Hydrophilic Agarose Macrobead Cultures Select for Outgrowth of Carcinoma Cell Populations That Can Restrict Tumor Growth

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Abstract
Cancer cells and their associated tumors have long been considered to exhibit unregulated proliferation or growth. However, a substantial body of evidence indicates that tumor growth is subject to both positive and negative regulatory controls. Here, we describe a novel property of tumor growth regulation that is neither species nor tumor-type specific. This property, functionally a type of feedback control, is triggered by the encapsulation of neoplastic cells in a growth-restricting hydrogel composed of an agarose matrix with a second coating of agarose to form 6- to 8-mm diameter macrobeads. In a mouse cell model of renal adenocarcinoma (RENCA cells), this process resulted in selection for a stem cell–like subpopulation which together with at least one other cell subpopulation drove colony formation in the macrobeads. Cells in these colonies produced diffusible substances that markedly inhibited in vitro and in vivo proliferation of epithelial-derived tumor cells outside the macrobeads. RENCA cells in monolayer culture that were exposed to RENCA macrobead-conditioned media exhibited cell-cycle accumulation in S phase due to activation of a G2/M checkpoint. At least 10 proteins with known tumor suppression functions were identified by analysis of RENCA macrobead-conditioned media, the properties of which offer opportunities to further dissect the molecular basis for tumor growth control. More generally, macrobead culture may permit the isolation of cancer stem cells and other cells of the stem cell niche, perhaps providing strategies to define more effective biologically based clinical approaches to treat neoplastic disease.

Introduction
We have recently reported the ability of various tumor cell lines to form tumor colonies following encapsulation in agarose–agarose macrobeads (1). Tumor colonies arise in the core of the macrobeads from a rare population of the 150,000 cells originally encapsulated. Approximately 99% of these cells undergo apoptosis within 1 week (1). When mouse renal adenocarcinoma (RENCA) cells are encapsulated in macrobeads, tumor colonies expand in size from a single cell to colonies containing several hundred or more cells. The initial rapid growth slows as the colonies enlarge to a relatively stable size. As such, encapsulated tumor colonies demonstrate aspects of a Gompertzian growth curve.

With regard to the growth patterns of tumors in vivo, Laird and colleagues (2, 3) and subsequently, Norton and colleagues and Speer and colleagues (4–7) have shown that both normal organs and tumors follow a similar Gompertzian (decremented exponential) growth curve that approaches an asymptote or falls off as they enlarge, consistent with feedback inhibition of growth. In the same way, the partial surgical removal of a tumor often induces tumor progression, analogous to partial resection of the liver which results in stimulation or “compensatory hyperplasia”. This is true for both laboratory models and in the clinic, suggesting that clinical reduction of tumor burden could actually promote tumor progression (8, 9). Building on these phenomena and the demonstrable loss of exponential growth of mammary tumors (10), but also taking into account other factors such as the internal structural

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heterogeneity ("anaplasia") of tumors generally (i.e., tumors as aggregates or "conglomerates" of growth centers) and the now-documented presence of cancer stem cells (11, 12) Norton and colleagues (4–6), as well as Speer and colleagues (7), have elaborated more detailed realistic and potentially clinically useful Gompertzian-based models that have helped to understand tumor growth and to optimize available antitumor therapy.

From the above demonstration of decremented exponential growth in tumors and the feedback inhibition that is implied by such models, it is not far to the hypothesis that the growth of a given tumor might be slowed or stopped by one or more biological signals indicating the presence of tumor mass, even when that mass is not actually present (8, 9, 13). Taken as a whole, the data are supportive of Prehn’s hypothesis that tumors, in fact, behave, “…as integrated organs rather than a collection of independent cells.” (13) As such, tumors respond to positive and inhibitory stimuli from both within and external to the tumor, at least in part in the same manner as normal organs and tissues.

We have previously reported the ability of agarose-encapsulated, insulin-producing porcine islets to provide protection from autoimmune and xenogeneic islet destructive responses for the treatment of type I diabetes (14–17). In the current article, we have taken advantage of the xenogeneic immunoprotection afforded by the agarose encapsulation to test the hypothesis that similarly encapsulated tumor cells are capable of providing growth-controlling signals to external tumor cells. The ability of encapsulated tumor cells to inhibit tumor growth would be in line with the notion of Prehn, namely, that tumor cells exhibit growth control analogous to developing organs as they reach their ultimate mass.

The encapsulation of mouse, feline, and human tumor cells of epithelial origin in an agarose matrix was performed to simulate tumor mass for both in vitro and in vivo studies. From observations on the behavior and molecular biology of such entrapped tumor cells, we have documented the existence of a novel growth regulatory mechanism that is neither species nor tumor-type specific.

Materials and Methods

Cell lines

The RENCA tumor cell line used for these experiments is a renal adenocarcinoma that arose spontaneously in Balb/c mice, originally obtained from the National Cancer Institute, (Bethesda, MD) and now available from ATCC (American Type Culture Collection) employing morphology and isoenzymology and/or Cytochrome C subunit I (COI) PCR assays for cell line authentication. RENCA cells were maintained in vitro (5% CO₂ + air at 37 °C) in tissue culture flasks (BD Biosciences) containing RPMI 1640 (Invitrogen) with 10% newborn calf serum (NCS; Invitrogen). Cell passages used were limited to no more than 20 from a frozen stock of these cells. DU145 (human prostate carcinoma), HCT116 (human colorectal carcinoma), J82 (human urinary bladder transitional cell carcinoma), and MCF7 (human mammary gland adenocarcinoma) were originally obtained from ATCC. DU145 cells were cultured in RPMI 1640 + 10% FBS. HCT116 cells were cultured in McCoy’s 5A + 10% FBS. J82 cells were cultured in MEM + 10% FBS + 1% NEAA, and MCF7 cells were cultured in MEM + 10% FBS.

RENSA macrobeads

RENSA agarose–agarose macrobeads were prepared essentially as previously described for islet macrobeads (18). Briefly, 100 μL of 0.8% agarose (HSB-LV; Lonza Copenhagen ApS, Vallensbak Strand) in MEM (Sigma-Aldrich) were mixed with 1.5 × 10⁵ RENCA cells. The agarose/cell suspension was expelled from a transfer pipette into mineral oil to form the core of the macro bead. Following washing with RPMI 1640 and overnight culture at 37 °C in 5% CO₂ and air, the core was rolled in approximately 1 mL of 4.5% agarose to apply an outer coat and transferred to mineral oil. For other cell lines, 100,000 to 250,000 cells per macrobead were used. Macrobeads were cultured in 90-mm Petri dishes (Nunc) at 10 macrobeads per 40 mL of RPMI 1640 with 10% NCS. Metabolic properties were evaluated using an MTT assay (Sigma-Aldrich).

Bioassays

For bioassays employing conditioned media, cells (15,000 per well) were seeded in 6-well plates in 4 mL fresh culture media or 5-day macrobead-conditioned media. For coculture bioassays, cells were seeded in 6-well plates (BD Biosciences), allowed to attach overnight, and cultured with macrobeads suspended in cell culture inserts (BD Biosciences). Cells were methanol-fixed, stained with 0.33% (w/v) neutral red and absorbance read at 540 nm with 630 nm as the reference wavelength. Inhibition was defined as the percent difference in Abs₅₄₀−₆₃₀nm between treated and untreated media.

Animals

Experiments were reviewed and approved by the IACUC of The Rogosin Institute. Balb/c mice were obtained from Charles River Laboratories. After a 7-day acclimatization period, tumors were started by injecting 2,500 RENCA cells under the capsule of the left kidney. Four days later, animals received 4 RENCA macrobeads via a small opening into the peritoneal cavity.

Image acquisition

Tumors were viewed with an Olympus model SZ6045 dissecting microscope using a bright-field illuminator (model SZH-ILLD, Olympus) and photographed with a Zeiss Axioscam MRc at 6× magnification using MRGrab v1.1 software (Carl Zeiss Vision GmbH). Formalin-fixed macrobead sections stained with hematoxylin and eosin (H&E) were viewed with an inverted microscope (Axiovert S 100) and photographed (Axiocam MRc) using MRGrab v1.1 software.

Time-lapse images of growing cells were obtained using an inverted phase contrast microscope (Olympus IX71), a Uni-blitz shutter and a Hamamatsu Orca ER B/W digital camera. Cells were maintained at 37 °C for microscopy by use of a Solent Scientific environmental chamber. Cells were kept in sealed 75-mm plastic flasks containing 8 mL of medium and...
5% CO₂ + air for up to 48 hours. Control experiments confirmed that growth continues normally in sealed flasks for at least 60 hours. Images were taken with exposure times of 100 milliseconds with 2 by 2 binning and interframe intervals of 5 minutes. Stored image stacks were converted to Cineplex time-lapse videos using Metamorph software (Universal Imaging) and further compressed to Quicktime videos with PremierePro (Adobe Systems, Inc.). All cells in the starting field were identified by location. Time-in-mitosis (as measured by time rounded up) and the fates of daughter cells were recorded visually by following each cell to the end of the video.

Isolation of RNA from RENCA monolayers

RENSA monolayers were maintained in RPMI 1640 supplemented with 10% NCS. For mRNA, trypsinized RENCA cells were seeded at 20,000 cells per 40-mm well (6-well plate) in 4 mL of RPMI 1640 + 10% NCS. Medium was replenished on day 2 and cells harvested on day 4 in late log growth. Total RNA was prepared using the RNeasy Mini Kit (Qiagen).

Gene microarrays and data analysis by GenMapp and MAPPFinder gene microarrays

RNA quality was confirmed with Total RNA Nano chips (Bioanalyzer 2100 (Agilent Technologies). Samples of total RNA meeting criteria (|rRNA > 30% of area, 28s/18s ratio > 1.5) were used for reverse transcription to cDNA. The standard target labeling assay (1–15 μg total RNA) was followed as per the Affymetrix Gene Chip Expression Analysis Technical Manual 1999. Biotin-labeled cRNA was then synthesized using the BioArray HighYield RNA transcript labeling kit (Affymetrix), purified by RNeasy Mini Kit (Qiagen), fragmented to a mean size of 35 to 200 base pairs and evaluated for quality by hybridization to Test 3 Chip (Affymetrix). Samples with background between 30 and 100, and scale factor between 1 and 3 were hybridized to Affymetrix Mu94v2 A, B and C chips in that order. All chips were scanned at the Genomics Resource Center of The Rockefeller University as per the Affymetrix Gene Chip Expression Analysis Technical Manual 1999.

The data set included 3 biological replicate samples each from RENCA monolayers cultured in fresh medium or RENCA macrobead-conditioned medium. Average probe intensities were globally scaled to a target intensity of 250. Data were normalized at the probe-level by GC-RMA using GeneTraffic software version 3.1.2. Data within each chip type were normalized separately and then combined in a single Excel worksheet. Fold change was calculated as log₂ (replete/fresh media). ANOVA P values were calculated for expression intensity from the 3 values per probe.

GenMapp 2.1.0 and MAPPFinder 2.0 software packages (www.GenMapp.org) were used and all probe sets measured in the experiment were used as input to GenMAPP. GO annotations were based on the GenMAPP Mm-Std_20051114.gdb database. The GenMAPP Criteria Builder was used to filter data using the following criteria: ANOVA greater than 100, log₂R greater than 0.20 for upregulated genes and log₂R greater than −0.20 for downregulated genes.

Flow cytometry

A Becton Dickinson FACScan flow cytometer was used with BD Cycle test PLUS DNA reagent kit to analyze RENCA cells. RENCA cells exposed to replete media from 105 to 117 days RENCA macrobeads or to fresh media were harvested from day 2 to 5 and 5 × 10⁷ to 1 × 10⁸ cells analyzed.

Protein isolation and mass spectroscopy

RENSA cells were conditioned to serum-free CDM4HEK293 media (Hyclone) + 4 mmol/L of l-glutamine and 1% P/S for 2 weeks following acclimation in SFM4HEK293 media (Hyclone) + 4 mmol/L of l-glutamine and 1% P/S according to manufacturer’s directions. RENCA macrobeads were cultured for 12 weeks in serum-free CDM4HEK293 media. Total protein was prepared from day 5 RENCA macrobead-conditioned media via acetone precipitation. Proteins were separated by 1D SDS-PAGE [1 mmol/L of 4%–12% Bis-Tris midl gel, MES-SDS Running buffer (Invitrogen) and stained using SimplyBlue Safe Staıın (Invitrogen)]. Visible bands were excised and submitted for protein identification by LC-ESI-MS + MS/MS (Protea Biosciences, Inc.). Briefly, in-gel digestion was performed with 500 ng trypsin in 50 mmol/L of ammonium bicarbonate buffer overnight. Peptide extraction was performed using 5% formic acid in 50% acetonitrile followed by rehydration with 50 mmol/L of ammonium bicarbonate buffer overnight. Three extraction cycles were performed per sample. Recovered peptides were lyophilized and reconstituted in 10 mmol/L of acetic acid and relyophilized followed by reconstitution in 0.1% formic acid in DI water. A Shimadzu LC-20AD HPLC and ACE 100 × 0.3-mm C18 capillary LC column by Advanced Chromatography Technologies was used for liquid chromatography at 35°C. ESI (electrospray ionization) mass spectrometry was performed on a Thermo LTQ XL Mass Spectrometer using Xcalibur 2.0SR2 acquisition software with electrospray ionization in positive ion mode with a spray voltage at 3.8 kV. The eight most intense ions with ion intensities above a threshold of 3,000 in each regular MS scan were subjected to MS/MS analyses. Database correlation analysis parameters included the use of ABI ProteinPilot software 3.0 with the Paragon search engine of the NCBI nr database (Mus musculus).

Results

In vitro and in vivo growth inhibitory effects of RENCA macrobeads

Shortly after the first appearance of macroscopically visible tumor colonies in the macrobeads at around 3 weeks post-encapsulation, 5-day conditioned media from RENCA macrobeads begins to show proliferation-suppressive activity. This antiproliferative effect increases over time from 0% at 2 to 3 weeks, to just under 20% by 5 weeks and ultimately slows the growth of RENCA cells in monolayer culture by up to 40% or more at 24 weeks postencapsulation (Fig. 1A). RENCA cell growth is increasingly inhibited when grown in macrobead-conditioned media from 5 to 7 days (Fig 1B). The inhibitory
effect plateaus by approximately 6 months postencapsulation and remains constant from 6 months to at least 2 years (data not shown).

To verify that the inhibitory capacity of the RENCA macrobead was a result of macrobead-secreted factors present in the conditioned media, and not a result of depleted media, several experiments were initiated. First, adherent RENCA cells, in fresh media, were exposed to RENCA macrobeads via a cell culture insert system in which macrobeads are suspended over freely growing RENCA cells by the use of a well insert. Using this well insert approach, RENCA cells demonstrated 39.7% growth inhibition, whereas cells exposed to conditioned media showed 44.8% growth inhibition (Fig. 1C). Secondly, concentrating macrobead-conditioned media followed by addition to fresh culture media (“replete media”) also inhibits the growth of RENCA monolayer cells. Analysis of conditioned medium has shown that cortisol, testosterone, estrogen, TGF-β, IFN-γ, and interleukin (IL)-2, IL-6, and IL-10 do not account for this activity (data not shown).

To determine whether the antiproliferative/antitumor effect of RENCA macrobeads was also present in vivo, Balb/c mice received 2,500 RENCA cells under the renal capsule in a series of experiments. Four days later, 4 RENCA macrobeads or 4 empty macrobeads were placed in the peritoneal cavity, whereas other tumor-induced mice received only sham surgery. Thirty days after implantation, the macrobead-treated mice had significantly smaller tumors (30%–60%) than those of control mice treated with empty macrobeads or surgery alone (Student’s t test: $P < 0.05$ and $P < 0.01$, respectively; Fig. 1D), suggesting that the growth inhibitory substance(s) produced by the macrobeads homed to the tumor site and inhibited growth.

Treatment of 54 dogs and cats with end-stage, naturally occurring tumors, including prostatic carcinoma, splenic hemangiosarcoma, and hepatocellular carcinoma in canines and gastrointestinal lymphoma, oral squamous cell carcinoma, and mammary carcinoma in felines, were confirmatory of this effect. Thirty-nine of the 51 evaluable dogs and cats in the study showed improvement in appetite and/or weight, activity and overall well being, irrespective of tumor state at implantation or tumor response to macrobead implantation and without any evident impairment of the immune system. Small group sizes, various tumor stages, and previous cancer therapies make statistical interpretation of efficacy difficult for every type of tumor treated. Nonetheless, significant examples of efficacy were found including a cat with GI lymphoma that received 5 RENCA macrobead implants and lived for 3 years as well as a cat with mammary carcinoma that received 4 implants and lived 8 years after initial diagnosis before eventually succumbing to recurrent disease. The most controlled group in this series with histologic confirmation of tumors and without previous cancer treatment were dogs with prostatic adenocarcinoma. These dogs ($n = 11$) received either 22 or 44 RENCA macrobeads per kilogram and survived a median of 177 days from diagnosis to elected euthanasia.
slow the growth of human epithelial cell lines including pros-
species or cell-type specific

Growth inhibition by encapsulated tumor cells is not
(survival days: 59*, 68, 76, 134, 177*, 255, 311, 328*, 340, 732; *
the hypothesis that RENCA cells exposed to conditioned

Growth inhibition by encapsulated tumor cells is not
species or cell-type specific

RENCA macro bead-conditioned medium was also able to
slow the growth of human epithelial cell lines including pro-
state DU145 cells (36%–40%; Fig. 2A), bladder J82 cells (17%–
43%; Fig. 2A) and colorectal HCT116 cells (43%–58%; Fig. 2A)
but had little effect on HeLa cells (0%–15%; not shown),
emphasizing that the effect operates across species lines, is not
specific to tumor type, at least within a series of epithelial-
derived tumors, and is independent of angiogenic or immu-
nologic mechanisms. Importantly, Figure 2A also dem-
strates that the inhibitory effect was not the result of
depleted conditioned media as the cell culture insert method
also produced equivalent inhibition (similar to Fig. 1C above).

Furthermore, other encapsulated tumor cell lines are also
able to inhibit freely growing tumor cells, across species lines
and tumor types, as illustrated by the inhibition of RENCA
cells (25%–35% inhibition) by encapsulated human bladder
J82 cells between the ages of 6 and 10 weeks (Fig. 2B). Encapsulated J82 cells also inhibit J82 monolayer cells
(16%–28% inhibition; Fig. 2B).

RENCA macrobeads induce cell-cycle alterations in
freely growing monolayer cells

To further investigate the inhibition of tumor cells in vitro,
cell growth was evaluated in RENCA cells exposed to reple-
t media (concentrated conditioned media brought up in fresh
media) by use of time-lapse videography (Fig. 3A). These
experiments revealed a 10% reduction in the number of cells
that entered mitosis during a 24-hour period between day 3
and 4 when cultured with replete media as compared with
fresh media (Fig. 3B). Of those cells entering mitosis, only 56%
went on to divide in replete media whereas 90% of cells
completed cell division when cultured in naive media. Non-
productive cell divisions, including those of cells that initiated
division, but ultimately apoptosed, were also significantly
increased when cells were grown in replete media (Fig. 3B).
The complete video of this experiment is available online as
Supplementary Figure 1.

These data suggest that RENCA cells exposed to conditioned
media undergo cell-cycle checkpoint delays with upregula-
tion of DNA damage surveillance and repair programs followed by
nonprogression of the cell cycle and death via apoptosis. In
fact, analysis of 48 hours of time-lapse videos, beginning on day
2 of growth, revealed an increase in normal cell cycle time from
24 hours for cells grown in fresh media to 30 hours for cells
grown in replete media. Cells that divided normally, whether in
fresh or conditioned medium, required equivalent time to
progress through mitosis (1.3 hours to 1.4 hours, respectively),
but cells with nonproductive divisions in conditioned medium
required significantly longer times (2.3±1.9 vs. 1.3±1.05 hours;
<0.001) before aborting mitosis or undergoing apoptosis.
During the 48-hour culture time, 52 cells produced 308 daugh-
ter cells in fresh media, whereas 51 cells produced 149 daugh-
ter cells in replete media. Furthermore, the frequency of cells
that entered abnormal cell divisions increased from 5% in
naive media to 26% in replete media.

Flow cytometry studies were initiated to further examine
the hypothesis that RENCA cells exposed to RENCA macro-
bead-conditioned replete media exhibited increased cell-cycle
times. When grown in replete media, the proportion of cells
that accumulated in the S phase of the cell cycle was increased
as compared with cells grown in fresh media (Fig. 3C).
Whereas 28% of the cells after 5 days in fresh medium were
in S phase, 45% of the cells grown in conditioned replete
medium were in S phase (Fig. 3D).

Given the above data that tumor cells cultured in macro-
bead-conditioned media exhibit an increase in S-phase time,
gen expression studies were performed to help elucidate
specific mechanisms of the cell-cycle alteration. Expression
analysis revealed a preponderance of upregulated S-phase
checkpoint genes (11/15 genes) in RENCA cells exposed to
conditioned media, as compared with RENCA cells grown in
fresh media (Table 1). GADD45 and various RAD homologues
associated with DNA damage response genes were signifi-
cantly upregulated (Table 1).
Figure 3. Cell-cycle study of RENCA cells responding to day 5 RENCA macrobead-conditioned replete media. A, beginning (top) images and ending (bottom) images of RENCA cells cultured in fresh media (left side) or RENCA macrobead-conditioned replete media (right side). Original magnification was 100×. The complete video is available online as Supplementary Figure 1. B, mitotic outcomes of RENCA cells exposed to either fresh or replete media as assessed with time-lapse videography. Inset shows individual cell-cycle times for productive mitosis collected over a 48-hour interval of log-linear growth between days 2 and 4 of culture. C, flow cytometry analysis of RENCA cell cycles following exposure to fresh or replete media. D, percentages of cells in S phase for RENCA cells exposed to either fresh or replete media. The accumulation of cells in S phase is evident.

### Table 1. S-phase checkpoint genes

<table>
<thead>
<tr>
<th>Probe Set</th>
<th>Fresh media</th>
<th>Normalized average intensity</th>
<th>Conditioned media</th>
<th>Log2R</th>
<th>ANOVA</th>
<th>Description</th>
</tr>
</thead>
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<tr>
<td>99005_at</td>
<td>365</td>
<td>310</td>
<td></td>
<td>-0.233</td>
<td>0.045</td>
<td>RAD1 homologue</td>
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<td>102878_at</td>
<td>129</td>
<td>119</td>
<td></td>
<td>-0.123</td>
<td>0.029</td>
<td>RAD52 homologue</td>
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<td>101429_at</td>
<td>418</td>
<td>1083</td>
<td></td>
<td>1.372</td>
<td>0.000</td>
<td>DNA-damage inducible 3</td>
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<tr>
<td>103944_at</td>
<td>65</td>
<td>114</td>
<td></td>
<td>0.806</td>
<td>0.007</td>
<td>RAD51-like 1</td>
</tr>
<tr>
<td>103457_at</td>
<td>52</td>
<td>389</td>
<td></td>
<td>0.627</td>
<td>0.005</td>
<td>RAD 54 like</td>
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<tr>
<td>102292_at</td>
<td>296</td>
<td>417</td>
<td></td>
<td>0.495</td>
<td>0.013</td>
<td>Growth arrest &amp; DNA damage 45</td>
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<tr>
<td>100459_at</td>
<td>236</td>
<td>317</td>
<td></td>
<td>0.426</td>
<td>0.042</td>
<td>RAD50 homologue</td>
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<td>92481_at</td>
<td>407</td>
<td>533</td>
<td></td>
<td>0.387</td>
<td>0.000</td>
<td>RAD53 homologue</td>
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<tr>
<td>165665_at</td>
<td>75</td>
<td>745</td>
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<td>0.372</td>
<td>0.017</td>
<td>RAD51 homologue</td>
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<tr>
<td>101885_at</td>
<td>2036</td>
<td>2539</td>
<td></td>
<td>0.319</td>
<td>0.006</td>
<td>Growth arrest specific 5</td>
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<td>92410_at</td>
<td>292</td>
<td>347</td>
<td></td>
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<td>0.047</td>
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<td>96102_1_at</td>
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<td>569</td>
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<td>0.147</td>
<td>0.004</td>
<td>RAD23b homologue</td>
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<td>94768_at</td>
<td>1742</td>
<td>1904</td>
<td></td>
<td>0.128</td>
<td>0.016</td>
<td>RAD21 homologue</td>
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Identification of released proteins

To overcome interference between macrobead-secreted proteins and bovine proteins found in conditioned media that had been supplemented with 10% normal calf serum, RENCA cells were adapted to serum-free culture media, encapsulated and cultured in serum-free media. RENCA macrobeads cultured in serum-free media displayed equivalent metabolic activity over time as compared with RENCA macrobeads cultured in serum-containing media, \( P > 0.05 \). \( n = 3–4 \) Lots per time point. B, RENCA macrobeads cultured in either serum-free or serum-containing media displayed similar inhibitory activity against RENCA cell growth, \( P > 0.05 \) except week 12 where \( P = 0.017 \). \( n = 2–5 \) Lots per time point. C, H&E micrograph of tumor colonies within RENCA macrobeads cultured in serum-containing media at 12 weeks of age. D, H&E micrograph of tumor colonies within RENCA macrobeads cultured in serum-free media at 12 weeks of age. Scale bars, 50 \( \mu m \).

Discussion

Our findings are consistent with the hypothesis that the growth of a given tumor and/or the tumor cells themselves can be slowed or stopped by one or more biological signals associated with the presence of tumor mass, even when that mass is not actually present. Tumor cells entrapped in the agarose matrix respond by producing a signal (or signals) that inhibit(s) the growth of freely growing tumor cells. More specifically, cells derived from epithelial tumors and encapsulated in the macrobead undergo a transformative process that results in the selection of at least 2 subpopulations of cells which give rise to tumor colonies. Although the mechanisms of the transformative process are not known, the formation of the tumor colonies in the growth-restrictive agarose environment is required to produce the tumor inhibitory effect as
Table 2. RENCA macrobead-secreted proteins with known tumor inhibitory function

<table>
<thead>
<tr>
<th>Accession number</th>
<th>Protein name (gene symbol)</th>
<th>Number of aa (UniProt)</th>
<th>Molecular weight, kDa (UniProt)</th>
<th>Description</th>
<th>Gene conservation (HomoloGene)</th>
</tr>
</thead>
<tbody>
<tr>
<td>gi</td>
<td>28916693</td>
<td>Gelsolin (GSN)</td>
<td>780 (secreted, Isoform 1: canonical)</td>
<td>86</td>
<td>Severs actin filaments to prevent toxic extracellular buildup during necrosis (19). Transfection of human bladder cancer cells reduced colony forming ability and tumorigenicity in vivo (20). Inhibits human pancreatic cancer cell invasion (21) and is associated with transition from adenoma to carcinoma in human colon carcinogenesis (22).</td>
</tr>
<tr>
<td>gi</td>
<td>13938049</td>
<td>Fibulin1 (FBLN1)</td>
<td>705 (secreted, Isoform D: canonical)</td>
<td>78</td>
<td>Extracellular matrix protein. FBLN1 transfection of human fibrosarcoma HT1080 cells inhibited growth in vivo, reduced angiogenesis, and increased apoptosis (23).</td>
</tr>
<tr>
<td>gi</td>
<td>84875537</td>
<td>Nucleolin (NCL)</td>
<td>707 (cytoplasm, nucleus)</td>
<td>77</td>
<td>Involved in rRNA transcription control, ribosome maturation and nucleocytoplasmic transportation (24). Endostatin receptor that mediates the antiangiogenic and antitumor activities of endostatin (25).</td>
</tr>
<tr>
<td>gi</td>
<td>3914939</td>
<td>Prosaposin (PSAP)</td>
<td>557 (secreted)</td>
<td>61</td>
<td>Assists in the lysosomal hydrolysis of sphingolipids (26). Identified as a tumor-secreted inhibitor of metastasis in which expression in human prostate cancer is correlated with metastases (27).</td>
</tr>
<tr>
<td>gi</td>
<td>46397639</td>
<td>Pigment epithelium-derived factor (PEDF)</td>
<td>417 (secreted)</td>
<td>46</td>
<td>A serine protease inhibitor of the serpin gene family with potent antiangiogenesis activity (28). Shown to have multimodal tumor inhibitory properties including antiproliferative, proapoptotic, and prodifferentiation activities in numerous tumor types (29–31).</td>
</tr>
</tbody>
</table>

(Continued on the following page)
<table>
<thead>
<tr>
<th>Accession number</th>
<th>Protein name (gene symbol)</th>
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<th>Description</th>
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</tr>
</thead>
<tbody>
<tr>
<td>gi</td>
<td>6679373</td>
<td>Serpine1 (Serbp1) Alternative name: plasminogen activator inhibitor-1</td>
<td>407 (cytoplasm, nucleus) Isoform 1: canonical)</td>
<td>45</td>
<td>May play a role in mRNA stability. Shown to inhibit orthotopic rat bladder tumor cells (32). Inhibits human prostate tumor growth in vivo (33). Also shown to function in a tumor-protective role by inhibiting chemotherapy-induced apoptosis (34).</td>
</tr>
<tr>
<td>gi</td>
<td>6678077</td>
<td>Secreted protein acidic &amp; rich in cysteine (SPARC) Alternative names: osteonectin, basement-membrane protein 40</td>
<td>302 (secreted)</td>
<td>34</td>
<td>Regulates cell growth through interaction with ECM. Inhibits in vitro growth of ovarian, breast, acute myeloid leukemia, and pancreatic cells (35–38).</td>
</tr>
<tr>
<td>gi</td>
<td>267133</td>
<td>Tissue inhibitor of metalloproteinase 2 (TIMP2)</td>
<td>220 (secreted)</td>
<td>24</td>
<td>Protease inhibitor shown to inhibit transfected melanoma cells in vivo (39), angiogenesis (40), and responses to growth factors (41).</td>
</tr>
<tr>
<td>gi</td>
<td>84794552</td>
<td>Phosphatidylethanol amine-binding protein 1 (PEBP1) Alternative name: raf kinase inhibitory protein</td>
<td>187 (cytoplasm)</td>
<td>21</td>
<td>Serine protease inhibitor. Decreased expression in human breast (42) and melanoma cancer metastases (43); anaplastic thyroid cancer (44); and transfected hepatocellular carcinoma growth suppression (45). Known metastasis inhibitor (46).</td>
</tr>
</tbody>
</table>
monolayer cultures do not inhibit tumor growth. It is the tumor colonies that produce some factor or factors that inhibit(s) the proliferation of other freely growing cancer cells both in vitro and in vivo.

Although there is considerable diversity among cancer tumor types with respect to gene changes and the signaling pathways and pathway interactions that flow from those gene changes and other supragenetic regulatory processes, the findings reported here indicate that there exists a growth regulatory system that is common to many epithelial-derived tumor types and that this regulatory system is conserved across species lines. This is evident from the in vitro studies in which encapsulated mouse RENCA cells were shown to inhibit the growth of various human tumor cell lines and also from encapsulated human J82 bladder tumor cells that were able to inhibit the growth of mouse RENCA cells. This conserved growth regulation is further illustrated by the in vivo studies in which mouse RENCA cell macrobeads significantly extended the survival of canine patients with prostate adenocarcinoma (median survival of 177 days) as compared with no treatment (21–30 days; ref. 49) or to total prostatectomy or subtotal intracapsular prostatectomy [<50 days (n = 10) and <2 weeks for 7/10 dogs, respectively; ref. 50]. The tumor inhibition appears to be the result of increased cell-cycle times in cells exposed to either macrobead-conditioned media or to RENCA macrobeads directly. In particular, the data demonstrate an increase in S-phase cycle time and a decrease in productive mitosis for freely growing tumor cells exposed to such conditions.

The use of serum-free media for the culture of the macrobeads has allowed 1-dimensional gel electrophoresis and mass spectroscopy of the bands which demonstrates the presence of at least 10 proteins from RENCA macrobead-conditioned media that have been previously reported to have tumor inhibitory properties (Table 2). These data suggest that multiple signals are involved in the inhibitory effect of the macrobeads. The fact that the proteins listed in Table 2 are all highly conserved across species is consistent with the lack of species specificity and the lack of tumor-type specificity of the macrobead growth-inhibitory effects.

Based on these findings, we believe that RENCA agarose–agarose macrobeads have promise as a form of anticancer therapy, by virtue of their ability to utilize conserved growth regulatory (feedback) pathway(s) and their lack of clinically significant toxicity to date. Using encapsulated tumor cells from one species to treat cancer in another species is attractive because strong immunologic (xenogeneic) barriers prevent cross-species seeding of unencapsulated cells. Beyond their potential therapeutic implications, the cancer macrobeads offer a unique opportunity to study the molecular biological basis of tumor cell proliferation control, in part, by virtue of their ability to select a subpopulation of cells that appear to have stem cell properties and to be responsible for enabling the founding and perpetuation of tumors (1). In addition, they provide a ready means for the in vitro preclinical testing of proposed therapeutic agents that, to be optimally effective, must show activity against such cells. A Phase 2 clinical trial in humans (FDA BB-IND-10091) commenced in 2010.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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References


Hydrophilic Agarose Macrobead Cultures Select for Outgrowth of Carcinoma Cell Populations That Can Restrict Tumor Growth


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