5-(Bis(3-(2-hydroxyethyl)-1H-indol-2-yl)methyl)-2-hydroxybenzoic acid (BHIMHA): showing a strategy of designing drug to block lung metastasis of tumors

This article was published in the following Dove Press journal:
Drug Design, Development and Therapy
16 February 2016
Number of times this article has been viewed

Abstract: Early metastasis is still the most recalcitrant factor in the treatment of lung cancer patients. By analyzing the structures and comparing the docking scores of the known pharmacophores, the authors of this paper designed 5-(bis(3-(2-hydroxyethyl)-1H-indol-2-yl)methyl)-2-hydroxybenzoic acid (BHIMHA) as a promising lead compound to develop metastasis inhibitors. In vitro, 5, 10, and 20 µM of BHIMHA concentration dependently inhibited the migration and invasion of A549 cells. In vivo, 0.4, 2.0, and 8.9 µmol/kg of BHIMHA dose dependently inhibited the metastasis of LLC (Lewis Lung Carcinoma) toward lung. In vivo, 2 µmol/kg of BHIMHA showed additional actions of slowing the growth of the primary tumor of C57BL/6 mice and S180 mice as well as inhibiting xylene-induced ear edema of the mice. Therefore, BHIMHA simultaneously blocked tumor metastasis toward lung, slowed the primary tumor growth, and limited the inflammation. These pharmacological actions were correlated with the inhibition of PKCα and NF-κB expression.

Keywords: migration, invasion, lung metastasis, anticancer, anti-inflammation, PKCα inhibitor

Introduction

Despite the advances in conventional and targeting therapies, lung cancer is one of the most aggressive human cancers. In the last decade, the early metastasis toward lung has been one of the most recalcitrant factors leading to failed treatment and high mortality rate of the cancer patients. Metastatic cascade of the cancer cells involves a series of events such as epithelial–mesenchymal transition, migration and invasion of cancer cells, the intravasation of the cancer cells toward the systemic circulation, the adhesion of the cancer cells onto endothelial cells, the extravasation of the cancer cells, the colonization of the cancer cells in distant organs, and the cancer cells-induced angiogenesis. Of the metastatic cascade, the migration and the invasion are the critical upstream events.

Protein kinase C (PKC) is a family of the kinases consisting of distinct isoforms of lipid-regulated serine/threonine kinases and exemplifies the specifically signaling molecules that link multiple cellular processes to cancer. The overexpression of PKC activated by phorbol esters promotes tumor development and is one of the biomarkers for cancer diagnosis. In respect of isoforms PKCα, PKCβIII, PKCγ, PKCη, PKCδ, and PKCθ, the regulation of PKCα relates to the proliferation, survival, differentiation, and motility of the cancer cells. As a regulator, the expression of PKCα is responsible for...
the invasion and metastasis of cancer cells. For instance, in vitro the downregulation of PKCα expression leads to the inhibition of the invasion and the migration of A549 cells, and in vivo not only is PKCα level is significantly higher in lung cancer tissue when compared to healthy lung tissue, but also the PKCα traffic in the nuclei is significantly increased when compared to the membrane. The knowledge means the downregulation of PKCα expression would be essentially considered in designing the drug capable of blocking the metastasis of tumors toward the lung.

Inflammatory component contributes to the proliferation, angiogenesis, and metastasis of tumors. Nuclear factor-κB (NF-κB) not only correlates with the onset and the progression of cancer but also correlates with cancer-related inflammation, ie, NF-κB leads to the cross-talk of inflammation with tumor growth, angiogenesis, and metastasis. The knowledge means the inhibition of NF-κB expression also would be considered in designing the drug capable of blocking the metastasis of tumors toward the lung.

It is well known that staurosporine is an inhibitor of PKC and limits the invasive and metastatic abilities of lung tumor cells. BPIC is an inhibitor of tumor growth and minimizes inflammatory response. MIAM upregulates sirtuin-3 and slows the growth of human hepatocellular carcinoma. Based on the pharmacophores from staurosporine, BPIC and MIAM, the drug could be capable of blocking the metastasis of tumors toward the lung. In this context, we surveyed the structural characteristics of staurosporine, BPIC, and MIAM; designed 5-(bis(3-(2-hydroxyethyl)-1H-indol-2-yl)methyl)-2-hydroxyl benzoic acid (BHIHMA); assayed the in vitro activities of BHIHMA inhibiting the proliferation, migration, and invasion of the cancer cells; tested the in vitro effect of BHIHMA on the expression of PKCs and NF-κB from cancer cells; measured the in vivo efficacy of BHIHMA blocking the metastasis of the tumors toward the lung; evaluated the in vivo potency of BHIHMA slowing the tumor growth; and estimated the in vivo action of BHIHMA preventing the mice from xylene-induced ear edema.

Materials and methods

General

The chemicals were purchased from Sigma-Aldrich Co. (St Louis, MO, USA) and were purified when necessary. Column chromatography was performed using silica gel (200–300 mesh, Qingdao Haiyang Chemical Co., Qingdao, People’s Republic of China). Purity of BHIHMA (>95%) and BHIHMA (>98%) were determined with TLC (thin-layer chromatography; Qingdao silica gel plates of GF254) and HPLC (CHIRALPAK AH-H column, 4.6×250 mm, Daicel Chemical Industries Ltd., Osaka, Japan), respectively.

1H-NMR (nuclear magnetic resonance) (800 MHz) and 13C-NMR (200 MHz) spectra were recorded on Bruker Avance II 800 MHz spectrometer (Bruker Daltonik, Billerica, MA, USA) with DMSO-d6 (dimethyl sulfoxide-d6) as the solvent and tetramethylsilane as internal standard. ESI/MS (electrospray ionization/mass spectroscopy) was tested on ZQ 2000 (Waters, Milford, MA, USA) and solariX FT-ICR mass spectrometer (Bruker Daltonik) consisting of an electrospray ionization/matrix-assisted laser desorption ionization dual ion source and 9.4 T superconductive magnet.

HaCaT-immortalized human epidermal cells (HaCaT cells), mice Lewis lung carcinoma cells (LLC cells), and human lung adenocarcinoma cell line A549 were purchased from ATCC, Manassas, VA, USA. Mice sarcoma S180 cells were purchased from the Animal Center of Peking University. Doxorubicin and bovine serum albumin (BSA) were purchased from Sigma. Roswell Park Memorial Institute-1640 (RPMI 1640) medium, Dulbecco’s Modified Eagle’s Medium (DMEM), and fetal bovine serum (FBS) were purchased from Gibco; products of Thermo Fisher Scientific, Waltham, MA, USA. The inserts (8.0 μm pores) of 24-well transwell were purchased from Corning Incorporated (Corning, NY, USA). Staurosporine, NF-κB antibody, and NF-κB inhibitor ammonium pyrrolidinedithiocarbamate (PDTC) were purchased from Beyotime Institute of Biotechnology (Nantong, People’s Republic of China). PKCα, PKCBII, PKCγ, PKCη, and PKCδ antibodies were purchased from Boster Biological Engineering Co., Ltd. (Wuhan, People’s Republic of China); PKC9 antibody was purchased from CST (Shanghai) Biological Reagents Company Limited, Shanghai, People’s Republic of China.

Male ICR mice and male C57BL/6 mice were purchased from the Animal Center of Peking University. The protocol was reviewed and approved by ethics committee of Capital Medical University. The committee assures that the welfare of the animals be maintained in accordance with the requirements of the Animal Welfare Act and in accordance with the NIH Guide for Care and Use of Laboratory Animals. All experiments were performed at least in triplicate. Data are presented as mean ± SD (standard deviation). The statistical analysis of all the biological data was carried out by use of analysis of variance (ANOVA) test. The P-value <0.05 was considered statistically significant.

Design of BHIHMA

As mentioned, the positive regulation of PKCα relates to the proliferation, survival, differentiation, motility, invasion,
and metastasis of the cancer cells. It was hypothesized that inhibiting PKCα expression would lead to the block of the metastasis of tumors toward the lung. The survey of structural characteristics of BPIC, MIAM, and staurosporine, the known PKC inhibitors, led to the design of BHI MHA as a novel inhibitor of PKCα (Figure 1). To theoretically ensure the rationality of the design, five members of PKC family, ie, PKCα (3iw4), PKCβII (3PFQ), PKCγ (2E73), PKCθ (1XJD), PKCη (3TXO), and PKCδ (3UEY), were selected from Protein Data Bank (PDB) as the candidates of the target protein, and BHI MHA was docked into their active sites. The theoretical investigations demonstrated that of the five proteins, only the docking of BHI MHA toward the active site of PKCα gave proper feature and higher docking score. In this case, the docking investigation of staurosporine toward the active site of PKCα was also performed, and the features and scores of BHI MHA and staurosporine are also shown in Figure 1.

Synthetic route of BHI MHA

As depicted in Figure 2, BHI MHA was prepared via a two-step procedure, ie, the Pictet–Spengler condensation of methyl-5-formyl-2-hydroxybenzoate and tryptophol, and the hydrolysis of 5-(bis(3-(2-hydroxyethyl)-1H-indol-2-yl)-methyl)-2-hydroxy benzoate (BHI MHAM).

Docking of BHI MHA into the active site of PKCs

In the molecular docking, the structure of PKCα (3iw4), PKCβII (3PFQ), PKCγ (2E73), PKCθ (1XJD), PKCη (3TXO), and PKCδ (3UEY) from PDB was treated as rigid and prepared by AutoDockTools 1.5, ie, merging nonpolar
hydrogens and assigning gasteiger charges and autodock elements. Then, energy-minimized 3D conformations of staurosporine and BHIMHA were treated as flexible and prepared by AutoDockTools 1.5, ie, merging nonpolar hydrogens, assigning gasteiger charges, finding root and aromatic carbons, detecting rotatable bonds, and setting torsions. The grid box dimensions were set to 22.5 Å × 30 Å × 30 Å using a grid spacing of 0.375 Å for two average structures.

To determine the probable binding conformations, AutoDock4, a very popular docking program with a high success rate, and energy-minimized 3D conformations of staurosporine and BHIMHA were used to dock into the active site of PKC\(\alpha\) (3iw4), PKC\(\beta II\) (3PFQ), PKC\(\gamma\) (2E73), PKC\(\delta\) (1XJD), PKC\(\eta\) (3TXO), and PKC\(\delta\) (3UEY). The global optimization was started with parameters of a population of 100 randomly positioned individuals. The maximum number of energy evaluation was increased to 2.5 × 10^7, and the maximum number of generation in Lamarckian genetic algorithm was increased to 2.7 × 10^5. A Solis and Wets local search was performed with a maximum number of 3,000. During each simulation, 256 runs were carried out for each agent. The resulting 256 conformations of staurosporine and BHIMHA were scored by the lowest binding energy and clustered using a root mean square tolerance of 2.0 Å. Lamarckian genetic algorithm was used to find the appropriate binding positions, orientations, and conformations of staurosporine and BHIMHA in the binding site.

**In vitro antiproliferation assay**

A549 cells, the carcinoma cells, were maintained in RPMI 1640 medium supplemented with 10% FBS, penicillin (100 U/mL), and streptomycin (100 μg/mL). HaCaT cells (control), the noncarcinoma cells, were maintained in DMEM medium supplemented with 10% FBS, penicillin (100 U/mL), and streptomycin (100 μg/mL). The cells were maintained in a humidified atmosphere of 5% CO\(_2\) at 37°C. The medium was renewed every 2 days. The proliferation of A549 cells and HaCaT cells was determined with MTT [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide] assay. In brief, A549 cells or HaCaT cells in the logarithmic growth phase were digested with 0.25% trypsin, added to RPMI 1640 medium or DMEM medium supplemented with 10% FBS, plated into 96-well plates (5 × 10^4 cells/well), and incubated for 6 hours. The cells were treated with BHIMHA (final concentration: 1, 10, 25, 50, and 100 μM), 25 μL solution of MTT in the medium (5 mg/mL) was added, and the plates were incubated at 37°C for 4 hours. The supernatant was discarded and to each well 100 μL of DMSO was added. The optical density (OD) value of BHIMHA-treated well was compared with that of DMSO-treated well. The proliferation of A549 cells or HaCaT cells was represented with the OD value. Each measurement was performed in triplicate. Data are presented as mean ± SD. The statistical analysis of all data was carried out by use of ANOVA test. The P-value <0.05 was considered statistically significant.

**A549 cell migration assay**

Transwell chamber with cell permeable membrane was used for migration assay of A549 cells as described previously.\(^{24}\) In brief, A549 cells (5 × 10^4 cells/chamber) and serum-free medium in the upper chamber were treated with BHIMHA (5, 10, and 20 μM). The medium with 10% FBS, the chemotactic factor, was added into the down chamber (600 μL/chamber), the transwell chamber was incubated at 37°C for 6 hours, and the cells that did not penetrate the membrane and at the bottom of the upper chamber were removed with cotton swabs. The cells were fixed with phosphate-buffered saline (PBS) containing 4%...
formaldehyde for 30 minutes and stained with 0.1% crystal violet for 15 minutes. The cells that penetrated through the membrane into the down chamber were counted under a light microscope at ×200 to take the count of the cells from six visual fields.

**A549 cell invasion assay**

Transwell chambers were used for A549 cell invasion assay as described previously.24 Membrane at the bottom of each chamber was coated with 50 µL matrigel and air-dried overnight. The chamber was blocked with BSA (2%, 50 µL/chamber), incubated at 37°C for 2 hours, and rinsed with PBS. A549 cells (2×10⁴ cells/chamber) and serum-free medium in the upper chamber were treated with BHIMHA (5, 10, and 20 µM). Into the down chamber, 600 µL of FBS (10%, control) was added. A549 cells were cultured at 37°C for 24 hours and the cells that did not penetrate the membrane were removed with cotton swabs. The cells penetrated through the membrane were fixed with 4% formaldehyde for 30 minutes and stained with 0.1% crystal violet for 10 minutes. The cells that penetrated through the matrigel were counted under a light microscope at ×200 to take the count of the cells from six visual fields. Data are presented as mean ± SD. The statistical analysis of the data was carried out by use of ANOVA test. The P-value <0.05 was considered statistically significant.

**Western blot assay**

A549 cells (2×10⁴/well) were seeded in 6-well plates overnight and treated with 5 µM of BHIMHA, 0.1 µM of staurosporine, and 1 µM of PDTC for 24 hours. The cells were harvested, washed twice with PBS, and at 4°C, lysed with ice-cold radioimmunoprecipitation assay buffer (1% NP-40 in 150 mM NaCl, 50 mM Tris, and 2 mM EDTA) for 30 minutes. The protein from the lysate was run on a sodium dodecyl sulfate–polyacrylamide gel, and the bands were transferred to polyvinylidene difluoride membrane. The membrane was blocked with 5% (w/v) BSA for 2 hours, washed thrice with Tris Buffered Saline Tween (TBST), and at 4°C incubated with primary antibodies PKCα, PKCβII, PKCγ or PKCζ or PKCδ or NF-κB (p65, 1:1,000) overnight. The membrane was washed and incubated for 1 hour (at room temperature) with IgG-horseradish peroxidase conjugated secondary antibody and washed thrice with TBST. Western blot chemiluminescence reagents were added for visualizing the bands of the proteins. The data are presented as mean ± SD. The statistical analysis of the data was carried out by use of ANOVA test. The P-value <0.05 was considered statistically significant.

**Tumor metastasis assay**

Tumor metastasis was studied with LLC model as described previously.25 In brief, subcutaneous tumors were implanted by injecting 1.5×10⁷ viable LLC cells in 0.2 mL of normal saline (NS) into the skin of the right armpit of 8-week-old C57BL/6 mice. Nine days after LLC cell injection, the tumor reached ~50 mm³ and the treatment was initiated. The mice were randomly divided to NS-treated group (oral dose: 10 mL/kg/d, 12 consecutive days) and BHIMHA-treated groups (oral dose: 8.9, 2.0, and 0.4 µmol/kg/d, 12 consecutive days). In mice, tumor weight and volume were recorded on 13th day and used to represent the activity. Twenty four hours after the last dosage, all mice were weighed, received ether anesthesia, killed, and immediately dissected to obtain the tumor samples for tests. Data are presented as mean ± SD (standard deviation). The statistical analysis of the data was carried out by use of ANOVA test. The P-value <0.05 was considered statistically significant.

**In vivo antitumor assay**

Male ICR mice were 10 weeks old at the beginning of in vivo antitumor assay.26 S180 cells were subcutaneously injected to form solid tumors. The cells obtained ascitic form from tumor-bearing mice were serially transplanted once per week. The subcutaneous tumors were implanted under mouse skin at the right armpit by injecting 0.2 mL of NS containing 2×10⁶ viable tumor cells. Twenty four hours after implantation, the mice were randomly divided to BHIMHA (oral dose: 0.4, 2.0 and 8.9 µmol/kg/d, 12 consecutive days) treatment groups (36 mice), doxorubicin (Dox, intraperitoneal dose: 2 µmol/kg/d, 12 consecutive days, 12 mice), or NS (oral dose: 10 mL/kg/d, 12 consecutive days, 12 mice). Mice were weighed daily. Twenty four hours after the last dosage, the mice were weighed and killed with ether anesthesia to obtain the organs and the tumor immediately. Data are presented as mean ± SD. The statistical analysis of the data was carried out by use of ANOVA test. The P-value <0.05 was considered statistically significant.

**In vivo xylene-induced ear edema assay**

Male ICR mice were 10 weeks old at the beginning of in vivo xylene-induced ear edema assay.21 The mice were randomly divided into BHIMHA-treated (oral dose: 0.4, 2.0 and 8.9 µmol/kg, 30 mice) groups, aspirin-treated (oral dose: 0.11 and 1.11 mmol/kg, 20 mice) groups, and NS orally treated group (10 mice). Thirty minutes after administration, 0.03 mL of xylene was applied to both anterior and posterior surfaces of the right ear of the mice. The left ear was used as a control. Two hours after xylene application,
the mice were anesthetized with ether and killed for removal of both ears. The circular punches were taken with bores of 7 mm in diameter and weighed. The increase in weight caused by xylene irritant was calculated by subtracting the weight of the untreated left ear punch from the weight of the treated right ear punch. Data are presented as mean ± SD. The statistical analysis of the data was carried out by use of ANOVA test. The P-value < 0.05 was considered statistically significant.

Results and discussion

The yield and chemical physical data of BHIMHAM

At 60°C, the 10-hour Pictet–Spengler condensation of a mixture of 1.66 g (10 mmol) of methyl-5-formyl-2-hydroxybenzoate, 3.22 g (20 mmol) of tryptophol and 0.17 g (1 mmol) of TsOH provided 1.08 g (93%) of BHIMHA as light gray powders.

FT-MS (m/e) 469.17775 [M–H]+ (calculated 469.18417). 

\[ \text{BHIMHA} \] 

\[ \text{H-NMR (800 MHz, DMSO-d_6)} \] 

\[ \delta = 10.53 \text{ (s, 2H), 7.53 (s, 1H), 7.50 (d, J=8 Hz, 2H), 7.30 (d, J=8 Hz, 2H), 7.26 (d, J=8.8 Hz, 2H), 7.03 (t, J=7.2 Hz, 2H), 6.97 (t, J=7.2 Hz, 2H), 6.92 (d, J=8.8 Hz, 1H), 6.07 (s, 1H), 3.53 (m, 2H), 3.49 (m, 2H), 2.85 (t, J=7.2 Hz, 4H).} \]

\[ \text{\textsuperscript{13}C-NMR (200 MHz, DMSO-d_6)} \] 

\[ \delta /\text{ppm} = 169.80, 160.35, 136.19, 136.02, 132.48, 132.76, 129.78, 128.52, 121.18, 118.88, 118.67, 117.63, 113.03, 111.59, 109.31, 61.97, 60.22, 28.37, 21.52, 21.23, 14.56. \]

These data demonstrate that via Pictet–Spengler condensation and hydrolysis, BHIMHA is successfully prepared in 73% yield and no rigorous conditions are required, which should benefit its preparation in large scale.

BHIMA fits the active site of PKCα

It is well documented that the interactions of a peptide ligand with the active site of PKCα involve a number of amino acid residues such as Val420, Asp467, Leu345, Phe350, and Val353. Accordingly, the interactions were disclosed with docking investigation and are shown in Figure 3. The hydrogen, Van der Waals, and hydrophobic interactions between BHIMHA and the residues Val420, Asp467, Leu345, Phe350, and Val353 of the active site are amplified and these are clearly shown in Figure 3A. The data mean that in the active site of PKCα the peptide ligand and BHIMHA have the same action. Similarly, the hydrogen, Van der Waals, and hydrophobic interactions between staurosporine and the residues Val420, Asp467, Leu345, Phe350, and Val353 of the active site are amplified and these are clearly shown in Figure 3B. The data also mean that in the active site of PKCα, the peptide ligand and staurosporine have the same action. Therefore, the docking investigations suggest that BHIMHA is worthy of the experimental assays.
BHIMHA did not affect the proliferation of A549 cells and HaCaT cells
The effect of BHIMHA (1, 10, 25, 50, and 100 µM) on the proliferation of A549 cells was examined with MTT assay. It was found that in the presence of 20 µM BHIMHA, the 48-hour incubation gave A549 cells 88.33%±0.98% of survival, and in the presence of 100 µM BHIMHA the 48-hour incubation gave A549 cells 58.44%±3.58% of survival. Therefore, 5, 10, and 20 µM of BHIMHA were used for in vitro experiments. The effect of BHIMHA (1, 10, 25, 50, and 100 µM) on the proliferation of HaCaT cells was examined with MTT assay. It was found that in the presence of 1, 10, 25, 50, and 100 µM BHIMHA the 48-hour incubations gave HaCaT cells 93.51%±6.35%, 90.24%±7.65%, 89.23%±7.35%, 86.55%±6.78%, and 80.6%±7.42% of survivals, respectively, demonstrating that the IC$_{50}$ value of BHIMHA against HaCaT cell proliferation is higher than 100 µM and has no cytotoxic action on noncarcinoma cells.

BHIMHA inhibited A549 cell migration
To show the effect of BHIMHA on the migration, the transwell migration assay of A549 cells was performed. The stained feature and the counted number of the migrated cells are represented with purple particles and are shown in Figure 4A and C. Figure 4A visualizes that 6 hours after the incubation 5, 10, and 20 µM of BHIMHA concentration dependently inhibits the migration of A549 cells. Figure 4C indicates that the migrating number of A549 cells treated with 5, 10, and 20 µM of BHIMHA gradually decreases.

Figure 4

Notes: (A) Migration A549 cells in transwell chambers treated with BHIMHA for 6 hours; (B) invasion A549 cells in matrigel-coated transwell chambers treated with BHIMHA for 24 hours; (C) concentration dependence of BHIMHA inhibiting migration of A549 cells; and (D) concentration dependence of BHIMHA inhibiting invasion of A549 cells; n=3.

Abbreviations: SD, standard deviation; BHIMHA, S-(bis(3-(2-hydroxyethyl)-1H-indol-2-yl)-methyl)-2-hydroxybenzoic acid.

BHIMHA inhibited A549 cell invasion
To show the effect of BHIMHA on the invasion, the transwell invasion assay of A549 cells was performed. The stained feature and the counted number of the invasion cells are represented with purple particles and are shown in Figure 4B and D. Figure 4B visualizes that 24 hours after incubation 5, 10, and 20 µM of BHIMHA concentration dependently
inhibits the invasion of A549 cells. Figure 4D indicates that the invasion number of A549 cells treated with 5, 10, and 20 µM of BHIMHA gradually decreases.

**BHIMHA inhibited the metastasis of LLC cells from planted tumor toward the lung of C57BL/6 mice**

The effect of 12 days’ treatment of BHIMHA on the metastasis of the planted LLC cells toward the lungs of C57BL/6 mice is represented with the number of metastatic tumor nodules. Figure 5A shows the dose (0.4, 2.0, and 8.9 µmol/kg/d)-dependent effect of BHIMHA on the number of the metastatic lung tumor nodule (labeled with blue ring). Figure 5B shows the number of the metastatic tumor nodules in the lungs of the mice orally receiving BHIMHA, and suggests that the inhibition is also a dose-dependent action.

**Effect of BHIMHA on PKCα expression from A549 cells**

As mentioned earlier that, the regulation of PKCα relates to the proliferation, survival, differentiation, and motility of the cancer cells, and positively regulates the invasion and the metastasis. For instance, the inhibition of PKCα expression leads to the inhibition of the invasion and migration of A549 cells. To clarify the reproducibility of the design, the action mode of BHIMHA inhibiting the migration, invasion, and metastasis of the cancer cells was explained with the expression of PKCα or PKCβII or PKCγ or PKCη or PKCδ or PKCθ from A549 cells by performing Western blot assay. Of the six antibodies, only PKCα gave useful result. Figure 5C and D show that 24 hours after the incubation, 5 µM of BHIMHA effectively decreases the intensity of PKCα band, correlating the downregulation of PKCα expression with the inhibition of the migration and invasion of A549 cells, and additionally with the metastasis of LLC tumor toward the lung. However, the downregulation of PKCα expression may result from quite different cellular mechanism. How BHIMHA inhibits PKCα expression and whether this inhibition is a result of directly affecting PKCα itself or a result of indirectly affecting the upstream kinases of PKCα remains to be known.

**Effect of BHIMHA on LCC tumor growth of C57BL/6 mice**

To clarify the effect of 12-day treatment of BHIMHA on the growth of the primary tumor of C57BL/6 mice planted with LLC cells, the tumor weights were measured. Figure 6 shows the weights of the primary tumors of the C57BL/6 mice receiving NS and BHIMHA (0.4, 2.0, and 8.9 µmol/kg/d), and suggests BHIMHA dose dependently slows the growth

Figure 5 In vivo BHIMHA dose dependently inhibits the metastasis of tumor toward lung.

Notes: (A) Representative photographs of LLC tumor metastasis lungs of the mice treated with NS and BHIMHA. (B) On LLC metastasis model BHIMHA inhibited metastasis of C57BL/6 mice, n=10. (C, D) Western blot assay showing the effect of BHIMHA on the expression of PKCα from A549 cells, n=3.

Abbreviations: NS, normal saline; LLC, Lewis lung carcinoma; PKCα, protein kinase Cα; BHIMHA, 5-(bis-(3-(2-hydroxyethyl)-1H-indol-2-yl)methyl)-2-hydroxybenzoic acid; staurosporine, (5S,6R,7R,9R)-6-methoxy-5-methyl-7-methylamino-6,7,8,9,15,16-hexahydro-5H,14H-17-oxa-4b,9a,15-triaza-5,9-methano-dibenzo[b,h]cyclopenta[e]jadisacen-14-one.
BHI MHA: a promising lead drug to block lung metastasis of LLC planted C57BL/6 mice and its minimal effective dose is 2 µmol/kg/d.

Effect of BHI MHA on tumor growth of S180 mice

The in vivo inhibition of 12-day treatment of BHI MHA to the primary tumor was further examined on S180 mouse model. Figure 7 shows the tumor weights of S180 mice treated with NS (10 mL/kg/d) and BHI MHA (0.4, 2.0 and 8.9 µmol/kg/d), ensures BHI MHA dose dependently slows the tumor growth of S180 mice and shows a minimal effective dose of 2 µmol/kg/d. Thus, either for LLC planted C57BL/6 mice or for S180 cells planted ICR mice 2 µmol/kg/d of oral BHI MHA effectively slows the tumor growth, and emphasizes that in addition to inhibiting the metastasis of tumor toward lung BHI MHA is capable of inhibiting the primary tumor to grow.

Effect of BHI MHA on IC R mice developing inflammation

The anti-inflammation activities of BHI MHA (0.4, 2.0, or 8.9 µmol/kg) were evaluated on xylene-induced ear edema. Figure 8A shows that oral BHI MHA dose dependently reduces xylene-induced ear edema of the mice and has a minimal effective dose of 2 µmol/kg. Besides, the potency of reducing ear edema of 8.9 µmol/kg of BHI MHA equals that of 0.11 mmol/kg of aspirin (P > 0.05), which means the anti-inflammation activity of BHI MHA is 12-fold higher than that of aspirin. In addition to inhibiting the metastasis of the tumor toward lung and inhibiting the growth of the primary tumors, BHI MHA also effectively inhibits the inflammatory response.

Effect of BHI MHA on NF-κB expression of A549 cells

Inflammation is a process of innate immunity in response to physical, physiological, and/or oxidative stress and correlates with the activation of NF-κB signaling pathway. To correlate the inhibition of inflammation with the downregulation of the expression of NF-κB from BHI MHA-treated A549 cells, the Western blot assay was performed, and PDTC was used as a positive control. Figure 8B and C show that 24 hours after incubation, 5 µM BHI MHA effectively decreases the intensity of NF-κB band. Thus, it would be proposed that by blocking NF-κB signaling pathway BHI MHA inhibits the mice to develop xylene-induced inflammation.

Conclusion

The survey of the structural characteristic of staurosporine, BPIC, and MIAM, as well as the docking toward the active sites of PKCs led to a rational design of BHI MHA capable of inhibiting the metastasis of tumor toward lung, slowing the tumor growth, and blocking inflammatory response. In vitro, BHI MHA concentration dependently inhibits the
migration and invasion of A549 cells. In vivo, BHIMHA dose dependently inhibits the metastasis of LLC toward lung, dose dependently inhibits the tumor growth of C57BL/6 mice and S180 mice, and dose dependently inhibits xylene-induced ear edema of ICR mice. These activities support the rationality of BHIMHA design. In vitro, BHIMHA downregulates the expressions of PKCα and NF-κB from the treated A549 cells and provides a clue to disclose the action mechanism.

Acknowledgments

This work was supported by Beijing Municipal Science and Technology Commission (Z141100002114049), PXM 2013014226000002, TJSHG (201310025008), IHLB (KZ201210025021), Beijing Nova Program (grant number XX2013039), National Natural Science Foundation (81172930, 81273379, 81373264, 81373265, and 81270046) and 863 program (2015AA020902), Beijing Natural Science Foundation (7132032).

Disclosure

The authors report no conflicts of interest in this work.

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