TUMORIGENESIS AND NEOPLASTIC PROGRESSION

Antiphospholipid Antibodies Promote Tissue Factor–Dependent Angiogenic Switch and Tumor Progression

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Progression to an angiogenic state is a critical event in tumor development, yet few patient characteristics have been identified that can be mechanistically linked to this transition. Antiphospholipid autoantibodies (aPLs) are prevalent in many human cancers and can elicit proangiogenic expression in several cell types, but their role in tumor biology is unknown. Herein, we observed that the elevation of circulating aPLs among breast cancer patients is specifically associated with invasive-stage tumors. By using multiple in vivo models of breast cancer, we demonstrated that aPL-positive IgG from patients with autoimmune disease rapidly accelerates tumor angiogenesis and consequent tumor progression, particularly in slow-growing avascular tumors. The action of aPLs was local to the tumor site and elicited leukocytic infiltration and tumor invasion. Tumor cells treated with aPL-positive IgG expressed multiple proangiogenic genes, including vascular endothelial growth factor, tissue factor (TF), and colony-stimulating factor 1. Knockdown and neutralization studies demonstrated that the effects of aPLs on tumor angiogenesis and growth were dependent on tumor cell-derived TF. Tumor-derived TF was essential for the development of pericyte coverage of tumor microvessels and aPL-induced tumor cell expression of chemokine ligand 2, a mediator of pericyte recruitment. These findings identify antiphospholipid autoantibodies as a potential patient-specific host factor promoting the transition of indolent tumors to an angiogenic malignant state through a TF-mediated pathogenic mechanism.


The identification of mechanisms that trigger the malignant transition of indolent, avascular tumors is an important goal for cancer prevention and treatment. The avascular state may occur as either an early stage in primary tumor development or minimal residual disease after treatment. The transition of avascular tumors to a malignant state depends on the induction of angiogenesis (angiogenic switch), but the mechanisms that drive this transition have not been well defined. It would be especially valuable to elucidate patient-specific host factors that modulate tumor angiogenesis and progression, but few such factors that are assessable in patients have been identified.

The antiphospholipid antibodies (aPLs) are a class of autoantibodies that recognize complexes of phospholipid-binding proteins bound to anionic phospholipids on the cell surface. Anionic phospholipids are absent from the surface of most cells but translocate to the surface on activation of endothelial cells, monocytes, and platelets, as well as during apoptosis and malignant transformation. The incidence of aPLs in the general population is approximately 1% to 5%. Chronic aPL elevation can lead to a thrombogenic state.

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known as antiphospholipid syndrome (APS), and aPL elevation also can occur transiently in association with viral and several other infections as well as in patients with other autoimmune disorders, such as lupus or rheumatoid arthritis. Numerous studies have shown a significantly increased frequency of circulating aPLs in cancer across multiple tumor types, compared with matched controls, with incidence generally 15% to 25%. Because the linkage of aPLs to coagulation is well-known, studies of the pathological implications of aPLs in cancer have focused on thromboembolic disorders, a common complication in many malignancies. Although comparatively few studies have examined aPLs as a potential risk factor for cancer development or progression, studies with relatively short-time follow-up found an association between aPLs and non–coagulation-related cancer mortality, and aPL-positive patients experienced a higher than expected frequency of non-Hodgkin lymphoma.

The potential for aPL as an angiogenic factor has long been evident. aPLs can induce the expression of vascular endothelial growth factor (VEGF) and tissue factor (TF) in monocytes and endothelial cells. TF, a central factor in coagulation cascade, can stimulate angiogenesis via either procoagulant function and platelet activation or, alternatively, the direct activation of protease-activated receptor-2 (PAR2; alias coagulation factor II receptor-like 1) and proangiogenic signaling in endothelial cells. Many tumor cells produce TF, and tumor cell–derived TF has been shown to promote tumor progression. Despite these studies, the effects of aPLs from patients on tumor cells and angiogenesis have not been previously examined. In this study, we present evidence that aPL activates proangiogenic gene expression in breast cancer cells, induces TF-dependent vascularization in small, slow-growing avascular tumors, and promotes tumor progression in multiple mouse models of breast cancer. Collectively, these findings indicate a potential protumor role for these autoantibodies.

Materials and Methods

Mice

All procedures were approved by the Institutional Animal Care and Use Committees of Albert Einstein College of Medicine (Einstein; New York, NY) and by the Animal Care and Use Review Office (US Department of Defense). All mice were housed in the pathogen-free barrier facility at Einstein. Nude mice in NCr background were obtained from Taconic Farms (Albany, New York) and bred at Einstein. Polyoma virus middle T antigen (PyMT) mice were kindly provided by Dr. Jeffrey W. Pollard (Albert Einstein College of Medicine, Bronx, NY) and backcrossed to mice with C57BL/6 background for >10 generations.

Cell Lines

MDA-MB-436 breast cancer cells were obtained from the National Cancer Institute Tumor Repository (Frederick, MD). The breast cancer lines MDA-MB-468 and MDA-MB-231, as well as the prostate cancer cell line PC3, were obtained from ATCC (Manassas, VA). Cells were cultured according to the supplier’s recommendations.

TF Knockdown

Four TF-knockdown shRNA constructs were selected from the shRNA library of The RNAi Consortium (vector backbone: pGIPZ). The target regions for three of the shRNA sequences, TFKD321 (5′-CTTCTATGGTGTGACATTGT-3′), TFKD322 (5′-CTGTATTACCATATTACAT-3′), and TFKD323 (5′-TGGAGCTACTGCAAATGCT-3′), were in the 3′ untranslated region of the TF (F3) gene; a fourth, 5′TFKD324 (5′-AAGTCTACACTGTTCAAAT-3′), was within the coding sequence of exon 2. The lentiviral expression vector carried turbo-green fluorescent protein (GFP). Virus-carrying turbo-GFP alone was used as a positive control for transfection. Selection and lentiviral packaging of the constructs, along with an empty vector control, were performed by the shRNA Core Facility at Einstein. Lentivirus carrying the above constructs was transfected into human breast cancer cells following the GIPZ lentiviral shRNA technical manual from Thermo Scientific (Waltham, MA). Lentivirally transduced cells were selected in 6 μg/mL of puromycin for 7 days, after which >90% of cells were GFP positive.

Human Breast Cancer Samples

Procedures using human samples were approved by the Institutional Review Board of Montefiore Medical Center (MMC)/Einstein (09-06-190X). Residual sera, collected for preoperative tests from patients with abnormal breast mass at MMC, were examined for aPL titer (described below). The diagnosis and staging of breast lesions was performed by the surgical pathologists (Y.W. and other on-service pathologists) at MMC.

Determination of aPL Titer

Serum samples were assayed for aPL by the Immunodiagnostic Laboratory at MMC using a commercial ELISA kit from BIO-RAD (Hercules, CA). The assays included the following: anti-cardiolipin (IgM, IgG), anti–β-2 glycoprotein I (IgM, IgG, IgA), and anti-phosphatidylserine (aPS; IgM, IgG). Samples with medium/high titer, according to criteria established by international consensus, were classified as aPL positive.

IgG Samples

Plasma samples, obtained from two patients with APS, were designated as aPL#1 and aPL#3. These samples, as well those from five healthy donors (referred to as control IgG herein), were prepared from discard plasma samples obtained during diagnostic or other unrelated clinical procedures (Institutional Review Board number 09-06-190X). Serum was prepared from

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plasma by incubation with 1 U/mL human thrombin (Sigma-Aldrich, St. Louis, MO) at 37°C for 30 minutes, followed by clot removal. Total IgG was isolated, as described previously, using protein G Sepharose 4, according to the manufacturer’s recommended procedure (Amersham Pharmacia Biotech, Piscataway, NJ). Purified IgGs were dialyzed against phosphate-buffered saline (PBS), and the protein concentration was determined by absorbance at 280 nm. A third sample of patient IgG with elevated aPL was obtained from LifeSpan Biosciences, Inc. (Seattle, WA) and designated aPL#2. aPL titers of all samples were determined as described above. IgGs from the three patients are referred to collectively as aPL+ IgGs.

Affinity-purified aPS/anti-cardiolipin IgG was prepared by modification of a published method. Briefly, phosphatidylserine and cardiolipin (Sigma-Aldrich) in 100% ethanol were mixed at a 2:3 molar ratio in a 50-mL conical tube, dried, and suspended in Tris-HCl buffer (pH 7.4). The tube was rotated at 37°C for 1 hour for liposome formation. aPL#1 serum was added, and the tube was rotated for 2 hours. Liposomes were then pelleted, washed with the same buffer, and resuspended in 20 mmol/L phosphate buffer containing 0.1% Tween 20 (pH 7.4). The affinity-purified IgGs were then isolated with protein G Sepharose 4 (Amersham Pharmacia Biotech). Affinity-purified IgG was approximately 14-fold enriched in aPL titer relative to unpurified IgG, as assessed by aPS activity.

Controls for Endotoxin Effects
IgGs from patients or normal controls were treated with Detoxi-Gel Endotoxin Removing Gel (Thermo Scientific, Rockford, IL) following the manufacturer’s recommended protocol. No significant differences were observed in the effect of human IgGs on tumor cells’ proangiogenic gene expression nor tumor growth and leukocytic infiltration in xenografted breast model between Detoxi-Gel endotoxin-treated and untreated human IgGs.

Cell Viability and Proliferation Assays
MDA-MB-436 cancer cells (1 × 10^5 per well) were plated in 96-well plates in RPMI 1640 medium for overnight growth. Cells were then treated with or without control or aPL+ IgG at the indicated concentrations for 24 hours and then assayed for viability or proliferation. To assay viability, cells were incubated for 4 hours with 0.5 μg/mL of MTT assay (Sigma-Aldrich). The reduction of MTT was measured using a standard method. To assay proliferation, BrdU incorporation was then assayed for viability or proliferation. To assay viability, cells were incubated with 1 mmol/L bromodeoxyuridine (BrdU; Sigma-Aldrich), and BrdU incorporation was determined by immunoassay according to the manufacturer’s instructions (Millipore-Calbiochem, Billerica, MA).

Gene Expression Test
Cells (5 × 10^5 per well) were placed in 6-well plates in RPMI 1640 medium containing 1% fetal bovine serum for overnight growth. Cells were then treated for the indicated times with 0.2 to 0.4 mg/mL aPL+ or control IgGs. RNA was extracted using TRIzol (Invitrogen/Life Technologies, Grand Island, NY) and reverse transcribed with the Superscript II system (Invitrogen/Life Technologies), according to the manufacturer’s instructions. Primers were obtained from Real Time Primers (Elkins Park, PA), and real-time quantitative PCR was performed on Applied BioSystems 7900HT (Applied Biosystems/Life Technology, Grand Island, NY). Glycer-aldehyde-3-phosphate dehydrogenase—normalized mRNA for specific gene expression was calculated by the 2^-ΔΔCt method using SDS software version 2.3 (Applied Bio-systems/Life Technologies).

In Vivo Assays
Xenografts
Breast cancer xenografts were established by inoculation of human cancer cells (5 × 10^5 cells per site in 0.1 mL RPMI 1640 medium) into the thoracic or abdominal mammary glands of female nude mice at 7 to 10 weeks of age. For coinjection experiments, the inoculum was first incubated for 1 hour on ice in 0.1 mL RPMI 1640 medium containing 20 μg total polyclonal human IgG from either healthy donors or patients; alternatively, 9.5 μg affinity-purified IgG was used. For TF neutralization, the incubation was in the presence of 10 μg anti-human TF antibody (American Diagnostica-Sekisui, Stamford, CT). For experiments using i.p. administration of human IgGs (rather than coinjection), one injection of 80 mg/kg body weight (bw) human IgG was administrated i.p. at day 14 after grafting. Mice were monitored every 7 days for the formation of visible tumors. Tumor incidence was calculated as the percentage of grafted glands bearing visible tumors. Tumor volumes were measured with a caliper for longest (L) and shortest (S) length and thickness (Th) by a blinded observer (Y.-Y.W., A.V.N., or E.Y.L.), and total volume was calculated as LxSxTh. Tumor volume in grafts of GFP-expressing tumor cells was determined weekly using a whole body Xenogen IVIS imaging system (Caliper Life Sciences, Hopkinton, MA). Background signal was measured using parallel mammary glands injected with 5 × 10^5 non-transduced cells.

PyMT Model
Female PyMT mice at 6 weeks of age were administered one i.p. injection of approximately 80 mg/kg bw human IgG. Tumors in all thoracic and abdominal mammary glands were analyzed for tumor stage by a blinded observer (Y.-Y.W., A.V.N., or E.Y.L.) using criteria established and described in detail in previous reports.

Determination of the Titer of Circulating Human aPLs in Mice
Four mice per group were injected i.p. once with approximately 80 mg/kg bw aPL#1 IgG or alternatively coinjected with 1 × 10^5 MDA-MB-436 cells and 20 μg aPL#1 IgG into each of two
Immunofluorescence

For quantitation of tumor blood vessel density, vessels were labeled in vivo as described previously. Briefly, lysine-fixable, Texas Red–conjugated dextran (mol. wt.: 70,000) (Life Technologies), 6.2 mg/mL in PBS, was administered i.v. (21 μg/gm bw) 5 to 7 minutes before tissue harvest. We demonstrated that this method precisely marked endothelium-lined tumor vessels in mice. Tissues were fixed with 10% formalin, divided into sections, and counterstained with DAPI, and the entire tumor area was imaged. Vessel density was measured as the ratio of Texas Red–positive area/DAPI-positive area using WCIF ImageJ software version 1.37a (NIH, Bethesda, MD). For size matching of tumors, the longest diameters of tumors were determined from the tagged image file format images using Photoshop (Adobe, New York, NY). To obtain fluorescent images of tumor-associated macrophages, lysine-fixable fluorescence isothiocyanate–dextran (mol. wt.: 70,000) (Life Technologies), 6.2 mg/mL in PBS, was administered i.v. (21 μg/gm bw) 2 hours before tissue harvest. We have previously demonstrated that this method specifically labels macrophages within the tumor.

To assess α-smooth muscle actin (SMA)–positive cells and their association with blood vessels, sections were simultaneously stained with a monoclonal anti–α-SMA antibody (Sigma-Aldrich) and rabbit anti–von Willebrand factor (Dako, Carpinteria, CA), followed by Alexa Fluor-488–conjugated goat anti-mouse IgG2α and Texas Red–conjugated goat anti-rabbit antibodies (Molecular Probes/Life Technologies). Confocal images were prepared by the Analytic Imaging Core Facility at Einstein.

In Vivo Matrigel Plug Assay

Human breast cancer cells (5 × 10^5) were incubated on ice with 20 μg control or aPL IgG, as in the coinjection procedure. BD Matrigel Matrix (Growth Factor Reduced; BD Biosciences, San Jose, CA) was added to a final concentration of approximately 70%, and the mixture was injected into mammary gland. To label blood vessels in the plug, Texas Red–conjugated dextran was administered before harvest, as described above. Plugs were fixed in 10% formalin or zinc fixative (BD Biosciences) at 4°C overnight and then paraffin embedded and divided into sections. Gross images of the plugs were obtained using a Zeiss STEMI (Zeiss, Thornwood, NY) at the Einstein Analytical Imaging Core Facility.

Statistical Analysis

The unpaired Student’s t-test was used to assess significance for all data sets, with the exception of tumor incidence, PyMT tumor stage, and aPL incidence in breast cancer patients, which were analyzed using a Fisher’s exact test.

Results

Occurrence of Elevated aPL Titer in Breast Cancer Patients

In view of previous reports of aPL occurrence in cancer across multiple tumor types, we examined aPL status in newly diagnosed breast cancer patients at MMC (Table 1). Eight patients diagnosed with carcinoma in situ and 13 patients with invasive carcinoma were tested for seven classes of autoantibody frequently detected in APS patients. Of the 21 total patients, 6 (29%) had elevated aPL according to international consensus laboratory criteria for APS. This value is consistent with previous reports of aPL prevalence in cancer. Notably, 6 (46%) of 13 invasive cases were aPL positive, in comparison to 0 of 8 patients with noninvasive tumors (P < 0.05), consistent with a potential association of aPL with invasive malignancy.

Promotion of Tumor Progression by Patient-Derived Antibodies

To investigate the potential role of aPLs in tumor progression, we used IgGs from patients with autoimmune elevation

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<th>Table 1</th>
<th>aPL Titer in Patients with Breast Lesions</th>
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<td>aPL status</td>
<td>Carcinoma in situ*</td>
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<tr>
<td>aPL⁺</td>
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<td>aPL⁺ (% total)</td>
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*Carcinoma in situ includes both ductal and lobular carcinoma in situ.
†Invasive carcinoma includes invasive ductal or lobular carcinoma.

P < 0.05 (Fisher’s exact test).
of circulating aPLs, including APS patients, because these samples are available in sufficient quantity for animal model studies. The serological findings for the aPL\(^+\) patients and healthy donors (referred to as control IgG hereafter) used in this study are shown in Table 2. Orthotopic models of human breast cancer were generated by xenografting human breast cancer cell lines into mammary fat pads of nude mice. To enhance our ability to detect potential effects of aPLs on tumor progression, we selected two human breast cancer cell lines, MDA-MB-436 and MDA-MB-468, which are known to exhibit slow growth in xenograft models, potentially because of limited angiogenesis.\(^{30,31}\) As controls, we examined cell lines that display rapid growth in vivo, including MDA-MB-231 breast cancer cells and PC3 prostate cancer cells.

In pilot studies, we observed that tumors remained undetectable or <2 mm in diameter for 4 weeks in >70% of MDA-MB-436 or MDA-MB-468 xenografts. This observation is consistent with previous reports that 90% of fat pad xenografts of MDA-MB-436 cells remain undetectable for at least 30 days after grafting.\(^{30}\) In initial experiments, mice grafted with MDA-MB-436 cells at 7 weeks of age were administered a single i.p. dose of either aPL#1 or control #1 polyclonal IgG (80 mg/kg bw) at 2 weeks after grafting. In control IgG-injected mice, tumors remained undetectable in >70% of grafts and mean tumor volume remained <5 mm\(^3\) at 5 weeks after grafting (Figure 1, A and B). In contrast, mice treated with aPL#1 IgG produced detectable tumors in >80% of grafts at 5 weeks, with a mean volume of >10 mm\(^3\) (Figure 1, A and B). Subsequently, the rate of tumor growth continued to increase in the aPL#1 IgG-treated, but not in the control IgG-treated, mice (Figure 1B). The volume of control tumors remained <8 mm\(^3\), consistent with a dormant phenotype,\(^{32}\) whereas aPL#1 IgG-treated tumors were >20 mm\(^3\) within 4 weeks after IgG injection (Figure 1B). The effect of aPL#1 IgG was specific to cell lines with suboptimal growth, because equivalent tumor growth was observed in control and aPL#1 IgG-injected mice grafted with more aggressive MDA-MB-231 breast cancer cells (data not shown) or PC3 prostate cancer cells (Figure 1C). To determine whether aPL levels achieved in the mouse model were representative of patient levels, we assessed human aPL titer in the circulation of injected mice. The circulating level of aPL#1 IgG at 1 week after injection was near the abnormal cutoff value in patients, and declined with a half-life of 9.3 days (Supplemental Figure S1), consistent with previous half-life estimates in mice for human IgG1 and IgG2,\(^{33,34}\) the major isotopes in patient aPL.\(^{35,36}\)

To extend these findings to a spontaneous breast cancer model, we used PyMT mice, in which mammary tumors are induced by the transgenic expression of the polyoma middle T oncogene specifically in mammary epithelial cells. Mammary tumors in this model progress through stages comparable to the human disease, including an onset of angiogenic switch and pulmonary metastasis.\(^{25,26}\) The reactivity of human aPL with murine antigens is indicated by the use of these antibodies in several studies to induce pathological effects in mice, similar to symptoms observed in patients with APS, including pregnancy loss, fetal injury, and arterial thrombosis.\(^{37-39}\) In \(\beta-2\) glycoprotein I, the phospholipid-binding protein that serves as the major pathogenic target of human aPL,\(^{30,41}\) the N-terminal domain to which aPL binds is perfectly conserved between mouse and human.\(^{51}\) This domain has been shown to mediate direct binding of human aPL to murine \(\beta-2\) glycoprotein I.\(^{41}\) In pilot studies, we observed that PyMT mice with C57Bl/6 background (used in this study) progressed to malignant stages at 8 to 9 weeks of age, representing an approximately 2-week delay relative to the earlier studies with this model in mixed genetic background.\(^{25}\) Consequently, we administered one i.p. injection of either control or aPL#1 IgG to the PyMT mice in this background at 6 weeks of age, when most tumors were at premalignant stages, and analyzed tumor stages at 9 weeks of age. In contrast to control IgG, a single treatment with aPL#1 IgG resulted in a sevenfold increase in the frequency of histological progression from early to late carcinoma within 3 weeks (Figure 1, D and E).

Tumor cells often display high levels of cell surface anionic phospholipids,\(^{43,44}\) potential targets of aPL, suggesting that local interaction between aPL and tumor cells may affect tumor progression. Because cell surface anionic phospholipids are expected to interact with the circulating protein annexin A5,\(^{45}\) we performed immunostaining to assess annexin A5 in xenografted tumors. We observed a marked localization of annexin A5 to tumor, compared with adjacent normal tissue (Figure 1F and Supplemental Figure S2A). High levels of annexin A5 were also observed in PyMT mammary tumors (Supplemental Figure S2A). These data are consistent with the presence of a cell surface target for aPL in tumors, although we cannot exclude that local expression of intracellular annexin A5 may also contribute to the observed staining.

To further distinguish local and systemic effects of aPL administration, we developed a coinjection protocol, in which breast cancer cells were preincubated for 1 hour on ice with 20 \(\mu\)g of aPL#1 or control IgG. Cells with aPL#1 IgG were then grafted ipsilaterally and cells with control IgG

| Table 2 | aPL Titer (IgG Isotype) in Serum of aPL\(^+\) Patients and Healthy Donors |
|---------|-----------------------------|-----------------------------|-----------------------------|
| Normal cutoff | aCL <23 | aPS <16 | \(\beta\)-2GPI <20 |
| aPL#1 | 30.6 | 78.5 | >154 |
| aPL#2 | 50 | 50 | 57 |
| aPL#3 | 12.9 | 100 | 152 |
| Control 1 | 0.34 | 1.18 | <5.0 |
| Control 2 | 2.71 | 1.98 | 0.84 |
| Control 3 | 1.75 | 3.4 | 0.21 |
| Control 4 | 1.73 | 0.63 | 0.14 |
| Control 5 | 2.19 | 0.92 | 0.28 |

Data are given as U/mL. aPL titer is determined by the Immunodiagnostic Laboratory at MMC. One unit is equivalent to 1 \(\mu\)g/mL of affinity-purified IgG.\(^{39}\)
contralaterally in the same mice (Figure 1G). Tumor volumes were determined at each injected site at weekly intervals. By using three different aPL-positive IgGs, matched with three different control IgGs, we consistently observed a significant increase in the volume of MDA-MB-436 tumors within 1 week after grafting at aPL-positive IgG-injected sites compared with control sites, at which growth was delayed and variable (Figure 1H and Supplemental Figure S2, B and C). After this initial
period of aPL+ IgG-stimulated growth, tumor size was maintained with little change for several weeks (Figure 1H and Supplemental Figure S2, B and C) until 5 to 6 weeks after grafting, at which time aPL+ IgG-treated sites produced rapidly expanding tumors, whereas control IgG-treated sites remained comparatively quiescent (Figure 1H). Although the magnitude of tumor growth showed some experimental variation, the distinct kinetics stimulated by aPL+ IgG, with respect to both initial growth and later tumor expansion, were a consistent finding and were verified with a second human breast cancer cell line, MDA-MB-468 (Figure 1I). To obtain further evidence for the tumor-accelerating effect of aPL, aPL was affinity purified from aPL#1 IgG using liposomes bearing a cardiolipin/phosphatidylserine mixture. Xenografts coinjected with affinity-purified aPL displayed stimulation of tumor growth comparable to that observed with whole aPL#2 IgG (Figure 1H). Taken together, these observations indicate that aPL can act locally at tumor sites to promote early tumor growth and long-term tumor progression.

Promotion of Tumor Angiogenesis by aPL+ IgG

No effect of aPL+ IgG on the proliferation or survival of cultured MDA-MB-436 cells was observed (Supplemental Figure S2, D and E), suggesting that it is unlikely that aPL+ IgG promotes tumor progression through a direct effect on tumor cell growth. We next considered whether the tumor-accelerating effect of aPL is associated with an onset of tumor angiogenesis, because previous studies have indicated that accelerated growth of MDA-MB-436 xenografts is angiogenesis dependent.30,46 We observed that tumors from aPL+ IgG-treated grafts were more vascularized than control tumors at 6 weeks after grafting (Figure 2A). Control grafts displayed blood vessels in the stromal region, but intratumoral vessels were at low density in comparison to

Figure 2  Effects of antiphospholipid antibodies (aPL)-positive IgG on tumor angiogenesis and leukocytic infiltration. A: Representative tumors imaged at 6 weeks after coinjection of MDA-MB-436 cells with normal (CON) or aPL+ IgG. B: Mice were injected i.p. with normal (CON) aPL#1 IgG 2 weeks after grafting MDA-MB-436 cells, and tumors were collected at 6 weeks after grafting for assessment of blood vessels and macrophages labeled with red and green fluorescent dextran, respectively. White arrows indicate labeled blood vessels; yellow arrow, tumor stroma. No vessels were observed within the imaged control tumor. C and D: The ratio of vessel density (Texas Red dextran intensity)/nuclear signal (DAPI) was compared between size-matched tumors in normal (CON) and aPL+ IgG-treated MDA-MB-436 grafts (n = 3 to 5). Tumor sizes, based on longest length, are displayed in the right half of each panel. E: MDA-MB-436 xenografts coinjected with control (CON) or aPL+ IgG were stained to detect smooth muscle actin. Arrows point to positively stained blood vessels in tumors. F and G: aPL+ IgG-induced leukocytic infiltration in xenografts. F: MDA-MB-436 xenografts from mice injected i.p. with control (CON) or aPL+ IgG (aPL) were stained to detect macrophages (F4/80) or neutrophils (Ly6G). G: Leukocytic infiltration in F was quantitated as the ratio of diaminobenzidine signal/nuclear counterstain. For F4/80, n = 3 (CON) or n = 4 (aPL); for Ly6G, n = 4 (CON) or n = 6 (aPL). *P < 0.05 versus controls (C, D, and G). Scale bars: 1 mm (A); 50 µm (B and F); 30 µm (E).
aPL$^+$ IgG-treated tumors (Figure 2B), suggesting increased angiogenesis in aPL$^+$ IgG-treated tumors. However, it could not be determined from this observation whether angiogenesis was promoted by aPL$^+$ IgG or was simply a consequence of the acceleration of growth. We, therefore, quantitated vessel density in size-matched tumors from grafts exposed to aPL$^+$ or control IgG, using a fluorescent labeling approach that we have validated for this purpose in previous studies.$^{27}$ The analysis revealed a pronounced effect of aPL$^+$ IgG on tumor angiogenesis independent of tumor size (Figure 2, C and D). Notably, the enhancement of angiogenesis by aPL$^+$ IgG was greater (>10-fold) for size-matched tumors <1.5 mm in diameter than for larger tumors, which displayed a less than threefold increase of vessel density (Figure 2, C versus D). This observation suggests that a proangiogenic effect occurs at an early stage of tumor acceleration by aPL. The ability of aPL to exert early effects on tumor progression is further indicated by the appearance of invasive histological features in small tumors in aPL$^+$ IgG-treated grafts, compared with size-matched control tumors in which capsules remain well defined and undisturbed (Supplemental Figure S3).

To further probe the effects of aPL$^+$ IgG on angiogenesis, we examined the coverage of tumor vessels with mural cells expressing SMA. Vessels in aPL$^+$ IgG-treated tumors displayed well-developed mural layers, with SMA organization indicative of parallel bundles of filaments forming a continuous perivascular sheath (Figure 2E), which has been observed as a characteristic feature of pericytes in normal vessels.$^{47}$ In control IgG-treated tumors, the vessels available for comparison were near the transition between tumor and stroma. These peripheral vessels were mostly surrounded by mural layers that were thinner and less developed than those of aPL$^+$ IgG-treated tumors (Figure 2E).

Because we and others have observed an involvement of tumor-associated macrophages in angiogenic switch,$^{26,48}$ we examined the effect of aPL$^+$ IgG on leukocytic infiltration. We observed more macrophages as judged by dextran labeling in aPL$^+$ IgG-treated tumors (Figure 2B). A quantitative analysis revealed a >10-fold increase in the density of F4/80$^+$ macrophages in aPL$^+$ IgG- compared with control IgG-treated tumors (Figure 2, F and G). Dense focal accumulations of Ly6G$^+$ neutrophils were also frequently observed in aPL$^+$ IgG-treated tumors, although this effect

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**Figure 3** Kinetics of antiphospholipid antibodies (aPL)-induced tumor angiogenesis. **A:** Representative gross images of Matrigel plugs harvested at 3, 6, and 9 days after grafting (3D, 6D, and 9D, respectively). Plugs were generated in mammary glands of nude mice by inoculating Matrigel-containing MDA-MB-436 cells and 20 µg of either control (CON) or aPL$^+$ (aPL) IgG. Arrows indicate vessel formation visible at the surface of the plug. **B:** Representative images of sections of Matrigel plugs containing either control or aPL$^+$ IgGs, harvested at day 6 after grafting. Arrows indicate blood vessels labeled with Texas Red dextran. **C:** Vessel density is quantitated as the ratio of Texas Red dextran intensity/DAPI nuclear counterstain (vessel/nuclei) in tumors harvested at 3D, 6D, and 9D. $n=3$ to 4 (3D), 6 (6D), or 4 (9D). **D:** Representative hematoxylin and eosin -stained sections of tumors in Matrigel plugs harvested at day 6 after grafting. The boxed areas in the left panels are shown at higher magnification on the right panels. Arrows indicate tumor vessels. *$P<0.05$ (C). Scale bars: 1 mm (A); 100 µm (B and D, left panels); 30 µm (D, right panels).
did not achieve statistical significance (Figure 2, F and G). IgGs treated with Detoxi-Gel to remove endotoxin gave similar results (data not shown), indicating that the effect of aPL is not due to the endotoxin contamination in aPL+ IgG. Taken together, these observations suggest that aPL has the potential to promote tumor progression by stimulating angiogenesis and inflammatory reactions at the tumor site.

Kinetics of Tumor Angiogenesis in Response to aPL+ IgG

To further examine the time course of aPL+ IgG-stimulated angiogenesis, it was necessary to examine time points within 1 week of aPL exposure. Because the xenograft provides limited material at these early time points, we chose to examine a Matrigel xenograft model. Matrigel plugs, in which MDA-MB-436 cells were combined with either control or aPL+ IgG, were implanted and harvested at 3, 6, and 9 days after grafting. Gross examination of the plugs (Figure 3A) revealed only minimal vessel development at 3 days. At 6 days, vessel development had clearly progressed in aPL+ IgG, but not in control IgG plugs. By day 9, vessel development was evident in control plugs, but remained more pronounced in aPL+ IgG plugs. Quantitative microscopic analysis of vessels labeled with dextran (administered before harvest27) confirmed these findings: vessel density was significantly elevated in aPL+ IgG-treated compared with control IgG-treated plugs at both 6 and 9 days after implant (Figure 3, B and C). These findings were confirmed by histological analysis, which demonstrated a high density of blood vessels in aPL+ IgG but not IgG-treated Matrigel plugs (Figure 3D).

Proangiogenic Gene Expression in aPL+ IgG-Treated Tumor Cells

We next sought to identify aPL-induced factors that may regulate angiogenesis and tumor progression. Because our studies indicate that aPL interacts locally with tumor (Figure 1, F, H, and I), and aPL is known to modulate gene expression in target cells,49 we considered that aPL might interact with cells in the tumor to promote angiogenesis. Previous studies have demonstrated that aPL can induce expression in macrophages of both VEGF-A and TF, central regulators of angiogenesis.21

![Figure 4](https://example.com/fig4.jpg)

**Figure 4** Effects of antiphospholipid antibodies (aPL)-positive IgG on gene and protein expression. A and B: Real-time quantitative PCR analysis of gene expression in cultured MDA-MB-436 cells treated with control (C) or aPL+ (A) IgG for 4 or 8 hours. For A, n is 5, 4, 4, 5, 9, 5, or 6 for angiomotin (AMOT), CCL2, CSF1, IL-8, tissue factor (TF), thrombospondin 1 (THBS1), or VEGF, respectively. For B, n is 4, 7, 8, or 4 for CCL2, CSF1, TF, or VEGF, respectively. C: Western blot analysis of cell lysates for MDA-MB-436 (30 µg/lane 436) or MDA-MB-231 (3 µg/lane 231) cells treated with control (C) or aPL#1 (A) IgG for 24 hours. Molecular weight markers are indicated on the left. D: IHC for TF in MDA-MB-436 tumors in mice injected i.p. with normal (CON) or aPL+ (aPL) IgG 2 weeks after grafting. Tumors were collected 7 weeks after IgG injection. Arrows indicate positive staining. E: IHC for TF in Matrigel plugs containing MDA-MB-436 cells and either control (CON) or aPL+ (APL) IgG. Plugs were harvested on day 3 or 6 after grafting (3D or 6D, respectively). Arrows in day 3 images indicate positively stained cells. F: TF protein expression in Matrigel plugs was quantitated as the ratio of TF stain/nuclear counterstain (hematoxylin) (n = 4). *P < 0.05 (A, B, and F). Scale bars: 50 µm (D); 30 µm (E).
However, the ability of aPL to act directly on tumor cells to promote a proangiogenic phenotype has not been examined. We, therefore, investigated the effect of aPL+ IgG treatment on the expression of a panel of proangiogenic and proinflammatory genes in human breast cancer cells. Treatment of MDA-MB-436 cells with aPL+ IgG for 4 hours, in comparison to control IgG, resulted in a significant elevation of the gene expression of TF and VEGF, as well as the macrophage growth factor colony-stimulating factor-1 (CSF1) (Figure 4A), which we and others have linked to tumor progression and angiogenesis.25,27,50 In contrast, the expression of other angiogenesis-related factors in human breast cancer cells, including thrombospondin 1, angiomotin, and IL-8 (alias CXCL8), was not significantly affected by aPL+ IgG (Figure 4A). At 8 hours of treatment, the expression of chemokine ligand 2 (CCL2), which has been implicated in tumor angiogenesis and monocyte recruitment,51–53 became significantly elevated, whereas expression of TF and VEGF returned to baseline levels (Figure 4B). The expression of CSF1 remained elevated at 8 hours (Figure 4B).

Regulation of TF Protein Expression by aPL+ IgG

We chose to evaluate TF as a potential mediator of aPL+ IgG effects on tumor angiogenesis and growth, because TF can potentially support angiogenesis via several pathways,
including procoagulant action and direct induction of proangiogenic gene expression. Protein expression analysis of MDA-MB-436 cells treated with aPL$^+$ IgG for 24 hours demonstrated up-regulation of an approximately 49-kDa species (Figure 4C), consistent with full-length TF protein.54 In comparison, MDA-MB-231 cells, which form aggressive, aPL$^+$ IgG-independent tumors when grafted to mammary fat pads (data not shown), displayed a high baseline level of TF expression (consistent with a previous report$^{55}$) that was little affected by aPL$^+$ IgG treatment (Figure 4C). The difference in TF mobility between the two cell lines is similar to previously observed variation attributable to variable glycosylation.$^{56}$

We next examined TF protein expression in tumors in vivo. Seven weeks after IgG injection of MDA-MB-436 xenograft-bearing mice, we observed focci of intense TF expression in tumors from aPL$^+$ IgG-treated mice compared with weaker diffuse staining in control IgG-treated mice (Figure 4D). To examine tumor TF expression at early time points, we used the Matrigel plug model. At 3 days after grafting, plugs containing aPL$^+$ IgG-treated tumor cells displayed a marked increase in TF-expressing cells compared with control IgG-treated plugs (Figure 4E). Most of the TF-expressing cells had the morphological appearance of tumor cells (Figure 4E). Quantitative analysis indicated aPL$^+$ IgG stimulation of plug TF expression at 3 and 6 days after grafting, although statistical significance was not achieved at 6 days because of variation (Figure 4F).

Role of Tumor Cell—Derived TF in aPL$^+$ IgG-Promoted Tumor Progression

We next sought to investigate the potential role of aPL$^+$ IgG-induced TF in tumor progression. When MDA-MB-436 cells were coinjected with both aPL$^+$ IgG and an anti-TF neutralizing antibody, tumor incidence and tumor size at 1 week after grafting were reduced by 67% to 71% compared with MDA-MB-436 coinjected with aPL$^+$ IgG alone (Figure 5, A and B). At 2 weeks, a similar reduction in tumor size was observed, although statistical significance at this time point was not achieved. Neutralization of TF also reduced aPL$^+$ IgG-mediated tumor vessel density by 60% compared with size-matched tumors treated with aPL$^+$ IgG (Figure 5, C and D), suggesting that TF may mediate the protumor effect of aPL$^+$ by promoting tumor angiogenesis.

Because the TF-neutralizing antibody recognizes both human and murine TF,$^{57}$ the TF dependency we observed potentially reflects a role of tumor cell— or host-derived TF. Host TF has been shown to have minimal impact on tumor growth in multiple murine models,$^{21}$ whereas tumor cell—derived TF has been shown to promote tumor progression in multiple murine cancer models.$^{20,21,58}$ To definitively address the role of tumor cell—derived TF, we used a gene-silencing approach. Lentiviruses carrying four different TF shRNA expression vectors, as well as an empty control vector, were used to infect MDA-MB-436 and MDA-MB-468 cells. Three of the shRNAs (TFKD321 to 323) were designed to target the TF 3′ untranslated region and one, TFKD324, was designed to target exon 2. In pilot experiments, all of the TF-shRNA—expressing cells displayed significant reduction of TF mRNA (Supplemental Figure S4A). We selected two shRNAs, TFKD321 and 324, for further analysis because of their differences in target site and inhibitory efficiency (approximately 60% for TFKD321 and >80% for TFKD324). Western blot analysis demonstrated a high efficiency of TF knockdown in MDA-MB-436 cells carrying either TFKD321 or TFKD324, with only minimal responses to aPL$^+$ IgG treatment in either knockdown line (Figure 5E). TF knockdown in MDA-MB-436 cells did not inhibit cell growth in vitro, and in fact TF-knockdown MDA-MB-436 cells showed increased viability by MTT assay (Supplemental Figure S4B). We then tested the effect of TF knockdown in tumor cells on aPL$^+$ IgG-induced tumor progression in vivo. MDA-MB-436 cells carrying TFKD321, TFKD324, or control vector were coinjected with aPL#1 IgG into mammary glands. Tumor formation and growth were monitored weekly and confirmed by whole body bioluminescent imaging for GFP, which is expressed by the lentiviral vector in transfected tumor cells (Supplemental Figure S4C). Consistent with our findings using anti-TF-neutralizing antibody, TF knockdown inhibited aPL$^+$ IgG-induced tumor progression. Throughout the 3-week period of observation after grafting, tumor sizes in TFKD324-bearing xenografts were reduced >85% relative to control ($P = 0.012$, $P < 0.001$, and $P = 0.004$ for weeks 1, 2, and 3, respectively) (Figure 5F). In xenografts with TFKD321-transduced cells, in which TF knockdown is less efficient, a trend of growth inhibition was observed, reaching approximately 50% inhibition at 3 weeks, although no statistical significance was achieved (Figure 5F). Consistent with these findings, vessel density in aPL$^+$ IgG-treated tumor xenografts containing TFKD324-bearing MDA-MB-436 cells was lower than that of aPL$^+$ IgG-treated control tumors (Figure 5G).

We next considered potential mechanisms downstream of TF that might mediate aPL$^+$ IgG-induced angiogenesis. Full-length TF, the form induced by aPL$^+$ IgG, is an integral membrane protein and may, therefore, act in an autocrine manner to trigger proangiogenic factor production in tumor cells, although it may also potentially be released in microparticles to affect neighboring tumor/stromal cells in a paracrine manner.$^{59}$ Although it is possible that tumor cell gene expression of VEGF and CSF1 is TF dependent, we have been unable to demonstrate such dependence using neutralizing anti-TF antibodies (data not shown). In contrast, a significant inhibition of CCL2 mRNA expression was observed in both TFKD321 and TFKD324 cells compared with control cells 8 hours after aPL$^+$ IgG treatment (Figure 5H). The results imply that aPL$^+$ IgG induces CCL2 production in tumor cells in a TF-dependent manner. These findings are consistent with studies showing that TF expression in
endothelial cells can promote angiogenesis via the up-regulation of CCL2 expression.60,61 One potential mechanism by which CCL2 might mediate aPL+ IgG-stimulated angiogenesis is via the recruitment of monocytes/macrophages into tumors.51 However, we observed no significant reduction in tumor-associated macrophages on TF depletion, using either TFKD321 or TFKD324 xenografts (data not shown). A second potential mechanism by which CCL2 may promote tumor angiogenesis is via the recruitment of pericytes or vascular smooth muscle cells, which form the mural cells that surround the endothelium in mature blood vessels. These closely related cell types, which both express SMA, are thought to play important supportive roles throughout the process of angiogenesis,62 and it has been suggested that the recruitment of these cells to sites of angiogenesis is mediated by the TF-dependent expression of CCL2.60 We used fluorescence microscopy to assess SMA+ cell association with xenograft tumor vessels in the presence or absence of tumor cell–derived TF. Because TF knockdown reduces vessel density in the xenografts (Figure 5G), we focused the analysis on vessels near the tumor periphery, because vessel density is generally higher in this region relative to the tumor center. Consistent with our IHC studies (Figure 2E), aPL+ IgG-treated xenografts derived from either parental or vector control MDA-MB-436 cells displayed a dense array of vessels with well-developed SMA+ mural layers. The SMA in the mural cells of these vessels appeared to be organized as parallel bundles forming a continuous perivascular sheath47 (Figure 5I and Supplemental Figure S5, A and D). In contrast, the SMA+ mural cell layers in the vessels of aPL+ IgG-treated, TF-deficient tumors were poorly developed, displaying thin bundles of actin stain that did not show parallel organization and typically formed incomplete perivascular sheaths (Figure 5I and Supplemental Figure S5, B–F). This deficiency in coverage by SMA+ cells was evident in both large and small vessels in TF-knockdown compared with control tumors (Supplemental Figure S5). In comparison, vessels in normal mammary tissue adjacent to TF-deficient xenografts displayed well-developed SMA+ mural cell layers similar to those observed in control MDA-MB-436 xenografts (Supplemental Figure S5, A and I). The overall density of SMA+ intratumoral cells was reduced by >70% in TF-knockdown xenografts relative to control xenografts (Supplemental Figure S5H). These findings indicate that the incorporation of mural cells during the aPL+ IgG-mediated development of tumor vasculature is dependent on aPL+ IgG-stimulated TF production in tumor cells.

Discussion

Dormant, microscopic tumors may occur as residual disease after treatment, and may also be prevalent as undiagnosed primary tumors in the general population.1 The individual variation with respect to progression of these tumors is potentially due to patient-specific mechanisms, including host characteristics. However, triggering mechanisms that convert precancerous tumors to malignancies are poorly understood,1,63 and few patient-specific characteristics have been identified as host factors that promote progression. Herein, we investigated antiphospholipid autoantibodies, which are prevalent in multiple human malignancies, as a potential patient-specific host mechanism promoting tumor progression. Our key findings are that aPL+ IgG promotes TF-dependent angiogenesis and tumor growth in an orthotopic breast cancer model, accelerates tumor progression in a spontaneous breast cancer model, and stimulates TF expression and the expression of multiple proangiogenic factors in tumor cells (Figure 6). These are the first data to show an effect of these autoantibodies on tumor cells, and, to our knowledge, the first studies to demonstrate tumor

![Figure 6 Hypothetical model of antiphospholipid antibodies (aPL)-triggered dormancy disruption. Binding of aPL to complexes of phospholipid-binding proteins bound to cell-surface anionic phospholipids on dormant tumor cells induces tumor cell production of proangiogenic factors, including trans-membrane tissue factor (TF) and secretory vascular endothelial growth factor (VEGF) and colony stimulating factor-1 (CSF-1). TF further regulates the production of the chemokine CCL2. These events result in the stimulation of local angiogenic switch and transition of the tumor from dormancy to malignancy.](https://ajp.amjpathol.org)
Antiphospholipid Antibodies Are Protumor

The mechanisms by which aPL binds to cell surfaces and initiates signal transduction are not well defined. Studies in monocytes and endothelial cells indicate potential contributions from a variety of cell surface proteins to aPL-mediated signaling, including annexin II, low-density lipoprotein receptor–related protein 8, and several members of the Toll-like receptor (TLR) family (TLR2, TLR4, and TLR8).78 Of these molecules, annexin II and TLR4 have been implicated in aPL induction of TF expression, severely affecting the intratumoral accumulation of SMA+ cells and resulting in the formation of tumor blood vessels with thin and incompletely developed SMA+ mural cell layers. These mural cells may support angiogenesis by both enhancing vessel stability and providing paracrine/juxtacrine cues to the underlying endothelial cells.77 Our study provides the first demonstration in vivo of a functional link between tumor cell TF and tumor recruitment of these potentially pivotal stromal elements.

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MDA-MB-436. It is conceivable, then, that aPL acts on dormant tumors, represented in our model by MDA-MB-436 cells, to activate an annexin II–based pathway of TF expression that is constitutively functional in cell lines, such as MDA-MB-231, that may be representative of advanced-stage tumors in which angiogenic switch has taken place. Consistent with this interpretation, we observed no effect of aPL+ IgG on either MDA-MB-231 or another fast-growing tumor, PC3.

Antiphospholipid autoantibodies include antibodies that recognize several different complexes of phospholipids and phospholipid-binding proteins. Antibodies recognizing complexes of cardiopatolin and β2-glycoprotein I (anti-β2GPI) are increasingly recognized as the major pathogenic entity in APS,40,41 and anti-β2GPI has been strongly linked to the induction of TF in monocytes in studies using monoclonal or affinity-purified antibodies.82,83 TF induction appears to be a robust property of these antibodies: a recent study of monocyte TF expression in response to IgG from patients bearing anti-β2GPI observed >10-fold stimulation by IgG from each of eight individuals, in comparison to 12 healthy donors.18 Notably, of the 13 patients with invasive breast cancer that we surveyed, 5 (38%) were above cutoff with respect to anti-β2GPI, and 5 more were between 50% to 100% of cutoff (data not shown). Therefore, although it remains possible that aPL in cancer patients will reveal functional differences with respect to protumor effects, compared with aPL from the three non-cancer patients used in this study, the available serological data, both in our and other studies,13,14 suggest the potential for aPL activation of TF-mediated angiogenic function in cancer settings. Investigation of protumor function in cancer patient sera will be the subject of future studies.

Antibodies that recognize tumor cells may, in principle, be capable of mediating anti-tumor responses, in addition to the protumor effects we have described. We did not observe anti-tumor effects of aPL in the immunocompetent PyMT model; however, it remains to be determined whether such effects might be observed in other settings. aPL antibodies, purified from APS patient serum, were reported to inhibit metastasis in a murine melanoma model.85 However, in human melanoma patients, autoimmune status (including the presence of antircardiopatolin antibodies) has not shown any consistent relationship with outcome.85,86 Bavituximab, a monoclonal antibody recognizing phosphatidylserine and β2GPI, displays an anti-tumor effect associated with the enhancement of anti-tumor immunity.87 The distinct function of bavituximab, compared with aPL+ IgG in our study, may be due to its distinct specificity. Bavituximab acts through binding to domain II of β2GPI,89 whereas pathogenic human aPL predominantly targets domain I at the N-terminus.89,90 Anti-domain I reactivity is strongly correlated with coagulopathy, with respect to both clinical manifestations in patients and in vitro studies of clotting time.90,91

The aPL prevalence we and others have documented in cancer patients presumably reflects chronic elevation of these autoantibodies, although this has not been verified by repeated sampling in most studies. However, in addition to chronic elevation, aPL may also become transiently elevated in infections.8,92 For example, parvovirus, which has an incidence of 30% to 60% in US women,93 is associated with elevated levels of anti-β2GPI.94 In the general population, the frequency of elevated aPL is approximately 1%.27 Because an unknown, but potentially large, fraction of these instances represent transient responses, the long-term risk of transient aPL elevation in cancer patients is difficult to determine and may be substantial. Transient aPL has only a limited association with coagulopathy,75 perhaps because of the low probability of a coincidence between transient aPL elevation and transient events that generate cell surface targets for the autoantibodies (the two-hit hypothesis96). In contrast, the findings in this study, as well as earlier reports describing phospholipid exposure in tumor cells,4,43,44 indicate that tumors are a potential source of chronically presented aPL targets, and therefore transient aPL responses may potentially be clinically significant in cancer.

It remains difficult to account for the transition of tumors to a proangiogenic, aggressive phenotype. Although multiple studies have identified host pathways, such as the promotion of host cell recruitment and VEGF mobilization by matrix metalloproteinase-9,97–100 as important for the promotion of tumor angiogenesis in animal models, there are few studies that can associate this transition with events or characteristics identifiable in patients. It is especially urgent to investigate such factors in breast cancer, in which many deaths are attributed to the transition of residual disease from a dormant to an aggressive state.1,101,102 Consequently, we have focused our studies on two cell lines chosen for their promise as potential models of dormancy, and it is notable that we observed effects of aPL+ IgG only on these lines, and not on lines that display no dormant tendencies. We demonstrated proangiogenic effects of aPL+ IgG in tumors whose size (<1.5 mm diameter) and vascular deficiency are consistent with previous descriptions of a dormant tumor phenotype.103 Escape from dormancy may also be a critical process in primary tumor development, as suggested by the estimate that >20% of women between the ages of 40 and 50 years have clinically undetectable breast cancer.104 Our studies indicate that elevation of patient aPL deserve consideration as a novel factor promoting angiogenic switch. Further studies are needed to assess the clinical significance of this mechanism and to further delineate the pathways linking these autoantibodies to tumor progression in experimental models.

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Supplemental Data

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References


36. Dostal-Johnson D, Rote NS, Branch DW: IgG1 and IgG2 are the predominant subclasses of antiphospholipid antibody in women with the lupus anticoagulant. Clin Immunol Immunopathol 1990, 54:309–319


68. Hjortoe GM, Petersen LC, Albrechtsen T, Sorensen BB, Norby PL, Mandal SK, Pendurthi UR, Rao LV: Tissue factor-factor
Villa-specific up-regulation of IL-8 expression in MDA-MB-231 cells is mediated by PAR-2 and results in increased cell migration. Blood 2004, 103:3029–3037


105. Antiphospholipid Antibodies Are Protumor

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