Re-activation of the p53 pathway inhibits in vivo and in vitro growth of hormone-dependent human breast cancer cells

YAYUN LIANG1, CYNTHIA BESCH-WILLIFORD2, INDIRA BENAKANAKERE1 and SALMAN M. HYDER1,3

1Dalton Cardiovascular Research Center, 2Department of Veterinary Pathology and
3Department of Biomedical Sciences, University of Missouri, Columbia, MO 65211, USA

Received April 4, 2007; Accepted May 9, 2007

Abstract. Mutations in wild-type p53 (wtp53) protein lead to loss of its tumor suppressor function in breast cancer cells, facilitating uncontrolled tumor growth. Consequently, procedures to repair defective p53 functions in tumor cells are being actively pursued. We sought to determine whether expression of wt p53 protein, or conversion of endogenous mutant p53 (mtp53) into a functional p53 protein with small molecule PRIMA-1, can override the tumor-promoting effects of naturally occurring mtp53 protein in hormone-responsive T47-D human breast cancer cells. We show that transfection of wt p53 gene into T47-D cells suppresses their proliferation in regular media, and inhibits estrogen-dependent cell proliferation in media containing dextran-coated charcoal treated serum. Growth inhibition was not due to the absence of estrogen receptor-α or progesterone receptors though receptor levels for estrogen receptor-α were drastically reduced in wt p53 expressing cells. Focused microarray analysis of wt p53 expressing cells revealed suppression of PCNA cell cycle regulatory mRNA and protein. Wild-type p53 transfected T47-D cells also failed to grow in vivo in estrogen supplemented nude mice. Furthermore, xenografts obtained with parental T47-D cells expressing mtp53 grew poorly in nude mice treated with PRIMA-1. PRIMA-1 treated tumors exhibited a low proliferation index, even though mice were estrogen-supplemented. PRIMA-1 treatment of tumor cells suppressed VEGF and induced expression of estrogen receptor-β though expression of estrogen receptor-α and progesterone receptors was unaffected. These data indicate that alteration of the p53 signal transduction pathway by re-expression of wt p53 protein in T47-D cells, or treatment of parental cells with PRIMA-1, can prevent in vivo and in vitro proliferation of T47-D breast cancer cells.

Introduction

p53 is a multifunction tumor suppressor protein that is a key player in the cellular response to DNA damage and other types of cellular stress (1,2). Activated wild-type p53 (wt p53) plays a key role in signaling pathways that lead to cellular stress-induced cell cycle arrest and/or apoptosis. Mutations in wt p53 are frequent in a wide range of cancers, including breast cancer, and this likely contributes to the uncontrolled growth phenotype associated with tumor cells (1,2). Many mutant p53 alleles in cancer cells express dysfunctional mutant p53 protein (mtp53) at a high level (3,4). Approximately 50% of all breast cancers carry point mutations in the p53 gene, and the majority of mtp53 alleles in breast cancer cells are defective in DNA binding as well as in cell cycle checkpoints and DNA damage-induced induction of apoptosis (5,6). Mutations in p53 or the p53 pathway are thought to play a key role in promoting tumor cell survival and tumor cell resistance to chemotherapeutic drugs (7,8).

PRIMA-1 is a small molecule that activates the mtp53 protein and restores wild-type functions of this protein (9,10). For example, PRIMA-1 stimulates expression of p21 and other p53-dependent promoters in several breast cancer cells that express mtp53 (10), and suppresses hormone-induced vascular endothelial growth factor (VEGF) expression (10). VEGF is associated with angiogenesis in tumor tissue (11,12). Because PRIMA-1 at least partially restores wt p53 function in several breast cancer cells, it seemed possible that co-expression of wt p53 and/or treatment with PRIMA-1 might have therapeutic potential as an anti-cancer/anti-angiogenic treatment for hormone responsive breast cancer cells that express mtp53 protein by suppressing the production of VEGF. The premise of this strategy is that wt p53 protein is essentially dominant over endogenous mtp53 in cancer cells. This study tests the hypothesis that growth of hormone-
responsive T47-D human breast cancer cell line can be suppressed by co-expression of wtp53 or by treatment with PRIMA-1. The results suggest that reactivation of the p53 mediated pathways may have potential as an anti-cancer strategy for human breast cancer and possibly other cancers.

Materials and methods

Materials. T47-D cells expressing mtp53 were from ATCC (Manassas, VA). Phenol red-free DMEM/F12 medium, phosphate-buffered saline, and 0.05% trypsin-EDTA were from Invitrogen Corporation & Life Technologies (Grand Island, NY) and fetal bovine serum (FBS) was from JRH Biosciences (Lenexa, KS). PRIMA-1 (p53-reactivation and induction of massive apoptosis) was from Tocris (Ellisville, MO). 17β-estradiol (1.7 mg/pellet, 60-day release) and placebo pellets were from Innovative Research of America (Sarasota, FL). sulforhodamine B (SRB) was from Sigma (St. Louis, MO), BD Matrigel Matrix was from BD Biosciences (Bedford, MA), anti-PCNA antibody was from Dako (Carpinteria, CA) and anti-ER-α and anti-ER-ß antibodies for Western blots were from Santa Cruz Biotechnology, Inc. (Santa Cruz, CA). The plasmid with the p53 gene cloned into pcDNA vector was a kind gift from Dr Xinhin Chen, University of Alabama.

Cell lines and cell culture. T47-D cells were stably transfected with a wt-p53 expression plasmid or vector (pcDNA), selected for neomycin resistance and tested for expression of functional p53 as described previously (10). Stably transfected T47-D cells were selected for loss of progesterin-dependent induction of VEGF induction (10). One such cell line, T47-D-p53-L (referred to as p53-L cells), and a control cell line, T47-D-PCDNA-A (referred to as PCDNA-A), carrying vector DNA, were selected for further study. Cells were maintained in phenol red-free DME/F12 medium supplemented with 10% FBS (T47-D) or the same medium supplemented with 200 μg/ml G418 for transfected clones. All cells were grown in 100x20 mm tissue culture dishes and harvested with 0.05% trypsin-EDTA.

Cell proliferation assay by SRB. A sulforhodamine B (SRB) assay was used to measure cell viability, as previously described (13,14). Briefly, T47-D, PCDNA-A, and p53-L cells were grown to 70% confluence, after which media were replaced with 5% FBS-DCC DMEM/F12 and incubated for 24 h to deplete any endogenous steroids. Cells were harvested by trypsin-EDTA treatment and 4x10³ cells/well were seeded 24 h to deplete any endogenous steroids. Cells were harvested with 0.05% trypsin-EDTA. In 100x20 mm tissue culture dishes and harvested with 0.05% trypsin-EDTA.

Preparation of nuclear extracts. A nuclear extract kit was used according to the manufacturer's instructions (Active Motif, Carlsbad, CA). Briefly, cells were grown in 100-mm dishes, washed with 3 ml ice-cold PBS containing phosphatase inhibitors and harvested by gentle scraping with a cell lifter. Cells were centrifuged for 5 min at 200 × g at 4°C, re-suspended in 500 μl 1X hypotonic buffer (Active Motif kit) and incubated for 15 min on ice. Detergent (25 μl) was added and the extract was centrifuged for 30 sec at 14,000 × g at 4°C. The pellet was re-suspended in 50 μl complete lysis buffer containing 1 mM DTT and 1% protease inhibitor cocktail and incubated on ice for 30 min with shaking. Samples were centrifuged at 14,000 × g for 15 min, and the supernatant was transferred to a microcentrifuge tube, aliquoted, and stored at -80°C. Supernatant was adjusted to the desired protein concentration prior to use.

Immunoblotting. Nuclear extracts (35 μg per lane) were separated on a NuPAGE 10% Bis-Tris Gel (Invitrogen, Carlsbad, CA). Electrophoresis was performed at 100 V for 2 h using NuPAGE MES-SDS Running Buffer. Separated proteins were electro-transferred to a polyvinylidene difluoride membrane (Bio-Rad Laboratories, Hercules, CA) at 35 V for 1.5 h. The blots were blocked for 1 h at room temperature (RT) in 5% non-fat dry milk in TBS containing 0.1% Tween-20 (TBS-T), and incubated with primary antibody for 2 h at RT at the indicated dilution: PCNA (clone PC10; 1:800 dilution), anti-PR (1:300 dilution of AB-52), anti-ERα (D-12; 1:200 dilution), anti-ER-ß (H-150; 1:200 dilution). The blots were washed 3 times with TBS-T, incubated with secondary antibody for 1 h at RT, and washed 6 times (8 min each) with TBS-T. Immuno-reactive bands were visualized using the ECL plus detection kit (Amersham, Pharmacia Biotech, Arlington Heights, IL). Membranes were stripped and re-blotted for β-actin (Sigma). Immuno-reactive protein signal was normalized to the actin signal in the same sample.

Animal xenograft tumor studies in nude mice with wt-p53 transfected cells. Female athymic nu/nu nude mice, 5-6-weeks old (18-22 g) were purchased from Harlan Sprague Dawley, Inc. (Indianapolis, IN). The mice were housed in a laminar air-flow cabinet under specific pathogen-free conditions. All facilities were approved by the American Association for Accreditation of Laboratory Animal Care in accordance with the current regulations and standards of the United States Department of Agriculture, the Department of Health and Human Services, and the NIH. All procedures were approved by IACUC.

Nude mice were inoculated in the dorsal flank with a 17β-estradiol pellet (1.7 mg/pellet; 60-day release) 48 h prior to injection with tumor cells. PCDNA-A and p53-L cells were harvested by trypsinization, washed twice with DMEM/F12, re-suspended in 100 μl DMEM/F12 and injected (1x10⁶ cells per mouse) subcutaneously in the dorsal flank. Tumors were measured every 3 days with a digital caliper and tumor volumes were calculated using the following formula: \((V = \frac{L \times W \times H}{2})\).
**PRIMA-1 treatment of xenografts obtained from parental T47-D cells expressing mtp53 protein.** Mice were inoculated with estradiol pellets as described above. After 48 h, 5x10⁶ T47-D cells were re-suspended in 100 μl DMEM/F12 medium mixed with Matrigel (50% v/v), and inoculated into both flanks subcutaneously. Animals were randomly assigned to two groups of 5 mice each. One group of mice received 25 mg/kg/day PRIMA-1 by i.p injection 2 h prior to injection of tumor cells (details in Fig. 5A). This treatment was continued for two additional days and then PRIMA-1 treatment was continued at 50 mg/kg/day by tail vein injection for 10 days. Experiments were terminated by sacrificing animals at 6 h after the final PRIMA-1 injection. Tumors were measured every 2 days with digital calipers and harvested into 4% paraformaldehyde for future studies. Tumor volume was calculated using the formula \((L \times W \times H) \times \pi/6\) (15).

**Histology and immunohistochemical analysis.** Immunohistochemical analysis was carried out for Ki67, VEGF, Factor VIII, PR, ER-α and ER-β. Tumor tissues was fixed overnight in 4% paraformaldehyde for immunohistochemistry. Tissues were processed for paraffin infiltration and embedding. Five-μm sections were mounted onto ProbeOn Plus microscope slides (Fisher Scientific Inc., Pennsylvania, PA), stained with hematoxylin-eosin and examined for cellularity by light microscopic analysis. For immunohistochemical analysis, sections were de-waxed in xylene, rehydrated through graded concentrations of ethanol, then rinsed in distilled water, and, if necessary, stored in phosphate-buffered saline (PBS) at 4˚C until use. Sections were subjected to heat-induced epitope retrieval in 10 mM citrate buffer (pH 6.0) (VEGF, Ki67, ER, PR) or proteinase K (20 μg/ml in TE buffer, pH 8.0) (Factor VIII antigen). Slides were treated with 3% hydrogen peroxide in absolute methanol (to inactivate endogenous peroxidase activity), washed in 3X PBS, incubated in blocking buffer with 5% bovine serum albumin for 20 min and treated with polyclonal antibody at room temperature for 60 min at the following dilution: anti-VEGF antibody [1:200 dilution of a rabbit anti-VEGF polyclonal antibody (sc-152); Santa Cruz Biotechnology, Inc.]; anti-Ki67 [1:200 dilution of rabbit anti-Ki67 polyclonal antibody (RB 1510-P); Lab Vision, Fremont, CA]; anti-factor VIII antibody [1:400 dilution of a rabbit anti-factor VIII polyclonal antibody (A0082); Dako]; anti-PR [1:50 dilution of a rabbit anti-PR polyclonal antibody, (A0098); Dako]; anti-ERα [1:300 dilution of a rabbit anti-ERα polyclonal antibody (sc-542); Santa Cruz Biotechnology, Inc.]; anti-ER-β [1:50 dilution of a mouse anti-ER-β monoclonal antibody (MAb974s); Serotec, Inc., Raleigh, NC]. Sections were then washed and sequentially incubated with a secondary antibody (biotinylated swine anti-mouse IgG and biotinylated swine anti-rabbit IgG; Dako), and a streptavidin-linked horseradish peroxidase product (Dako) for 30 min, also at room temperature. Alternatively, some sections were incubated with EnVision™, a horseradish peroxidase labeled polymer conjugated with anti-rabbit antibodies (Dako). Bound antibodies were visualized following incubation with 3, 3’-diaminobenzidine tetrahydrochloride (0.05% with 0.015% H₂O₂ in PBS) solution (Dako) for 3-5 min. Sections were counterstained with Mayer's hematoxylin, dehydrated, cleared and coverslipped for microscopic examination.

**Proliferation index.** The number of Ki67 positively staining nuclei was counted per ~1000 tumor nuclei in the most intensely stained areas of tumor sections. We measured areas that were uniform (2049 μ²) among samples. The proliferation index was then defined as Ki67 positive nuclei/Ki67 positive +Ki67 negative nuclei x 100.

**RNA isolation and synthesis of cDNA.** Total RNA was isolated using Ultra spec RNA reagent (Biotex Products, Oxon, UK). Total RNA was used as a template for biotinylated probe synthesis using TrueLabeling-AMP Kit (SuperArray, Inc., Bethesda, MD) according to the manufacturer's instructions.

**Hybridization and chemiluminescent detection.** Microarray membranes were hybridized using the Oligo GEArray Kit (SuperArray, Inc.) according to the manufacturer's instructions. The chemiluminescent detection kit (SuperArray, Inc.) was used to conform proper probe hybridization. Membranes were washed, rinsed, incubated with ECL chemiluminescent substrate (Amersham, Arlington Heights, IL), after which the membranes were exposed to radiographic film. All images were stored electronically and uploaded into an integrated (web-based) GEArray Expression Analysis System at the SuperArray web site. Data analysis and normalization was based on quartile analysis as described by SuperArray Inc.

**Statistical analysis.** Statistical significance was measured using Student’s t-test or One-way analysis of variance (ANOVA) with repeated measure over time. When necessary, it was assumed that ANOVA was non-parametric. Values are reported as mean ± SE. For samples with significant F-ratio (p<0.05), the Student-Newman-Keuls multirange test was employed (SigmaStat).

**Results**

Stable T47-D transfectants that co-express wtp53 and mtp53 were selected by screening for neomycin resistance, the presence of wtp53 protein and for inducibility of p21 as previously described (10). Stable transfectants were also screened for progestosterone-stimulated induction of VEGF and expression of p53 was found to suppress hormone-induced VEGF in several clones (10). A cell line that lacks progesterone-inducible VEGF, p53-L, and a control cell line carrying empty vector, PCDNA-A, were selected for further study. Cell proliferation studies were carried out on p53-L cells in the presence of 10% FBS using a sulforhodamine B (SRB) dye-binding assay (13,14). Control cells included PCDNA-A cells and parental T47-D cells. The results showed that p53-L cells grew 1.5- to 2-fold less efficiently than T47-D or PCDNA-A cells in the presence of 10% FBS. These data suggest that stable co-expression of wtp53 protein suppresses in vitro proliferation of T47-D cells in the presence of serum.

The expression of human cell-cycle related genes were compared in p53-L and PCDNA-A cells using a human gene array profiling system (SuperArray). The results showed that PCNA mRNA was the most extensively down-regulated (2.3-fold) in p53-L compared with expression in PCDNA-A cells, and correspondingly PCNA protein was 2-fold lower in
p53-L cells than in PCDNA-A and T47-D cells (Fig. 2A and B).

Estrogen stimulates proliferation of many breast cancer cell lines including T47-D cells (16) thus, the effect of exogenous estrogen on growth of p53-L, PCDNA-A and T47-D cells was examined by measuring cell proliferation as described above. The results showed that exogenous estradiol-17-ß (0.1 nM to 1 μM) strongly stimulated growth of PCDNA-A and T47-D cells in a dose-dependent manner, but did not stimulate the growth of p53-L cells (Fig. 3A), despite continued expression of estrogen receptor-α ER-α in these cells albeit at 3-4-fold lower level in p53-L cells than in PCDNA-A and T47-D cells. The level of ER-ß was similar in p53-L, PCDNA-A and T47-D cells (Fig. 3B).

The proliferative capacity of p53-L cells was also examined in nude mice treated with estradiol pellets (note that T47-D cells do not grow as xenografts in nude mice in the absence of exogenous estrogen). p53-L and PCDNA-A cells were injected subcutaneously in the left and right dorsal flanks of nude mice, and tumor volume was measured by caliper every three days for 30 days. The results showed that mice injected with PCDNA-A cells formed large tumors, which grew for ~10 days before beginning to regress (Fig. 4). Tumors of similar size formed with similar efficiency in mice injected with T47-D parental cells (not shown). In contrast, mice injected with p53-L cells formed very small tumors by day 10, which regressed slowly from 10 to 30 days post-injection (Fig. 4). These data demonstrate that co-expression of wtp53 suppresses growth of T47-D cells in vivo in nude mice supplemented with exogenous estrogen.

The studies described above demonstrate that co-expression of wtp53 and mtp53 in T47-D cells suppresses cell growth in vivo and in vitro. This result suggests that cancer cell growth might also be suppressed by restoring wild-type function to mtp53. This approach was tested here using PRIMA-1, a previously characterized small molecule activator of mtp53 (9). For this experiment, estrogen pellets were implanted into nude mice, the mice were treated with PRIMA-1 for 2 h, and then tumor cells were mixed with Matrigel and injected subcutaneously in the left and right dorsal flanks of nude mice.

Figure 1. Growth of p53-L, PCDNA-A and T47-D cells. p53-L, PCDNA-A and T47-D cells (4x10³ cells/well) were seeded into a 96-well plate and incubated overnight in 100 μl DMEM/F12 + 10% FBS. Media was replaced and cells were incubated for the indicated period of time with an additional media change on day 3. Cells were fixed in situ and cell viability was determined from SRB absorbance (see Materials and methods). Asterisks indicate values that are significantly different than control (p<0.05, ANOVA).

Figure 2. (A) Gene expression analysis in PCDNA-A and p53-L cells. Gene expression was analyzed using a human SuperArray microchip for cell cycle genes as described in Materials and methods. Signal intensity was normalized using an interquartile approach and the GEArray expression analysis suite from SuperArray. Changes in expression ≥2.0-fold were assessed. Only PCNA was found to reduce by >2-fold in the p53L cells (circles). (B) Western blotting of PCNA in T47-D, PCDNA-A and p53-L cells. Cells were grown to 80% confluence in the presence of 10% FBS, washed with cold PBS and harvested. Nuclear extract was prepared and analyzed by Western blotting using antibody to PCNA. Quantification of data is shown in bar graph to the right. Values were normalized to β-actin.
Fig. 5A). PRIMA-1 dosing continued for 3 days at 25 mg/kg and then for 10 days at 50 mg/kg as shown in Fig. 5A. Tumor volume was measured every other day for 14 days (Fig. 5B). The results show that PRIMA-1 strongly inhibits tumor growth in nude mice injected with T47-D cells (Fig. 5B). Treated animals showed no toxic side effects of PRIMA-1 as also reported by others (9) and maintained normal body weight over the duration of the experiment (Fig. 5C). Very small tumors were detected at the site of injection in animals treated with PRIMA-1 (Fig. 5D). These data show that PRIMA-1 strongly inhibits growth of T47-D as xenograft tumors in nude mice.

Because PRIMA-1-treated mice injected with T47-D cells developed small tumors (Fig. 5B and D), representative tumors were excised and subject to histological analysis. The results showed tumor cells in both PRIMA-1 treated mice and control mice; however, the cellularity of tumors from PRIMA-1 treated animals was sparse (Fig. 6A, left panel). There were fewer Ki67-positive cells in tumors from PRIMA-1-treated mice than in controls (Fig. 6A, middle panel). The proliferation index was 37±5 (mean ± SEM) in PRIMA-1 treated samples compared with 57±4 in the control samples; p<0.05, Student’s t-test). The number of apoptotic cells was similar in PRIMA-1-treated and control animals as determined using TUNNEL assay (not shown). These data suggest that PRIMA-1 mainly inhibited proliferation of T47-D cells in vivo under the protocol tested. Furthermore, VEGF expression was severely inhibited in tissue samples from PRIMA-1-treated animals (Fig. 6A, right panel). Analysis of Factor VIII expression and tumor vascularity was inconclusive due to insufficient tissue. PRIMA-1 did not alter the ER-α or progesterone receptor status of tumor tissue, but it significantly stimulated expression of ER-β in tumor cells isolated from treated animals (Fig. 6B).

Discussion

The present study demonstrates that over-expressed wtp53 is dominant over mtp53 in T47-D cells, and that co-expression of wtp53 and mtp53 or activation of endogenous mtp53 is a viable strategy to inhibit tumor cell growth. Importantly, activation of the p53 pathway by one of these two approaches inhibits proliferation of T47-D cells in cell culture in the presence of serum-derived growth factors or exogenous estrogen, and inhibits growth of T47-D xenograft tumors in vivo in nude mice.

The results presented here show that co-expression of wtp53 protein inhibits growth of T47-D cells in culture in the presence of 10% FBS. The mechanism of this effect is not known. It is possible that wtp53 protein binds to mtp53, thereby preventing adverse biological impacts of the mutant protein. This result is consistent with recent observations from human clinical trials, showing that expression of wtp53 protein can cause regression of human breast disease many of which contain the mtp53 protein initially (17,18). Alternatively, co-
expression of wtp53 in T47-D cells may directly regulate genes involved in cell cycle arrest and/or apoptosis, thus indirectly blocking the adverse growth-promoting effects of mtp53. Focused microarray studies in T47-D cells co-expressing wtp53 demonstrated decreased transcription of proliferation genes including PCNA. These data suggest that T47-D cells have an intact p53 pathway, which could potentially be exploited for therapeutic benefit. One caveat is that the present observations may not be generalized to all human breast cancer cells or to other human cancer cell types. For example, additional downstream effectors of cell cycle control may be present or absent in cell lines other than T47-D. Additional studies are needed to address this possibility. Nevertheless preliminary studies showed that co-expression of wtp53 prevents progesterone-dependent induction of VEGF not only in T47-D cells but also in two other cell lines containing mt53, namely BT-474 and HCC-1428 cells, suggesting that restoration of the p53 pathway is achievable in many different breast cancer cell types following an attempt to restore this function with wtp53 expressing plasmid or use of small molecules such as PRIMA-1 (10).

Co-expression of wtp53 in T47-D cells also blocked estrogen-dependent cell growth. Cell growth was inhibited despite normal expression of ER-ß though there was a 3- to 4-fold reduced expression of ER-α. Previous studies suggest that wtp53 may also interact with and inhibit downstream signaling by ER-α (19-21), though the role of p53 in influencing ER-ß mediated signal transduction pathway remains unknown. Alternatively, wtp53 may suppress other signal transduction pathways (i.e., MAPK or the PI3 kinase) which in turn influence ER-dependent signaling (22).

This study also shows that co-expression of wtp53 in T47-D cells inhibits growth of xenograft tumors in nude mice. There are several possible explanations for this result. First, co-expression of wtp53 could induce inhibitors of tumor or endothelial cell growth and/or inhibitors of angiogenesis.

---

**Figure 5.** (A) PRIMA-1 dosing regimen is shown schematically. Dose and time of each injections or PRIMA-1 is indicated. Time of inoculation with E2 pellet and tumor cells is also indicated. (B) Xenograft tumor volume in PRIMA-1-treated and control nude mice. Five- to 6-week old nude mice were inoculated with 17ß-estradiol pellet. Forty-eight hours later, mice were injected subcutaneously in both dorsal flanks with 5x10^6 T 47-D cells in 100 μl DMEM/F12 containing Matrigel. PRIMA-1-treated mice were dosed as shown in (A). Tumors were measured every 2 days and tumor volume was calculated using the formula (L x W x H) x π/6. Asterisks indicate values that are significantly different than control (p<0.05 using t-test). (C) Average body weight of PRIMA-1-treated and control animals. (D) Selected images of xenograft tumors in PRIMA-1-treated and control animals at the end of the experiment as shown in (A).
Second, wtp53 could promote apoptosis of T47-D cells in xenograft tumors. Third, co-expression of wtp53 inhibited proliferation and induced cell cycle arrest in T47-D cells in xenograft tumors. These possibilities are not mutually exclusive and all three mechanisms could contribute to suppression of xenograft tumors in nude mice. Additional studies will explore and possibly discriminate between one or a combination of these effects.

Activation of endogenous mtp53 in T47-D cells by PRIMA-1 also strongly inhibited, but did not completely suppress growth of T47-D cells in xenograft tumors. These possibilities are not mutually exclusive and all three mechanisms could contribute to suppression of xenograft tumors in nude mice. Additional studies will explore and possibly discriminate between one or a combination of these effects.

Activation of endogenous mtp53 in T47-D cells by PRIMA-1 also strongly inhibited, but did not completely suppress growth of T47-D cells in xenograft tumors. These possibilities are not mutually exclusive and all three mechanisms could contribute to suppression of xenograft tumors in nude mice. Additional studies will explore and possibly discriminate between one or a combination of these effects.

a survival factor for breast cancer cells. Since PRIMA-1 activates mtp53 into its wt counterpart, it is possible that PRIMA-1 activated p53 protein blocks the production of VEGF at transcriptional level as has been shown by others. As an alternative for lack of tumor growth, nascent blood vessels at the site of tumor cell injection might increase the local concentration of PRIMA-1, ultimately causing apoptosis and/or cell cycle arrest. More information on the pharmacokinetic and pharmacodynamic properties of PRIMA-1 in vivo is needed to evaluate these possibilities. Future experiments will test alternative PRIMA-1 dosing strategies and their effects on growth of T47-D xenograft tumors in nude mice. Such studies are feasible because to date, PRIMA-1 has not been associated with any adverse toxicity in nude mice.

Immunohistochemical analysis also showed that tumors in PRIMA-1-treated mice contain a smaller fraction of Ki67-positive cells, indicating a lower rate of cell proliferation. In contrast, the rate of apoptosis was similar in tumors in PRIMA-1-treated and control mice. Since the tumors were collected at the end of the experiment it is possible that initially there was a higher apoptotic rate reducing the number of cells in tumor overall. Thus PRIMA-1 may have effects both in terms of increasing apoptosis as shown by others and in reducing cellular proliferation. It is also possible that the rate of apoptosis would increase in tumors of mice exposed to a higher concentration of PRIMA-1, leading to complete suppression of xenograft tumors. Thus, future studies will include PRIMA-1 dose escalation and alternative routes of PRIMA-1 delivery.

Interestingly, expression of ER-ß increased in tumors in mice treated with PRIMA-1 (Fig. 6B), while expression of ER-α and PR did not change. While the function of ER-ß in breast cancer cells remains controversial, it has been implicated as a negative regulator of cell growth and proliferation. A recent report has also indicated a role for ER-ß in blocking angiogenesis in T47-D by reducing VEGF and ER-ß has been shown to interact with the VEGF promoter. In addition, it has been shown that ER-ß negatively impacts the
proliferative effects of ER-α by altering ER-α mediated gene expression in breast cancer cells (32,33). Thus it is possible that increased expression of ER-β during PRIMA-1 treatment may lead to loss of progression of breast cancer cells though the cells are continuously exposed to estrogen. The mechanism by which PRIMA-1 stimulates expression of ER-β in breast cancer cells, and how such an increase influences ER-α mediated proliferative response remains to be explored.

In conclusion, our results show that the p53 pathway negatively regulates growth of T47-D cells in vivo and in vitro, and that activation of the p53 pathway suppresses the transformed phenotype of these cells. In cell culture, this effect is observed in the presence of estradiol. An important outcome of this study is that endogenous mtp53 can be re-activated by small molecule p53 activators such as PRIMA-1, and such compounds may have therapeutic potential for preventing or treating at least a subset of human breast tumors.

Acknowledgements

This research was supported by NIH grant CA-86916; Dept of Defense Breast Cancer Research Program W81XWH-05-1-0416; PDF0600723 and BCTR0600704 from Susan G Komen Breast Cancer Foundation, and by funds from the Research Animal Diagnostic Laboratory. We would like to thank Dr Xinbin Chen from the University of Alabama for helpful discussions and Dr Jennifer Schnell and Ms. Jill Gruenkemeyer for excellent technical assistance. S.M.H. is the Zalk Missouri Professor of Tumor Angiogenesis.

References