Abstract. Axillary bromhidrosis is sweat excreted by apocrine glands in the armpits, mouth corners and other parts. The clinical manifestation includes excessive sweating and heavy odor, leading to the growth of bacteria and skin disease. The present study investigated the mechanism underlying the effect of paeoniflorin (PF) in the treatment of bromhidrosis. PF was injected into the feet of rats, and the foot skin was dissected for histological analysis. Primary human sweat gland cells (hSGCs) were isolated from patients with bromhidrosis. After 24 h treatment with PF or 3-methyladenine, the production of reactive oxygen species (ROS), autophagy, apoptosis, proliferation and cell cycle distribution were determined. PF induced nuclear pyknosis in rat SGCs. In vitro PF treatment inhibited cell proliferation with a 25% inhibitory concentration of 9.530 µM. Treatment with 9.530 µM PF for 24 h significantly increased apoptosis, ROS production and autophagy in hSGCs. PF promoted LC3B and Beclin 1 expression, but inhibited p62, phosphorylated (p)-PI3K and p-Akt expression. 3-methyladenine treatment reversed PF-induced changes in hSGCs. PF-induced inhibition of hSGC proliferation was associated with ROS production, apoptosis, and autophagy. These findings provide a basis for treating bromhidrosis.

Introduction
Axillary bromhidrosis is a common dermatological condition among the general population, and its overall worldwide prevalence is reported to be 1-3% (1,2). Bromhidrosis has a negative effect on a person’s quality of life (3). The treatments for bromhidrosis include surgery, physical therapy and botulinum toxin A injection (4,5).

Bromhidrosis results from hyperactivity of axillary sweat gland (SG)s, and most treatment methods focus on removing the axillary SG and decreasing axillary sweating (3,6). However, there is no radical cure for bromhidrosis and the development of novel drugs or therapies for managing bromhidrosis remains important. The etiology of bromhidrosis is multifactorial (7) and is associated with infection (8) as well as metabolic and neurological dysfunction (9). However, the exact etiology of bromhidrosis is unclear.

Paeoniflorin (PF), an agent isolated from Paeoniae alba, is the primary active component in Shaobei injection (10). PF has been shown to promote production of intracellular Ca²⁺ in salivary gland cells (11) and exert a neuroprotective effect (12) by inducing autophagy-associated pathways (13). In addition, PF regulates the metabolism of amino acids, cholesterol and fat (14-16) and inhibits proliferation of fibroblast-like synoviocytes (17) and pulmonary artery smooth muscle cells (SMCs) (18). To the best of our knowledge, however, the effects of PF on SG cells (SGCs) and its mechanism in treating bromhidrosis have not been reported.

In clinical practice, patients with axillary bromhidrosis who had been injected with Shaobei under their armpits reported that axillary bromhidrosis was notably decreased. Therefore, the present study aimed to determine the effect of PF on bromhidrosis both in vivo and in vitro. The effect of PF on SG morphology, as well as proliferation, apoptosis and autophagy of SGCs was determined.

Materials and methods
Animals and treatment. The protocols for all animal experiments were approved by the Institutional Animal Care and Use Committee of the First Affiliated Hospital of Guangdong
Pharmaceutical University. Male Sprague-Dawley rats (n=25; age, 3-4 weeks; weight, 210±15 g) were purchased from the Experimental Animal Center of Sun Yat-sen University and housed at 27°C, relative humidity 45%, light/dark cycle of 12 h. Rat chow and water were available ad libitum. PF (2.5 mg/kg body weight) was injected into the upper right claw via foot injection. At 6, 24, 48 and 72 h post-treatment (n=5/group), animals were sacrificed by overdose of pentobarbital sodium (135 mg/kg; R&D Systems, Inc.). The foot skin was dissected and prepared for histological examination. All experiments were performed in accordance with the Chinese regulations on the use and breeding of experimental animals (19).

**Histological examination.** The foot skin samples were cut into pieces, fixed with 4% paraformaldehyde at room temperature for 48 h (cat. no. p1110; Beijing Solarbio Science & Technology Co., Ltd.), then dehydrated, made transparent, and embedded in paraffin. Next, the embedded tissues were cut into serial sections (5 µm thickness) using a microtome. After baking the sections of tissue at 60°C for 2 h, they were immersed in xylene, ethanol (100, 95, 80 and 70%), and pure water, dehydrated with gradient ethanol and mounted with neutral resin as previously reported (20). The photos were captures at the magnifications of x20 and x400 by an Olympus light microscope (cat. no. BX51; Olympus Corporation) and the data were analyzed using Olympus Stream software (Olympus Corporation).

**Cell isolation and culture conditions.** Primary human SGCs (hSGCs) were isolated from the alar skin of five patients with bromhidrosis (three men and two women; age, 25-45 years) recruited between June and September 2020 at The First Affiliated Hospital of Guangdong Pharmaceutical University (Guangzhou, China). Bromhidrosis was diagnosed according to previously reported criteria (21). The protocols for all experiments involving humans were approved by the Institutional Review Board and Ethics Committee of The First Affiliated Hospital of Guangdong Pharmaceutical University Witten (approval no. 202182). Written informed consent was obtained from each of the five patients prior to enrollment in the study. Primary hSGCs were isolated as previously reported (22). The pieces of foot skin were cleaned of subcutaneous fat, washed with precooled PBS, then incubated with dispase (0.8 U/ml; Roche Diagnostic Ltd.) at 37°C for 16-20 h. The dermis was obtained and subsequently digested with collagenase type IV (2.5 mg/ml; Gibco; Thermo Fisher Scientific, Inc.) at 37°C for 1 h. The hSGCs were then carefully collected from the lysate using a Transferpettor under a microscope. Next, the hSGCs were cultured in DMEM-F12 (Gibco; Thermo Fisher Scientific, Inc.) supplemented with FBS, 1% penicillin/streptomycin (Beutoye Institute of Biotechnology), 10X Insulin Transferrin Selenium, 2 mM l-glutamine (Gibco; Thermo Fisher Scientific, Inc.), 2 nmol/ml triiodothyronine (Sigma-Aldrich; Merck KGaA), 10 ng/ml recombinant human epidermal growth factor (Invitrogen; Thermo Fisher Scientific, Inc.) and 0.4 mg/ml hemisuccinate hydrocortisone (Sigma-Aldrich; Merck KGaA) at 37°C and 5% CO₂. Cells at passage 3-4 were used for further experiments.

**Immunofluorescence assay.** Isolated hSGCs were identified using cytokeratin 8 (CK8) and α-smooth muscle actin (SMA) immunofluorescence assays. Primary cells were harvested, resuspended in DMEM-F12 and then placed 2x10⁵ cells into 24-well plates with slides (14 mm in diameter; Costar; Corning, Inc.) at 37°C for 24 h. Next, cells attached to slides were fixed with 4% Paraformaldehyde at room temperature for 30 min, treated with 0.5% Triton X-100 (Beyotime Institute of Biotechnology) at room temperature for 5 min, then incubated with antibodies against CK8 (1:100; cat. no. ab353280; Abcam) and α-SMA (1:200; cat. no. ab23535; Abcam) at 4°C overnight. This was followed by incubation with secondary goat anti-mouse/rabbit IgG antibody labeled with Alexa Fluor 594/488 (both 1:200; cat. no. ab150113; Abcam) at 37°C for 1 h. Cell nuclei were stained with DAPI (10 µM) at 37°C for 10 min. Images of the stained cells were captured using a fluorescence microscope (magnification, x20 and x400; Lionheart LX; BioTek Instruments Inc.).

**hSGC treatment.** Following identification and passaging, hSGCs at passage 3-4 were treated with 0, 2, 4, 8, 16, 32 and 64 µM PF (Sigma-Aldrich; Merck KGaA) at 37°C in a 5% CO₂ atmosphere for 24 h. For inhibition of autophagy, PF and/or 5 mM 3-methyladenine (3-MA; Sigma-Aldrich; Merck KGaA) were added to hSGCs at 37°C for 24 h. Each condition was replicated three times.

**Cell proliferation analysis.** The inhibitory effect of PF on hSGC proliferation was examined using Cell Counting Kit-8 (CCK-8) assay (5 mg/ml; Beyotime Institute of Biotechnology) for 2 h according to the manufacturer’s instructions. The absorbance of each well was read at 450 nm using a microplate reader (BioTek Instruments, Inc.). The 50 and 25 inhibitory concentration (IC) were calculated based on CCK-8 assay results. Untreated hSGCs were used as the control.

**Measurement of intracellular reactive oxygen species (ROS) levels.** Intracellular ROS levels were determined using a CellROX® Green assay kit (Invitrogen; Thermo Fisher Scientific, Inc.) following treatment with PF, hSGCs were incubated with CellROX® probe at 37°C and 5% CO₂ for 30 min, followed by three washes with DMEM (Gibco; Thermo Fisher Scientific, Inc.) without FBS. The fluorescence intensity of each group of cells was measured using a BD FACS Calibur™ flow cytometer (BD Biosciences) and analyzed by FlowJo software v10.0. N-Acetyl-Cysteine (1 mM) and tert-butylhydroperoxide (200 µM; BD Biosciences) were used as negative and positive controls, respectively.

**Cell apoptosis.** Cell apoptosis was determined by flow cytometry and Hoechst 33258 staining. In brief, hSGCs (2x10⁵ cells/ml) were placed into 24-well plates (Costar; Corning, Inc.) and incubated with PF (9.53 µM) for 24 h at 37°C in a 5% CO₂. The cells were then harvested by Trypsin, fixed with 70% ethanol at room temperature for 30 min and stained using Hoechst 33258 DNA intercalating dye (Beyotime Institute of Biotechnology) or Annexin V-FITC/PI fluorescent double staining solutions (Beyotime Institute of Biotechnology).
An Olympus fluorescent microscope (magnification, x400) was used to record images of the Hoechst 33258-stained cells. Cell apoptosis, as indicated by Annexin V/PI fluorescent double staining, was analyzed using a FACS Calibur™ flow cytometer (BD Biosciences) and the data were analyzed using FlowJo 10.07 software (FlowJo LLC).

Cell cycle distribution analysis. The effect of PF on hSGC cell cycle distribution was analyzed using flow cytometry (BD Biosciences). hSGCs were treated with PF (9.530 µM) at 37°C for 24 h, and then the cells were incubated with Trypsin and 4 mL of blocking reagent DMEM-F12 (Gibco; Thermo Fisher Scientific, Inc.) was added at 37°C for 10 sec. The cells washed using PBS and fixed with 70% ethanol at 4°C for 30 min prior to 50 µg/ml PI/RNase staining solution (Sungene Biotech Co., Ltd.) at room temperature for 20 min. A FACS Calibur™ flow cytometer was used for cell cycle analysis and analyzed by FlowJo software v10.0.

Western blot analysis. The total cellular proteins were extracted from PF-treated and control hSGCs using lysis buffer (Beyotime Institute of Biotechnology) and protein determination by BCA Protein Assay kit (Pierce; Thermo Fisher Scientific, Inc.). Following protein quantification, an aliquot of total protein (30 µg) from each extract was separated by 10% SDS-PAGE (Invitrogen; Thermo Fisher Scientific, Inc.) and the protein bands were electro-transferred onto PVDF membranes (MilliporeSigma). Then, 5% skimmed milk (Beyotime Institute of Biotechnology) was used to block the membrane at room temperature for 30 min, followed by primary antibody incubation at 4°C overnight. After washing with TBS- 0.05% Tween-20 (Sigma-Aldrich; Merck KGaA) and polysorbate buffer (Invitrogen; Thermo Fisher Scientific, Inc.), the membrane was incubated with the secondary HRP Goat anti-Rabbit IgG (1: 20000; BOSTER, cat. no. BA1054) at room temperature for 1 h. Next, the membranes were incubated with anti-LC3B (1:1,000, cat. no. ab192890, Abcam), anti-Beclin 1 (1:1,500, cat. no. ab210498, Abcam), anti-P62 (1:1,000, cat. no. ab109012, Abcam), phosphorylated (p)-PI3K (1:1,000, cat. no. ab182651, Abcam), PI3K (1:2,000, cat.no.ab140307,Abcam), Akt (1:500, cat.no.ab8805, Abcam),p-Akt (1:1,500, cat. no.ab38449, Abcam), and GAPDH (1:1,000, cat. no. ab8245, Abcam) primary antibodies at 4°C overnight. GAPDH served as a control protein. The immunostained protein bands were visualized using an enhanced chemiluminescence kit (Beyotime Institute of Biotechnology) according to the manufacturer's protocol. The optical density was analyzed by Image-Pro Plus 6.0 (Easybio Technology Co., Ltd.).

Statistical analysis. All data were analyzed using GraphPad Prism 6.0 software (GraphPad Software, Inc.) and results are expressed as the mean ± standard deviation (n=3). One way analysis of variance followed by post hoc Tukey's test was performed to compare multiple groups. P<0.05 was considered to indicate a statistically significant difference.

Results

PF alters the histology of skin and SGs. Treatment with PF for 24, 48 or 72 h significantly induced nuclear pyknosis in SGs (Fig. 1). At 48 h, SGCs were invisible and small numbers of inflammatory cells were present. In addition, the cytoplasm of glandular epithelial cells from rats treated with PF appeared to be loose compared with cytoplasm in control cells. Treatment with PF for a short period of time (6 h) did not change the histology of skin and SGs.

PF inhibits proliferation of hSGCs. CK8 and α-SMA are specific markers of SGCs (23,24). To identify SGCs, localization was confirmed by immunohistological analysis of these maker proteins. Fluorescent images of cells treated with CK8 and α-SMA showed that hSGCs had been successfully isolated from human alar skin (Fig. 2A). Primary hSGCs were treated with a series of PF concentrations; PF inhibited hSGC proliferation in a dose-dependent manner (Fig. 2B). The IC<sub>50</sub> and IC<sub>25</sub> values for PF were 41.013 and 9.530 µM, respectively. After considering the toxicity of PF to human skin (25), 9.530 µM PF was subsequently selected for use in treating hSGCs.

PF promotes intracellular ROS production and apoptosis. Treatment with 9.530 µM PF for 24 h
significantly increased ROS production (Fig. 3A; 15.62±1.52 vs. 39.44±2.42%) and significantly promoted hSGC apoptosis (Fig. 3B and C). PF treatment for 24 h increased the percentage of apoptotic hSGCs from...
6.35±0.67 to 32.01±2.78%. These results suggested PF exerted a cytotoxic effect on hSGCs.

**PF alters cell cycle distribution.** Flow cytometric analysis of cell cycle distribution showed that PF treatment for 24 h did not affect the cell cycle distribution of hSGCs. The percentage of hSGCs at G0/G1 phase following PF treatment was 59.05±5.91 vs. 56.95±1.54% for control hSGCs (Fig. 4).

**3-MA inhibits PF-induced ROS production and apoptosis in hSGCs** Previous reports have demonstrated that PF promotes autophagy (26,27). Here, 3-MA, an autophagy inhibitor, partially suppressed ROS production and apoptosis in hSGCs that had been treated with PF (Fig. 5A and B); these results were confirmed by Hoechst 33258 staining (Fig. 5C). These findings showed that PF-induced ROS production and apoptosis in hSGCs may be associated with promotion of autophagy.

**Expression of autophagy-associated proteins following PF and 3-MA treatment.** To verify whether autophagy was associated with the molecular mechanism underlying PF-induced cellular changes in hSGCs, expression levels of autophagy-associated proteins were detected. PF alone significantly upregulated the levels of LC3B and Beclin 1 expression, but decreased p62 expression in hSGCs (Fig. 6A and B). By contrast, administration of autophagy inhibitor (3-MA) partially reversed the PF-induced changes in LC3B, Beclin 1 and p62 expression (Fig. 6B). In addition, PF-induced inhibition of p-Pi3K and p-Akt expression was also partially reversed by 3-MA (Fig. 6B). These protein expression profiles indicated the involvement of autophagy in PF-induced hSGC cytotoxicity.

**Discussion**

To the best of our knowledge, the present study is the first to report the inhibitory effect of PF on SGC proliferation. The in vivo results showed that PF inhibited SGC proliferation, while in vitro experiments showed that PF suppressed SGC proliferation and promoted apoptosis, autophagy and ROS production.

Cao et al (13) and Sun et al (28) showed that PF administration attenuates 1-methyl-4-phenylpyridinium-induced production of cytosolic free Ca2+ in PC12 cells and that levels of LC3-II protein are upregulated during this process (13,28). Chen et al (29) reported that pretreatment with PF restored advanced glycation end product -modified bovine serum albumin-induced decreases in cell viability and p62 expression, but enhanced the expression of LC3-II in human umbilical vein endothelial cells (29). These studies suggested that PF promotes autophagy in cells.

Additionally, PF is reported to inhibit proliferation of different types of cell, including fibroblast-like synoviocytes (17), pulmonary artery (18) and vascular SMCs (30) and several types of tumor cell (31) by regulating various pathways,
including the NF-κB pathway. In accordance with these findings, the present study showed that PF promoted autophagy and apoptosis in hSGCs cells by increasing LC3B and Beclin 1 expression and downregulating p62 expression. The PF-induced inhibition of hSGC proliferation was consistent with nuclear pyknosis observed in SGs of rat foot skin in vivo. In addition, PF-induced inhibition of hSGC proliferation was associated with downregulation of the pro-proliferative PI3K/Akt pathway. Accordingly, the present results showed that PF inhibited proliferation of hSGCs, potentially by inducing autophagy.

Zhao et al (32) showed that PF attenuates α-naphthyl isothiocyanate-induced ROS production in rats. In certain types of cell, activation of the PI3K/Akt pathway is ROS-dependent (33-35). However, the overproduction of ROS can damage cellular DNA and decrease cell membrane stability (36,37). The interaction between ROS release and autophagy is complex and multifarious (38). ROS-induced autophagy and its mechanisms are commonly studied in cancer cells (38-40). In salivary gland cells, PF promotes production of intracellular Ca²⁺ (11); this effect can be triggered by ROS production, which trigger short-term autophagy in cells (41). The present study showed that ROS production promoted by PF in hSGCs was associated with apoptosis and autophagy; all these factors were altered by inhibition of autophagy. These findings suggest a complex mechanism by which PF inhibits hSGC proliferation. In summary, PF treatment suppressed SGC proliferation and promoted apoptosis, autophagy and ROS production, suggesting that PF may be useful for managing bromhidrosis. However, the exact mechanism by which PF reduces bromhidrosis was not fully clarified and requires further investigation.

To the best of our knowledge, the present study is the first to demonstrate that PF inhibits hSGC proliferation by promoting autophagy, ROS production and apoptosis. The in vivo anti-proliferative effect of PF on hSGCs was also confirmed. The present data revealed the effects of autophagy, ROS production, cell proliferation and apoptosis in the management of bromhidrosis by PF.

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Availability of data and materials

The datasets generated and/or analyzed during the present study are available from the corresponding author on reasonable request.

Authors' contributions

YX, HH and PL conceived, designed and performed the experiments. HH, PL and HL analyzed and interpreted data. YX and HL wrote and revised the manuscript. YX and HL confirm the authenticity of all the raw data. All authors have read and approved the final version of the manuscript.

Ethics approval and consent to participate

The protocols for all animal experiments were approved by the Institutional Animal Care and Use Committee of the First Affiliated Hospital of Guangdong Pharmaceutical University (approval no. 202182). All patients provided written informed consent. The protocol for the clinical study was approved by the Research Ethics Committee of the First Affiliated Hospital of Guangdong Pharmaceutical University (approval no. gyfykydw039). All procedures were performed in compliance with Ethics Committee regulations and Animal Research: Reporting of In Vivo Experiments guidelines.

Patient consent for publication

Not applicable.

Competing interests

The authors declare have no competing interests.

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