Abstract. BC (BC), as the most common malignancy in women worldwide, is associated with high morbidity and mortality. However, chemoresistance and toxicity are the main causes that limit the success of treatment in aggressive BC cases. Thus, there is a vital need to identify and develop novel therapeutic agents. Frankincense, pine needle and geranium essential oils have been reported to play critical biological activities in cancer. However, to the best of our knowledge whether frankincense, pine needle and geranium essential oils have any effect on the progression of BC in MCF-7 cells remains unclear. In the present study, we assessed the possible effects of frankincense, pine needle and geranium essential oils on cell viability, proliferation, migration and invasion as well as the possible mechanisms. MCF-7 cells were treated with oils, and associations with BC were investigated. In the present study, we clearly revealed that frankincense, pine needle and geranium essential oils suppressed cell viability, proliferation, migration and invasion in human BC MCF-7 cells. Further data demonstrated that frankincense, pine needle and geranium essential oils induced apoptosis, but did not affect cell cycle progression. Consistent with the in vitro activities, frankincense essential oil was effective in inhibiting tumor growth and inducing tumor cell apoptosis in a human BC mouse model. In addition, these 3 essential oils modulated the activity of the AMPK/mTOR signaling pathway.

Introduction

BC (BC) is the most common malignant disease among women worldwide, and is associated with high morbidity and mortality (1). Surgery combined with adjuvant therapy was the main treatment strategy in BC patients. In addition, endocrine and targeted therapies were gradually applied to clinical therapy (2,3). However, in the treatment of aggressive BC cases, chemoresistance and toxicity are the leading causes for failure. Thus, it is important to define and develop new therapeutic agents, which may bind to BC cells specifically and effectively.

Frankincense, the gum resin derived from *Boswellia* species, contains active ingredients. The oil has been demonstrated to modulate critical biological activities including anti-rheumatism, anti-inflammatory (4,5), antibacterial, antifungal and anticancer activities (6-9). Frankincense oil is prepared by the steam distillation of frankincense gum resin. Based on the biological function of frankincense, it possibly possesses anticancer characteristics. Pine needle (*Pinus densiflora* Siebold & Zucc.), is usually utilized as a herbal medicine, tea bag infusion and health supplement in East Asian countries, such as Korea and China (10). It is beneficial in the therapy of patients with coronary heart disease (CHD), neurodegenerative disorders and carcinoma. Moreover, it was also reported that extracts from pine needle inhibited apoptosis of the normal cells induced by a hydroxyl radical (11). As a central material, geranium essential oil has been used in the cosmetic, perfume, aromatherapy and food industries. In addition, the oil famous for its antibacterial, antioxidative and anti-inflammatory properties, has been used as a traditional drug for a long time (12-15).

In conclusion, the present study indicated that frankincense, pine needle and geranium essential oils were involved in the progression of BC cells possibly through the AMPK/mTOR pathway.
pathway involving essential oils in BC cell proliferation, invasion and apoptosis development.

Materials and methods

Cell culture. MCF-7 cells were obtained from the Dalian Institute of Chemical Physics, Chinese Academy of Sciences (Dalian, China). Cells were seeded in RPMI-1640 medium (HyClone, Logan, UT, USA) supplemented with 10% fetal bovine serum (FBS; Gibco, Grand Island, NY, USA) and 1% penicillin-streptomycin (Invitrogen, Carlsbad, CA, USA). The cells were maintained in a humidified incubator at 37°C with 5% CO₂.

Materials. Stock solutions of frankincense, pine needle and geranium essential oils were obtained from the Hualin Natural Health Cosmetics Company (Beijing, China). Dimethyl sulfoxide (DMSO) at a ratio of 1:2 (v/v) was used as a vehicle of the oils. Subsequently, the oils were diluted with complete medium to a series of different concentrations. The frankincense injection was prepared by diluting the stock solution of frankincense by mixing it with DMSO in a 1:2 ratio, and then diluting this mixture to 1:1,000 (v/v) with phosphate-buffered saline (PBS). The frankincense smear was prepared by diluting the stock solution of frankincense by 1% (v/v) with grape seed oil.

Western blotting. SDS buffer (60 mM Tris-HCl with a pH value at 6.8, 10% glycerol, 2% SDS, with 5% 2-mercaptoethanol) was utilized to store the lysate of the cells and tissues. The 4-12% sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) gels were used to separate the cell lysate. Proteins were transferred to polyvinylidene fluoride (PVDF) membranes (Invitrogen), and the membranes were blocked with Tris-buffered saline plus 0.1% Tween-20 (TBST) containing 5% skim milk for 2 h. The primary antibodies specific to phospho-ERK, total-ERK, phospho-4E-BP1, phospho-mTOR, total-mTOR, phospho-AMPK, total-AMPK, poly(ADP-ribose) polymerase (PARP) (Asp214) (Cell Signaling Technology, Beverly, MA, USA), Bcl-2 (Abgent, Inc., San Diego, CA, USA) were immunoblotted. All bands were washed in TBS with Tween-20 (TBST) and immunoblotted with peroxidase-conjugated anti-mouse or anti-rabbit secondary antibodies, respectively. The bands were detected using a chemiluminescence (ECL) Western Blotting kit and exposed to film. The β-actin antibody (BIOSS, Beijing, China) was used as an internal marker for control. All experiments were carried out thrice.

Cell cycle analysis. For DNA content analysis, MCF-7 cells were treated with different oils. The cells were gathered, washed and resuspended after 48 h. Ethanol (75%) was used to fix cells overnight. The cells were centrifugated and exposed to RNase (100 µl) for 30 min at 37°C. Propidium iodide (PI) (400 µl) was used to stain DNA for 30 min without light. DNA contents were analyzed using BD Biosciences Accuri C6 flow cytometer (BD Biosciences, Franklin Lakes, NJ, USA) and FlowJo software. DNA histograms in the G0-G1, S and G2-M phases were defined and the ratio in each phase was calculated.

Apoptosis assay. Cell apoptosis assays were carried out with the use of an Annexin V-FITC apoptosis detection kit. Specifically, cells (1x10⁶ cells/well) were treated with or without oils for 48 h, collected, washed and resuspended. Then, the cells were treated with a mixture of Annexin V-FITC and PI for 15 min in the dark at room temperature. Subsequently, binding buffer (400 µl) was added. The cells were then analyzed by the BD Biosciences Accuri C6 flow cytometer.

Cell Counting Kit-8 (CCK-8) assay. To evaluate cell proliferation, cells (1x10⁵ cells/well) were plated in 96-well plates containing 100 µl full medium, and incubated with different oils at various time-points. CCK-8 (10 µl) was added to the plates and incubation followed at 37°C in 5% CO₂ for 4 h. The spectrometric absorbance was determined by microplate spectrophotometer (Multiscan MK3; Thermo Fisher Scientific, Waltham, MA, USA) at 450 nm.

Focus formation assay. Cells (5x10² cells/well) were trypsinized to a single-cell suspension and seeded into 6-well plates. The cultures were plated in the corresponding media with different oils until the appearance of foci from transformed cells was evident (8 days after incubation). Crystal violet (0.2%) was utilized to stain the colonies and images were captured with a Nikon digital camera (Nikon Corporation, Tokyo, Japan).

Wound healing assay. Wound healing assay was performed to evaluate the migratory ability of the tumor cells. Briefly, MCF-7 (1x10⁴) cells were cultured into a 12-well plate with various oil treatments and increased to 70-80% confluence. The sterilized P200 pipette tip passed through the cell monolayer causing a wound. The cells were observed to migrate to the wound at different times. The migrated cells to the injured area were photographed at different time-points under an inverted microscope. In each well, at least 8 regions of each condition were captured randomly at a magnification of x100. The experiment was conducted in triplicate.

Cell invasion assay. A cell invasion assay was performed using Transwell inserts with polycarbonate membranes of 8.0-µm pore size (Corning Inc., Corning, NY, USA) with ECMatrix gel (Chemicon, Temecula, CA, USA) to form a continuous thin layer. In brief, 4x10³ cells (with different oil treatments) and serum-free medium were inoculated into the upper chamber. Culture medium containing 10% FBS was used as a chemical attractant in the lower chamber. Cells were incubated at 37°C in incubators for 24 h. Then, the invaded cells under the membrane were fixed with methanol, stained with Wright-Giemsa, photographed (magnification, x400) and cells were counted in 5 random areas. Each experiment was performed thrice.

In vivo assays for tumor growth. To further determine whether frankincense regulated tumor growth, in vivo tumorigenesis was performed. MCF-7 cells (2x10⁵) were injected into the left and right dorsal flank of 4-5 week-old female nude mice (purchased from the Animal Center Dalian Medical University), respectively. When mice exhibited a palpable tumor (~1 week after tumor cell inoculation), they were randomly divided into 7 groups (n=5 animals/group): the WT
group, treated with nothing; the DMSO and negative group, treated with DMSO; the base oil and negative group, treated with grape seed oil smear; the injection group, treated with frankincense subcutaneous injection (1:1,000 v/v, 0.1 ml); the external smear group, treated with frankincense smear (0.1 ml); the combination group, treated with frankincense subcutaneous injection combined with frankincense smear; the PTX and positive group, treated with PTX subcutaneous injection (10 mg/kg) (16). The treatment was administered every 4 days for 12 days. The tumor dimensions were gauged every 4 days for 12 days (17). The estimated tumor volume (mm$^3$) was calculated using the following formulas: Tumor volume = length x width$^2$ x 0.52. All mice were kept under specific pathogen-free conditions for air filtration. All experiments with animals complied with the standards in the guidelines of the University Animal Care and Use Committee of Dalian Medical University. Finally, the tumors were fixed in 10% formalin and embedded in paraffin for immunohistochemistry.

**Tunel assay.** TUNEL analysis was used to determine the apoptosis of tumor cells induced by essential oils. In short, the paraffin section was cut into 5-µm of thickness, dewaxed and rehydrated. Apoptotic cells were detected using an *in situ* cell death assay kit. After the terminal deoxynucleotidyl transferase reaction, the labeled end of the incision was identified.

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**Figure 1.** Frankincense, pine needle and geranium essential oils suppress the cell viability and proliferation of MCF-7 cells. (A-C) MCF-7 cells were seeded in a 96-well tissue culture. The cells were subjected to serial dilutions of frankincense, pine needle and geranium essential oils treatment (WT, cells treated with nothing; DMSO, cells treated with DMSO; oil, cells treated with oil; PTX, cells treated with paclitaxel). Cell viability was determined at 48 h after essential oil treatment by CCK-8 assay. (D) Morphological changes of MCF-7 cells following frankincense, pine needle and geranium essential oil stimulation. MCF-7 cells were subjected to frankincense, pine needle and geranium essential oil, respectively (1:3,000; 1:3,300; and 1:6,000 v/v). Images were captured at 48 h after treatment. (E) Cells were seeded in each well of a 6-well plate, subsequently exposed to and incubated with frankincense/pine needle/geranium essential oil (1:3,000; 1:3,300; and 1:6,000 v/v). After 8 days of culture, the surviving colonies were counted with crystal violet staining. Data are presented as the mean ± SD from 3 independent experiments (*P<0.05, **P<0.01).
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by alkaline phosphatase-based immunohistochemistry, with fast red as the substrate. Stained slides were washed and sealed with an aqueous mounting medium.

Statistical analysis. Student's t-test was applied to detect statistically significant differences for non-paired replicates. One-way analysis of variance (ANOVA) was used to compare replicate means. P-value $<0.05$ was considered to indicate a statistically significant result ($P<0.05$, $P<0.01$). Error bars represent the mean ± standard deviation (SD) unless specified otherwise.

Results

Frankincense, pine needle and geranium essential oils suppress MCF-7 cell viability and proliferation. To determine whether these three essential oils affected MCF-7 cell viability, the number of viable MCF-7 cells was determined following various dilutions (frankincense, 1:1,000-1:4,000; pine needle, 1:1,000-1:4,000; geranium, 1:1,000-1:8,000) of oil exposure. As shown in Fig. 1A–C, the cell viability was decreased in a dose-dependent manner when MCF-7 cells were treated with oil. In addition, no viable MCF-7 cells remained after 48 h treatment with 1:1,000 dilution of oil. Based on the CCK-8 assay, IC$_{50}$ values (the 50% inhibitory concentration of frankincense/pine needle/geranium essential oil) for MCF-7 cells were 42.8 µg/ml, 90.2 µg/ml and 73.9 µg/ml, respectively (data not shown). Moreover, to determine whether oil suppresses cell viability, MCF-7 cells were subjected to morphological evaluation assessment. As shown in Fig. 1D, MCF-7 cells underwent significant morphological changes, such as detaching from tissue culture plates and shrinking following oil exposure (with IC$_{50}$ dilution treatment for 48 h). Furthermore, the colony formation assay revealed that the colony size and colony

Figure 2. Frankincense, pine needle and geranium essential oils suppress the migration and invasion of MCF-7 cells. (A) The ability of migration was compared in frankincense, pine needle and geranium essential oil treatments and control groups based on wound healing ($P<0.05$). (B) In vitro ECMatrix gel analysis was performed to compare the cell invasion between 3 essential oil-treated cells and control groups ($P<0.05$). Values are presented as the mean ± SD from 3 independent experiments.
forming capacity of MCF-7 cells were strongly reduced in the presence of the oils compared to the controls (Fig. 1E). These results indicated that frankincense, pine needle and geranium essential oils played a role in BC cell proliferation.

Frankincense, pine needle and geranium essential oils regulate aggressiveness of MCF-7 cells. To examine whether these 3 essential oils exerted a direct regulation on the response to BC aggressiveness, MCF-7 cells were treated with different essential oils. Upon frankincense, pine needle and geranium essential oil treatments, the migratory capability was significantly decreased as determined by wound healing assay (Fig. 2A). Concomitantly to the aforementioned experiments, we investigated the potential role of these 3 essential oils in mediating the invasive abilities of MCF-7 cells. The results of the Transwell invasion assay revealed that the invaded cell number/field was much lower than that in the control group (Fig. 2B). Collectively, these results demonstrated that frankincense, pine needle and geranium essential oils regulated the aggressiveness of BC cells.
Frankincense, pine needle and geranium essential oils regulate apoptosis and the cell cycle of MCF-7 cells. To explore the potential role of these 3 essential oils in modulating the apoptosis and cell cycle, MCF-7 cells were treated with oils (frankincense, 1:3,300; pine needle, 1:3,000; geranium essential oil, 1:6,000) for 48 h, respectively. Cell apoptosis and the cell cycle were analyzed by flow cytometry. As shown in Fig. 3A, the number of apoptotic cells was increased in groups treated with the oils, compared with the WT and DMSO control groups. Cleavage of PARP was involved in DNA repair following environmental stress (18). We also detected the expression of cleaved PARP to evaluate the apoptotic activity of MCF-7 cells treated with the oils. Cleaved PARP levels were increased in the oil-treated cells (Fig. 3B). We also determined the expression of the Bcl-2 protein, and found that the oils induced a decrease in the expression of Bcl-2. Furthermore, we analyzed the cell cycle of MCF-7 cells and found that the percentage of cells arrested at the G0/G1, S and G2-M phases were not significantly altered (Fig. 3C). Collectively, these results indicated that frankincense, pine needle and geranium essential oils induced apoptosis but not cell cycle arrest of MCF-7 cells.

Frankincense essential oil modulates tumor growth in a xenograft mouse model. Finally, we examined the tumorigenicity of frankincense essential oil in MCF-7 cells by in vivo experiments. Administration with frankincense subcutaneous injection, frankincense smear, frankincense subcutaneous injection combined with frankincense smear significantly suppressed tumor growth in nude mice implanted with established MCF-7 tumors (Fig. 5A). Moreover, the tumor growth events such as mTOR/p70S6K (19). In addition, mTOR was demonstrated to be a central controller of cell proliferation, growth and survival (20). In the present study, we assessed the activity of the AMPK signaling and its downstream target by treatment of MCF-7 cells with frankincense, pine needle and geranium essential oils. Western blot analysis revealed that the levels of phosphorylated-AMPK were increased in MCF-7 cells treated with these 3 essential oils (Fig. 4A and B). Concomitantly, the degrees of phosphorylation of mTOR and its downstream effector 4E-BP1 were markedly decreased in MCF-7 cells treated with these 3 essential oils. Conversely, there was no change in the total amount of AMPK and mTOR protein, demonstrating a true decrease in phosphorylation status. These results revealed that the antiproliferative, anti-invasive and induced-apoptosis effect of frankincense, pine needle and geranium essential oils on MCF-7 cells involved AMPK-initiated mTOR inhibition.
rate of the combined group was slower than that of the injection group and the smear group. Furthermore, TUNEL assay revealed a significantly higher number of TUNEL-positive cells that were detected in the experimental group, compared with the control group. In addition, the results demonstrated that the number of apoptotic cells in combination group was higher than that in the other experimental control groups (Fig. 5B). Collectively, in vivo studies confirmed that frankincense suppressed tumor growth and induced apoptosis.

Discussion

In the present study, we clearly demonstrated that frankincense, pine needle and geranium essential oils suppressed cell viability, proliferation and invasion in human BC cell line MCF-7. In addition, we determined that the frankincense, pine needle and geranium essential oils induced apoptosis, but did not affect cell cycle progression. The frankincense essential oil was also effective in inhibiting tumor growth and inducing tumor cell apoptosis in human BC mouse model. We demonstrated that frankincense, pine needle and geranium essential oils suppressed cell progression through the AMPK/mTOR pathway.

Essential oils, which are distilled from flowers, leaves, stems, the bark or roots of a specific plant, contained terpenes, aldehydes, esters, alcohols and other chemical molecules. It has been demonstrated that essential oils have an antibacterial and anti-inflammatory effect. Moreover, research has reported that essential oils also have a certain anticancer effect. Wu et al demonstrated that essential oils from Angelicae dahuricae and Inula japonica increased the sensitivity of BC cell line MCF-7/ADR to doxorubicin (21). Essential oils distilled from the leaves and flowers of Callistemon citrinus from the western Himalayas gave rise to the antiproliferative effect on human lung carcinoma cell line A549 and rat glioma C-6 cells via induction of apoptosis (22). Thymoquinone decreased proliferation and accelerated apoptosis in ID8-NGL (mouse ovarian cancer cells) tumors after 10 and 30 day-treatment (23). It has been observed thymoquinone mediated cell cycle arrest and apoptosis in BC and hepatocellular carcinoma (24,25). Boswellic acids (major components of frankincense) were reported to possess antitumor activity due to their cytostatic and pro-apoptotic properties in many human cancer cell lines containing meningioma (26), leukemia (27), hepatocellular carcinoma (28), melanoma, fibrosarcoma (29), colon (30) and prostate cancer (31-33). Moreover, the essential oil of frankincense inhibited proliferation and modulated apoptosis of human cancer cell lines both in vitro and in vivo (17,34,35). According to a study by Jeong et al, apoptosis, oxidative cell damage, induced by exposure to hydroxyl radical was inhibited by the extracts from pine needle (11). In the present study, we demonstrated that frankincense, pine needle and geranium essential oils reduced MCF-7 cell viability in a dose-dependent manner. Moreover, frankincense, pine needle and geranium essential oils strongly reduced colony size and colony forming capacity of MCF-7 cells. The treatment of frankincense, pine needle and geranium essential oils was responsible for the altered migratory and invasive phenotype of MCF-7 cells in vitro. Our results further indicated that the frankincense, pine needle and geranium essential oils induced apoptosis. However, the oils did not affect cell cycle progression. Furthermore, we observed that the essential oil of frankincense inhibited tumor growth and induced apoptosis in vivo. These results clearly revealed that frankincense, pine needle and geranium essential oils could play an important role in many biological functions of BC cells such as proliferation, invasion as well as apoptosis.

Further analysis of molecular mechanisms for cancer cell progression may provide more data concerning novel molecular targets of frankincense, pine needle and geranium essential oil treatments for BC. AMPK is a central cellular energy-sensing system that constructively takes part in the interaction between metabolism and cancer progression by regulation of the mTOR pathway (36). The activation of AMPK directly phosphorylates and activates TSC2 by increasing its GAP activity and inhibiting mTOR signaling (37). The serine/threonine kinase mammalian target of rapamycin (mTOR) functions as a major regulator of cellular growth and survival, and resides in two multiprotein complexes, mTORC1 and mTORC2 (38).
mTORC1 regulates phosphorylation of p70 S6 kinase 1 (S6K1) and eukaryotic initiation factor 4E (eIF4E) binding protein 1 (4EBP1) (20,39). It was reported that aspirin decreased viability and anchorage-independent growth of mutant PIK3CA BC cells through AMP-activated protein kinase (AMPK) activation and mTORC1 inhibition (40). Furthermore, in BC, 17-β-oestradiol (E2) directly activated AMPK through interaction of its α-subunit with estrogen receptors, implying its roles in cell proliferation (41). Knockdown of AMPK inhibited glucose metabolism and proliferation of TNBC cells (42). Based on the aforementioned, we analyzed the correlation of the frankincense, pine needle and geranium essential oil-mediated AMPK and mTOR signaling pathway. We revealed that these 3 essential oils notably regulated the activity of the AMPK/mTOR pathway in human BC cells MCF-7. These results demonstrated that frankincense, pine needle and geranium essential oil-modulated BC cell progression was, at least in part, AMPK/mTOR-dependent.

In conclusion, our results demonstrated that frankincense, pine needle and geranium essential oils have an antitumor effect, which could be mediated by the AMPK/mTOR pathway. The novelty of the present study was that the frankincense, pine needle and geranium essential oils may be a promising treatment for BC. However, the present study still had some shortcomings. To further confirm our present findings, similar experiments using the other BC cell lines should be performed. In addition, it is unclear whether the oils have an effect on BC patients. Thus, further investigation is warranted to analyze the effects and accurate mechanisms of frankincense, pine needle and geranium essential oils in BC.

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