AKT serine/threonine protein kinase modulates bufalin-triggered intrinsic pathway of apoptosis in CAL 27 human oral cancer cells

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Abstract. Bufalin has been reported to induce apoptosis in a variety of cancers but little is demonstrated in oral squamous cell carcinoma (OSCC) cells. The present study investigated the inhibition of proliferation, cell cycle arrest and apoptotic effects of bufalin in CAL 27 human oral cancer cells. Bufalin inhibited the growth of CAL 27 cells in a concentration-dependent manner and an IC₅₀ value of bufalin was about 125 nM for 24 h treatment using the MTT assay. Moreover, the cell cycle distribution was arrested at the G₀/G₁ phase in CAL 27 cells after bufalin exposure. Upon bufalin stimulation, the expression of Bcl-2 was significantly decreased while that of cytochrome c, Apaf-1 and AIF was increased compared to the control group by western blot analysis. An increase in the expression of the active form of caspases was found in bufalin-treated cells, and the caspase activities were also elevated. Bufalin-triggered apoptosis was blocked by specific inhibitors of caspase-9 (z-LEHD-fmk) and caspase-3 (z-DEVDF-fmk), respectively. In contrast, CAL 27 cells overexpressing constitutively active AKT (CAL 27/CA-AKT) were exposed to bufalin at different concentrations, and cell growth remained unchanged. Bufalin exhibited minimal apoptotic effects on CAL 27/CA-AKT cells. Taken together, bufalin induced G₀/G₁ phase arrest and provoked the intrinsic apoptotic pathway via AKT activation in CAL 27 cells. Our data suggest that bufalin could be potentially efficacious in the treatment of oral cancer in the future.

Introduction

Oral squamous cell carcinoma (OSCC) is the most common head and neck cancer which exhibits frequent lymph node metastasis and local invasion, causing poor prognosis (1,2). The addiction to betel, tobacco, and alcohol is found to be highly correlated with the risk of HNSCC (3) and the studies in this area might lead to new approaches in the prevention and treatment of this important group of human cancers (4).

Apoptosis involves a cascade of molecular changes such as morphology changes, chromatin condensation, and DNA fragmentation (5,6). Abnormal regulation of apoptosis leads to many human disorders including autoimmune disease and cancer. Thus, understanding the mechanisms of apoptosis is an important strategy for treatment of cancer (6,7). Several gene products have been demonstrated to be critical in the regulation of apoptosis (8,9). For example, caspases are synthesized as proenzymes and become activated by cleavage. Caspase activation is often regulated by various cellular factors, including members of the Bcl-2. The Bcl-2 protein family is divided into two functional subfamilies: pro-apoptotic subfamilies (Bax and Bid) and anti-apoptotic proteins (Bcl-2 and Bcl-xL) (8,10). The family members translocate to the mitochondria and mediate the membrane potential to induce cytochrome c release. Cytosolic cytochrome c is further involved in caspase activation. The caspase cascade is a key pathway in apoptotic signal transduction, and can be divided into two types of subfamilies: upstream initiator caspases (caspase-8 and -9) and downstream effector caspases (caspase-3, -6 and -7), which directly induce the final events of apoptosis (6,11,12).

Bufalin, a cardioactive C-24 steroid, is the major component of the traditional Chinese medicine Chan-Su obtained from the skin and parotid venom glands of the toad (13-15). A previous study has shown that bufalin is applied for a treatment of heart failure and used for a variety of biological activities, such as blood pressure stimulation and antineoplastic activities (14). In addition, bufalin processed biological functions as inhibitors of...
Na+/K+-ATPase and topoisomerase II, leading to protein-linked DNA double-strand breaks (14,16). It has been reported that the topoisomerase inhibitor, etoposide and adriamycin are widely prescribed anticancer drugs which inhibit cell proliferation and induce apoptosis in numerous cancer cell lines (17,18). Our earlier study showed that bufalin is found to inhibit cell growth at G1/G0 phase of the cell cycle and to induce apoptosis in a dose-dependent manner, which was associated with both mitochondrial-regulated and death receptor-initiated pathways (19). Furthermore, bufalin has been found to inhibit Bel-2 and c-myc in human leukemia cells (20) and to induce apoptosis of human prostate cancer cells in part with Fas stimulation, cytochrome c release and caspase activation (15). Additionally, bufalin has also been found to induce apoptosis by upregulating the expression of Bax (21) and suppressing orthotopic transplantation tumor in nude mice in human hepatocellular carcinoma cells (22). Therefore, bufalin is thought to be a valuable anticancer drug.

There is limited information, however, on the cellular and molecular mechanisms underlying the bufalin-induced apoptosis in human oral cancer cells. The purpose of this study was to define the biological and therapeutic effects of bufalin-treated human oral cancer cells for the first time. This study was designed to: i) evaluate the cytotoxicity effects of bufalin; ii) characterize the effect of bufalin on the cell cycle; iii) investigate the apoptotic effects of bufalin by analyzing the protein expression in CAL 27 human oral cancer cells in vitro.

Materials and methods

Chemicals and reagent. Dulbecco’s modified Eagle’s medium (DMEM), L-glutamine, fetal bovine serum (FBS), penicillin/streptomycin and Trypsin-EDTA were purchased from Gibco/Life Technologies (Carlsbad, CA, USA). 4,6-diamidino-2-phenylindole dihydrochloride (DAPI), dimethyl sulfoxide (DMSO), propidium iodide (PI), bufalin, Triton X-100 and anti-β-actin antibody were obtained from Sigma-Aldrich Corp. (St. Louis, MO, USA). Caspase-3 inhibitor (z-DEVD-fmk, Cat. 264155), caspase-9 inhibitor (z-LEHD-fmk, Cat. 218761), anti-p-AKT (Ser473) (Cat. 07-310), anti-AKT (Cat. 05-591), caspase-9 inhibitor (z-LEHD-fmk, Cat. 218761), anti-p-AKT (Ser473) (Cat. 07-310), anti-AKT (Cat. 05-591) and Immobilon Western Chemiluminescent HRP substrate (Cat. WBKLS0500) were bought from Merck Millipore (Billerica, MA, USA). Tdt-mediated deoxyuridine triphosphate nick end labeling (TUNEL) assay kit (in situ Cell Death Detection Kit, Fluorescein) was purchased from Roche Diagnostics (Boehringer Mannheim, Mannheim, Germany). These primary antibodies (anti-cyclin D1, anti-p-BAD, anti-BAD, anti-Bcl-2, anti- cytochrome c, anti-APaf-1 and anti-AIF) and horseradish peroxidase (HRP) conjugated secondary antibodies for western blot analysis were obtained from Santa Cruz Biotechnology Inc. (Santa Cruz, CA, USA). The primary antibodies (anti-caspase-9 and anti-caspase-3) were obtained from Cell Signaling Technology (Danvers, MA, USA).

Cell culture and bufalin treatment. CAL 27 (CRL-2095) cell line was purchased from the American Type Culture Collection (Manassas, VA, USA). Cells were grown in DMEM supplemented with 10% FBS, 100 U/ml penicillin and 100 µg/ml streptomycin at 37°C in a humidified atmosphere of 5% CO₂ atmosphere incubator. Cells were treated with bufalin for indicated concentrations as in each experiment. Equivalent volume of 0.1% DMSO was used as vehicle control.

DNA construct and transfection. Constitutively active AKT was a gift from Dr Way (Department of Biological Science and Technology, China Medical University) and subcloned into pcDNA3 (Invitrogen/Life Technologies). CAL 27 cells were transfected with either an empty vector (pcDNA3), or a constitutively active AKT construct (pcDNA3-CA-AKT) using Arresti-In transfection reagent and performed according to the manufacturer’s instructions (GenDiscovery Biotechnology, Taipei, Taiwan). Cells were selected in neomycin 48 post-transfection (23,24). Viable stably transfected cells expressing constitutively active AKT were analyzed by western blotting as described elsewhere (5,25).

Cell viability assay. The effects of bufalin on cell viability were determined using the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT, Sigma-Aldrich Corp.) assay. Briefly, 1x10⁴ cells per well were seeded in 96-well culture plates. After overnight incubation, the cells were treated with different concentrations of bufalin (0, 50, 100, 150 or 200 nM) for 24 h. The cells were treated with 50 µl of 5 mg/ml MTT for 4 h at 37°C and the resulting formazan crystals were dissolved in dimethyl sulfoxide (DMSO). The absorbance was measured by microplate spectrophotometer (Bio-Tek Instruments Inc., Winooski, VT, USA) at 570 nm. Results were expressed as percentage of the controls, which were arbitrarily assigned 100% viability. Viability assays were performed in triplicate from three independent experiments. The 50% inhibitory concentration (IC₅₀) of bufalin was calculated as described previously (26,27).

Determination for cell morphology. Cells (2x10⁵ cells/well) were maintained in 24-well plates and then were treated with 0, 50, 100 and 200 nM of bufalin. Cell morphological examination was determined utilizing a phase-contrast microscope as previously described (19). Chromatin condensation was detected using the DAPI staining method as previously described (19,28). CAL 27 cells were incubated with 100 nM bufalin for 0, 12, 24 and 48 h. After that, cells were fixed gently by putting 70% ethanol, stained with DAPI, and then photographed using a fluorescence microscope.

Analysis for cell cycle progression by flow cytometry. Cells (2x10⁵ cells/well) in 24-well plates were exposed to different concentrations of bufalin for 24 h. For determination of cell cycle phase and apoptosis, cells were then collected, fixed in 70% ice-cold ethanol overnight, washed in PBS once, and resuspended in PBS containing 40 µg/ml PI, 0.1 mg/ml RNase A and 0.1% Triton X-100 in dark room for 30 min. Cell cycle distribution and apoptotic nuclei were determined by flow cytometry (BD Biosciences, FACSCalibur flow cytometer, San Jose, CA, USA). The percentages of cells in G₀, S, and G₂/M phases were analyzed using CellQuest Pro Software (BD Biosciences) as described previously (29,30).

Assessment of apoptotic cells by TUNEL staining. TUNEL staining was performed according to the manufacturer’s instructions (Roche Diagnostics). Cells (2x10⁵ cells/ml) in 24-well plates were treated without or with 125 nM bufalin for 24 h.
Cells were harvested and immediately incubated with terminal deoxynucleotidyl transferase (TdT) enzyme at 37°C for 1 h. Following TUNEL staining, all samples were washed once with PBS and resuspended in 0.5 ml of PBS containing PI (10 µg/ml) and DNase free-RNase A (200 µg/ml). TUNEL positive cells were analyzed by flow cytometry. The fluorescence intensity was quantified by BD Pro CellQuest software. TUNEL assays were performed in triplicate from three independent experiments as described previously (19,27).

Assay for caspase-3 activity. Caspase-3 colorimetric assay kit (R&D Systems, Inc., Minneapolis, MN, USA) was aped according to the manufacturer’s recommendations. In brief, cells (1x10⁷ cells/flask) in T75 flasks were incubated with 125 nM bufalin for 24 h. Cells were harvested and lysed in a cold Lysis buffer (provided in the kit). Cell lysates (50 µg protein) were incubated with caspase-3 specific substrate (Ac-IETD-pNA) for 1 h at 37°C. The caspase activity was determined by measuring cleavage of chromogenic caspase substrates (pNA) at OD₄₀₅ in a microplate spectrophotometer (BioTek Instruments Inc.) (27,31).

Western blot analysis. Cells (1x10⁷ cells/flask) were seeded in T75 flasks and incubated with or without 125 nM bufalin for 24 h. Total proteins were prepared and determined as previously described (25,28). Briefly, the protein concentration was measured using a BCA assay kit (Pierce Chemical, Rockford, IL, USA). Equal amounts (40 µg) of proteins were boiled for 5 min, separated by 12% SDS-PAGE, and then electro-transferred to Immobilon-P transfer membrane PVDF. The transferred membranes were blocked for 1 h with Tris-buffered saline/Tween-20 containing 5% non-fat dry milk and incubated with primary antibodies at 4°C overnight. Membranes were washed three times with Tris-buffered saline/Tween-20 for 10 min and incubated with secondary HRP-conjugated antibody (5,25). Signals of the blots were detected using an enhanced cheniluminescence (ECL) kit and developed in Kodak Bio-MAX MR film (Eastman Kodak, Rochester, NY, USA). The band density was quantified using National Institute of Health (NIH) ImageJ 1.45 program (28). Blots were probed with β-actin antibody used as the loading control.

Statistical analysis. All the statistical results were expressed as the mean ± SD of triplicate samples. Statistical analyses of data were done using one-way ANOVA followed by Student’s t-test, and p<0.05 was considered significant.

Results

Effects of bufalin on the proliferation and viability of CAL 27 cells. Firstly, we investigated the effects of bufalin treatment on the growth of CAL 27 cells. As shown in Fig. 1, bufalin significantly reduced the cell viabilities of CAL 27 cells after 24 h exposure in a concentration-dependent manner. IC₅₀ values were calculated to be about 125 nM for 24 h. Thus, 125 nM was applied for all subsequent experiments. To determine whether the observed decrease in cell viability is associated with apoptosis, we examined the nuclear morphology under a phase-contrast microscope. In the control cells no significant changes were seen in cell nuclei or cell membrane integrity, whereas CAL 27 cells treated with 125 nM of bufalin for 24 h showed various extent of cell shrinkage, volume reduction, apoptotic body formation and cell blebbing (Fig. 1B). These results suggest that bufalin exhibited a significant apoptosis-inducing effect on CAL 27 cells.

Assay for caspase-3 activity. Caspase-3 colorimetric assay kit (R&D Systems, Inc., Minneapolis, MN, USA) was applied according to the manufacturer’s recommendations. In brief, cells (1x10⁷ cells/flask) in T75 flasks were incubated with 125 nM bufalin for 24 h. Cells were harvested and lysed in a cold Lysis buffer (provided in the kit). Cell lysates (50 µg protein) were incubated with caspase-3 specific substrate (Ac-IETD-pNA) for 1 h at 37°C. The caspase activity was determined by measuring cleavage of chromogenic caspase substrates (pNA) at OD₄₀₅ in a microplate spectrophotometer (BioTek Instruments Inc.) (27,31).

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Cell cycle analysis of CAL 27 cells after exposure to bufalin. To elucidate whether growth inhibition by bufalin is associated with apoptosis, we determined apoptotic features by measurement of the amount of cells in the sub-G₀/G₁ phase in CAL 27 cells in vitro. The results from flow cytometric assay using PI staining revealed that treatment with bufalin resulted in increased accumulation of G₀/G₁ phase in CAL 27 cells and this effect is dose-dependent (Fig. 2A and B). Also, we found that bufalin increased sub-G₀/G₁ population (apoptosis) in CAL 27 cells (Fig. 2A). These results suggest that bufalin inhibited proliferation of CAL 27 cells through G₀/G₁ phase arrest and induction of apoptotic cell death.
Apoptotic feathers in bufalin-treated CAL 27 cells. Cells were treated with 125 nM bufalin for 24 h, then stained with DAPI and analyzed under a fluorescence microscope (Fig. 3A). In addition, the average percentage of apoptotic cells (TUNEL positive cells) increased from 5% of the control to 40% by DAPI/TUNEL double staining (Fig. 3B). In the present study, we investigated the possible mechanisms of bufalin-induced apoptosis in CAL 27 cells. Since caspasas are known to play a pivotal role in mediating various apoptotic signaling (8,10), we measured the activity of effector caspase (caspase-3) in bufalin-treated cells. Fig. 3C shows that exposure of CAL 27 cells to 125 nM of bufalin led to increased levels of activated caspase-3. Taken together, we concluded that 125 nM bufalin decreased the percentage of viable CAL 27 cells through apoptotic cell death. Moreover, caspasas are central regulators of the apoptotic pathway.

Bufalin inhibits the growth of CAL 27 cells via the AKT signaling pathway. We investigated the effects of bufalin treat-
ment on the growth of CAL 27/pcDNA or CAL 27/CA-AKT cells. Results in Fig. 4A indicated that CAL 27/pcDNA cells were inhibited in growth and viability was reduced by bufalin in a concentration-dependent manner. To further determine if the observed decrease in cell viability is associated with apoptosis, we investigated the nuclear morphological changes CAL 27/pcDNA cells. Fig. 4C demonstrates that the control cells were not significantly changed in cell nuclei and cell membrane integrity, whereas bufalin-treated CAL 27/pcDNA cells showed various extent of chromatin condensation, nuclear fragmentation and destruction of cell membrane integrity after a 24-h incubation particularly with 125 nM bufalin. Typical apoptotic nuclei were observed as early as 24 h in CAL 27/pcDNA cells after treatment with bufalin. Characteristically morphological changes of apoptosis were also observed under a microscope, including cell shrinkage, volume reduction, chromatin condensation, cell blebbing and formation of membrane embedded apoptotic bodies (Fig. 4C). Strikingly, bufalin had minimal apoptotic effects (alteration of cell viability) on CAL 27/CA-AKT cells (Fig. 4B and C).

Bufalin causes cell cycle arrest at G₀/G₁ phase in CAL 27/pcDNA but not in CAL 27/CA-AKT cells. To elucidate whether growth inhibition by bufalin is associated with apoptosis via the AKT signaling pathway, we explored the amount of cells in the G₀/G₁ phase in CAL 27/pcDNA and CAL 27/CA-AKT cells. The results from flow cytometric assay using PI staining revealed that treatment with 50-150 nM bufalin resulted in increased accumulation of G₀/G₁ phase in CAL 27/pcDNA cells and this effect is dose-dependent (Fig. 5A and B). We also observed the sub-G₁ population in CAL 27/pcDNA cells (Fig. 5A). In contrast, bufalin did not significantly affect the cell cycle arrest and had minimal apoptotic effects on CAL 27/CA-AKT (Fig. 5A and B). Based on these observations, we propose that bufalin-induced G₀/G₁ phase arrest and apoptotic death in CAL 27 cells is carried out through the AKT signaling pathway.

Bufalin triggers apoptosis in CAL 27/pcDNA but not in CAL 27/CA-AKT cells. To confirm whether AKT expression-modulated bufalin-induced apoptosis in CAL 27 cells, CAL 27/pcDNA and CAL 27/CA-AKT cells were used to investigate DAPI staining, TUNEL assay and caspase-3 activity. Cells were treated with 125 nM bufalin for 24 h and then stained with DAPI. Data in Fig. 6A show that bufalin stimulated chromatin condensation in CAL 27/pcDNA but no effect was found in CAL 27/pcDNA cells. In addition, the average percentage of apoptotic cells (TUNEL positive cells) increased from 5% of the control to 40% (Fig. 6B). In the present study, we investigated the possible mechanisms of bufalin-induced apoptosis in CAL 27 cells and found that AKT signaling might be involved in this event. Caspase-3 is known to play a pivotal role in mediating various apoptotic signaling (8,10), and then we measured the activity of effector caspase-3 in bufalin-treated cells. As illustrated in Fig. 6C, exposure of CAL 27 cells to 125 nM bufalin led to increased caspase-3 activity, but no significant effect occurred in CAL 27/CA-AKT cells after bufalin incubation.

Effects of caspase-9 and caspase-3 inhibitors on bufalin-induced apoptosis in CAL 27, CAL 27/pcDNA and CAL 27/CA-AKT cells. To further investigate the involvement of intrinsic caspase signals in bufalin-induced apoptosis, both of z-DEVD-fmk (a caspase-9 inhibitor) and z-LEHD-fmk (a
Figure 5. Bufalin induces concentration-dependent cell cycle arrest in CAL 27/pcDNA or CAL 27/CA-AKT cells. Cells were treated with bufalin at different concentrations (0, 50, 100 and 150 nM) for 24 h. (A) Respective profiles from flow cytometry and (B) the results are mean ± SD and representative of three independent experiments. *p<0.05 versus the control group (0 nM).

Figure 6. Bufalin-induced caspase-3-dependent apoptosis is mediated by AKT signaling in CAL 27/pcDNA or CAL 27/CA-AKT cells. The both types of cells were exposed to 125 nM bufalin for 24 h. (A) cells were stained with DAPI and analyzed under a fluorescence microscope followed by (B) TUNEL staining and flow cytometric analysis. The TUNEL positive cells were regarded as apoptotic. (C) Effect of bufalin on caspase-3 activity of CAL 27/pcDNA or CAL 27/CA-AKT cells. The results are mean ± SD and representative of three independent experiments. *p<0.05 versus the control group.
caspase-9 inhibitor) blocked intracellular apoptotic proteases and attenuated bufalin-reduced viability and caused cell death in CAL 27 and CAL 27/pcDNA cells (Fig. 7A and B). Importantly, cells overexpressing CA-AKT had minimal effect on bufalin-induced cell death as can be seen in Fig. 7A and B. These results suggest that bufalin-induced apoptosis in CAL 27 cells was associated with the mitochondria-mediated caspase-9 and caspase-3-dependent pathway.

 Bufalin activates the mitochondrial apoptotic pathway involving the regulation of Bcl-2 family members through AKT signaling in CAL 27 cells. To confirm if AKT signal was overexpressed in CAL 27 cells, our data (Fig. 8A) showed that bufalin down-regulated level of AKT only when phosphorylated at threonine 308 but no alteration was observed in AKT expression in treated cells. Phosphorylated serine/threonine protein kinase AKT has been reported to modu-
late Bad phosphorylation at Ser136 (p-Bad) and cyclin DI (32,33). We found that the decreased expression of cyclin DI and p-Bad occurred in CAL 27 and pcDNA/CAL 27 cells after exposure to bufalin. However, no dramatic effect on the levels of cyclin DI and p-Bad was observed in caAKT/CAL 27 cells (Fig. 8A). Our results demonstrated the downstream of AKT signaling (cyclin DI and p-BAD) was decreased in bufalin-treated CAL 27 cells. Upon apoptotic signals, pro-apoptotic Bcl-2 members, such as BAD, was activated; in contrast, Bcl-2 can prevent this occurrence. The imbalance of expression of pro- and anti-apoptotic proteins is associated with the ultimate fate of cells (33,34). To assess whether mitochondrial pathway is involved in bufalin-induced apoptosis, we evaluated the expression levels of BAD and Bcl-2 by western blot analysis. As shown in Fig. 8B, BAD protein levels increased, whereas Bcl-2 protein levels decreased after exposure of CAL 27 cells to 125 nM bufalin for 24 h. Thus, bufalin treatment increased the ratio of BAD/Bcl-2, which is in favor of the occurrence of apoptosis and leads to the release of cytochrome c from mitochondria. In contrast, CAL 27 cells overexpressing constitutively active AKT (CA-AKT) had minimal effect on bufalin-induced cell death. Once released, cytochrome c combines with apoptotic protease activating factor-1 (Apaf-1) and procaspase-9 to form the apoptosome in the presence of ATP, resulting in the activation of caspase-9 and caspase-3 (8,10). Next, we detected the expression of cytochrome c, Apaf-1 and AIF in bufalin-treated cells. The expression of cytochrome c, Apaf-1 and AIF significantly increased after 24 h bufalin treatment (Fig. 8B). These results suggest that bufalin treatment-induced mitochondria-dependent apoptosis is mainly through AKT-regulated signaling (Fig. 9).

Bufalin, a digitalis-like molecule from an animal source, functions as a Na+-K+-ATPase inhibitor and causes an increase in intracellular calcium in cancer cells (13-16). Just like several antitumor drugs including etoposide, adriamycin, and genistein, bufalin is also known as an inhibitor of topoisomerase II (17,18). In addition, bufalin has been shown to induce apoptosis in a variety of human tumors, including colorectal carcinoma, melanoma, hepatoma, breast carcinoma, and gastric carcinoma (19-22). However, the molecular mechanisms responsible for the pro-apoptotic effects of bufalin in human oral cancer cells remain elusive.

Apoptotic characteristics include the appearance of the sub-G1 population and chromatin condensation and elevated pro-apoptotic protein expression levels, the cytochrome c release and caspase cascade activation (6,10,12). In the current study, we provided evidence that bufalin induced apoptosis in CAL 27 cells in a dose-dependent manner through cell cycle arrest at G0/G1 phase (Fig. 2), enhanced chromatin condensation by DAPI staining (Fig. 3A), up-regulation of pro-apoptotic proteins (Fig. 8A), the cytochrome c (Figure 8B), and intrinsic caspase activation (Fig. 3C). Significant apoptotic death was observed using TUNEL assay, by bufalin in CAL 27 cells (Fig. 3B). Our results demonstrated that bufalin also serves as an apoptotic inducer in vitro.

Caspases participate in the execution of apoptosis (8,11). There are two major caspase-dependent pathways. One is the death receptor induced caspase activation pathway, which results in caspase-8 activation. The other is the mitochondrial apoptotic pathway, dependent on the release of cytochrome c from mitochondria to the cytosol. Released cytochrome c binds with Apaf-1 and activates caspase-9, which then activates caspase-3 (6,12). Our study revealed that bufalin failed to activate the death receptor-mediated caspase-8 pathway in CAL 27 cells (data not shown). In contrast, we observed that the process of bufalin-induced apoptosis is involved the activation of caspase-9 and -3, and that the treatment with specific inhibitors of caspase-9 and -3 significantly prevented the bufalin-induced cell apoptotic effects (Figs. 7 and 8B). Thus, our results demonstrated that the bufalin-induced apoptosis is carried out through the mitochondrial-dependent response. Additionally, cytochrome c-mediated apoptosis is controlled prominently by the members of Bcl-2 family. BAD and Bcl-2 have been identified as major regulators. BAD possesses proapoptotic ability, while Bcl-2 blocks apoptosis. Therefore, the balance between the levels of Bcl-2 and BAD is important in determining cell survival or death (8,10). Our finding shown in the present study demonstrated that bufalin-treatment increased the ratio of BAD/Bcl-2, suggesting that the increase of the ratio of BAD/Bcl-2 might be the key factor of bufalin-induced apoptosis (Fig. 8A).

Next, we investigated whether the possible mechanism of bufalin-induced apoptosis is through activation of the AKT signaling pathway in CAL 27 cells (Figs. 4-8). AKT is a serine-threonine kinase and regulates cancer cell progression. It has been demonstrated that AKT is over-activated or over-expressed in many human malignancies (5,25). Understanding the control of the AKT signaling pathway is potentially important for developing therapeutic inhibitors. CAL 27 cells overexpressing constitutively active AKT (CA-AKT) is used for evaluating
bufalin-induced cell death. Data demonstrated that CA-AKT diminished bufalin-induced cell apoptosis. Hence, our study is the first report regarding AKT signaling contributed to bufalin-triggered mitochondrial apoptotic death in CAL 27 cells in vitro.

In summary, our results provided further insight into bufalin-induced apoptosis and deepen our knowledge on the mechanisms of anticancer activity of bufalin in CAL 27 cells. Bufalin causes cell cycle arrest at the G2/M phase. The bufalin-induced apoptosis is dependent on the mitochondria-mediated caspase activation and involvement of the regulation of Bcl-2 and BAD (Fig. 8). An exciting finding in this study is that constitutively active AKT had no significant effect on bufalin-induced apoptosis. These data provide a hint toward clarification of the mechanisms of bufalin-induced apoptosis, but it might be a long way before unveiling the complete mechanisms underlying bufalin-induced apoptosis in tumor cells. Moreover, other signaling components such as FOXO3a, p27kip1, and c-Myc might be also involved in bufalin-induced apoptosis (35,36). We believe that bufalin has important antitumor properties and is a promising chemotherapeutic agent for the treatment of human oral cancer in the future. Continued examination of AKT and other signaling pathways will be important in further delineating the cell death mechanisms. Work ongoing in our laboratory is addressing these issues.

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