Melanoma Prevention Using Topical PBISe

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Abstract

Malignant melanoma is the deadliest form of skin cancer, known for its drug resistance and high metastatic potential. Deregulated PI3 and mitogen activated protein (MAP) kinase pathways promote early melanocytic lesion development and confer drug resistance. No agent exists to target these deregulated pathways to prevent cutaneous noninvasive melanocytic cells or invasive melanomas from developing into more aggressive widely disseminated metastatic disease. In this study, a selenium containing isosteric analogue of PBIT [S,S'-1,4-phenylenebis(1,2-ethanediyl)bis-isothiourea] called PBISe [Se,Se'-1,4-phenylenebis(1,2-ethanediyl)bis-isoselenourea] is shown to moderate these 2 major signaling pathways to prevent cutaneous melanocytic lesion or melanoma development. Topical application of PBISe retarded melanocytic lesion development in laboratory-generated skin by 70% to 80% and in animal skin by approximately 50%. Mechanistically, prevention of lesion development occurred due to decreased Akt3 signaling, which increased MAP kinase pathway activity to inhibitory levels. The combined effect of targeting these pathways led to decreased cell proliferation and increased apoptotic cell death thereby preventing melanoma development. Thus, topically applied PBISe treatment has potential to prevent noninvasive melanocytic lesion and invasive metastatic melanoma development in skin. Cancer Prev Res; 4(6); 935–48. ©2011 AACR.

Introduction

Malignant melanoma remains the most deadly and invasive form of skin cancer leading to more than 80% of all skin cancer deaths (1, 2). Despite the use of surgery to excise early noninvasive melanocytic lesions to prevent progression to metastatic disease, incidence and mortality rates for metastatic melanoma continue to rise (2, 3). Accumulating evidence has shown protective effects of sunscreens from damaging UV rays, which otherwise can potentially induce the development of melanomas from melanocytes located at the base of the skin epidermis (4). However, the utility of sunscreens for protecting skin from sunburn damage is raising concerns that its use might encourage sun exposure, which could have the opposite effect and increase rates of skin cancers (5).

No agent targeting Akt3 or V600E-B-Raf signaling involved in early melanoma development is available clinically to prevent the disease or retard early stage noninvasive melanocytic lesion cells from progressing into metastatic melanoma (6). Preventing melanoma development in its early stages using chemopreventive agents is a viable strategy to decrease premalignant lesions from progressing into lethal advanced stages in humans. If an agent of this type were available, it could have significant clinical potential to decrease mortality rates and reduce costs associated with treatment and management of advanced-stage disease (7). Therefore, novel compounds are needed that could be used to augment existing preventive strategies and one approach is the development of topical agents, which could moderate key pathways involved in development of melanocytic lesions or melanoma in skin.

Development of melanomas from melanocytes is a multistage complex process involving changes in expression and activities of a number of genes and signaling pathways regulating cellular differentiation, growth, senescence, survival, and migration (8). B-Raf is a member of the mitogen-activated protein (MAP) kinase pathway whose activity is deregulated in melanomas by mutation to a constitutively active form called V600E-B-Raf in approximately 90% of benign nevi or normal moles (9, 10). However, the presence of V600E-B-Raf alone does not cause melanoma as this aberrant protein activates the downstream MAP kinase pathway to inhibitory levels, leading to cellular senescence (10–16). Therefore, very few V600E-B-Raf containing moles ever develop into melanoma and remain in a senescent stage (14, 15). Other factors such as loss of tumor suppressors PTEN, p16INK4a or activation of oncogenes such as Akt3 are needed to moderate V600E-B-Raf activity and the downstream
MAP kinase pathway to drive tumor progression (14–17). Elevated Akt3 activity, occurs in approximately 70% of melanomas and inhibits V600E-B-Raf protein activity by phosphorylating negative regulatory sites, which moderates protein activity to levels promoting melanoma development rather than driving cells into senescence (10, 15, 17–20). Therefore, therapeutics targeting Akt3 activity has potential to increase MAP kinase activity to inhibitory levels to promote cell senescence. Furthermore, melanoma cells containing V600E-B-Raf treated with PBIT, a specific inhibitor of this mutant protein, were found to develop resistance to the drug by activating Akt3 signaling (21). Finally, combined targeting of Akt3 and V600E-B-Raf has been reported to synergistically inhibit melanoma (22). Thus, agents regulating these 2 signaling pathways could have significant potential to prevent melanocytic lesion development (10, 15, 17).

PBIs (S,S'-1,4-phenylenbis(1,2-ethanediyl)bis-isoselenourea) is a selenium containing analogue of the iNOS inhibitor PBIT that has been evaluated as a therapeutic agent for treatment of systemically spread melanoma; however, its chemopreventive potential has not been examined (23–25). In this study, the chemopreventive efficacy of topically applied PBISe has been tested on laboratory-generated skin reconstructions containing melanocytic lesions and xenografted cutaneous melanoma tumors in mice (26, 27). Mechanistically, PBIs decreased Akt3 signaling resulting in increased MAP kinase activity to inhibitory levels, which promoted cell senescence and apoptosis (23). Thus, topically applied PBIs could have significant clinical potential for preventing early melanocytic lesion development in skin.

Materials and Methods

PBIs and PBIT synthesis

Bromide salts of PBIs (Mol. Wt. 538.80) and PBIT (Mol. Wt. 444.44) were synthesized, and identity as well as purity verified by nuclear magnetic resonance (NMR) and mass spectrometry as described previously (24). Aliquots of 10 mmol/L PBIs and PBIT (>95% purity by HPLC) stock solutions made in PBS were stored at −20°C for in vitro use. For in vivo experiments PBIs and PBIT were dissolved in dimethyl sulfoxide (DMSO) to make a 48× stock solution and diluted to 120 μL in acetone prior to use.

Cell lines and culture conditions

WM35 radial growth phase melanocytic lesion cell line expressing green fluorescence protein (GFP) and WM115 vertical growth phase cells were grown as described previously (28). Normal human primary melanocytes were cultured in 1× MCDB (Molecular, Cellular, and Developmental Biology) 153 (Sigma), 2% FBS, 10% chelated FBS (Hyclone), 100 nmol/L endothelin 3 (ET3; VWR), 10 ng/mL stem cell factor (SCF; R&D), 20 pmol/L cholera toxin (Sigma), 4.5 ng/mL bFGF (Promega) and 2 mmol/L l-Glutamine (Mediatech) as described previously (29). Normal human FF2441 fibroblasts and GFP-expressing human metastatic melanoma cell lines UACC 903 and 1205 Lu were cultured in high glucose Dulbecco’s modified Eagle’s medium (DMEM) with glutamax (Invitrogen) supplemented with 10% FBS. Passage 2 to 5 human foreskin keratinocyte cells were isolated and cultured in Epilife E-medium (a serum-free HEPES based medium) containing 1X HKGS consisting of bovine pituitary extract, bovine insulin, hydrocortisone, bovine transferrin, and human epidermal growth factor (EGF; Cascade Biologics) as detailed previously (17).

Creation of laboratory-generated skin

Laboratory-generated skin was made from human cell lines (average size measurements: length- 14 mm, breadth-21 mm, height- 1 mm) by suspending normal human FF2441 fibroblast cells in 10% reconstitution buffer, 10% DMEM (Mediatech), 2.4 μL/mL of 10 mol/L NaOH, and 80% collagen I (Becton Dickinson) at a cell density of 3.75 × 10^5 cells/mL on ice followed by incubating 1.5 mL aliquots in 12-well culture plates at 37°C tissue culture incubator for 3 hours to form a dermal matrix (17). One milliliter aliquot of E-medium was added to each well containing a dermis and allowed to grow for 2 days. A mixture of GFP-expressing WM35 or UACC 903 cells and normal human keratinocytes at a ratio of 1:10 were resuspended in 1 mL E-medium and added on top of the dermal matrix to produce a keratinized layer containing noninvasive melanocytic lesions or invasive melanomas. After 2 days of incubation, skin reconstructions were transferred onto wire grids to form a complete keratinized layer for 7 to 8 days in a tissue culture incubator (17). During this period, developing skin reconstructions were fed via diffusion from E-medium (replaced on alternate days) below the wire grids.

Topical drug treatment of laboratory-generated skin

Skin reconstructions containing noninvasive melanocytic lesions or invasive melanoma tumor nodules that were similar in number and size were grouped into control vehicle PBS or 5, 10, 15, and 20 μmol/L PBIs treatment groups (n = 3 skin reconstructions in each group) and exposed to each agent applied once per day for 8 days. At the end of treatment, 6 images from each skin reconstruct were photographed using a Nikon SMZ 1500 fluorescent microscope (Nikon Instruments) to quantitate number and area of melanoma tumor nodules expressing GFP using IP Lab software (BD Biosciences). Average area occupied by melanoma tumor nodules from each treatment group was analyzed by fixation with 10% paraformaldehyde (Electron Microscopy Science) followed by storage in 0.5 mol/L EDTA. Skin reconstructions were trimmed into strips, frozen in optimum cutting temperature (OCT) compound and sectioned.

Histological and morphological characterization of laboratory-generated skin

Morphology and architecture of skin reconstructions prior to, and at the end of each treatment regime were analyzed by fixation with 10% paraformaldehyde (Electron Microscopy Science) followed by storage in 0.5 mol/L EDTA. Skin reconstructions were trimmed into strips, frozen in optimum cutting temperature (OCT) compound and sectioned.
Formalin-fixed paraffin-embedded sections were stained with hematoxylin and eosin (H&E) to examine skin architecture, whereas frozen sections were used to photograph GFP-expressing tumor nodules.

**Melanoma tumor xenograft studies using topical PBISe treatment**

Animal experiments were conducted according to protocols approved by Institutional Animal Care and Use Committee at Penn State University. Twelve, 4 weeks old female athymic nude mice having an average weight of 20 g (Harlan Sprague Dawley) were injected with $2.5 \times 10^5$ cells in 200 μL DMEM-containing 10% FBS into the subcutaneous skin layer on the right and left flanks above the rib cages to conduct tumor kinetics studies. After 24 h, mice were randomly assigned to 3 groups ($n = 4$ mice per group) before starting topical treatment. Vehicle control (Acetone-120 μL), 0.095 μmoles PBISe or PBIT (corresponding to 15 μg selenium each side or 1.5 ppm selenium/animal) in 120 μL vehicle were topically applied at the site of tumor cells injection every day for 29 days. Average tumor size and body weights from each treatment group were measured and plotted against days.

For mechanistic studies, $5 \times 10^6$ UACC 903 cells were injected into the subcutaneous skin layer of nude mice. Mice were treated with vehicle, PBIT or PBISe, as detailed above. Size and time matched tumors were harvested at days 9, 11, 13, and 15 to assess changes in cell proliferation and apoptosis (23). Cell proliferation and apoptosis rates in tumor cells were measured in formalin-fixed, paraffin-embedded tumor sections using purified mouse anti-human Ki-67 (PharMingen) and TUNEL (terminal deoxynucleotidyl transferase–mediated dUTP nick end labeling) staining (Roche Diagnostics). A minimum of 6 different tumors with 4 to 6 fields per tumor was analyzed and results reported as the average ±SEM.

**Analysis of caspase-3/7 activity in tumor lysates**

A small portion of flash frozen tumor was pulverized into powder followed by isolation of protein lysates using protein lysis buffer [600–800 μL per 50 mg powder, 50 mmol/L Tris-HCl, pH 7.5 containing 0.1% Triton X-100, 1 mmol/L EDTA, 1 mmol/L EGTA, 50 mmol/L sodium fluoride, 10 mmol/L sodium β-glycerol phosphate, 5 mmol/L sodium pyrophosphate, 1 mmol/L activated sodium orthovanadate, protease inhibitor cocktail from Sigma and 0.1% (v/v) 2-mercaptoethanol]. Whole tumor lysates were centrifuged at 10,000 × g thrice to remove cell debris. Protein concentrations were quantitated using Bio-Rad protein assay reagent (Bio-Rad laboratories) and analyzed for caspase-3/7 activity. Caspase-3/7 activity was determined by incubating 100 μg of protein lysate with caspase-3/7 substrate as described previously (23, 30).

**Toxicity assessments**

Four to six weeks old female nude mice (Harlan Sprague Dawley) were treated with either vehicle control, PBISe or PBIT ($n = 5$) as described in tumor kinetics studies. At the end of treatment, blood was collected from each sacrificed animal in a plasma separator tube with lithium heparin (BD Microtainer) following cardiac puncture and analyzed for SGOT (AST—aspartate aminotransferase), SGPT (ALT—alanine aminotransferase), alkaline phosphatase, glucose, blood urea nitrogen, total protein and creatinine levels to ascertain possible liver, heart, kidney, and pancreas related toxicity. Harlan has provided values of serum enzymes and metabolites representing range of normal values for female athymic nude mice. A portion of vital organs—liver, heart, kidney, intestine, pancreas, and adrenal—from each animal was formalin-fixed and paraffin-embedded to examine for toxicity-associated changes in cell morphology and tissue organization following H&E staining (23).

**Introduction of siRNAs in to UACC 903 cells by nucleofection**

$1 \times 10^6$ UACC 903 cells were nucleofected (Amaya Nucleofector-I) with 100 and 200 picomoles of scrambled control or Akt3 targeting siRNA, using reagent-R and K-17 program as described previously (15). Scramble and Akt3 siRNAs (Invitrogen) sequences used in this study were reported previously (15, 18). After 2 days recovery in DMEM supplemented with 10% FBS, cells were transferred to media containing no serum and 2 hours later, stimulated with 10% FBS containing DMEM for 60 m. Cell lysates collected and analyzed by Western blotting to measure the expression of Akt3, pAkt (S473), pErk-1/2 (T202/Y204), and α-enolase, which served as a control for protein loading. The intensities of protein bands were quantitated using Image-J and normalized against α-enolase.

**In vitro drug treatment and collection of cell lysates**

$1.5 \times 10^6$ GFP-expressing cells WM35, WM115, or UACC 903 in 10 mL DMEM containing 10% FBS were grown in a 100 mm culture dish for approximately 36 hours. Exponentially growing cells were treated with 3, 5, 9, 10, or 15 μmol/L PBISe in 10 mL DMEM supplemented with 10% FBS for 6, 12, and 24 hours. Total cell lysates from adherent and floating cells were collected from PBS and PBISe treated plates. Floating cells were transferred to a 50 mL conical tube, centrifuged at 1,500 rpm for 5 m and cell pellet washed twice with PBS before being lysed using protein lysis buffer. Adherent cells were washed twice with PBS and incubated with 100 μL protein lysis buffer for 30 m on ice. Lysates collected from adherent and nonadherent cells were then combined, centrifuged at 10,000 rpm for 10 m at 4°C, and clear supernatant transferred to a prechilled 1.5 mL centrifuge tube (31). Protein concentration in the samples was quantitated using the bichinchoninic acid (RCA) assay (Thermo Scientific).

**Western blotting**

Fifteen to thirty micrograms of protein derived from cells treated with different agents were analyzed by Western blotting on NuPAGE Gels (Invitrogen). Following electrophoresis, samples were transferred to polyvinylidene
Caspase-3/7 activity in the cell lysates collected for Western blot analysis was determined using Apo-ONE (Promega Corporation). In brief, 20 to 30 μg of protein in 40 μL lysis buffer was incubated with caspase-3/7 substrate (R110-Z-DEVD dissolved in caspase-3/7 assay buffer) for 2 hours at 37°C with constant shaking in a light protected container. Amount of R110 released was determined using a SPECTRA max-M2 plate reader ( Molecular Devices Corporation) set at 485 nm excitation and 520 nm emission wavelengths. Average values of relative fluorescence units from triplicate wells were plotted as a bar graph with ±SEM (23, 30, 32).

**Cell viability, IC50, proliferation, and cell-cycle analysis**

The viability of cells was measured using the MTS Assay (Promega Corporation). A total of 5 × 10^{3} FF2441 or melanoma cells/well in 100 μL of DMEM containing 10% FBS were grown in a 96-well plate for 36 or 72 hours to reach approximately 70% confluence and treated with either PBS vehicle, PBIT or PBISe for 24 hours and number of viable cells compared with PBS controls. Similarly, 20 × 10^{3} melanocytes/well in 100 μL of melanocyte medium were grown in a 96-well plate for 72 hours to reach approximately 70% confluence and treated with 1 to 100 μmol/L PBISe for 24 hours. IC_{50} values for each compound were determined for each cell line from 3 independent experiments using GraphPad Prism version 4.01 and averages represented with ±SEM (GraphPad software).

Cellular proliferation rate was measured by seeding 5 × 10^{3} cells in a 96-well plate, followed by treatment with PBISe, PBIT, or PBS for 24 hours. Cells were labeled with BrdU, using a Cell Proliferation ELISA kit, 4 hours prior to end of treatment (Roche Diagnostics; ref. 18). Amount of BrdU taken up by cells was measured by incubating with a peroxidase conjugated anti-BrdU binding antibody and read in a 96-well plate reader at 370 nm and compared to PBS treated control cells. Values were determined from 3 independent experiments.

Cell-cycle analysis was undertaken by growing 1.5 × 10^{6} melanoma cells in 100-mm culture dishes for 36 hours, followed by treatment with PBISe or PBIT for 24 hours. Adherent and nonadherent cells were collected and stained using 1 mL of 100 μg/mL propidium iodide (Sigma), 20 μg/mL of RNase A (Roche Diagnostics), and 3 μg/mL of Triton X-100 dissolved in 0.1% (w/v) sodium citrate for 30 m at 4°C (33). Stained cells were analyzed using the FACScan analyzer (BD Biosciences). Data were processed utilizing ModFit LT software (Verity Software House).

**Statistical analysis**

Statistical analysis was undertaken using the 1-way ANOVA followed by Dunn’s multiple comparisons test or Student’s t test. Results were considered significant at P < 0.05.

**Results**

**PBISe but not control PBIT inhibits melanocytic lesion and melanoma cell growth in culture**

PBISe and PBIT are iNOS inhibitors containing selenium and sulfur, respectively, which are shown diagrammatically in Figure 1A (23, 34). PBISe but not PBIT was effective at reducing the growth of noninvasive melanocytic lesion WM35 and WM115 cells. IC_{50} values of PBISe for WM35 and WM115 cells were 9.5 and 9.7 μmol/L, respectively (Fig. 1B). Prior studies have shown antimeLANoma activity of PBISe against invasive metastatic melanoma UACC 903 and 1205 Lu cells (23). In contrast, PBIT had no effect on cell viability at these concentrations suggesting a possible cancer preventive potential for PBISe (Fig. 1B). Specificity of PBISe for killing melanocytic lesion and melanoma cells but not normal melanocytes or fibroblast cells was determined by comparing the IC_{50} values to normal human epidermal melanocytes (NHEM) and fibroblasts (FF2441) that had a value of 30.7 and 17 μmol/L, respectively (Fig. 1C). Fibroblasts and melanocytes had an IC_{50} value that was 1.7 to 3.8 fold higher than that of melanocytic lesion or melanoma cell lines. Thus, PBISe can inhibit survival of cultured noninvasive melanocytic lesion as well as invasive melanoma derived cell lines and has a lesser effect on normal cells.

**PBISe is effective at preventing melanocytic lesion development in laboratory-generated skin**

The potency of PBISe for killing cutaneous noninvasive melanocytic lesion or invasive melanoma cells was examined by seeding these cells into laboratory-generated skin and topically treating with PBISe (17). Skin reconstructs used for this study contain GFP-tagged WM35 melanocytic lesion or UACC 903 melanoma cells and the amount of GFP following treatment was quantified. This approach is an accepted organotypic skin melanoma model for evaluating the efficacy of topically added chemopreventive agents (17). PBISe decreased total area occupied by GFP following treatment was quantified. This approach is an accepted organotypic skin melanoma model for evaluating the efficacy of topically added chemopreventive agents (17). PBISe decreased total area occupied by GFP following treatment was quantified. This approach is an accepted organotypic skin melanoma model for evaluating the efficacy of topically added chemopreventive agents (17). PBISe decreased total area occupied by GFP following treatment was quantified.
Figure 1. PBISe inhibits melanoma cells growth in culture. 
A, structures of PBIT and PBISe. PBISe is an isosteric analogue of PBIT and was synthesized by replacing sulfur with selenium. B, PBISe inhibits growth of cultured melanoma cells. Cell lines representing noninvasive melanocytic lesions (WM35 and WM115) were treated with increasing concentrations of PBIT or PBISe for 24 hours. Cell viability was determined by MTS assay and IC\textsubscript{50} values calculated using GraphPad Prism. Results of 3 independent experiments were plotted; bars, mean ± SEM. C, PBISe has negligible effect on normal cells at concentrations killing melanocytic and melanoma cells. Human normal epidermal melanocytes, fibroblasts, and melanocytic lesion or melanoma cell lines (WM35, WM115, 1205 Lu, and UACC 903) were treated with PBISe for 24 hours and IC\textsubscript{50} values compared; bars, average IC\textsubscript{50} from 3 independent experiments ± SEM.
Figure 2. Topical application of PBISe inhibits growth of melanoma tumors in laboratory-generated skin reconstructs. A and B, Topically applied PBISe inhibited the growth of melanocytic lesions and melanoma tumors developing in skin reconstructs. Laboratory-generated skin containing melanocytic lesions or melanoma tumors were treated with PBISe or vehicle for 8 days and sizes of area occupied by developing GFP tumors quantified. (** and *** = P < 0.05, One-Way ANOVA) C, PBISe causes negligible damage to the cells present in skin. H&E stained skin reconstructs containing melanocytic lesion of melanoma cells were treated topically with PBISe and compared to untreated controls. No change in skin morphology or morphology of keratinocytes or fibroblasts was observed.
negligible effect (Figs. 2A and B; \( P < 0.05 \), One-way ANOVA).

Skin architecture and morphology of constituent cells containing radial WM35 and invasive UACC 903 melanoma tumors were compared in H&E stained sections from PBISe versus control treated skins (Fig. 2C). Similar histological and morphological features were observed under all treatment regimes showing an intact keratinized layer and dermal fibroblasts having a similar size, shape, and distribution, suggesting that PBISe does not damage skin or the cells present in it (Fig. 2C). Thus, PBISe can decrease melanocytic lesion and melanoma tumor development in laboratory-generated skin with negligible effect on skin architecture or morphology of fibroblasts or keratinocytes.

Topical application of PBISe is not toxic to mice and inhibits melanoma tumor development in animal skin

To determine the inhibitory efficacy of PBISe on tumor inhibition in animal skin, \( 2.5 \times 10^5 \) cutaneously invasive UACC 903 cells were subcutaneously injected above the left and right rib cages of 4 to 6 weeks old female nude mice, and skin above the tumor treated topical with PBISe, PBIT, or vehicle acetone. Noninvasive WM35 melanocytic lesion cells lines could not be used for this study since they do not form tumors in animal skin. Animals containing cutaneous UACC 903 tumors were exposed to PBISe and consistently found to have approximately 50% smaller lesions starting at day 17 compared to those treated with either PBIT or vehicle control (Fig. 3A; \( P < 0.05 \); 2 tailed Students’ t test).

No statistically significant differences in animal body weights between PBISe, PBIT, or control treated groups were observed, suggesting a lack of systemic toxicity at drug concentrations used (Fig. 3B). This was subsequently confirmed by examining serum biomarkers for blood parameters indicative of major organ related toxicity at the end of treatment regime, which showed negligible differences between groups and fell within normal parameter for this mouse species (Fig. 3C; ref. 35). Thus, PBISe can retard cutaneous tumor formation in animals when applied topically without significant systemic toxicity.

PBISe exposure triggered apoptosis in melanoma cells growing in culture and in xenografted tumors

The mechanism preventing cutaneous melanocytic lesion development following PBISe treatment was established by analyzing apoptosis and proliferation rates using lysates derived from cultured melanocytic lesion or melanoma cells or xenografted tumors treated with agents. Since elevated caspase-3/7 activity in cultured cells is an indicator of apoptotic cell death, levels in cells exposed to PBISe or PBIT were measured using Apo-ONE homogenous caspase-3/7 activity kit. A dose-dependent increase in caspase-3/7 activity in cultured noninvasive WM115 and invasive 1205 Lu cells was observed only when cells were treated with PBISe but not with PBIT (Fig. 4A). For both cell lines, higher concentrations of PBISe caused significant cell death, which consequently reduced caspase-3/7 activity. Similar results were observed for protein lysates derived from cultured noninvasive WM35 and invasive UACC 903 cells treated with PBISe and analyzed for caspase-3/7 activity (Fig. 4B). Analysis of caspase-3/7 activity in protein lysates collected from size and time matched tumors harvested at days 11, 13, and 15 likewise showed a 1.5 to 2.2-fold increase in apoptosis (Fig. 4C; \( P < 0.05 \); Student’s t test). In contrast, PBIT or acetone vehicle treatment did not increase apoptosis rates. Confirming these observations, PBISe treatment increased number of TUNEL positive apoptotic cells observed following immunohistochemical analysis of xenografted size and time matched tumors isolated at days 11, 13, and 15 in a dose-dependent manner (Fig. 4D). Compared to controls, a consistent increase in TUNEL positive cells was observed following PBISe treatment (Fig. 4D; \( P < 0.05 \); Student’s t test). Thus, PBISe increases apoptosis rates in melanoma cells growing in culture or in animals.

PBISe decreases cellular proliferation rates and halts cell-cycle progression in cultured melanocytic lesion as well as in melanoma cells

Since topically applied PBISe had potential to prevent noninvasive melanocytic lesion as well as invasive melanoma cell growth in laboratory-generated skin and decrease xenografted tumor development, effect of PBISe exposure on proliferation rates and cell-cycle progression was examined. Compared to PBS vehicle or PBIT, a significant decrease in proliferation measured by BrdU incorporation was observed at 1 to 5 \( \mu \)mol/L PBISe for noninvasive melanocytic lesion WM35 or WM115 cells (Fig. 5A). In contrast, metastatic melanoma 1205 Lu cells, required approximately 14 \( \mu \)mol/L PBISe to decrease cell proliferation by approximately 60% (Fig. 5A).

Basis for decreased cellular proliferation following PBISe treatment was examined next by analyzing percentage of cell population in the different phases of the cell cycle. Compared to control PBS or PBIT, PBISe treatment increased the sub-G0/1 WM35 cell population and decreased S-phase cell population in a dose-dependent manner (Fig. 5B). At 10 to 15 \( \mu \)mol/L PBISe, a 7 to 16 fold increase in sub-G0/1 WM35 cells were observed compared to PBIT indicating increased apoptosis (Fig. 5C). Compared to controls, a significant decrease in S-phase cells was observed at 5 to 15 \( \mu \)mol/L PBISe (Fig. 5D). Thus, PBISe treatment inhibited noninvasive melanocytic lesion and metastatic melanoma cell proliferation by decreasing the S-phase cell population and increasing the sub-G0/1 component indicating elevated apoptotic cell death.

Targeting Akt3 using siRNA or PBISe increased MAP kinase activity reducing the proliferative potential and promoting apoptosis of cultured cells

Western blotting was used to identify the protein signaling events decreasing cell survival following Akt3 inhibition in invasive metastatic cells using siRNA or following PBISe treatment. Compared to UACC 903 cells nucleofected with scrambled siRNA, Akt3 siRNA led to 2 to 5 fold higher levels of pErk1/2 in a dose-dependent manner.
Tumor kinetics (UACC 903)

A

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<th>Tumor volume (mm³)</th>
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<td>17</td>
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<td>19</td>
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50% tumor reduction (P < 0.05 t-test)

B

Animals body weight

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C

Analysis of serum enzymes and metabolites

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<td>SGOT (AST) (Units/L)</td>
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<td>171.9 ± 30.2</td>
<td>218.6 ± 27.2</td>
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<td>SGPT (ALT) (Units/L)</td>
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<td>48.6 ± 6.7</td>
<td>76 ± 31.5</td>
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<td>Alkaline phosphatase (ALP) (Units/L)</td>
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<td>81.3 ± 15.2</td>
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<td>Blood glucose (mg/dL) (108 – 232)</td>
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<td>Creatinine (mg/dL) (0.2 – 0.31)</td>
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<td>Blood urea nitrogen (BUN) (mg/dL) (18 – 35.7)</td>
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<td>17.3 ± 1.9</td>
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<td>Total protein (g/dL) (3.6 – 6.1)</td>
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<td>5.3 ± 0.18</td>
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<td>Metabolic activity: liver and kidney function</td>
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</table>

Figure 3. Topically applied PBISe inhibit development of cutaneous melanomas with negligible systemic toxicity. A, topically applied PBISe decreases development of subcutaneous melanocytic lesions in mice. Following injection of UACC 903 cells subcutaneously into mice, area above sites was topically treated daily with acetone, PBISe, or PBIT and sizes of developing tumors measured on alternate days. (* P < 0.05, Student t-test) B, daily topical treatment with PBISe did not significantly alter animal body weight. Body weights of animals treated topically with acetone, PBISe, or PBIT were measured on alternate days to establish possible toxicity. C, daily topical treatment with PBISe did not significantly alter blood biomarkers indicative of major organ related toxicity. Blood collected from animals treated topically with PBISe, PBIT, or acetone was collected from animals at day 29 and analyzed for enzyme activities of liver (SGOT, SGPT), kidney (alkaline phosphatase), and heart (SGOT, SGPT, and alkaline phosphatase). Concentrations of metabolites (glucose, creatinine, urea nitrogen, and total protein) in serum were also measured. Control values for this mouse species are listed below each factor in brackets for comparison.
Figure 4. PBISe induces apoptosis to inhibit survival of melanocytic lesion and melanoma cells. A and B, PBISe treatment increased caspase-3/7 activity in cultured melanocytic lesion and melanoma cell lines. Cell lines representing noninvasive melanocytic lesions (WM35 and WM115) and invasive melanomas (1205 Lu and UACC 903) were treated with increasing concentrations of PBIT or PBISe for 24 hours and caspase-3/7 activity from culture media or cell lysates measured. Bar graph represents fold increase over PBS control. C and D, tumors from mice treated with PBISe showed increased caspase-3/7 activity in tumor lysates and had more TUNEL positive cells compared to PBIT treated controls. Protein lysates collected from PBISe or PBIT-treated size and time matched tumors harvested at days 11, 13, and 15 were incubated with R110-conjugated caspase-3/7 substrate (R110-Z-DEVD) for 1 hour and released Rhodamine-110 measured in a plate reader. TUNEL positive cells from formalin-fixed paraffin-embedded size and time matched tumors from PBISe or PBIT were scored for percentage of positive cells from a minimum of 3 tumors (3 to 5 fields/tumor); bar, average ± SEM (* and ** P < 0.05, Student t-test).
Figure 5. PBISe treatment of cultured melanoma cells increases the subG0/G1 and decreases the S-phase cell population. A, PBISe inhibits the proliferative potential of cultured melanocytic lesion and melanoma cells. A, BrdU ELISA kit was used to assess the proliferative potential of cells following treatment with PBISe, PBIT, and PBS; bar mean ± SEM. B, C, and D, PBISe treatment decreased population of cells in S-phase and increased sub-G0/G1 phase of the cell cycle. Cultured noninvasive WM35 cells were treated with increasing concentrations of PBIT, PBISe, or PBS for 24 hours. Adherent and detached cells were collected, stained with propidium iodide, and cell-cycle analyzed using a FACScan.
Figure 6. PBISe targets Akt3 signaling to regulate noninvasive melanocytic and invasive melanoma cell survival. A, inhibition of Akt3 signaling increases MAP kinase pathway signaling. SiRNA mediated inhibition of Akt3 protein levels led to increased pErk1/2 levels in UACC 903 cells. UACC 903 cells were nucleofected with 100 and 200 picomoles of siRNA targeting Akt3 or a control scrambled siRNA and cell lysates collected after 2 hours serum starvation followed by 1 hour serum stimulation. Expression of pAkt and pErk1/2 were measured by Western blotting and the band intensities measured using Image-J software. Protein levels were normalized against α-enolase and the pErk1/2 to pAkt ratio measured. B and C, PBISe treatment of noninvasive melanocytic lesion and invasive melanoma cell decreased Akt3 signaling and increased MAP kinases pathway activity to inhibitory levels. PBISe treatment in all cell lines decreased the pAkt and downstream pPRAS40 levels, which consequently increased levels of pErk1/2. Combined targeting of these pathways promoted decreased cell growth indicated by lowered cyclin-D1 and increased p21 levels as well as increasing rates of apoptosis observed as increased levels of cleaved caspase-3 and PARP.
Akt3 activity thereby decreasing the inhibitory effect on increasing pErk1/2 (Fig. 6B and C). Thus, PBISe reduced p21 levels increased consistently in a dose and time-dependent manner following PBISe treatment corresponding to p21 levels increased consistently in a dose and time-dependent manner (Fig. 6B). Similar effects on Akt and Erk signaling pathways were observed when invasive UACC 903 cells were treated with PBISe (Fig. 6C). For WM35, WM115, and UACC 903 cell lines, cyclin D1 levels decreased and p21 levels increased consistently in a dose and time-dependent manner following PBISe treatment corresponding to increasing pErk1/2 (Fig. 6B and C). Thus, PBISe reduced Akt3 activity thereby decreasing the inhibitory effect on V600EB-Raf, which led to high inhibitory MAP kinase pathway activity down regulating cyclin D1 and increasing p21 levels thereby promoting apoptosis and cell senescence.

Discussion

Despite skin cancer prevention programs, availability of UV protecting sunblocks, and surgical procedures for removing preinvasive melanocytic skin lesions, incidence of metastatic melanoma and mortality rates resulting from disseminated metastatic disease continue to rise (1, 36). Therefore, additional agents to prevent the disease from developing or progressing past its earliest stages, which could be added to creams, lotions, or sunblocks, are needed. Preventive agents of this type would have potential to decrease disease incidence and mortality rates (6).

Chemopreventive effects of inorganic and organic selenium derivatives have been evaluated for colon, lung, prostate, and esophageal cancers (37–39). Selenium containing compounds can induce the activities of phase-II enzymes and inhibit phase-I enzymes to prevent the progression as well as reduce growth of tumor cells (23). Selenium has also been shown to increase the therapeutic efficacy of drugs in combination treatment regimes, and to increase the potency of chemopreventive and therapeutic agents when selenium is substituted for sulfur in drugs by increasing rates of cellular apoptosis (23). Furthermore, low selenium levels have been reported in the serum of patients suffering from cancers including melanoma (40, 41). Various preclinical and clinical studies have shown anticancer activity of inorganic and organic selenium containing agents (37, 39, 40, 42) and the mechanistic basis of tumor inhibition was studied (43–45). Our study shows that substitution of sulfur in PBIT with selenium provided the compound with novel properties to enhance its anticancer activity. Promoting enhanced anticancer activity by replacing sulfur in various chemotherapeutic agents with selenium has been reported (37, 42). PBISe could be metabolized in to alkyl selenol or incorporated in to proteins to confer this activity, but the possibility was not explored in this study.

While selenium-containing compounds can enhance the cancer inhibitory activity of agents, recent preclinical and clinical studies have shown that administration of selenium compounds might have no effect or induce serious side effects (46, 47). For example, combining selenium and vitamin E in a cancer prevention trial (SELECT), the largest phase III randomized placebo-controlled study, found that oral administration of selenomethionine did not prevent prostate cancer and might increase rates of diabetes (47). Another study found that dietary supplementation of an anti-oxidant mixture containing selenium increased melanoma risk in women (46). Lack of chemopreventive efficacy of selenomethionine for inhibiting tumor growth has also been reported in preclinical rodent models (43, 44).

Although several concerns and controversies abound regarding the use of selenium for preventing cancers, the present study shows a strategy to improve the efficacy of pharmaceutical agents by substituting sulfur with selenium. In addition, several recent studies have shown that the dose and form of selenium are the key factors that influence the outcome of selenium treatment (43, 44). Therefore, care must be taken when deciding the dose and form of selenium for clinical use. Methylselenocysteine was effective at inhibiting tumor growth compared to selenomethionine suggesting requirement for particular structural features necessary for anticancer activity (48). Based on our observations, PBISe topical treatment successfully retarded the development of early WM35 as well as metastatic UACC 903 in laboratory-generated skin reconstruct models. These observations suggest PBISe might be useful for inhibiting the progression of dysplastic nevi in to premalignant lesions as well as metastatic melanomas. Also, where a cutaneous melanoma has been removed, PBISe might prevent recurrence which would be a scenario analogous to the use of tamoxifen as a breast cancer prevention agent where any risks incurred by using the agent are outweighed by the high risk of cancer recurrence (49).

Prior studies have shown that intraperitoneal administration of PBISe but not PBiT retarded melanoma tumor growth without causing systemic toxicity (23). This study extends this initial discovery showing that daily topical application of PBISe retards cutaneous noninvasive melanocytic lesion or invasive melanoma development. Topical PBISe decreased tumor development in laboratory-generated skin reconstructs by 70% to 80% and the development of tumors in the skin of animals by approximately 50% compared to controls, thereby demonstrating the chemopreventive potential of PBISe.

Similar to our data, prior studies have shown that substituting sulfur with selenium could increase the potency of pharmaceutical agents to inhibit oncogene function (37, 45). Similar to other selenium containing chemotherapeutic agents, PBISe triggered cellular apoptosis by inhibiting Akt3/PRAS40 signaling, thereby increasing caspase-3/7 activity in melanoma cells (30). Recent studies have also shown that selenium can decrease active and total Akt
protein levels by destabilization of this protein in cells; however, decreased total Akt levels were not observed in this study (50). PBIsE can also trigger cell death mediated by activation of caspases. Caspases are activated during the process of cell apoptosis and caspase-3 is a key factor responsible for either partial or complete triggering of this cascade. PBIsE elevated caspase-3/7 activity of cells by inducing cleaved caspase-3 and PARP. TUNEL assay carried out on tumors taken from mice treated with PBIsE showed a 2-fold increase of apoptosis compared to controls, indicating reduced tumor volume is, in part, due to increased cellular apoptosis.

In addition to triggering apoptosis and increasing the sub-G0/G1 cell population, PBIsE decreased cell proliferation by inhibiting the proportion of cells in the S-phase of the cell cycle (23). Mechanistically, this occurred by PBIsE regulating MAP kinase signaling through inhibition of the Akt3 pathway. By decreasing the Akt3 pathway activity, mutant \(V_{\text{600E}}\) B-Raf could not be phosphorylated by Akt3 to decrease the activity of the MAP kinase pathway (23). Akt3 has been shown to phosphorylate \(V_{\text{600E}}\) B-Raf to release the senescence block induced by high MAP kinase signaling pathway activity, which in turn can promote early melanoma development (8, 31).

In conclusion, topical application of PBIsE inhibited cutaneous noninvasive melanocytic lesion development and invasive melanoma development in skin reconstructs and retarded the growth of subcutaneous xenografted tumors with negligible systemic toxicity. Since no targeted chemopreventive agents are available for melanoma, PBIsE is a promising candidate for preventing early melanocytic lesion development with insignificant side effects. Thus, PBIsE has potential for use as a topical chemopreventive agent to retard development of cutaneous melanocytic lesions or melanoma development.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

Acknowledgments

The authors thank Drs. Sung Jin Huh, Arati Sharma, and Mitchell Cheung for technical assistance with animal experimentation and Drs. Melissa Tran, Samina Alam, and Craig Meyers for help with the laboratory-generated skin technique.

Grant Support

This work was supported by the American Cancer Society (RSG-04-053-01-GMC), NIH CA-127892-01A, NIH NCI contract (NO2-CA-56603), and The Zimmerman Foundation for Melanoma Research.

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Received August 17, 2010; revised January 21, 2011; accepted February 21, 2011; published OnlineFirst March 2, 2011.

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