Abstract. The aim of the present study was to investigate the cytotoxic effects of bufalin on SCC-4 human tongue cancer cells. Cell morphological changes and viability were examined using phase contrast microscopy and flow cytometry, respectively. The results indicated that bufalin induced morphological changes and reduced total viable cells. Apoptotic cell death was analyzed by DAPI staining and DNA gel electrophoresis; the results revealed that bufalin induced cell apoptosis. Levels of reactive oxygen species (ROS), Ca\textsuperscript{2+}, nitric oxide (NO) and mitochondrial membrane potential (ΔΨ\textsubscript{m}) were measured by flow cytometry, and bufalin was observed to increase Ca\textsuperscript{2+} and NO production, decrease the ΔΨ\textsubscript{m} and reduce ROS production in SCC-4 cells. In addition, western blotting was performed to detect apoptosis-associated protein expression. The results demonstrated that bufalin reduced the expression of the anti-apoptotic protein B-cell lymphoma 2 (Bcl-2) and increased the expression of the pro-apoptotic protein, Bcl-2-associated X protein. However, bufalin treatment also increased the expression of other apoptosis-associated proteins such as apoptosis-inducing factor and endonuclease G in SCC-4 cells. Based on these findings, bufalin may induce apoptotic cell death via mitochondria-dependent pathways in human tongue cancer SCC-4 cells.

Introduction

Oral cancer is one of the major causes of cancer-associated mortalities in the global human population and the total number of patients with oral cancer who are younger than 40 years old has markedly increased in the last decade (1). In Taiwan, a report by the Department of Health in 2014 (2) demonstrated that there are 36.8 mortalities per 100,000 people associated with oral cancer every year. During the early stages of oral cancer patients are mainly treated with surgery, which is occasionally coupled with radiotherapy; however, as it becomes more advanced a combination of surgery, radiotherapy and chemotherapy is applied, which can have adverse effects on the patients' quality of life (3,4). Currently, there are a number of clinically used chemotherapeutic drugs that have been approved for the treatment of oral cancer, including cisplatin, cetuximab and endothelial growth factor receptor (EGFR) inhibitors (5). However, the development of more effective treatments and drugs with lower toxicities is required in order to improve patient prognosis and reduce the number of oral cancer-associated mortalities.

Bufalin, a small molecular compound, was isolated from chansu, a traditional Chinese medicine (TCM) that was obtained from the skin and parotid glands of the Chusan Island toad (Bufo gargarizans) (6). The extract from chansu has been used as a TCM to treat various types of cancers in the Chinese population (7). A number of studies have revealed that bufalin induced biological activities such as inhibiting cell proliferation, and inducing cell differentiation and apoptosis.
in various cancer cells, including human hepatocellular carcinoma BEL-7402 cells (8), leukemia HL-60 (9) and U937 (10) cells, gastric cancer MGC803 cells (11), prostate cancer DU145 and PC3 cells (12), and glioma cancer cells (13). Bufalin also induced G0/G1 phase arrest and apoptosis in human bladder cancer T24 cells through mitochondria signaling pathways (14). In addition, bufalin inhibited human non-small lung cancer A549 cell proliferation via the vascular EGFR (VEGFR1/VEGFR2/EGFR) tyrosine-protein kinase-protein kinase B/p44/p42/p38-nuclear factor-κB signaling pathways (15). A previous study reported that the inhibition of the Janus kinase-signal transducer and activator of transcription 3 signaling pathway enhances bufalin-induced apoptosis in colon cancer SW620 cells (16).

Previous studies have demonstrated that bufalin induced cell death through cell cycle arrest and inducing apoptosis in many human cancer cell lines, there has yet to be a report investigating bufalin-induced cell death via the induction of apoptosis in human tongue cancer SCC-4 cells (2,17). In addition, the molecular mechanism underlying bufalin-induced cell death in SCC-4 cells is unknown. In the present study, the SCC-4 cell line was used as an in vitro tongue cancer model to investigate the effects of bufalin treatment. The present study reported that bufalin induced cell cycle arrest and induced cell apoptosis in SCC-4 cells via endoplasmic reticulum stress and caspase- and mitochondria-dependent pathways.

Materials and methods

Chemicals and reagents. Bufalin of 99% purity, 4',6-diamidino-2-phenylindole, dilactate (DAPI), dimethyl sulfoxide (DMSO), propidium iodide (PI) and Trypsin-EDTA were obtained from Sigma-Aldrich; Merck KGaA (Darmstadt, Germany). A stock solution of bufalin (10 mM) was prepared in DMSO and further diluted in culture medium. DMSO was used as vehicle control in all experiments. Dulbecco’s modified Eagle’s medium (DMEM/F12 (1:1) medium, fetal bovine serum (FBS), L-glutamine and penicillin-streptomycin were purchased from Gibco; Thermo Fisher Scientific, Inc. (Waltham, MA, USA). Primary antibodies and peroxidase-conjugated secondary antibodies were obtained from Santa Cruz Biotechnology, Inc. (Dallas, TX, USA). Fluoro-3/AM, DiOC₆(3), H₂DCF-DA and DAF-FM were obtained by Invitrogen (Carlsbad, CA, USA).

Cell culture. The SCC-4 human tongue cancer cell line was obtained from the Food Industry Research and Development Institute (Hsinchu, Taiwan) and cultured in DMEM/F12 (1:1) medium supplemented with 10% FBS, 100 µg/ml streptomycin, 100 units/ml penicillin, and 2 mM L-Glutamine at 37°C incubator with 5% CO₂ (18).

Cell morphology examinations, total viability and cell cycle assays. SCC-4 cells (1x10⁶ cells/well) were cultured in 12-well plates with DMEM/F12 (1:1) medium supplemented with 10% FBS, 100 µg/ml streptomycin, 100 units/ml penicillin, and 2 mM L-Glutamine at 37°C incubator with 5% CO₂ (18).

Cell morphology examinations, total viability and cell cycle assays. SCC-4 cells (1x10⁶ cells/well) were cultured in 12-well plates with DMEM/F12 (1:1) medium for 24 h. Cell were pretreated with or without inhibitor [1 µM cyclosporine A, an inhibitor of ΔΨm or 1 mM N-acetyl cysteine (NAC), a general ROS scavenger; both from Sigma-Aldrich; Merck KGaA, Darmstadt, Germany] for 4 h at 37°C, and then incubated with bufalin at a final concentration series of 100, 200, 300, 400 and 500 nM, or 0.5% DMSO only as a vehicle control for 48 h at 37°C. Plated cells were examined and photographed under a contrast phase microscope at x200 magnification to analyze cell morphological changes. Cells were harvested and stained with PI (4 mg/ml) at room temperature, followed immediately by flow cytometry (FACSCalibur™; BD Biosciences, San Jose, CA, USA) to perform total viability assays or cells were analyzed for cell cycle distribution as previously described (19).

DAPI staining for chromatin condensation examination. SCC-4 cells (2x10⁵ cells/well) were cultured in 6-well plates and treated with bufalin (100, 200, 300, 400 and 500 nM) or 0.5% DMSO only as a vehicle control for 24 and 48 h at 37°C. Cells were fixed in 3% methanol in PBS at room temperature for 20 min and were then stained with DAPI solution (2 µg/ml) at 37°C for 30 min. Cells were photographed using a fluorescence microscope as previously described (19). The ratio of nuclei condensation of cells to total cells was calculated; 150 cells/field in at least 3 fields from each well were counted. The analysis software to quantify the level of DNA damage was TriTek CometScore™ Freeware version 1.5 (TriTek Corp., Sumerduck, VA, USA).

DNA fragmentation assay by Comet assay and DNA gel electrophoresis. SCC-4 cells (2x10⁵ cells/well) were cultured in 6-well plates for 24 h and incubated with bufalin (100, 300 and 500 nM), 1.25 µM H₂O₂ or 0.5% DMSO only as a vehicle control for 48 h at 37°C. All samples were collected for the Comet assay as described previously (20). SCC-4 cells (1.5x10⁶ cells/dish) were cultured in 10-cm dishes for 24 h and incubated with bufalin (100, 300 and 500 nM) or 0.5% DMSO only as a vehicle control for 48 h at 37°C, then cells were extracted using the Tissue and Cell Genomic DNA Purification kit (GMbiolab Co., Ltd., Taichung, Taiwan) as described previously (20). A total of 2 µg DNA from each treatment group was loaded onto 0.5% agarose gels (at 100 V for 40 min) in TBE buffer (89 mM Trisboric acid and 2 mM EDTA, pH 8.0) for electrophoresis. Ethidium bromide was used for visualization and gels were photographed by fluorescence microscopy.

Measurements of reactive oxygen species (ROS), intracellular Ca²⁺, mitochondrial membrane potential (ΔΨm) and nitric oxide (NO) production. The levels of ROS, Ca²⁺ and ΔΨm in SCC-4 cells following exposure to bufalin were measured by a flow cytometric assay. Briefly, SCC-4 cells (1x10⁶ cells/well) in 12-well plates were treated with bufalin (300 nM) or 0.5% DMSO only as a vehicle control for various time periods (0-9 h for ROS and NO measurements; 0-48 h for Ca²⁺ and ΔΨm measurements) at 37°C. Cells were centrifuged at 453 x g for 5 min at 25°C and resuspended with 500 µl DCFH-DA (10 µM) for ROS (H₂O₂) measurements, with 500 µlDiOC₆(3) (4 µmol/l) for ΔΨm measurements, as with 500 µl Fluoro-3/AM (2.5 µg/ml) for intracellular Ca²⁺ concentration measurements and with 500 µl DAF-FM (10 µM) to measure NO production. All samples were maintained in the dark for 30 min at 37°C and then analyzed by flow cytometry as described previously (21).

Western blot analysis. Western blotting was performed to detect the levels of apoptotic proteins. SCC-4 cells
(5x10⁶ cells/dish) were cultured in a 10 cm dish for 24 h and were then incubated with 0.5% DMSO only as a vehicle control or 300 nM bufalin for 0, 12, 24, 36 and 48 h at 37˚C. The protein was lysed by PRO-PREP Protein Extraction Solution (iNtRON Biotechnology, Sungnam, Korea), and total protein was measured using a Bio-Rad protein assay kit (Bio-Rad Laboratories, Inc., Hercules, CA, USA) as described previously (21). Proteins (20 µg) were separated by electrophoresis 12% (v/v) SDS-PAGE and then transferred onto polyvinylidene difluoride membranes (EMD Millipore, Billerica, MA, USA). Membranes were blocked for 1 h at room temperature with 5% non-fat milk in PBST (0.1% Tween-20 in 1X PBS), then hybridized with the following primary antibodies at 4˚C overnight. The primary antibodies supplied by Santa Cruz Biotechnology, Inc. (Dallas, TX, USA) were used at a 1:1,000 dilution; endonuclease G (Endo G; sc-365359), apoptosis-inducing factor (AIF; sc-13116), cytochrome (Cyt) c (sc-13560), X-box binding protein (XBP)-1 (sc-7160), DNA damage inducible transcript 3 (GADD153; sc-7351), inositol-requiring enzyme (IRE)-1α (sc-25563), activating transcription factor (ATF)-6α (sc-58323), ATF-6β (sc-30596), glucose-regulated protein, 78 kDa (GRP78; sc-13968), calpain-1 (sc-58323) and tumor necrosis factor (TNF)-related apoptosis-inducing ligand (TRAIL; sc-80393). The primary antibody supplied by EMD Millipore was used at a 1:500 dilution for BH3 interacting-domain death agonist (Bid; AB1730). The primary antibodies supplied by Cell Signaling Technology, Inc. (Danvers, MA, USA) were used at a 1:1,000 dilution for B-cell lymphoma 2 (Bcl-2; 2870), Bcl-2-associated X protein (Bax; 2772), death receptor (DR) 5 (8074), DR4 (42533), PARP (9532), caspase-3 (9669) and caspase-9 (9508). The primary antibody supplied by Sigma-Aldrich (Merek KGaA) was used at a 1:10,000 dilution for β-actin (A5316). This was followed by incubation with the secondary antibody at room temperature (25˚C) for 1 h. The secondary antibodies supplied by GeneTex (Irvine, CA, USA) were used at a 1:10,000 dilution for horseradish peroxidase (HRP)-conjugated goat anti-rabbit IgG (GTX213110), HRP-conjugated donkey anti-goat IgG (GTX26885) and HRP-conjugated rabbit anti-mouse IgG (GTX213112). Protein bands were visualized by chemiluminescence signals, which were enhanced using electrochemiluminescence detection (Amersham ECL™; GE Healthcare, Chicago, IL, USA) (21,22). ImageJ version 1.49o software (National Institutes of Health, Bethesda, MD, USA) was used to quantify changes in protein expression by densitometry analysis and using β-actin as the loading control.

Statistical analysis. All values are presented as the mean ± standard deviation of three independent experiments. Differences between groups were analyzed by one-way analysis of variance and Dunnett or Tukey test for multiple comparisons (SigmaPlot for Windows version 12.0; Systat Software, Inc., San Jose, CA). P<0.05 was considered to indicate a statistically significant difference.

Results

Bufalin induces cell morphological changes, decreases total viability and induces G2/M phase arrest in SCC-4 cells. Following exposure to various concentrations of bufalin for 48 h, SCC-4 cells were examined to determine any cytotoxic effects (Fig. 1). Bufalin induced cell morphological changes (cell shrinkage, cytoplasmic and nuclear condensation or pyknosis) and reduced the total cell viability in a dose-dependent manner; a half-maximal inhibitory concentration (IC₅₀) of 300 nM was observed at 48 h (Fig. 1A and B). In addition, bufalin induced G2/M phase arrest in SCC-4 cells (Fig. 1C). Therefore, the findings of the present study indicated that bufalin may induce cytotoxic effects on SCC-4 cells via cell morphological changes and by decreasing the number of viable SCC-4 cells in vitro.
and 500 nM) for 24 and 48 h, SCC-4 cells were stained with DAPI and photographed using fluorescence microscopy; the results are presented in Fig. 2. Treatment with bufalin at 100-500 nM for 24 and 48 h, induced brighter DAPI fluorescence in SCC-4 cells when compared with control cells (Fig. 2A and B). Brighter fluorescence indicates nicked DNA and chromatin condensation, which were observed in SCC-4 cells in a dose-dependent manner (Fig. 2C). These findings indicate that bufalin may induce early and late apoptosis in SCC-4 cells, characterized by the presence of irregular and fragmented nuclei, and shrunken cells.

Bufalin induces DNA fragmentation and DNA damage in SCC-4 cells. It has been previously revealed that DNA fragmentation (DNA ladder) is one of the hallmarks of apoptotic cell death (23). Following SCC-4 cell treatment with bufalin (0, 100, 300 and 500 nM) for 48 h, total DNA was isolated from each treatment group and run on DNA agarose gel electrophoresis (Fig. 3A). Bufalin induced a typical DNA ladder for all 3 examined doses in SCC-4 cells. In addition, the results of the Comet assay indicated that bufalin-induced DNA damage based on the production of Comet tails (Fig. 3B); these effects were observed in a dose-dependent manner (Fig. 3C).

Bufalin induces the production of ROS and Ca^{2+}, and decreases the levels of ΔΨm and NO in SCC-4 cells. In order to further investigate whether bufalin-induced cell apoptosis is involved in the production of ROS, Ca^{2+} and NO, or the dysfunction of the ΔΨm, SCC-4 cells were treated with 300 nM bufalin for various time periods. All samples were collected and analyzed by a flow cytometric assay (Fig. 4). The results revealed that bufalin decreased the production...
of ROS following 1-9 h of treatment (Fig. 4A) and the ΔΨm following 6-48 h of treatment (Fig. 4B). However, 6-9 h of bufalin treatment increased NO production (Fig. 4C) and increased Ca^{2+} production following 12-48 h of treatment (Fig. 4D). Cells were also pretreated with 1 µM cyclosporine A, an inhibitor of ΔΨm, for 4 h at 37˚C and then co-treated with or without 300 nM bufalin to analyze cell viability. The results demonstrated that pretreatment with cyclosporine A increased cell viability when compared with bufalin only treatment (Fig. 4E). These findings indicate that ROS, NO, Ca^{2+} and ΔΨm may be involved in bufalin-induced apoptotic cell death in SCC-4 cells in vitro.

**Bufalin alters apoptosis-associated protein expression in SCC-4 cells.** The present study also investigated whether bufalin-induced cell apoptosis alters the expression of apoptosis-associated proteins in SCC-4 cells. Cells were treated with bufalin (300 nM) for 0, 12, 24, 36 and 48 h, then proteins were measured and quantitated by western blotting (Fig. 5). The results demonstrated that following 48 h
of incubation, bufalin markedly increased the expression of Endo G, Bcl-2, Bax, poly(ADP-ribose) polymerase, AIF, Cyt c (Fig. 5A), TRAIL, DR5, DR4, caspase-3 and -9; (C) XBP-1, GADD153, IRE-1α, ATF-6β, ATF-6α, GRP78 and Calpain 1. Bufalin treatment for 36 h increased the expression of DR5 and caspase-9 (Fig. 5B); however, following 48 h their expression decreased when compared with 0 h of treatment. In addition, bufalin treatment markedly reduced the expression of Bcl-2, Bid (Fig. 5A), calpain 1 and ATF-6β (Fig. 5C). These results indicated that bufalin may induce apoptosis in SCC-4 cells via mitochondria-dependent signaling pathways.

Discussion

Previous studies have demonstrated that bufalin induces cell death via cell cycle arrest and inducing apoptosis in many human cancer cells (2,24,25); however, currently there is a lack of information available on bufalin-induced apoptosis in human tongue cancer SCC-4 cells in vitro. Thus, the present study investigated the cytotoxic effects of bufalin on human tongue cancer SCC-4 cells. The results indicated that: i) Bufalin induced cell morphological changes and reduced the total number of viable cells (Fig. 1); ii) bufalin induced chromatin condensation (Fig. 2), DNA fragmentation (apoptotic cell death) and DNA damage (Fig. 3); iii) bufalin increased NO and Ca2+ levels; however, treatment decreased the levels of ROS and ΔΨm (Fig. 4); and iv) bufalin increased the expression of pro-apoptotic proteins, including Bax, Endo G and AIF, and decreased the expression anti-apoptotic proteins such as Bcl-2 and calpain 1 (Fig. 5).

The present study indicated that bufalin reduced the total percentage of viable cells in a dose-dependent manner, which is similar to findings reported previously in other cell
lines (26-28); the results of the present study also identified an IC_{50} of 300 nM following 48 h treatment in SCC-4 cells. In addition, bufalin induced G2/M phase arrest in SCC-4 cells, which is also in agreement with previous reports (11,29-31). It has been previously established that anticancer drugs, such as paclitaxel have been used for treating patients with cancer, also known to induce G2/M phase arrest (32). Apoptosis has been demonstrated to be one of the potential mechanisms of anticancer activity and it is also a fundamental process in the development of various cell types (33,34). Therefore, the present study further investigated the induction of apoptosis in SCC-4 cells in vitro. DAPI staining and DNA gel electrophoresis were performed to confirm that bufalin induced cell apoptosis in SCC-4 cells. The results demonstrated that bufalin significantly induced cell apoptosis in SCC-4 cells in dose-dependent manner.

Apoptosis pathways can be divided into two types: The death receptor (extrinsic) pathway and the mitochondrial (intrinsic) pathway (35). The death receptor pathway may function via receptors including the Fas cell surface death receptor, DR5 and DR4, which then activate caspase-8, followed by caspase-9 and -3 in order to induce apoptosis or it may trigger mitochondrial dysfunction to induce the intrinsic cell apoptotic pathway (36,37). The western blotting results revealed that bufalin promoted the expression of TRAIL and DR4 following 48 h, and DR5 following 36 h of treatment, indicating that bufalin may have induced cell apoptosis via the death (extrinsic) receptor pathway. DR4 and DR5 proteins are involved in apoptosis and are the membrane receptors for TRAIL (38). In addition, other membrane receptors for TRAIL are also involved in inducing apoptosis (39). Bufalin was also observed to increase the protein expression of caspase-3 following 48 h, and caspase-9 following 36 h, which supports the involvement of both pathways.

Flow cytometric analysis indicated that bufalin increased the production of Ca^{2+} and NO, and decreased the levels of ΔΨ_{m} and ROS in SCC-4 cells in a time-dependent manner. SCC-4 cells were also pretreated with 1 mM NAC for 4 h at 37°C and then co-treated with or without bufalin. As expected, the general ROS scavenger NAC did not significantly decrease bufalin-induced ROS generation and did not increase the total number of viable cells (data not shown). ROS have an important role in cancer cell death; under starvation or stress conditions, the levels of ROS are increased for the induction of autophagy (40). These findings indicate that bufalin did not induce cell apoptosis via the ROS signaling pathway. Ca^{2+} uptake into the mitochondrial matrix has been demonstrated to be critical for cellular function (41). The present study revealed that bufalin promoted Ca^{2+} release. In addition, bufalin significantly reduced the ΔΨ_{m}, and pretreatment with cyclosporine A, followed by bufalin treatment increased the total number of viable SCC-4 cells when compared with bufalin treatment only. Anticancer drugs may induce cell apoptosis through the dysfunction of mitochondria or by decreasing ΔΨ_{m} (42,43), and mitochondria are involved in the stimulation of apoptosis in the intrinsic signaling pathway (44). Thus, the results of the present study suggest that bufalin may induce cell apoptosis via a mitochondria-dependent signaling pathway in SCC-4 cells. Western blot analysis revealed that bufalin increased the expression of the pro-apoptotic protein Bax and reduced the expression of the anti-apoptotic protein Bcl-2. Previous studies determined that the ratio of Bax/Bcl-2 affects the ΔΨ_{m}, and in turn the initiation of apoptosis (45,46). Mitochondrial control of apoptosis is thought to primarily involve the ΔΨ_{m} and membrane permeability (47). Thus, if the levels of ΔΨ_{m} are reduced, cytochrome c, AIF or Endo G may release from the mitochondria, triggering apoptosis.

In conclusion, to the best of our knowledge, the present study demonstrated for the first time that bufalin treatment inhibits the progression of SCC-4 human tongue cancer cells in vitro. Bufalin induced cell morphological changes, reduced the number of viable cells, and induced G2/M phase arrest and apoptosis. Bufalin may have triggered cell apoptosis via a mitochondria-dependent signaling pathway in SCC-4 cells, which is summarized in Fig. 6. Apoptosis may also have been induced via the extrinsic pathway, which involves DR4, DR5, TRAIL and various caspases, or via intrinsic pathways that lead to the release of AIF and Endo G in order to induce apoptosis.

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