSP1 null mice underwent laser Doppler analysis of normal hind limbs (D). After obtaining baseline perfusion data, DEA/NO (1 \( \mu \)l/g weight of 100-mM stock) was given through rectal bolus. Data sampling was then done every 2.5 minutes. Results represent the mean ± SD of four separate animals of each strain.
Tumor volume was calculated by the formula: volume = \( W^2 \times L/2 \), where \( W \) = shortest diameter and \( L \) = longest diameter. For each cell line, six mice per group were used.

**Tumor Explant Assay**

Tumor biopsies that had reached 1 cm\(^3\) were harvested from WT, TSP1 null, and Cr(Cnr)athymic<nu>fBf nude mice and were explanted in type I collagen as described [15]. Explants were incubated in the presence of DMEM with FCS and the indicated treatment agents for 7 days, and cell migration through the extracellular matrix was measured. Results represent the mean ± SD of at least three separate experiments.

**Immunohistochemistry**

Tumor explants were prepared as described. On postexplantation day 7, tumor biopsies were removed from the constructs, and the collagen gel was digested with type II collagenase (Worthington, Lakewood, NJ). The mixture was centrifuged, supernatant was aspirated, and cells were resuspended in basal medium without serum. Cells were attached to charged glass slides (Fisherbrand Superfrost Slides; Fisher Scientific, Pittsburgh, PA) by cytopinning at 1000 rpm for 5 minutes and then fixed and rehydrated through 80%, 70%, and 50% alcohol. Slides were rinsed with PBS and then incubated with 3% H\(_2\)O\(_2\) for 30 minutes at room temperature. Sections were then blocked with 100% normal goat serum in PBS for 30 minutes at room temperature, washed, and incubated 1 hour with a rat antimouse monoclonal CD31 antibody (clone MEC 13.3; BD Pharmingen, Franklin Lakes, NJ) at 1:100 dilution. Slides were washed and incubated with a rat antimouse IgG secondary antibody (ALX-211-052; Alexis Biochemicals, San Diego, CA) diluted at 1:5000 for 45 minutes, washed with PBS, and incubated in prediluted streptavidin–HRP conjugate (BD Pharmingen) for 45 minutes at room temperature. Color was developed by DAB substrate kit (BD Pharmingen). Slides were counterstained using Mayer's hematoxylin for 2 minutes, dehydrated, and mounted. CD31-positive cells were counted in 10 randomly selected high-power fields for each tumor explant type.

**Endothelial and Tumor Cell Coculture Experiments**

Bovine aortic endothelial cells (4 × 10\(^4\) cells/well) were plated in 24-well culture plates in DMEM + 10% FCS and allowed to reach confluence. MDA-MB-435 cells or TSP1-transfected MDA-MB-435 cells (clone TH-26, 5000 cells/well) were plated onto Transwell inserts (Falcon, Becton Dickinson, NJ) in RPMI 1640 medium and allowed to adhere. Inserts were then placed in wells containing BAEC and incubated for 1 week in DMEM at 37°C and 5% CO\(_2\). Bovine aortic endothelial cells were then fixed with 1% glutaraldehyde, and images were acquired in phase-contrast illumination with a microscope (Model IX 70; Olympus, Center Valley, PA).

**Statistics**

All experiments were replicated at least three times. Results are presented as mean ± SD with analysis of significance using Student’s \( t \) test or one-way or two-way analysis of variance (ANOVA) with Tukey’s post hoc test where indicated using Origin software (version 7; OriginLab Corp., Northampton, MA), with significance taken at \( P \) values < .05.

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**Figure 3.** Differential BOLD response and effect of TSP1 in tumor and normal leg. Blood oxygen level–dependent MR scans for WT and TSP1 null mice bearing B16F10 melanoma tumors 1000 mm\(^2\) in size were obtained from \( T_2^* \)-weighted MSME sequences. Positive (A) and negative (B) BOLD MRI signal changes are presented in tumor and normal leg as a function of time after NO challenge.

**Results**

**Vascular and Tumor-Derived TSP1 Limit Tumor Angiogenic Responses**

Expression of TSP1 either by tumor cells or by the host is known to limit melanoma tumor growth in mice [40,41]. To assess whether tumor and host TSP1 differentially regulate NO-stimulated tumor angiogenic responses, we used a previously published ex vivo assay that quantifies vascular outgrowth and invasion through a threedimensional type I collagen matrix [42]. Explants from B16F10 melanomas grown in wild-type and TSP1 null animals were embedded into three-dimensional type I collagen matrices, and vascular cell invasion and migration through the matrix were quantified (Figure 1A). Vascular outgrowth was consistently greater in melanoma explants from TSP1 null animals. Treating with a slow-releasing NO-donor diethyltriamine NONOate (DETA/NO) enhanced the vascular cell
responses to a greater extent in tumor explants from wild type mice to essentially match that of melanoma explants from TSP1 null mice (Figure 1A, left panel). Conversely, addition of exogenous TSP1 suppressed the outgrowth of the melanoma explants from the TSP1 null mice to match that of explants from wild type mice (Figure 1A, right panel).

MDA-MB-435 was previously considered to be a breast carcinoma cell line but was recently confirmed to be derived from the M14 melanoma cell line [43]. Parental MDA-MB-435 cells express very low levels of TSP1 [44] and form metastatic tumors with strong angiogenic potential when grown as xenografts in athymic mice [40]. Growth of the parental cells and TSP1-transfected clones as subcutaneous xenografts in Cr(Cnr) nude mice essentially reproduced our previously published growth curves after mammary fat pad injection (Figure 1A, right panel). TH26 express high levels of intact TSP1 that suppresses tumor growth, whereas TH50 cells overexpress a C-terminally truncated TSP1 that lacks this activity [40].

Consistent with their in vivo growth rates, tumor explants of these three clones showed markedly different vascular cell outgrowth in three-dimensional cultures. Maximum cell outgrowth occurred in explants of the parental cell line and TH50 clone with minimal enhancement by NO (Figure 1C). A control vector transfected showed slightly less vascular outgrowth. In sharp contrast, the TH26 clone demonstrated no basal vascular cell response and decreased outgrowth after stimulation with exogenous NO (51.3 ± 4.3% less outgrowth compared to MDA-MB-435 explants). Immunohistochemical detection of endothelial cells by anti-CD31 staining of cypopsins recovered from the outgrowths verified that overexpression of TSP1 in the TH26 explants inhibited endothelial cell outgrowth (Figure 1D). Thus, TSP1 expressed either in the tumor vasculature or by the tumor can limit vascular outgrowth in this tumor explant angiogenesis assay.

Coculture experiments corroborated these findings. Postconfluent monolayers of BAECs are known to spontaneously form sprouts and tubes in culture that express characteristic proangiogenic genes [45]. Quiescent confluent BAECs grown under Transwell chambers containing MDA-MB-435 cells exposed the BAECs to diffusible factors secreted by the tumor cells that significantly enhance sprouting and branching of the BAEC (Figure 1E, center panel). In contrast, BAEC exposed to factors released by the TSP1-expressing clone TH26 cells maintained their quiescent morphology (right panel), indicating that the level of TSP1 secreted by TH26 cells effectively

Figure 4. Nitric oxide challenge simultaneously decreases MDA-435 tumor blood flow and increases normal tissue blood flow. Age- and sex-matched athymic nude mice were injected with an MDA-MB-435 tumor cell line to the lateral thigh. When lesions reached 1 cm in size, laser Doppler analysis of tumor blood flow was performed (A, B). After obtaining baseline perfusion data, therapeutic agents were given through rectal bolus (DEA/NO 1 μl/g weight of 100-mM stock). Data sampling was then done every 2.5 minutes. Results represent the mean ± SD of five separate experiments. Representative images are presented. Age- and sex-matched Cr(Cnr) athymic<nu> mice underwent laser Doppler analysis of normal hind limb perfusion (A, B). After obtaining baseline perfusion data, DEA/NO (1 μl/g weight of 100-mM stock) was given through rectal bolus. Data sampling was then done every 2.5 minutes. Results represent the mean ± SD of four separate animals. Representative images are presented. Scale bar, 10 mm.
blocks the branching and sprouting response stimulated by proangiogenic factors released by this melanoma cell line.

**Endogenous TSP1 in the Tumor Vasculature Does Not Limit NO-Driven Perfusion in B16F10 Melanoma Tumors**

Wild type and TSP1 null mice bearing B16F10 tumors underwent BOLD MRI to assess acute changes in tumor perfusion induced by systemic NO treatment. Average-intensity analysis of the BOLD signal integrated over the indicated areas of interest (Figure 2, A and B) demonstrated the expected net increase in BOLD signal in normal tissue, with the increase being greater in the TSP1 null mice (Figure 2C). In contrast, the average BOLD signal decreased approximately 5% in the B16F10 tumors of both wild type and TSP1 null mice (Figure 2C). This decrease is consistent with decreased blood flow into the tumor due to a steal effect resulting from the peripheral vasodilation by the administered NO. Laser Doppler analysis of normal hind limb perfusion in C57BL/6 wild type and TSP1 null mice after an NO challenge confirmed the data obtained by BOLD MRI (Figure 2D). Increased blood flow was detected in TSP1 null limbs within 2.5 minutes after administering the NO donor, and increased flow relative to wild type limbs was maintained for the duration of measurement.

Further analysis of the BOLD MRI data selecting areas showing positive versus negative signals confirmed our previous finding in healthy mouse skeletal muscle that endogenous TSP1 limits the increased BOLD MRI signal in the normal tissue in response to NO.

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**Figure 5.** Thrombospondin 1 expression in MDA-MB-435 melanomas tempers vasoactive tumor blood flow responses to vasodilator or vasopressor challenge. Age- and sex-matched athymic nude mice were injected with equal numbers of either parental MDA-MB-435, clone TH26, or clone TH50 cells to the lateral thigh. When lesions reached 1 cm in size, laser Doppler analysis of tumor blood flow was performed. After obtaining baseline perfusion data, therapeutic agents were given [epinephrine, 0.002 μg/g weight through i.p. injection (A, B); DEA/NO, 1 μl/g weight through rectal bolus (C, D)]. Data sampling was then done every 2.5 minutes. Results represent the mean ± SD of five separate experiments. Representative images are presented. See Movie W2 for a real-time movie of tumor blood flow. Scale bar, 10 mm.
challenge in areas with positive signal but did not influence those with negative response (Figure 3A). In the tumors, however, neither positive nor negative BOLD MRI signals differed between wild type and TSP1 null mice (Figure 3B). Thus, the Thbs1 genotype of vascular cells recruited into these melanomas does not significantly influence their acute response to NO, and the net effect of a systemic NO challenge (Figure 2C) is to acutely decrease blood oxygen levels in the tumor secondary to increasing flow in normal peripheral tissue (Figure 2, C and D; see Movie W1 for a real-time movie of tumor blood flow).

Tumor TSP1 Limits NO-Stimulated Alterations in Tumor Blood Flow and Perfusion

Consistent with the results in C57BL/6 mice (Figures 2 and 3), normal vascular beds in contralateral hind limbs of the athymic mice demonstrated increased blood flow as determined by laser Doppler after a vasoactive challenge with exogenous NO (Figure 4, A and B). To address whether overexpression TSP1 by the tumor can control tumor blood flow, athymic nude mice were injected with MDA-MB-435 tumor cells engineered to express variable amounts of TSP1. Parental MDA-MB-435 cells express very low levels of TSP1 [44]. On the basis of laser Doppler imaging, epinephrine moderately but rapidly increased blood flow in these tumors, which was sustained over 30 minutes (Figure 5, A and B). Relative to the parental tumors, tumors formed by the TSP1 overexpressing clone TH26 demonstrated attenuated changes in tumor perfusion after vasoactive challenge using epinephrine (Figure 5, A and B; see Movie W2 for a real-time movie of tumor blood flow). Epinephrine did not significantly increase blood flow in these tumors. The TH50 clone overexpresses a C-terminal truncated form of TSP1, lacking residues 1073 to 1152 of the mature protein and shown previously to not inhibit MDA-MB-435 tumor growth [40]. The epinephrine response of these tumors was also attenuated relative to the parental cells and returned to baseline by 30 minutes.
435 tumors. The immediate decrease in tumor perfusion returned
examined NO responses in these tumors. Tumors formed by TH50
ceptor that mediates its antagonism of NO signaling [36], we also
cause the C-terminal domain of TSP1 is recognized by the CD47 re-
with changes persisting over the 30 minutes measuring period. Be-
an NO challenge than that seen in parental MDA-MB-435 tumors
435 tumors resulted in a rapid 50% decrease in tumor blood flow over
clone TH50 tumor cells (2.5 × 10⁴/well) were plated in RPMI and
or TSP1-sensitive cGMP signaling. MDA-MB-435, clone TH26, and
Figure 6. MDA-MB-435 melanoma cells do not demonstrate NO-
sensitive cGMP signaling. MDA-MB-435, clone TH26, and
cell TH50 tumor cells (2.5 × 10⁴/well) were plated in RPMI
conditioned medium (0.01% in basal smooth muscle
–
1% in basal smooth muscle
–
M) was
increased tumor blood flow has been noted with exogenous NO ad-

Conversely, administering NO to mice bearing parental MDA-MB-
35 tumors resulted in a rapid 50% decrease in tumor blood flow over
the first 10 minutes that returned toward baseline by 30 minutes (Fig-
ure 5, C and D). Thrombospondin 1-expressing TH26 tumors dem-
strated less than half as much decrease in tumor perfusion after
an NO challenge than that seen in parental MDA-MB-435 tumors
with changes persisting over the 30 minutes measuring period. Be-
cause the C-terminal domain of TSP1 is recognized by the CD47 re-
ceptor that mediates its antagonism of NO signaling [36], we also
examined NO responses in these tumors. Tumors formed by TH50
cells exhibited a similar response to NO as the parental MDA-MB-
435 tumors. The immediate decrease in tumor perfusion returned
to baseline by 30 minutes (Figure 5, C and D; see Movie W3 for a
real-time movie of tumor blood flow). Therefore, this truncated TSP1
lacks the ability to functionally limit acute NO responses in vivo.

**MDA-MB-435 Tumor Cells Lack NO–cGMP Signaling**

A number of human tumors have been reported to have decreased
levels of or lack TSP1 expression [46–49]. Although NO signaling
stimulated by tumor-produced angiogenic factors under such condi-
tions should be increased in vascular cells [15], it is not known
whether NO and TSP1 can similarly regulate cGMP signaling in
tumor cells. Analysis of parental MDA-MB-435 cells and the TH50
and TH26 clones demonstrated no significant change in cGMP levels
after treatment with either exogenous NO or TSP1 (Figure 6A). Analy-
sis of several additional cancer cell lines including B16F10 melanoma,
PC3 prostate carcinoma, and SKOV3 ovarian carcinoma cells demon-
strated a similar lack of NO-cGMP signaling (data not shown). There-
fore, cGMP signaling responses to NO in these tumors are restricted to
the tumor vasculature.

**MDA-MB-435 Clone TH26–Conditioned Medium Limits
NO-Driven cGMP Accumulation in VSMC**

To assess the ability of the truncated TSP1 secreted by TH50 cells
to regulate vascular cGMP signaling, porcine VSMCs were treated
with conditioned medium from parental MDA-MB-435, TH26, and
TH50 cells. cGMP levels were determined by immunoassay (A). Results represent the mean ± SD of three separate experiments. Porcine femoral artery VSMCs were plated
(2.5 × 10⁴/well) in 12-well plates in growth medium, weaned of
serum over 48 hours, incubated in the indicated concentrations
of tumor-conditioned medium (0.01% in basal smooth muscle
medium + 0.1% BSA) for 15 minutes, and DEA/NO (10
μM) ± TSP1 (2.2 nM), and cGMP levels were determined by
ELISA (B). Results represent the mean ± SD of three

**Discussion**

The use of NO and other vasoactive agents for controlling tumor
blood flow remains controversial. Although some evidence indicates
that NO does not acutely and directly alter tumor blood flow
[2,5,50], others have reported a diminution of tumor blood flow
after NO synthase inhibition in vivo [19,20,50–54]. Conversely, in-
creased tumor blood flow has been noted with exogenous NO ad-
ministration [17,51]. Some of these disparities may be attributed to
variations in imaging modalities and techniques, vasoactive agents
used, and the specific tumor models. We used two distinct and com-
plementary noninvasive imaging modalities, BOLD MRI and laser
Doppler, which are highly reproducible in both animals and people.
Analysis of a syngeneic murine melanoma (B16F10) and a human
xenograft melanoma (MDA-MB-435) demonstrate that NO does
not show significant vasodilator activity for the tumor vasculature.
Regardless of murine strain or tumor cell line studied, administration
of exogenous NO resulted in a rapid decrease in tumor blood flow as
documented by both BOLD MRI and laser Doppler. Decreased
blood flow in the abnormal vascular network of the tumor was par-
alleled by a concurrent increase in blood flow in adjacent and distant
normal vascular networks (Figures 2C, 3A, and 4, A and B), estab-
shing that NO drives a redistribution of blood away from the tumor
vessels through active dilation of normal vascular networks through a
steal effect. Conversely, constriction of peripheral vasculature using
epinephrine acutely increased blood flow into these tumors, presum-
ably by increasing the mean pressure in arteries supplying the tumor.
Therefore, the tumor behaves in both cases as a passive resistance
whose flow is inversely related to that in the host periphery.

The presence of TSP1 in the tumor vasculature and surrounding
normal tissues of the host animal had minimal effects on acute regu-
lation of tumor blood flow. In contrast, overexpression of TSP1 by
the tumor cells significantly attenuated blood flow changes after NO
treatment (Figure 5, C and D), supporting a role for TSP1 in mod-
ulating tumor blood flow responses. Intravenous treatment with the
type 1 repeats of TSP1 was inferred to alter blood flow in pancreatic
tumors [39]. Assessed by intravenous injection of fluorescein-labeled
dextran and histologic analysis of excised tumors, treated animals
had decreased total vessel area and less fluorescence signal in CD31-
positive vessels. Although suggesting a decrease in vessel filling and
blood flow, these findings may represent primarily a loss of perfused
vessels rather than a change in flow, which was not directly assessed.
The type 1 repeats can block NO signaling in vascular cells through
CD36 [15,55], but our previous studies in ischemic injury models in-
dicate that CD36 signaling does not significantly contribute to acute
regulation of blood flow [38]. Thus, the antitumor activity of
TSP1-based drugs targeting CD36 may occur exclusively through in-
hibiting angiogenesis rather than flow, the latter being controlled
through CD47.

At the cellular level, vascular cells are the major target for this acute
activity of NO in tumors because most tumor cells do not demon-
strate NO-driven cyclic nucleotide signaling. MDA-MB-435 cells and
cloned from the same parent did not demonstrate an accumula-
tion of intracellular cGMP after treatment with NO. This lack of
response is not unique to MDA-MB-435 cells. Several other com-
mon tumor cell lines including B16F10, PC3, and SKOV3 cells
failed to demonstrate a cGMP flux after NO treatment (data not
shown). Tumor blood vessels have also been found incapable of pro-
ducing NO in response to acetylcholine [56]. Thus, tumors generally
lack NO-responsive cGMP signaling.

Consistent with our data, increased tumor blood flow has been
found in animal models after challenge with one [3,57,58] or more
vasoconstrictors [59] alone or in combination with β-blockers such as
propranolol [60]. The same response has been confirmed in pa-
tients [32,61]. However, the increased tumor perfusion induced by
 systemic treatment with vasoconstrictors was relatively brief. Pro-
longed increases in tumor blood flow could only be achieved through
continuous infusion of the vasoconstrictor [3]. In a xenograft soft
tissue tumor model, we found moderate enhancement of tumor
blood flow after a single i.p. dose of epinephrine. As with NO, blood
flow changes to epinephrine were tempered in tumors that overex-
press TSP1. In all cases, tumor blood flow tended to return to base-
line within 30 minutes. Although similar agents can modify tumor
blood flow in people, their use may be limited by increases in heart
rate and blood pressure [61].

Tumor blood flow, therefore, is a passive response to alterations in
flow and resistance in the nontumor vascular networks and system of
the host both regionally and centrally. If vascular resistance beyond
the tumor decreases, as with administration of exogenous NO, blood
rapidly drains from the tumor. Alternatively, increasing peripheral vas-
cular resistance in normal vascular networks with a centrally acting
hypertensive agent redistributes blood into the tumor vasculature.

Acutely, the tumor vasculature is resistant to vasoactive challenge and
registers no direct response.

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Movie W1. Syngeneic B16 melanoma perfusion responses to NO in wild type versus TSP1 null mice. Each movie shows an initial light image followed by a pretreatment Doppler image and a time lapse movie (500 milliseconds = 2.5 minutes) of Doppler scans taken over 30 min after administration of DEA/NO as a rectal bolus.

Movie W2. Perfusion responses to NO in a M14 (MDA-MB-435) human melanoma xenograft overexpressing full-length thrombospondin-1 (clone TH26). Time lapse movies (500 milliseconds = 2.5 minutes) of Doppler scans taken over 30 min after administration of DEA/NO as a rectal bolus.

Movie W3. Perfusion responses to NO in a M14 (MDA-MB-435) human melanoma xenograft overexpressing C-terminal–truncated thrombospondin-1 (clone TH50). Time lapse movie (500 milliseconds = 5 minutes) of Doppler scans taken over 40 min after administration of DEA/NO as a rectal bolus.