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TITLE: Prevention of the Angiogenic Switch in Human Breast Cancer

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**15. ABSTRACT**

Our overall goal is to determine if human breast cancer can be prevented from becoming angiogenic when it is still at a microscopic size (< ~ 1 mm³). From Feb. '05 – Feb. '06, we have made the following progress: (1) We have cloned three different human breast cancers that undergo the angiogenic switch at predictable times. (2) We have found that the angiogenic switch time is modified by host stroma: two-fold earlier for tumor in the mammary fat pad, compared to tumors implanted in subcutaneous tissue. (3) We have found that the angiogenic switch is preceded by repression of stromal expression of thrombospondin-1. Angiogenic tumor cells continue to secrete a novel thrombospondin-1 repressing factor. This protein has been purified and partially sequenced. (4) For one of the breast cancers, the angiogenic switch can be detected at a microscopic size by a significant increase in bFGF in platelet alpha granules. (5) We have determined that the BRCA1 gene (breast cancer susceptibility gene), appears to regulate a ratio of thromobospondin-1 to VEGF in breast cancer cells. The lower the thrombospondin-1/VEGF ratio, the sooner the tumor cells will spontaneously switch to the angiogenic phenotype and grow large tumors in SCID mice.

**16. SECURITY CLASSIFICATION OF:**

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<th>a. REPORT</th>
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**19a. NAME OF RESPONSIBLE PERSON**

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I. INTRODUCTION:

The purpose of this research is to determine if human breast cancer (and possibly other human tumors) can be prevented from becoming angiogenic when they are still at a microscopic size of less than approximately 1 millimeter diameter. We are developing angiogenesis-based biomarkers that will recognize the presence of such an early tumor long before it is detectable by any conventional method, such as magnetic resonance imaging, CAT scan, or ultrasound. We are trying to determine the mechanism of the switch to the angiogenic phenotype in human breast cancer.

With support from the Innovator Award we have recently discovered that a subpopulation of angiogenic human breast cancer cells can reversibly switch to the non-angiogenic, dormant phenotype. This spontaneous reversal of the angiogenic phenotype is not unique to human breast cancer. To date we have found it in other human cancers as well. Approximately 5–15% of angiogenic breast cancers can spontaneously turn off their own angiogenic activity. We wish to learn whether administration of endogenous angiogenesis inhibitors can increase the percentage of angiogenic tumor cells that reverse their angiogenic phenotype. Furthermore, it may be possible to administer oral non-toxic drugs which increase the expression of endogenous angiogenesis inhibitors. Because of a recent discovery supported by this Innovator Award, we wish to determine if the BRCA1 gene is regulating the angiogenic phenotype in human breast cancer.

The translational goal of this project is two-fold:

(i) To use angiogenesis-based biomarkers to detect recurrent breast cancer years before detection by conventional methods and before symptoms, so that the biomarker itself could be employed to guide therapy by angiogenesis inhibitors of little or no toxicity. In other words, our long-term goal is to liberate the management of cancer from dependence on anatomical location.

(ii) Women with the mutated breast cancer gene may be monitored by periodic tests of angiogenesis-based biomarkers. Rising biomarkers could be treated as described above. If the biomarkers returned to normal, this could possibly obviate the current medical practice of prophylactic bilateral mastectomy and oophorectomy.
II. BODY:

Task 1: **Median time of switching to the angiogenic phenotype for human breast cancers.**

The timing of the switch to the angiogenic phenotype has been determined for three human breast cancers, and for other types of cancer as well (colon cancer, liposarcoma). These switching times and the percentage of tumors that switch to the angiogenic phenotype have been highly reproducible in many replicative experiments over the past 2 years. We have now turned our attention to understanding the mechanisms of the switch to the angiogenic phenotype.

**A. Research Accomplishments:**

To determine whether the breast cancer susceptibility gene, BRCA1, controls the angiogenic switch through regulation of the angiogenesis regulators VEGF and thrombospondin-1. (With Sandra Ryeom Ph.D.)

This is a new direction within Task 1. It is based on our finding of a possible novel mechanism for the breast cancer susceptibility gene (BRCA1), during support by this Innovator Award.

Through the analysis of families at high risk for breast and ovarian cancer, the breast cancer susceptibility genes were identified in 1996 and named BRCA1 and BRCA2 (1). Despite the identification of these genes, their roles in tumorigenesis are still not clearly defined. BRCA1 has been implicated in a number of different functions including DNA repair and recombination, cell cycle control and transcription. One mutant BRCA1 copy in the germline predisposes an individual to cancer with the second allele consistently lost in tumors thereby leaving no functional copy of BRCA1 (1) (Figure 1).

While many studies have investigated BRCA1 function both upstream and downstream of DNA damage responses (1), there has been little work studying the role of BRCA1 as a regulator of angiogenesis which is necessary for expansion of tumor mass beyond 1 mm³. One of the most widely studied pro-angiogenic proteins is vascular endothelial growth factor (VEGF), a critical endothelial cell mitogen *in vivo* (2). Studies have demonstrated that VEGF levels are 7-fold higher in breast tumors compared to normal adjacent breast tissue with VEGF levels directly regulated by estrogen (3). BRCA1 has been postulated to negatively regulate estrogen receptor signaling (1). Therefore, loss of BRCA1 function may lead to constitutive activation of estrogen receptor signaling, upregulation of VEGF levels, and ultimately loss of estrogen receptor expression (a hallmark of BRCA1-null tumors). Estrogen has also been shown to downregulate expression of thrombospondin-1, an endogenous angiogenesis inhibitor (4).

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1. Estrogen → thrombospondin-1 → angiogenesis

2. BRCA1 (mutated, low copy number) or BRCA1 (extremely high copy number) → thrombospondin-1 → angiogenesis (tumor growth

Fig. 1. Model of estrogen and BRCA1 regulation of thrombospondin-1 and angiogenesis.
We hypothesize that loss of BRCA1 function may also lead to constitutive inhibition of thrombospondin-1 expression, further promoting the angiogenic switch. The angiogenic switch is the change in tumors from a dormant, poorly vascularized harmless tumor, to a highly vascularized, rapidly expanding, angiogenic tumor (5). One of the characteristics of the angiogenic switch is the reversal in the ratio of pro- and antiangiogenic regulators such as VEGF and thrombospondin-1 from a ratio of low VEGF and high thrombospondin-1 to the opposite (5).

**New Findings:**

(i) We established two different human breast cancer cell lines with either over-expression of wild-type BRCA1, or with mutant (i.e., no functional copies of BRCA1) lines (with the generous collaboration of Dr. Mary-Claire King’s lab in Seattle).

<table>
<thead>
<tr>
<th>Cell Line Name</th>
<th>Tissue type</th>
<th># of functional BRCA1</th>
</tr>
</thead>
<tbody>
<tr>
<td>HCC1937</td>
<td>Human breast ca.</td>
<td>0</td>
</tr>
<tr>
<td>HCC1937+ BRCA1</td>
<td>Human breast ca.</td>
<td>many (&gt;1000)</td>
</tr>
<tr>
<td>UWB</td>
<td>Human ovarian ca.</td>
<td>0</td>
</tr>
<tr>
<td>UWB + BRCA1</td>
<td>Human ovarian ca.</td>
<td>many (&gt;1000)</td>
</tr>
</tbody>
</table>

(ii) We established the presence of non-angiogenic and angiogenic subpopulations in each of these cell lines by injection of all four tumor cell lines in SCID mice and we then determined the time until each cell line underwent the angiogenic switch.

(iii) We determined the *in vitro* production of the angiogenic regulators VEGF, thrombospondin-I and bFGF in the four different cell lines. We characterized each cell line by their ratio of thrombospondin-1 to VEGF and found that their time to the angiogenic switch correlated with their thrombospondin-1 to VEGF ratio, i.e., the lower the ratio, the sooner the cells undergo the angiogenic switch in SCID mice.

**Reportable Outcomes:**

We are currently preparing a manuscript describing our findings on the regulation of the angiogenic switch by the breast cancer susceptibility (BRCA1) gene.

**Conclusions and Experiments Planned:**

Our future studies will examine the role of wild-type BRCA1 as a putative regulator of VEGF and thrombospondin-1. Our initial expectations would have predicted that the human breast cancer cells (HCC1937) and the human ovarian cancer cells (UWB1.289) with no functional copies of BRCA1 would undergo the angiogenic switch sooner than the cells over-expressing wildtype BRCA1.
However, our most recent preliminary data suggests a new hypothesis: that the function of BRCA1 may best fit a bell shaped curve) with the number of functional copies of BRCA1 regulating angiogenesis and subsequently expansion of tumor mass (Figure 2). The mutant breast and ovarian cancer cells have zero functional copies while the BRCA1 over-expressing cells have hundreds to thousand-fold higher copy numbers than wildtype or heterozygous cells. Therefore, we postulate that at either end of the spectrum of BRCA1 copy number, tumorigenesis will occur. The timing of the angiogenic switch may be a direct consequence of the ratio of thrombospondin-1 / VEGF levels. In other words, the sooner expression of thrombospondin-1 decreases, and VEGF expression increases, the sooner the angiogenic switch will occur. Our future studies will address the validation of this hypothesis.

B. Research Accomplishments: (Task 1 Continued):

Studies on the molecular mechanism of the switch to the angiogenic phenotype in human breast cancer. (With George Naumov, Ph.D, and Eugene Lifshits, M.D.).

New findings:
Experiments were carried out to validate the model of quantifying the switch to the angiogenic phenotype in human breast cancer.

(i) We isolated and characterized the in vivo growth of the angiogenic and non-angiogenic subpopulations of human breast cancer cells (MDA-MB-435) (Figure 3). Approximately 70% of non-angiogenic tumor cells from this line.
reproducibly switch to the angiogenic phenotype at a median of 100 days. In contrast, for a different human breast cancer (MDA-MB-415), only 30% of non-angiogenic tumors undergo the angiogenic switch at a median of 500 days.

(ii) We worked out the minimum number of human tumor cells that need to be inoculated into a mouse to produce a non-angiogenic microscopic tumor in mice, i.e., $1 \times 10^6$ tumor cells is sufficient to produce a dormant tumor in mice. Inoculation of higher numbers of tumor cells, i.e., $2 \times 10^6$, $3 \times 10^6$, or up to $5 \times 10^6$ tumor cells, carried over excessive concentrations of vascular endothelial growth factor (VEGF), so that a few early “non-angiogenic” tumors were sparsely vascularized for approximately the first 15 to 30 days after implantation. Beyond that time period, these sparse new vessels could no longer be sustained, and the resulting microscopic tumor implants returned to their original non-angiogenic size ($\sim 1 \text{ mm}^3$). In contrast, when not more than $1 \times 10^6$ tumor cells were inoculated subcutaneously, microscopic dormant tumors formed that were always completely non-angiogenic until the predicted time for the angiogenic switch.

(iii) We found that the angiogenic switch for breast cancer could be modified by the host stroma. For example, non-angiogenic human breast cancers that switched to the angiogenic phenotype at a median of 150 days in the mammary fat pad, did not undergo the angiogenic switch until $\sim 500$ days (median) when implanted subcutaneously. Our preliminary data by Randy Watnick shows that angiogenic human breast cancers repress thrombospondin-1 significantly more efficiently in the mammary fat pad than in subcutaneous tissue (see below).

(iv) We performed microarray analysis that compared gene expression profiles of angiogenic to non-angiogenic human breast cancer cells (MDA-MB-436). For example, a few of the more significant gene expression changes during the switch to the angiogenic phenotype are listed in the table below, as a ratio of angiogenic/non-angiogenic expressions.

Microarray Results for human breast cancer MDA-MB-436.

<table>
<thead>
<tr>
<th>Gene Description</th>
<th>Expression ratio (Angiogenic/Non-angiogenic)</th>
</tr>
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<tbody>
<tr>
<td>Heat shock protein, HSP27</td>
<td>26.97</td>
</tr>
<tr>
<td>Quinolinate Phosphoribosyltransferase, QPRT</td>
<td>4.30</td>
</tr>
<tr>
<td>Bone marrow stromal cell antigen 2, BST2</td>
<td>4.22</td>
</tr>
<tr>
<td>Interferon alpha inducible protein 27, IFI27</td>
<td>3.28</td>
</tr>
<tr>
<td>Cytochrome P450, CYP1B1</td>
<td>3.11</td>
</tr>
<tr>
<td>Spondin 2, SPON2</td>
<td>3.06</td>
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</table>

(v) We have developed SCID immunodeficient mice transplanted with GFP- (green fluorescent protein) expressing bone marrow. These mice will be
used to address the role of bone marrow-derived cells in the angiogenic switch of human breast cancer.

(vi) We have continued to make progress in understanding a newly discovered phenomenon, i.e., that the angiogenic switch is not permanently unidirectional (originally demonstrated by Taturo Udagawa in the Folkman lab). A subpopulation of angiogenic tumor cells (10-20% of tumor cells) can spontaneously switch in the reverse direction and become non-angiogenic. We proved this by cloning single tumor cells from highly angiogenic tumors that had already switched to the angiogenic phenotype, and expanding them briefly in vitro and then implanting them subcutaneously. A major goal for the next year will be to understand the mechanism of reversal of the angiogenic switch. Do any of the natural angiogenesis inhibitors operate by this mechanism? Can novel non-toxic drugs be developed that reverse the angiogenic switch?

(vii) In collaboration with Professor Douglas Wallace (University of California, Irvine), we have sequenced and compared mitochondrial DNA from angiogenic and non-angiogenic human breast cancer cells (MDA-MB-436). Very significant tumor-type specific mitochondrial DNA mutations have been identified in non-angiogenic tumors that are not found in their angiogenic counterparts.

Reportable Outcomes:


Conclusion and Plans:
We plan to utilize the prolonged dormancy period of the MDA-MB-415 human breast cancer cells (30% undergo the angiogenic switch at a median of ~500 days), to study changes in the platelet proteome during the angiogenic switch (See Task 3 below). This is possible because the tumors remain at a microscopic size for long periods of time (up to 500 days). We expect that this particular model will be very useful for the discovery of angiogenesis-based biomarkers of human dormant non-angiogenic breast cancers.

We plan to characterize and compare the gene expression profiles of non-angiogenic and angiogenic MDA-MB-415 breast cancer cells. A separate analysis will be performed under hypoxic and normoxic conditions. We anticipate that these analyses will identify key molecular players in the angiogenic switch. The identified gene products will be qualitatively and quantitatively assessed in vivo during the angiogenic switch in all of our breast cancer models.
(viii) We have already identified a potentially important mediator of the angiogenic switch... heat shock protein 27 (HSP27). HSP27 is overexpressed 27-fold in angiogenic human breast cancer MDA-MB-436 cells, in contrast to the non-angiogenic tumor cells. HSP27 has been identified as an intra-cellular target of endostatin, thrombospondin-1, fumagillin, and its synthetic derivative TNP-470 (6). This gene expression difference was also evident when comparing these breast cancer cells under normoxic and hypoxic conditions (preliminary data). We plan to investigate in vivo changes in intratumoral and stromal HSP27 during the angiogenic switch in collaboration with Professor Lars Akslen of the Department of Pathology at the University of Bergen, Norway. Professor Akslen was on sabbatical in the Folkman lab during the past year.

(ix) We will explore the possibility of horizontal genetic transfer, as discovered by Dr. Lars Holmgren and J. Folkman (7, 8), between angiogenic and non-angiogenic tumor cells. We will test whether non-angiogenic tumor cell phagocytosis of apoptotic bodies from angiogenic tumor cells will increase the frequency and shorten the time-span of the angiogenic switch.

Future plans include in vitro studies where apoptotic GFP-expressing angiogenic switch breast cancer cells are fed to non-GFP-expressing non-angiogenic cells. If some of the non-angiogenic tumor cells initiate GFP expression, this would provide evidence for horizontal DNA transfer between the two populations in vitro.

Apoptotic bodies from angiogenic GFP-expressing cells will also be inoculated into non-GFP-expressing dormant tumors grown in vivo. If the non-angiogenic tumors acquire GFP-expression and induce a switch to the angiogenic phenotype, this would provide in vivo evidence for horizontal transfer of genetic material between angiogenic and non-angiogenic tumor cells. The reverse experiments would also be performed.

**Task 2. Molecular and genetic studies of the mechanism of the stability of the non-angiogenic phenotype.**

**Introduction:**
In the Folkman lab George Naumov has previously demonstrated that one form of tumor dormancy results from an inability of a microscopic tumor to induce angiogenesis. Such human tumors, grown in SCID immunodeficient mice, can remain non-angiogenic, and dormant at a microscopic size of \(~ 1 \text{ mm}^3\), for months to more than a year, before they switch to the angiogenic phenotype at a predictable time. The presence of dormant tumors provides an excellent tool to study the regulation of tumor progression as well as the role of the surrounding normal tissue in this process. We hypothesized that the inability of dormant breast cancers to induce neovascularization is due to an imbalance in their production of pro- and antiangiogenic factors.
Key Research Accomplishments:

**Stromal production of thrombospondin-1 as a regulator of angiogenesis in human breast cancer.** (With Randolph Watnick, Ph.D., and Soo-Young Kang, Ph.D.)

(i) We have found that non-angiogenic human tumor cells express relatively high levels of thrombospondin-1, a potent endogenous inhibitor of angiogenesis. In contrast, thrombospondin-1 decreases significantly prior to the switch to the angiogenic phenotype. Furthermore, the c-Myc oncoprotein, which negatively regulates the expression of thrombospondin-1, is expressed in greater amounts in angiogenic human tumor cells, as compared to non-angiogenic dormant tumor cells.

(ii) In the first year of this grant we established that two breast cancer cell lines that form non-angiogenic, microscopic, dormant tumors in mice express very high levels of thrombospondin-1, and very low levels of c-Myc or phosphorylated c-Myc. Conversely, two breast cancer cell lines that are able to form rapidly growing angiogenic tumors in mice, express very low levels of thrombospondin-1 and high levels of c-Myc.

(iii) We have now determined that angiogenic breast cancer cell lines are able to repress the expression of thrombospondin-1 in surrounding stromal fibroblasts in the tumor bed. In contrast, dormant non-angiogenic tumor cell lines actually stimulate thrombospondin-1 expression in stromal fibroblasts. We are currently in the process of identifying and purifying the thrombospondin-1 repressing factor that the angiogenic cells secrete as well as the thrombospondin-1 stimulating factor that the non-angiogenic cells secrete. Interestingly, the expression and/or secretion of the thrombospondin-1 repressing factor appears to be downstream of c-Myc because inhibition of Myc activity in the angiogenic tumor cells results in the loss of the thrombospondin-1 repressing activity.

(iv) Based on this new experimental data, we can now hypothesize that all breast cancers must pass through the dormant non-angiogenic phase prior to gaining the capacity to becoming an expanding tumor mass. Based on our current data we postulate that this transition or “angiogenic switch” is triggered by the increased expression of c-Myc, which, in turn, represses the expression of thrombospondin-1 and allows the tumor to become angiogenic. It is the angiogenic tumor that can be metastatic, expand its mass, and produce symptoms and metastases.

(v) Also during the past year we ectopically expressed a Myc-estrogen receptor gene in dormant breast cancer cells. This fusion protein is constitutively expressed, but is only activated in the presence of tamoxifen, because as the estrogen moiety is a mutant that only binds to tamoxifen. Furthermore, we ectopically expressed a dominant-negative Myc-estrogen receptor gene in the angiogenic breast cancer cells.
We have recently found that activation of Myc in dormant tumor cells leads to the repression of thrombospondin-1 both in the tumor cells and in mammary fibroblasts that were co-cultured with these cells. Meanwhile, the activation of the dominant-negative Myc in the angiogenic tumor cells induced the stimulation of thrombospondin-1 expression, both in the tumor cells and in mammary fibroblasts co-cultured with these cells.

**Conclusions and Future Plans:**

We have now inoculated these cells into immunodeficient mice to determine whether activation of the Myc-estrogen receptor construct will force tumor cells to an earlier angiogenic switch. We will also inoculate tumor cells that express the dominant negative construct into immunodeficient mice to determine whether disruption of Myc activity will reverse the angiogenic phenotype to a microscopic, nonangiogenic, dormant tumor phenotype. Next, we will also try to determine the genetic and molecular mechanisms that regulate the expression of c-Myc in order to determine how the angiogenic switch is triggered.

**Task 3: Development of novel angiogenesis-based biomarkers in platelets to detect non-angiogenic, microscopic-sized dormant tumors before or just after the angiogenic switch, but before tumors can be detected by palpation in mice (i.e., less than 50 mm³).**

**Introduction:**

We first reported that platelets harbor more than 25 angiogenesis regulatory proteins (9). We subsequently found (10), that some of these proteins can be taken up by platelets in the presence of a microscopic implanted tumor in mice, even in the absence of coagulation. Therefore, platelets can detect the release of angiogenesis stimulators as well as inhibitors from tumors.

**Key Research Accomplishments:**

**The platelet angiogenesis proteome as a potential biomarker of the non-angiogenic phenotype of human breast cancer.** (With Giannoula Klement, M.D., David Cervi, Ph.D., and Nandita Bhattacharya, Ph.D.)

This project is led by Giannoula Klement, M.D. and David Cervi, Ph.D. They have assembled a team of collaborating scientists so that together they can rapidly validate mass spectroscopy as a general method of analyzing the platelet angiogenesis proteome as a detector of microscopic human cancer, including breast cancer.

(i) We discovered (with Tai-Tung Yip (Ciphergen), that platelet-associated PF4 is a reliable biomarker for early tumor detection across a spectrum of human tumor xenografts and that the platelet-associated PF4 is a reliable biomarker for early tumor detection across a spectrum of human tumor xenograft. (Manuscript in preparation, see “Reportable Outcomes,” below).
(ii) We found (with George Naumov) that among the initial angiogenesis regulatory proteins sequenced in this study (Figure 4) human breast cancer, (MDA-MB-436) induced an elevated plasma bFGF and CTAPIII (connective tissue activating protein III) in mice bearing tumors that were non-angiogenic. This breast cancer also induced an elevated platelet VEGF>CTAPIII>PDGF>PF4 (platelet factor 4) profile in mice bearing tumors that were angiogenic and grossly visible.

(iii) By conducting a time course experiment with human breast cancer xenograft clone MDA-MD-436, it was discovered that significant increases in bFGF and PDGF were found in platelets, but not in plasma, as early as day 32 after implantation of breast cancer that was non-angiogenic and microscopic. (Figure 5).

(iv) By conducting a time course experiment with human breast cancer clone MDA-MB-415, we found (with George Naumov) that significant increases in platelet-associated bFGF and PF4 (platelet factor 4) were evident by day 30 after subcutaneous implantation of non-angiogenic tumor cells in SCID immunodeficient mice (Figure 6a). The plasma angiogenesis proteome revealed an elevation of PDGF and PF4 and decreased bFGF at day 30, while VEGF and did not increase until day 120 (Figure 6b).
Reportable Outcomes:

A manuscript entitled, “The platelet-associated PF4 is a reliable biomarker for early tumor detection across a spectrum of human tumor xenografts” has been prepared for submission. The study reports that the changes observed with PF4 are reproducible and that it is consistently elevated early when non-angiogenic tumors are at a microscopic size.
Fig. 7: Identification of the platelet- and plasma-derived candidate proteins.

Figure. 8 a & b.: SELDI-ToF MS/MS of digested candidate biomarker, 1351.76Da.
Conclusion and Future Plans.

Angiogenesis regulatory proteins have been identified in platelets that change their concentrations in the presence of microscopic implanted human tumors in SCID mice. In addition to these proteins, several other signature proteins of tumor onset and progression are under investigation.

We plan to identify these proteins as well, and to determine whether they can be used in a multi-factorial analysis that incorporates the angiogenesis proteome: (i) to detect the presence of microscopic (non-angiogenic) human breast cancer; (ii) to detect the switch to the angiogenic phenotype, and possibly (iii) to distinguish among different tumor types.

Platelet factor 4 is emerging as a putative general biomarker of the non-angiogenic phenotype of microscopic dormant human tumors, and especially breast cancer.

V. OVERALL CONCLUSIONS.

a. Summary. This Progress Report reveals advances of our knowledge in the three major areas of the research program: (i) The molecular mechanisms by which human breast cancer switches from non-angiogenic, in situ cancer, to
angiogenic cancer capable of rapid growth and metastasis; (ii) The mechanisms by which the primary tumor must repress endogenous angiogenesis inhibitors normally in its local tumor bed, in order to induce angiogenesis; and (iii) The study of platelets as a vehicle for selecting and collecting angiogenesis regulatory proteins elaborated by the earliest microscopic breast cancers.

b. “So What” Section. All three aspects of this research are amenable to translation into clinical practice for the improvement of the treatment of breast cancer. For example, the thrombospondin-1 repressor protein, discovered by Randolph Watnick to be secreted by tumors to prepare future metastatic sites, could become the target for the development of an antibody that would neutralize the repressor protein. Therefore, whenever a breast cancer was removed, a simple test for the presence of this repressor protein, would alert the oncologist to administer the anti-repressor antibody, which would prevent future metastases from becoming angiogenic. This would be a non-toxic approach to prevent tumor recurrence.

Alternatively, as we come to understand the platelet angiogenesis proteome in depth, it is becoming ever more obvious that the detection of tumor angiogenesis in microscopic breast cancer, may someday permit the “treatment of a biomarker itself,” years before any symptoms of recurrent cancer appear and long before any conventional imaging methods could detect the anatomical location. In fact, the platelet study is teaching us that anatomical location of cancer may not be necessary to successfully treat early disease, just as anatomical location is no longer necessary to treat infection with antibiotics, or to treat atherosclerosis with statins.

VI. REFERENCES.


VII. APPENDICES:

A Model of Human Tumor Dormancy: An Angiogenic Switch From the Nonangiogenic Phenotype

George N. Naumov, Elise Bender, David Zurakowski, Soo-Young Kang, David Sampson, Evelyn Flynn, Randolph S. Watnick, Oddbjorn Straume, Lars A. Akslen, Judah Folkman, Nava Almog

Background: Microscopic human cancers can remain dormant for life. Tumor progression depends on sequential events, including a switch to the angiogenic phenotype, i.e., initial recruitment of new vessels. We previously demonstrated that human tumors contain tumor cell populations that are heterogeneous in angiogenic activity. Here, we separated angiogenic from nonangiogenic human tumor cell populations and compared their growth. Methods: Severe combined immunodeficient (SCID) mice were inoculated with nonangiogenic human MDA-MB-436 breast adenocarcinoma, KHOS-24OS osteosarcoma, or T98G glioblastoma cells. Most of the resulting tumors remained microscopic (<1 mm diameter), but some eventually became angiogenic and enlarged and were used to isolate angiogenic tumor cells. Angiogenic and nonangiogenic tumor cells were inoculated into SCID mice, and time to the development of palpable tumors was determined. Cell proliferation was assayed in vitro by growth curves and in vivo by staining for proliferating cell nuclear antigen or Ki67. Microscopic tumors from both tumor cell populations were examined for histologic evidence of vascular development 14 days after inoculation in mice. Expression of the angiogenesis inhibitor thrombospondin-1 was examined by immunoblotting. Results: Nonangiogenic tumors of each tumor type developed palpable tumors after means of 119 days (range: 53–185 days) for breast cancer, 238 days (184–291 days) for osteosarcoma, and 226 days (150–301 days) for glioblastoma. Angiogenic cells developed palpable tumors within 20 days after inoculation. However, nonangiogenic and angiogenic cells of each tumor type had similar proliferation rates. Fourteen days after tumor cell inoculation, tumors from angiogenic cells showed evidence of functional vasculature. In contrast, nonangiogenic tumors remained microscopic in size with absent or nonfunctional vasculature. Thrombospondin-1 expression was statistically significantly lower (by five- to 23-fold, depending on tumor type) in angiogenic than nonangiogenic cells. Conclusions: This model provides a conceptual framework and a reproducible in vivo system to study unresolved central questions in cancer biology regarding the initiation, reversibility, and molecular regulation of the timing of the angiogenic switch. [J Natl Cancer Inst 2006;98:316–25]
genetically modified tumor cells [i.e., Ras-transfected dormant cells (17–19)].

To better understand the pathogenesis and timing of the angiogenic switch in human tumors, we developed a novel approach in which we created separate populations of nonangiogenic and angiogenic tumor cells in vivo without using molecular alterations. We used this approach to study three different human cancer types in severe combined immunodeficient (SCID) mice: breast cancer, osteosarcoma, and glioblastoma. For angiogenic and nonangiogenic populations of each tumor type, we compared proliferation rates in vitro and in vivo and determined ratios of proliferation to apoptosis. The angiogenic switch in vivo was characterized by examining changes in the vasculature.

**METHODS**

**Establishment of Nonangiogenic and Angiogenic Human Cancer Cell Lines**

Human breast adenocarcinoma (MDA-MB-436), osteosarcoma (KHOS-240S), glioblastoma (T98G), and human embryonic kidney (293T) cell lines were obtained from the American Type Culture Collection (ATCC, Manassas, VA) and confirmed to be free of mycoplasma. The breast carcinoma, osteosarcoma, and glioblastoma cell lines were defined as nonangiogenic based on: 1) the absence of angiogenic activity, as evidenced by repulsion of existing blood vessels and/or absence of microvessels within the tumor; 2) their growth to only approximately 1 mm in diameter or less in vivo, at which time further expansion stopped; 3) the apparent absence of “tumor take” for at least 130–238 days, until the emergence of the angiogenic phenotype; and 4) the fact that the tumors remained harmless to the host until they switched to the angiogenic phenotype.

In mice injected with nonangiogenic human tumor cells (i.e., the cell lines as originally obtained from ATCC), occasional tumors grew for each of the three cancer types following a prolonged dormancy period. These vascularized tumors were removed under sterile conditions, and tumor cell lines were established from them using standard tissue culture techniques. The cell lines derived from the nonangiogenic (i.e., dormant) tumors that had switched to the angiogenic phenotype were called “angiogenic” because they produced large angiogenic tumors (up to 2000 mm$^3$) within 1 month of their subsequent inoculation into mice. For breast cancer, a single cell–derived clone, designated clone A1, was also isolated from the angiogenic subpopulation. This clone formed occasional angiogenic tumors following a prolonged dormancy period.

All cell lines were maintained in tissue culture (37 °C, 5% CO$_2$, humidified atmosphere). Breast cancer MDA-MB-436 cells were maintained in Dulbecco’s modified Eagle medium (DMEM; Invitrogen, Carlsbad, CA) supplemented with 10% fetal bovine serum (FBS; Invitrogen). Osteosarcoma KHOS-240S cells were maintained in Minimum Essential Medium (MEM; Invitrogen) supplemented with 10% FBS and 0.1 mM nonessential amino acids. Brain tumor T98G cells were maintained in MEM supplemented with 10% FBS, 0.1 mM nonessential amino acids, and 1 mM sodium pyruvate. For in vivo experiments, subconfluent monolayers of tumor cells were harvested by trypsinization and resuspended in DMEM to a final concentration of 2.5 × 10$^7$ cells/mL.

**In Vitro Tumor Cell Proliferation**

In vitro growth curves were generated by plating 5 × 10$^4$ cells in 60-mm dishes in triplicate. The cells were counted daily over a 7-day period. In an additional experiment, cells were collected during the exponential growth phase for subsequent histologic characterization (staining for Ki67).

**Anchorage-Independent Growth In Vitro**

For all three cancer types, the ability of nonangiogenic and angiogenic cells to undergo anchorage-independent growth was compared. In brief, 10$^4$ cells were suspended in 5 mL of 0.4% agar and plated in triplicate on 10-cm plates that had been pre-coated with 4 mL of 0.6% agar. An immortalized and transformed human kidney cell line (293T) was used as a positive control. Plates were monitored every other day, and when colonies became evident (after 3 weeks), plates were incubated for 8 hours with 2 mL of thiazolyl blue tetrazolium bromide (MTT) (Sigma, St. Louis, MO) to visualize viable cell colonies. The number of colonies was then counted from 10 digital images of random high-power fields for each plate.

**Subcutaneous Tumor Growth**

SCID male mice aged 6–8 weeks (Massachusetts General Hospital, Boston) were used for in vivo studies and were cared for in accordance with the standards of the Institutional Animal Care and Use Committee (IACUC) under a protocol approved by the Animal Care and Use Committee of the Children’s Hospital Boston. Mice were anesthetized using a 2% isoflurane (Baxter, Deerfield, IL) inhalation oxygen mixture. Suspensions of 5 × 10$^6$ human breast cancer (MDA-MB-436), osteosarcoma (KHOS-240S), or glioblastoma (T98G) cells in 0.2 mL of DMEM were then inoculated subcutaneously into the lower-right quadrant of the flank of each mouse. For in vivo characterization of the dormancy period, we inoculated mice as follows: 21 mice were inoculated with nonangiogenic breast cancer cells, 11 mice with angiogenic breast cancer cells, and 12 mice with nonangiogenic clone A1 tumor cells; 18 mice with nonangiogenic osteosarcoma cells and 11 mice with angiogenic osteosarcoma cells; five mice with nonangiogenic glioblastoma cells and five mice with angiogenic glioblastoma cells. Mice were monitored every other day for palpable tumors at the site of tumor cell inoculation. Tumors could be detected by palpation in these shaved SCID mice once the tumors reached approximately 50 mm$^3$. Once tumors became palpable, tumor size was measured twice a week. Mice were killed by cervical dislocation when the tumor size reached approximately 1200 mm$^3$. To assess tumor growth kinetics, tumors were measured from the time that they were first palpable until they reached 500 mm$^3$. Mice that did not form palpable tumors were monitored for tumor growth for up to 350 days after tumor cell inoculation. The number and proportion of angiogenic tumors that had switched from dormancy was compared between all three cancer types at day 160 after tumor cell inoculation. Representative mice were killed at various time points for histology of microscopic and macroscopic tumors (for example, at 14, 47, and 77 days after tumor cell inoculation).

Additional mice were inoculated with 5 × 10$^6$ MDA-MB-436 breast cancer cells for morphologic and histologic analysis as follows: 25 mice were inoculated with angiogenic, 30 mice with...
nonangiogenic, and 30 mice with nonangiogenic clone A1 breast cancer cells. Tumor cell proliferation and apoptosis were analyzed at the following intervals: 4–7 days (10 mice) and 29–40 days (10 mice) after inoculation with angiogenic tumor cells; 4–14 days (15 mice) and 40–70 days (10 mice) after inoculation with nonangiogenic tumor cells, and 4–15 days (15 mice) and 40–70 days (10 mice) after inoculation with nonangiogenic clone A1 breast cancer cells.

Histology and Immunohistochemistry

Tumor tissue was excised from mice killed by cervical dislocation, rinsed in ice-cold phosphate-buffered saline (PBS), fixed in 4% paraformaldehyde, and embedded in paraffin. Representative 4-μm tumor cross-sections were cut from paraffin-embedded tissue, and sets of four adjacent sections were stained as follows: 1) To identify proliferating cells, section 1 was stained with a 1:150 dilution of PC10 monoclonal antibody (DAKO, Carpinteria, CA) against proliferating cell nuclear antigen or with a 1:50 dilution of the monoclonal antibody against Ki-67 (M7240, clone MIB-1; DakoCytomation, Copenhagen, Denmark) after antigen retrieval for 20 minutes in citrate buffer in a microwave at 500 W; 2) To evaluate tissue quality and tumor morphology, section 2 was stained with hematoxylin and eosin; 3) To assess fragmented DNA, section 3 was labeled by the terminal deoxynucleotidyl transferase biotin–dUTP nick-end labeling (TUNEL) technique (ApopTag kit, Intergen, Purchase, NY) according to the method of Gavrielli et al. (20); 4) To identify endothelial cells, section 4 was stained with a 1:250 dilution of anti-CD31 antibody (Pecam; BD Biosciences, San Diego, CA) after antigen retrieval by Proteinase K treatment. Necrotic areas were avoided for all histologic analyses.

In Vivo Permeability Assay

A modified Miles assay (21) was performed to investigate differences in vascular permeability in size-matched nonangiogenic and angiogenic tumors. Five mice per group were inoculated with 5 × 10^6 angiogenic, nonangiogenic, or nonangiogenic clone A1 human breast cancer cells. The modified Miles assay was performed at day 14 after tumor cell inoculation. Mice were anesthetized by intraperitoneal injection of Avertin (150 mg/kg of body weight) and were then inoculated intravenously with filtered Evans blue solution (100 μL of 1% in PBS per mouse), which was allowed to circulate systemically for 30 minutes while the mice were maintained at 30 °C. All mice were then perfused systemically with PBS (pH 7.4) for 3 minutes at a pressure of 120 mm Hg from an 18-gauge cannula that was inserted into the aorta via an incision in the left ventricle. The right atrium was then incised to create a route for the saline to escape. Patches of the dorsal skin of each mouse were harvested to permit visualization of dye leakage from blood vessels to subcutaneous tumor tissues, to the skin overlying the tumor, and to control skin from the contralateral flank (this perfusion ensured that no dye remained in the vascular lumen). For each mouse, three types of tissues (tumor, overlying and control skin) were excised, weighed, and placed in 1 mL of formamide (Sigma) for 5 days at room temperature to extract all dye from the tissues. The extracted dye was quantified by measuring optical density at 620 nm of the formamide at the end of the 5-day incubation. All optical densitometry values were normalized to tumor (or skin) tissue weight.

ELISA Quantification of Secreted VEGF_{165}, bFGF, and Thrombospondin-1

Nonangiogenic and angiogenic tumor cells from each of the three cancer types were plated in triplicate (5 × 10^6 cells/15 cm plate) in 15 mL of DMEM supplemented with 10% FBS. After 24 hours, the medium was replaced with 15 mL of serum-free medium. The medium was then collected, and enzyme-linked immunosorbent assay (ELISA) kits were used to determine concentrations of human vascular endothelial growth factor (VEGF_{165}; R&D, Minneapolis, MN), basic fibroblast growth factor (bFGF; R&D), and thrombospondin-1 (CYT Immune Science, College Park, MD) according to the manufacturers’ protocols. For each sample, blank values (i.e., those for serum-free media) were subtracted, and mean results were normalized per 10^4 cells. This assay was performed in triplicate.

Protein Extraction and Immunoblot Analysis

Angiogenic and nonangiogenic breast cancer cells (MDA-MB-436) were grown in 10-cm tissue culture plates in DMEM supplemented with 10% FBS. When cells reached 80% confluence, half the plates were treated with the phosphoinositide 3-kinase (PI3K) inhibitor LY294002 (Calbiochem, San Diego, CA) at a final concentration of 10 μM in DMEM with 0.1% FBS for 12 hours. The remaining plates were untreated. For immunoblotting, cells were harvested by mechanical scraping into 4 °C PBS. Cell pellets were then lysed in 50 mM Tris–HCl (pH 7.4), 150 mM NaCl, 1% NP40, 1 mM sodium orthovanadate, 5 mM NaF, 20 mM β-glycerophosphate, and complete protease inhibitor (Roche, Indianapolis, IN). Fifty micrograms of protein was loaded in the wells of a 4%–12% precast Bis–Tris gel (Bio-Rad, Hercules, CA). Membranes were blocked with 5% nonfat milk and incubated with primary antibodies to c-Myc (Abcam, Cambridge, UK), phospho-Myc (Cell Signaling Technology, Beverly, MA), thrombospondin-1 (LabVision, Fremont, CA), and β-actin (Abcam, Cambridge, MA). Antibody binding was detected by incubating blots with horseradish peroxidase–conjugated secondary antibodies, either goat anti–mouse (c-Myc, thrombospondin-1, and β-actin) or goat anti–rabbit (p-Myc) (Jackson Immunoresearch Labs, West Grove, PA). Bands were visualized with enhanced chemiluminescence reagent (Pierce, Rockford, IL).

Statistical Analysis

The time to development of a palpable tumor (i.e., after tumor cell inoculation) was used as the endpoint of the dormancy period in survival analyses. For each of the three tumor types, times to palpable tumor for the angiogenic, nonangiogenic, and nonangiogenic clone A1 (breast cancer) cells were compared using the Kaplan–Meier product limit method (22). Confidence intervals around estimated survival curves were calculated using Greenwood’s formula (22). In a subgroup analysis, we compared time to palpation for angiogenic breast carcinoma, osteosarcoma, and glioblastoma tumor cell populations. The log-rank test was used to compare the statistical significance of differences between survival curves (23). For tumors that switched to the angiogenic phenotype, we compared the median times from initial tumor palpation (i.e., from a size of approximately 50 mm^3) until the tumor reached a size of 500 mm^3 using the nonparametric
Mann–Whitney U test. The Student’s t test was used to compare angiogenic and nonangiogenic tumor cell lines with respect to continuous data including in vitro tumor cell doubling times, anchorage-independent growth, vascular permeability, and ELISA. Two-tailed P values less than .05 were considered statistically significant. Statistical analyses were performed with the SPSS package (version 13.0 SPSS, Chicago, IL).

**RESULTS**

**In Vitro Growth of Nonangiogenic and Angiogenic Tumor Cells**

In vitro assessment of doubling times of all three cancer types showed no statistically significant difference between the nonangiogenic and angiogenic cell lines (Fig. 1, A–C). During the exponential tumor cell proliferation phase, approximately 90% of both nonangiogenic and angiogenic breast cancer cells expressed comparable levels of Ki-67 protein (data not shown).

We then analyzed the differences in anchorage-independent growth between nonangiogenic and angiogenic tumor cells for all three tumor types (Fig. 2). Both nonangiogenic and angiogenic subpopulations of the breast carcinoma, osteosarcoma, and glioblastoma formed colonies 3 weeks after plating in soft agar with no statistically significant differences (all \( P > .05 \)) in the number of colonies formed.

**Subcutaneous Tumor Growth**

To investigate in vivo tumor growth, we inoculated mice with either angiogenic or nonangiogenic cells from each cancer type and monitored them for as long as 350 days. Additional mice were inoculated with the nonangiogenic clone A1 breast cancer cells. The number and proportion of angiogenic tumors that had switched from dormancy within a period of 160 days after tumor cell inoculation was determined for all mice. For breast carcinoma, palpable tumors developed in all seven mice inoculated with angiogenic cells, in four of the five mice inoculated with nonangiogenic cells, and in three of the eight mice inoculated with nonangiogenic clone A1 cells. For osteosarcoma, angiogenic tumors developed in all six mice inoculated with angiogenic cells, in one of the eight mice inoculated with nonangiogenic cells, and in three of the eight mice inoculated with nonangiogenic cells. For glioblastoma, angiogenic tumors developed in all five mice inoculated with angiogenic...
cells and in one of the five mice inoculated with nonangiogenic cells (Fig. 1, D–F).

Kaplan–Meier analysis was used to compare the time between inoculation of tumor cells and detection of palpable tumors for nonangiogenic and angiogenic cell populations of all three cancer cell lines (Fig. 3). This analysis accounted for those mice in which tumor cells switched to the angiogenic phenotype and formed tumors as well as those mice that remained free of palpable tumor for a prolonged time and were censored at various time points during the dormancy period for histological analysis (as denoted by tick marks along the Kaplan–Meier curves in Fig. 3). The mean times between tumor cell inoculation and the development of palpable tumors (i.e., tumors of approximately 50 mm³) were 19 days (95% confidence interval [CI] = 16 to 22 days) in mice (n = 11) inoculated with angiogenic breast cancer cells, 119 days (95% CI = 150 to 301 days) for mice (n = 5) inoculated with nonangiogenic breast cancer cells, 238 days (95% CI = 184 to 291 days) for mice (n = 18) inoculated with nonangiogenic osteosarcoma cells and 234 days (95% CI = 199 to 269 days) in mice (n = 12) inoculated with nonangiogenic clone A1 breast cancer cells (log-rank P < .001 for comparison with angiogenic cells). Mean times to develop palpable tumors were 21 days (95% CI = 18 to 25 days) in mice (n = 11) inoculated with angiogenic osteosarcoma cells and 238 days (95% CI = 184 to 291 days) for mice (n = 18) inoculated with nonangiogenic osteosarcoma cells (log-rank P < .001). Finally, mean times to palpable tumor were 20 days (95% CI = 16 to 24 days) for mice (n = 5) inoculated with angiogenic glioblastoma cells and 226 days (95% CI = 150 to 301 days) for mice (n = 5) inoculated with nonangiogenic glioblastoma cells (log-rank P = .002). By contrast, comparison of time to develop a palpable tumor for angiogenic tumor cell populations of the three cancer types showed no differences (P = .15). Similar dormancy times were observed for the breast and glioblastoma tumors when inoculated orthotopically in the mammary fat pad or brain (George Naumov, unpublished data).

To determine whether tumors from nonangiogenic cells, once they emerge from dormancy, have growth kinetics similar to those of tumors from angiogenic cells, we assessed the kinetics of tumor growth in mice that developed palpable tumors from both angiogenic and nonangiogenic cell lines of each cancer type (including for breast cancer, clone A1). For this analysis, we determined the time from the detection of a palpable tumor to the development of tumors either lacked vasculature (CD31 negative, Fig. 4, H, M) or CD31 positive (Fig. 4, K) and were filled with red blood cells (Fig. 4, I). By contrast, all angiogenic tumors either lacked vasculature (CD31 negative, Fig. 4, H, M) or contained sparse microvessels that lacked red blood cells (Fig. 4, I, N). At this early time point, nonangiogenic tumors either appeared white or contained microvessels mainly at the periphery of the lesion by gross examination (Fig. 4, C, D). By contrast, all angiogenic tumors appeared red and vascularized by gross examination (Fig. 4, A). Histologic examination of the nonangiogenic tumors at day 14 after inoculation showed that a minority of tumors (<10%) were completely avascular with no evidence of intratumoral endothelial cells, as demonstrated by absence of CD31 staining (Fig. 4, M); most tumors showed histologic evidence of undeveloped (CD31 positive) microvasculature that lacked lumens, had no red blood cells (Fig. 4, I), and possibly contained a few blind vascular sprouts that were CD31 positive (Fig. 4, N). Macroscopic tumors that had spontaneously switched

Fig. 3. Kaplan–Meier analysis of time from tumor cell inoculation to palpable tumor detection. The persistence of mice free of palpable tumors was represented using the Kaplan–Meier analysis. Mice were inoculated with either angiogenic or nonangiogenic tumor cell lines: (A) Breast cancer MDA-MB-436 (angiogenic, n = 11 mice; nonangiogenic, n = 21; nonangiogenic clone A1, n = 12), (B) Osteosarcoma KHOS-24OS (angiogenic, n = 11; nonangiogenic, n = 18). (C) T98G glioblastoma (angiogenic, n = 5; nonangiogenic, n = 5). A drop in the curve indicates the occurrence of a palpable tumor at the corresponding time since tumor cell inoculation. Tick marks denote mice euthanized for histologic analysis that had not developed a palpable tumor at that time (i.e., censored data). Log-rank tests for differences between angiogenic and nonangiogenic cell populations yielded P < .001 for breast cancer and osteosarcoma and P = .002 for glioblastoma.

Histologic Differences Between Nonangiogenic and Angiogenic Tumors

We analyzed tumors for evidence of blood vessels on day 14 after tumor cell inoculation, when mice inoculated with either the angiogenic or nonangiogenic tumor cells had only microscopic tumors (Fig. 4, A, C, D), and when tumors reached 1000–2000 mm³ (Fig. 4, B, E). Fig. 4 shows results for breast cancer cells; results for osteosarcoma and glioblastoma were similar (data not shown). At day 14 after inoculation, angiogenic tumors (i.e., derived from angiogenic cell lines), although still less than 1 mm in diameter, contained functional blood microvessels that were CD31 positive (Fig. 4, K) and were filled with red blood cells (Fig. 4, F). By contrast, at the same time point nonangiogenic tumors either lacked vasculature (CD31 negative, Fig. 4, H, M) or contained sparse microvessels that lacked red blood cells (Fig. 4, I, N). At this early time point, nonangiogenic tumors either appeared white or contained microvessels mainly at the periphery of the lesion by gross examination (Fig. 4, C, D). By contrast, all angiogenic tumors appeared red and vascularized by gross examination (Fig. 4, A). Histologic examination of the nonangiogenic tumors at day 14 after inoculation showed that a minority of tumors (<10%) were completely avascular with no evidence of intratumoral endothelial cells, as demonstrated by absence of CD31 staining (Fig. 4, M); most tumors showed histologic evidence of undeveloped (CD31 positive) microvasculature that lacked lumens, had no red blood cells (Fig. 4, I), and possibly contained a few blind vascular sprouts that were CD31 positive (Fig. 4, N). Macroscopic tumors that had spontaneously switched
Fig. 4. Histologic analysis of tumors from nonangiogenic and angiogenic human MDA-MB-436 breast cancer cells. Photomicrographs of representative tumors are shown at day 14 (panels A, C, D) and day 47 or 77 (panels B, E) after inoculation for both nonangiogenic and angiogenic tumor variants (human breast cancer). Panels A and B represent microscopic and macroscopic tumors generated from angiogenic tumor cell lines, respectively. Panels C and D represent microscopic tumors, generated from a nonangiogenic tumor cell line, that differ in morphologic appearance. Panel E represents a macroscopic tumor, generated from a nonangiogenic cell line, that has spontaneously switched to the angiogenic phenotype from tumor dormancy. For each tumor, representative hematoxylin and eosin–stained sections (F–J) and CD31-stained sections (K–O) in high power field of views (×400) are shown. At day 14 after inoculation, angiogenic tumors (although still less than 1 mm in diameter) contained functional blood microvessels that were CD31 positive (K) and were filled with red blood cells (F). At the same time point, size-matched nonangiogenic tumors either lacked vasculature (absence of CD31 stain [M] and red blood cells [H]) or contained microvessels (N) that were not filled with red blood cells (I). Ratios of tumor cell proliferation to apoptosis were determined by immunohistochemical staining for Ki67 and terminal deoxynucleotidyl transferase biotin–dUTP nick-end labeling (TUNEL). These ratios were statistically significantly lower for microscopic than macroscopic tumors from the nonangiogenic cell line (P = .002) but not for the angiogenic line or nonangiogenic clone A1. Bars represent means based on at least 10 independent mice, with upper 95% confidence intervals.
to the angiogenic phenotype in mice inoculated with nonangiogenic tumor cells showed histologic evidence of vessels with open lumens (Fig. 4, O) that were filled with red blood cells (Fig. 4, J), similar to the microvasculature of tumors derived from angiogenic cell lines (Fig. 4, G, L).

Next, we analyzed breast tumor cell proliferation and apoptosis to determine whether tumor dormancy resulted from tumor cell quiescence. Histologic studies of tumors generated from the angiogenic cell line revealed similar ratios of tumor cell proliferation to apoptosis in tumors generated up to 14 days after inoculation and in tumors that had reached a size of more than 1000 mm³ (around day 40). For tumors generated from the angiogenic breast cancer cell line, the mean ratios of Ki-67 stain to TUNEL assay results, in arbitrary units, were 7.6 (95% CI = 4.8 to 10.4) for 14-day microscopic tumors and 6.3 (95% CI = 3.8 to 8.8) for 40-day macroscopic tumors (not statistically significantly different, \( P = .43 \)) (Fig. 4, P). Similar ratios were observed in tumors derived from the nonangiogenic clone A1 cells, both during early dormancy (4–14 days after inoculation) and after the switch to the angiogenic phenotype, i.e., past day 70 of inoculation (early dormancy, 6.8 [95% CI = 4.6 to 8.9]; angiogenic, 5.8 [95% CI = 3.8 to 7.8]; \( P = .53 \)). However, the ratios in nonangiogenic tumors during early dormancy were statistically significantly lower than those in tumors that had switched to the angiogenic phenotype (early dormancy, 3.1 [95% CI = 2.1 to 4.1], angiogenic, 5.6 [95% CI = 4.5 to 6.6]; \( P = .002 \)). More than 50% of tumor cells were proliferative and more than 10% underwent apoptosis in all microscopic and macroscopic tumors analyzed. These findings indicate that nonangiogenic tumors, even before they have switched from dormancy to the angiogenic phenotype, have high proliferation rates and therefore are not quiescent (i.e., are not in a G0 state). These results are consistent with those of previous reports (13,17,24). Moreover, establishment of angiogenic and expanding tumors appeared to be independent of the tumor cell proliferation/apoptosis ratio.

Vascular Integrity and Functionality

To compare vascular integrity and functionality between angiogenic and nonangiogenic tumors, we performed a modified Miles assay. Vascular permeability was assessed within size-matched nonangiogenic and angiogenic tumors, as well as in the skin overlying the tumor and in control skin from the contralateral flank of the mouse (Fig. 5). We assessed permeability by measuring Evans blue dye that had leaked from the vasculature after removing intravascular dye by systemic perfusion of the mice with PBS (Fig. 5, B). Both macroscopic analysis (Fig. 5, A) and quantification of extracted dye indicated that angiogenic tumors retained the dye, and nonangiogenic tumors remained dye free. Vascular permeability within the skin that remained in contact with the tumor was comparable to that of skin obtained from the contralateral flank of the mouse. Thus, the observed differences in vascular permeability between angiogenic and nonangiogenic tumors appeared to be localized to the neoplastic tissue and did not affect the permeability of adjacent nonneoplastic tissue (i.e., skin).

Differences in Expression of Pro- and Antiangiogenic Proteins Between Nonangiogenic and Angiogenic Tumor Cells

Levels of bFGF and VEGF₁₆₅ (both of which are proangiogenic) and thrombospondin-1 (which is an angiogenesis inhibitor) secreted by tumor cells were quantified in the media of cells grown in vitro (Fig. 6). In all three cancer types, angiogenic tumor cells secreted much higher levels of bFGF than nonangiogenic cells (13-fold difference for breast cancer cells, 79-fold for osteosarcoma, and fourfold for glioblastoma; \( P < .001 \) for all three cancer types, Student's t test). All tumor cells secreted substantial levels of VEGF₁₆₅. However, whereas angiogenic osteosarcoma cells secreted 66-fold higher levels of VEGF₁₆₅ than nonangiogenic tumor cells (\( P < .001 \)), levels of VEGF₁₆₅ secreted by nonangiogenic and angiogenic tumor cells were similar for both breast carcinoma and glioblastoma. In all three cancer types, nonangiogenic subpopulations secreted statistically significantly higher levels of thrombospondin-1 than angiogenic cells (23-fold difference for breast cancer cells, sixfold for osteosarcoma, and fivefold for glioblastoma and breast carcinoma; \( P = .011 \), osteosarcoma and glioblastoma; \( P < .001 \)).

Repression of Thrombospondin-1 Activity via the PI3K–c-Myc Pathway Correlates With the Switch From Dormancy to the Angiogenic Phenotype

Because nonangiogenic cells secreted higher levels of thrombospondin-1 than angiogenic cells, we hypothesized that thrombospondin-1 expression plays a role during tumor dormancy.
than nonangiogenic cells. This finding confirmed the results obtained by measuring secreted thrombospondin-1. PI3K has previously been demonstrated to induce a signal transduction cascade leading to the phosphorylation of c-Myc and subsequent repression of thrombospondin-1 (19). Therefore, we sought to determine whether this pathway was active in the angiogenic cell line. We treated both angiogenic and nonangiogenic tumor cells with the PI3K inhibitor LY294002. Treatment with this compound caused thrombospondin-1 levels within angiogenic cells to increase but had no effect on the levels in nonangiogenic cells (or those in nonangiogenic clone A1 cells) (Fig. 7, B). Therefore, we conclude that the PI3K signaling pathway is responsible for the repression of thrombospondin-1 and is differently regulated in angiogenic and nonangiogenic tumor cells.

**DISCUSSION**

The results presented here show that the switch to the angiogenic phenotype in human cancer cells is highly reproducible in frequency and proportion of otherwise microscopic and dormant
tumors. The model for human tumor dormancy that we have developed here is novel because the angiogenic tumor cell lines had spontaneously switched from the nonangiogenic cell lines without the use of in vitro selection (by a particular tumor cell marker) or artificial genetic alterations (15,17–19). Our results suggest that the onset and extent of angiogenesis are critical determinants of tumor progression and growth. Also, our results indicate that the establishment of angiogenic and expanding tumors in vivo (i.e., the angiogenic switch) is independent of the ratio of tumor cell proliferation to apoptosis.

Histologic examination of microscopic nonangiogenic breast cancers revealed two categories of such tumors: 1) completely avascular tumors and 2) tumors containing empty lumens, without red blood cells, which we assume represent blind sprouts. Neither category of nonangiogenic tumors showed evidence of vascular permeability, most likely because of the absence of vasculature or presumably insufficient vascular sprouting. These categories are likely to represent early stages leading to the angiogenic switch. The gross difference between the nonangiogenic tumors and angiogenic tumors (i.e., white versus red tumors) is most likely due to the reactive hyperemia that accompanies the onset of blood flow after the angiogenic switch is completed in a previously hypoxic tumor.

In the experiments reported here, we used $5 \times 10^6$ tumor cells in each inoculation. We observed sparse microvessels in the nonangiogenic tumors up to 40 days after inoculation. These structures did not appear to be functional microvessels, as evidenced by lack of vascular lumens and absence of red blood cells. After 40 days, these microvascular structures could not be sustained, and they disappeared until the angiogenic switch. After the completion of this study, further work (data not shown) indicated that these sparse microvessels could be induced by excessive VEGF or other proangiogenic proteins carried over in the original tumor cell inoculum. Therefore, we decreased the inoculum by decrements from $5 \times 10^6$ to 500 tumor cells. We found the optimum inoculum to be $10^6$ tumor cells; this inoculum produced nonangiogenic, microscopic, dormant tumors that were completely avascular until the time of the angiogenic switch. Therefore, in future studies, we suggest that the model we now report could be optimized by using approximately $10^6$ tumor cells as the initial inoculum.

Although our findings indicate that the lack of angiogenesis is a rate-limiting step for tumor expansion in the dormant state, there may also be other processes involved in human tumor dormancy, such as differentiation programs, tumor cell survival, or immune response of the host. Although the precise mechanism and timing of tumor dormancy have not yet been completely elucidated, our novel approach provides a means to determine the molecular events and sequential steps of the angiogenic switch. The kinetics of individual stages might be of substantial practical importance for identifying an optimal therapeutic window in the treatment of early cancer.

In other animal models, the growth of angiogenic tumors has been associated with reduced expression of endogenous angiogenesis inhibitors, such as angioptatin (24), endostatin (25), tumstatin (26), pigment epithelium–derived factor (27), and thrombospondin-1 (28). Although it is likely that other endogenous angiogenesis inhibitors play a role in the switch to the angiogenic phenotype for other cancer types, in these experiments we focused on the differences in thrombospondin-1 expression between nonangiogenic and angiogenic tumor cells. In our experiments, thrombospondin-1 appeared to be a major negative regulator of angiogenesis. That is, thrombospondin-1 expression had to be reduced in the angiogenic tumor cells for the angiogenic switch to take place. This finding further strengthens Noël Bouck’s hypothesis that the onset of tumor angiogenesis is associated with a substantial decrease in thrombospondin-1 levels (29). Previous reports have shown that transfection of thrombospondin-1 into human breast cancer cell lines decreased angiogenesis, tumor growth, and metastasis in immunocompromised mice (19,30). Immunohistochemical analysis of human breast cancer tissue has shown marked increases in levels of thrombospondin-1 mRNA in stromal cells immediately adjacent to ductal carcinomas in situ compared with normal breast tissue (31). These findings raise the possibility that increased expression of thrombospondin-1 in stromal cells and cancer cells may inhibit angiogenesis induced by VEGF and bFGF.

Our dormancy model of breast cancer shows that thrombospondin-1 expression is regulated by the PI3K–c-Myc signaling pathway. This finding is consistent with those of previous reports (19,32–34). Western blot analysis demonstrated that the angiogenic MDA-MB-436 breast cancer cells expressed higher levels of c-Myc and lower levels of thrombospondin-1 than the nonangiogenic cell line or the nonangiogenic single cell–derived clone A1. Furthermore, when the activity of PI3 kinase was inhibited in the angiogenic cells, levels of p-Myc decreased and levels of thrombospondin-1 increased compared with levels in the nonangiogenic cells. We conclude that the PI3K–c-Myc pathway plays a key role in the regulation of thrombospondin-1 and may contribute to the observed differences in growth in vivo.

In our study, most of the nonangiogenic glioblastoma and osteosarcoma tumors never switched to an angiogenic phenotype during the lifetimes of the mice. By contrast, most of the nonangiogenic breast tumors did undergo the angiogenic switch. Therefore, the amount of time until the angiogenic switch appears to be distinct for a given tumor type and to differ among tumor types. We speculate that most human microscopic cancers can survive for prolonged periods by keeping the balance of pro- and antiangiogenic proteins in a steady state that favors dormancy.

The human tumor dormancy model has a few limitations. One limiting factor is that experiments characterizing the timing of the angiogenic switch can be time-consuming, lasting up to 1 year or longer. Consequently, up to this point, the mechanism of the angiogenic switch is still unclear and under investigation. Nonangiogenic tumors are invisible until the angiogenic switch, unless they are imaged using luciferase or green fluorescent protein.

Our results provide the basis for further analysis of dormant nonangiogenic tumors. Among the questions that it can be used to address are the following: 1) What are the determinants and mechanism for the “clock” that governs the angiogenic switch? Why is the timing of the switch to the angiogenic phenotype so predictable for a given tumor type? For example, why do 100% of angiogenic breast cancers become palpable at a median of 19 days, when less than 20% of nonangiogenic (clone A1) breast cancers switch to the angiogenic phenotype after approximately 1 year? In vivo growth of these single-cell clones demonstrates heterogeneity in the dormancy clock and will be used for subsequent molecular studies of this process. 2) What is the sequence of the different stages of the angiogenic switch, i.e., lumen formation, initiation of flow, and hyperemia? 3) Is the angiogenic switch reversible? Can angiogenic tumor cells revert to nonangiogenic cells? If not, why is it possible to isolate nonangiogenic tumors?
tumor cells from angiogenic tumors, such as the nonangiogenic clone A1 of breast cancer?

During the dormancy period, nonangiogenic tumors had high proliferation rates and were therefore not quiescent (i.e., were not in a G0 state). Moreover, the angiogenic switch appeared to be independent of the tumor cell proliferation/apoptosis ratio. Therefore, angiogenesis and proliferation/apoptosis represent different programs in tumor progression. These models can permit further studies of the angiogenic switch, especially the different stages and timing of the process. Also, this approach may be suitable for the study of sensitive angiogenesis-based biomarker assays and novel treatment strategies. We, along with others, are currently developing an angiogenesis-based panel of blood and urine biomarkers that can be quantified and used to detect microscopic tumors before or during the angiogenic switch. If this approach is feasible, microscopic tumors could possibly be treated years before they become symptomatic or their anatomical site is detectable. The use of this animal model has taught us that in the future, it may be possible to liberate the management of cancer from dependency on anatomical site.

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NOTES

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