Research Article

Honokiol Eliminates Human Oral Cancer Stem-Like Cells Accompanied with Suppression of Wnt/β-Catenin Signaling and Apoptosis Induction

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Honokiol, an active compound of Magnolia officinalis, exerted many anticancer effects on various types of cancer cells. We explored its effects on the elimination of cancer stem-like side population (SP) cells in human oral squamous cell carcinoma SAS cells. The sorted SP cells possessed much higher expression of stemness genes, such as ABCG2, ABCC5, EpCAM, OCT-4, CD133, CD44, and β-catenin, and more clonogenicity as compared with the Non-SP cells. After 48 h of treatment, honokiol dose dependently reduced the proportion of SP from 2.53% to 0.09%. Apoptosis of honokiol-treated SP cells was evidenced by increased annexin V staining and cleaved caspase-3 as well as decreased Survivin and Bcl-2. Mechanistically, honokiol inhibited the CD44 and Wnt/β-catenin signaling of SP cells. The Wnt signaling transducers such as β-catenin and TCF-4 were decreased in honokiol-treated SP cells, while the β-catenin degradation promoting kinase GSK-3α/β was increased. Consistently, the protein levels of β-catenin downstream targets such as c-Myc and Cyclin D1 were also downregulated. Furthermore, the β-catenin-related EMT markers such as Slug and Snail were markedly suppressed by honokiol. Our findings indicate honokiol may be able to eliminate oral cancer stem cells through apoptosis induction, suppression of Wnt/β-catenin signaling, and inhibition of EMT.

1. Introduction

Oral squamous cell carcinoma (OSCC) is the most common type of head and neck cancer [1], which ranks the sixth worldwide for cancer-related mortality [2]. It accounts for almost 3% of cancer cases in the world [3]. In Taiwan, OSCC has the highest rate of increase among male cancers. Despite the improvements in surgical and radiotherapy, the five-year survival rate for oral cancer has remained unchanged at about 50% over the past 30 years [4]. Recently, targeted therapy has been useful primarily in early malignancies; however, it becomes ineffective eventually, due to the nonredundant tyrosine kinase receptors and activation of signaling through several pathways in advanced cancers [5]. As the available therapies are limited by the resistance and severe adverse side effects, more effective therapeutics are urgently needed to combat this life-threatening disease.

The cancer stem cell (CSC) hypothesis asserts that malignant tumors are maintained exclusively by a small subpopulation of cells that give rise to phenotypically diverse cancer...
cells [6–8]. In recent years, the CSC hypothesis has been studied in many types of cancers including OSCC, and there is growing evidence suggesting CSC is responsible for the tumor resistance to chemotherapy and radiation therapy [9, 10]. Targeting CSC has thus been a potential strategy to improve the treatment outcome of cancers.

The CSCs typically overexpress ATP-Binding cassette (ABC) transporters to pump out drugs and thus result in the resistance to chemotherapeutic agents [11]. This overexpression of ABC transporters could facilitate CSCs isolation by flow cytometry on the basis of their ability to pump out the fluorescent DNA binding dye such as Hoechst 33342. With the capacity to pump out the Hoechst 33342, the signals of CSCs fell to the "side" of the bulk positively stained cells in FACS (fluorescence activated cell sorter) analysis plots and were referred as side population (SP) cells [12]. The SP cells from OSCC have been shown to possess phenotypes of CSC such as elevated levels of ABCG2, CD44, Oct-4, and so forth, higher clonogenicity, and tumorigenicity of xenografts [1, 10]. We therefore used SP cells as a model to evaluate natural products with potential targeting of CSC.

Honokiol is an active component isolated from the bark of the traditional Chinese medicine Magnolia officinalis. It had been shown to exert various biological effects, including muscle relaxation, anti-inflammation, antioxidants activity, and various protecting effects against hepatotoxicity, neurotoxicity, thrombosis, and angiopathy [13]. Interestingly, honokiol had been reported to induce apoptosis or growth inhibition in a variety of cancer cell lines including breast, lung, colon, and prostate cancer [14–17] and inhibit tumor growth in xenograft animal models [14, 15, 17–21]. In head and neck cancer, the anticancer effects of honokiol also had been demonstrated both in vitro and in vivo [22, 23]. It is worthy and interesting to investigate if honokiol could eliminate the CSC in such type of cancer. Here, we studied the effects of honokiol on the cancer stem-like SP cells isolated from SAS human OSCC cells. In addition to apoptosis induction and CD44 surface marker expression, the Wnt/β-catenin signaling pathway, epithelial-mesenchymal transition (EMT) markers [24–26], and epithelial cell adhesion molecule (EpCAM) [27] which were known to play crucial roles in regulation of CSC properties were also examined.

2. Materials and Methods

2.1. Chemicals and Reagents. Honokiol (purity > 98%) was purchased from Sigma. Primary antibodies Survivin, Bcl-2, caspase-3, β-catenin, TCF-4, GS-3α/β, c-Myc, Cyclin D1, CD44, Slug, Snail, and β-actin were purchased from Cell Signaling Technology. Honokiol was dissolved in dimethyl sulfoxide (DMSO) and further diluted in sterile culture medium immediately prior to use. The final concentrations of DMSO in cell cultures were all less than 0.05%. A TRIzol RNA isolation kit was obtained from Life Technologies, and primers for RT-PCR, dNTP, reverse transcriptase, and Taq polymerase were obtained from Gibco BRL. N2 supplement, human recombinant bFGF, and EGF were all purchased from Invitrogen.

2.2. Cell Culture. The human oral squamous cell carcinoma cell line SAS (JCRB0260) was purchased from the JCRB Cell Bank (Japanese Collection of Research Bioresources Cell Bank). This cell line was established from a poorly differentiated squamous cell carcinoma of the tongue. SAS oral cancer cells were maintained as monolayers in Dulbecco’s modification Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (HyClone) and 1% penicillin/streptomycin (Mediatech). Cells were cultured at 37°C in a water-jacketed 5% CO₂ incubator.

2.3. Side Population Analysis and Purification Using Flow Cytometry. The side population (SP) cells were analyzed and sorted by Hoechst 33342 (Sigma) staining and FACS Aria III cell sorter (BD). SAS cells were detached from the dishes with Trypsin-EDTA (Invitrogen) and suspended at 1 × 10⁶ cells/mL in Hank’s balanced salt solution (HBSS) supplemented with 3% fetal calf serum and 10 mM HEPES. These cells were then incubated at 37°C for 90 minutes with 2.5 µg/mL Hoechst 33342, either alone or in the presence of 50 µM verapamil (Sigma), a nonspecific inhibitor of ATP-binding cassette transporter (ABC transporter). The diminishment of SP cells in the presence of verapamil was used to define the flow cytometry gate for sorting SP cells. After 90-minute incubation, the cells were centrifuged immediately for 5 minutes at 300 × g, 4°C and resuspended in ice-cold HBSS. The cells were kept on the ice to inhibit efflux of Hoechst dye and 1 µg/mL propidium iodide (BD) was then added to discriminate dead cells. Finally, these cells were filtered through a 40 µm cell strainer (BD) to obtain single suspension cells for the analysis and sorting on FACS Aria III flow cytometer.

2.4. Sphere Culture. After sorting, SAS side population cells were seeded with a density of 500 cells/well in 6-well ultra-low attachment plates (Corning) in DMEM/F-12 culture medium supplemented with N2 supplement, 10 ng/mL human recombinant bFGF, and 10 ng/mL EGF. After culture for 14 days, spheres were harvested for the subsequent assays of SP cells.

2.5. Colony Formation Assay. SAS SP or Non-SP cells were plated at about 200 cells per well in 6-well coated plates and cultured in the medium described in Section 2.4. for 10 days. After most cell clones had increased to >50 cells, they were washed with PBS, fixed in methanol for 15 min, and stained with crystal violet for 15 min at room temperature. After washing out the dye, colonies with >50 cells were counted as positive colonies.

2.6. Assessment of the Proliferation of SAS SP and Non-SP Cells after Treatment with Honokiol. The SAS SP and Non-SP cells were seeded into 96-well plates in the medium described in Section 2.4. at a density of 3000 cells/well. After 24 h, the medium was replaced with fresh growth medium containing various doses of honokiol or vehicle only and the cells were incubated for another 48 hours. At harvest, cells were fixed with trichloroacetic acid (TCA) by gently adding it to each well to a final concentration of 10% with subsequent incubation for 1 hour at 4°C. The plates were then washed...
5 times with tap water and air dried. The dried plates were
stained with 100 μL of 0.4% (v/v) Sulforhodamine B (SRB)
prepared in 1% (v/v) acetic acid for 10 minutes at room
temperature. The plates were rinsed quickly 4 times with 1% acetic
acid to remove unbound dye and then air dried until no mois-
ture was visible. The bound dye was solubilized in 20 mmol/L
Tris base (100 μL/well) for 5 minutes on a shaker. Optical
densities were read on a microplate reader (Molecular De-
vices) at 562 nm. The optical density is directly proportional to
the cell number over a wide range.

2.7. Flow Cytometric Assessment of Stemness Markers Expres-
sion. After being cultured in the medium described in
Section 2.4. for 14 days, the SAS SP and Non-SP cells were
harvested and incubated with phycoerythrin- (PE-) conju-
gated antibodies against Oct-4 (Santa Cruz), human CD133/2
(Milenyi Biotec, recognizes epitope 2 of the CD133), and
CD44 (Cell Signaling), respectively, at 4°C for 30 minutes.
After washing the cells of excess reagents, the stained cells
were immediately analyzed by a FACSCalibur flow cytomter
(BD). Approximately 10,000 counts were made for each
sample. Anti-IgG (isotype) were used to detect the non-speci-
fic binding background activity in this assay.

2.8. Flow Cytometric Assessment of Apoptosis Using Annexin
V/PI Staining. Apoptosis was determined by using a com-
mercially available Annexin V apoptosis detection kit (BD)
and flow cytometry. The SAS SP cells were seeded in 6-well
plate (2 × 10⁵ cells/well) in medium described in Section 2.4.
and incubated for 24 h before treatment. Following treatment
with different doses of honokiol for 48 h, the SAS SP cells were
harvested, washed twice with 2 mL of ice-cold phosphate-
buffered saline (PBS), and then incubated with 100 μL of
HEPES buffer containing 2 μL of fluorescein–isothiocyanate–
(FITC-) conjugated annexin V (10 μg/mL) and 2 μL of pro-
pidium iodide (PI, 100 μg/mL) for 15 minutes. After washing
the cells of excess reagents, 400 μL of binding buffer was
added. The stained cells were immediately analyzed with a
FACSCalibur flow cytomter. Approximately 10,000 counts
were made for each sample.

2.9. β-Catenin/Tcf/Transcription Reporter Assay. SAS SP cells
were plated in 6-well plates, grown to 80%–90% confluence,
and transiently transfected with the plasmids of TOPflash.
TOPFlash has 3 copies of the Tcf/Lef binding sites in the
upstream of a thymidin kinase (TK) promoter and the fire-
fly luciferase gene. All transfections were performed with
Lipofectamine and 1.8 μg of TOPFlash plasmids. To normalize
transfection efficiency, cells were cotransfected with 0.2 μg
of the internal control reporter encoding Renilla reniformis
luciferase driven under the TK promoter. After transfection,
cells were incubated in the medium described in Section 2.4.
with or without honokiol (0–10 μM) for 48 h and then lysed
with reporter lysis buffer at harvest. Luciferase activity was
determined by using the Dual-Luciferase Assay System kit
(Promega) according to the manufacturer’s protocol. The
experiments were performed in triplicate, and the results
were expressed in relative luciferase units (RLU), normalized
to transfection efficiency as described above.

2.10. RT-PCR Analysis of Stemness Genes Expression. Trizol
reagent was used to extract the mRNAs from the SAS
SP and Non-SP cells according to the manufacturer’s rec-
commended protocol. Two μg of RNA was added to RT-
PCR reactions containing primers at a concentration of
0.5 μM. After a 42°C/60 min reverse transcription step, 30
cycles of PCR amplification were performed at 94°C for
30 sec, 58°C for 50 sec, and 72°C for 30 sec. PCR products
were run on 1.5% agarose gels for identification. Primers
used were for ABCG2, forward: 5'-CATCAACCTTTCC-
GGGGTGA-3' and reverse: 5'-TGTGAGATTGACCAA-
CACACCA-3' for ABCG2, forward: 5'-CCAAGCTGA-
C CCCAAAATGAAAA-3' and reverse: 5'-TGGATG-
TCTTTGCCCCCTTCTTCTTCT-3' for EpCAM, forward:
5'-CTGCCAAATGTTTGGTGATG-3' and reverse: 5'-ACG-
GTGTTGATCTCCTTCT-3' for Oct-4, forward: 5'-GGA-
GAGCAACTCCGATGG-3' and reverse: 5'-TTGATGTC-
GGGACTCCTC-3' for CD133, forward: 5'-CATAAA-
GCTGGACCATATTGG-3' and reverse: 5'-CCTGTGCT-
TGTTATGTTT-3' for CD44, forward: 5'-CTCTGCTC-
CTTCTAGTACA-3' and reverse: 5'-GTTAGCGGAG-
GTAGTCTGTA-3' for GAPDH, forward: 5'-ACACA-
GTCAATGCCATCAC-3' and reverse: 5'-TCCACCACC-
CTGTTGCTGA-3'.

2.11. Western Blotting. After treatment with various concen-
tration of honokiol for 48 h, the cell lysates were prepared
using ReadyPrep Protein Extraction Kit (Bio-Rad) according
to instructions provided. Total cell lysates (20 μg) were se-
parated electrophoretically by a 10% polyacrylamide SDS-PAGE
gel and transferred onto a polyvinylidene fluoride membrane
using the Bio-Rad Mini-Protein transfer system. The mem-
brane was cropped and further incubated overnight at 4°C
with specific antibodies against Survivin, Bcl-2, caspase-3,
active β-catenin, TCF-4, GSK-3β, c-Myc, cyclin D1, CD44,
Slug and Snail, and β-actin, respectively. After incubation
with primary antibodies, the cropped membranes were
washed with TBST 3 times and then incubated with horse-
ardish peroxidase-labeled secondary antibody for 45 min-
utes at room temperature. After washing with TBST 3 times,
final detection was performed with enhanced chemilumi-
nescence (ECL) Western blotting reagents (Minipore) and
BioSpectrum Imaging System (UVP, Upland, CA, USA).

2.12. Statistical Analysis. The significant difference between
control and experimental groups was analysed using
\( t \)-test (* \( P < 0.05; ** P < 0.01 \).

3. Results

3.1. SAS SP Cells Possessed Characteristicsof Cancer Stem Cell.
To evaluate the effects of honokiol on the cancer stem cells, we
sorted and characterized the SP cells from human SAS
oral cancer cells as described. As shown in Figure 1(a), a
small proportion (2.7%) of SP was detected in the left-down
side of the flow cytometry histogram, which could be dim-
inished in the presence of verapamil (50 μM). After 7 days of
Figure 1: SAS SP cells possessed the properties of stem cells. (a) Spheroid morphology of SP cells and fibroblast-like shape of Non-SP cells after 7 days of culture. The SP and Non-SP cells were sorted by Hoechst 33342 staining as indicated in the flow cytometry histogram and the gated region for dye excluding SP cells was identified by verapamil as described in Section 2. (b) SP cells had markedly higher clonogenicity than the Non-SP cells. The colonies formed were counted as described in materials and methods. (c) The mRNA expressions of multiple drug resistance genes (ABCG2, ABCC5) and epithelial cell adhesion molecule (EpCAM) in SP cells were much higher than those in Non-SP cells (GAPDH expression used as loading control). (d) The SP cells expressed much higher mRNA levels of stemness markers Oct-4, CD133, CD44, and β-catenin than the Non-SP cells (GAPDH expression used as loading control). (e) Flow cytometric analysis of stem cell markers expression in SP and Non-SP cells. The histograms display the expression of Oct-4, CD133, and CD44 in SP and Non-SP cells (I-SP, I-Non-SP; isotype control of SP and Non-SP cells, resp.; N: Non-SP cells; S: SP cells; the I-SP and I-Non-SP in the histograms of Oct-4 and CD44 are almost overlapped.).
culture, the sorted SP cells displayed sphere-like morphology in contrast to the flattened shape of Non-SP cells (Figure 1(a)). Besides, the SP cells appeared to have higher clonogenicity than the Non-SP cells (Figure 1(b)). In accordance with the lowered Hoechst blue-red intensity, the mRNA expression of ATP-binding cassette transporters such as ABCG2 and ABCC5 was much higher in SP cells as compared with that of Non-SP cells (Figure 1(c)). Recently, elevation of epithelial cell adhesion molecule (EpCAM) was reported to enhance tumor sphere formation and tumor initiation [27]. Incompatible with this report, we found the SP cells also possessed much higher level of EpCAM mRNA than the Non-SP cells (lower part of Figure 1(c)). We next examined the expression of stemness markers in SP cells. As expected, the mRNA levels of Oct-4, CD133, and CD44 in SP cells were much higher than those of Non-SP cells (Figure 1(d)). This phenomenon was further confirmed by flow cytometry analysis, in which the peaks of SP cells markedly shifted apart from those of the Non-SP cells to the high intensity (right) side (Figure 1(e)). All these data indicated the CSC properties of SAS SP cells.

3.2. Honokiol Eliminated the SP Cells and Suppressed Their Clonogenicity. The sensitivities of SP and parental SAS cells to honokiol are quite different. The SP cells were markedly more sensitive than the parental SAS cells. After 48 h of treatment, honokiol suppressed 21% parental SAS cell growth at dose of 10 μM. However, it inhibited 78% of SP cell growth at the same dose (Figure 2(a)). We then examined the percentage of SP in honokiol-treated SAS cells. After 48 h of incubation with honokiol at doses of 2.5, 5, and 10 μM, the SP cells were dose dependently eliminated from 2.53% to 1.45%, 0.43, and 0.09%, respectively (Figures 2(b) and 2(c)). Accordingly, the clonogenicity of SAS SP cells was also markedly suppressed by honokiol at doses of 5 and 10 μM (Figure 3).
3.3. Honokiol-Induced Apoptosis of SAS SP Cells. To investigate the mechanism of action underlying the SP cell elimination, we examined the apoptosis in honokiol-treated SP cells. After treatment with honokiol for 48 h, quantitative analysis of annexin V-FITC/PI staining by flow cytometry showed that the early apoptotic fraction (annexin V positive and PI negative) was increased from 0.59% of control to 8.48% and 11.5% at doses of 5 and 10 μM, respectively. Simultaneously, the late apoptotic fraction (annexin V positive and PI positive) was increased from 0.46% of control to 6.16% and 22.7%, respectively (Figure 4(a)). This honokiol-induced apoptosis of SP cells was further confirmed by the dose-dependent cleavage of caspase-3 and downregulation of antiapoptosis proteins such as Bcl-2 and Survivin (Figure 4(b)).

3.4. Honokiol Inhibited the Wnt/β-catenin Signaling and EMT Markers in SAS SP Cells. The Wnt/β-catenin pathway had been reported to play a crucial role in oral CSC and considered as an important target for screening therapeutics [26]. To more understand the molecular mechanisms responsible for the effects of honokiol against SAS SP cells, we analyzed the Wnt signaling pathway in honokiol-treated SP cells. Western blot results showed that honokiol not only decreased the Wnt signaling transducers such as β-catenin and TCF-4 (T cell transcription factor-4) but also increased GSK-3α/β (glycogen synthase kinase), a β-catenin degradation promoting kinase. Consequently, the protein products of β-catenin downstream genes such as c-Myc and Cyclin D1 were downregulated by honokiol (Figure 5(a)).

Since it is known that Wnt/β-catenin pathway could mediate EMT (epithelial-mesenchymal transition), we thus examined the EMT markers in SP cells after treatment with honokiol. As shown in Figure 5(c), the protein levels of mesenchymal markers such as Slug and Snail were markedly suppressed, while β-catenin and its downstream Cyclin D1 were inhibited.

4. Discussion

OSCC is an important public health issue worldwide. To date, there is no effective treatment for patients with local relapse and distant metastasis. It remains a highly lethal and disfiguring disease [1, 28]. Recent studies showed that the resistance of OSCC to conventional chemotherapy or radiation therapy might be due to the presence of CSC [28]. Elimination of CSC would be an effective way to improve the clinical outcome of OSCC [10]. Searching the CSC targeting agents thus had been of great importance.

The "side population" (SP) is a well-known method to isolate cancer stem-like cells [12], although not all SP cells isolated from different cell lines possess CSC properties [29]. The SP from other OSCC cells had been found to possess CSC phenotypes and may play an important role in tumorigenesis [1]. Consistently, in this study, the SP isolated from SAS human OSCC cells was characterized with CSC properties as described. Moreover, Lin et al. recently reported that the expression of EpCAM in SAS OSCC cells was elevated when cultivated the attached cells into the tumorsphere [27]. In parallel with this, our results showed the mRNA level of EpCAM in SAS SP cells was much higher in comparison with that of Non-SP cells. Therefore, it is rational to use the SAS SP cells for investigating the effects of honokiol on CSC elimination.

According to recent studies, phytochemicals and herbs would be potential sources of therapeutics for CSC elimination. For example, resveratrol, curcumin, sulforaphane, and...
so forth had been reported to suppress the cancer stem-like cells [30]. Recently, Ponnurangam et al. found honokiol in combination with radiation could suppress the colonosphere formation and DCLK1+ and CD133+ populations of colon cancer cells, indicating the effects of honokiol on CSC inhibition [31]. In agreement with the study by Ponnurangam et al., we demonstrated the effects of honokiol on the elimination of CSC-like SP in OSCC cells. Ponnurangam et al. suggested honokiol targeted CSC by inhibiting the γ-secretase complex and the Notch signaling pathway in colon cancer [31]. In OSCC cells, our results showed honokiol suppressed the Wnt/β-catenin pathway and EMT markers of the CSC-like SP cells. The precise molecular mechanisms underlying the CSC targeting effects of honokiol may be complicated and variant in different types of cancer cells. Further investigation is needed to more precisely clarify the mechanisms of action.

The Wnt pathway is an important target for developing novel therapeutics targeting on CSC [26]. In OSCC cells, activation of Wnt pathway was shown to play a crucial role in the proliferation of CSC-like cells [28]. According to our results, the elimination of SP cells by honokiol may highly attribute to the inhibition of Wnt/β-catenin pathway, the most intensively studied and well-characterized Wnt pathway [26]. The level of β-catenin is tightly regulated by ubiquitin-proteasome-mediated degradation, which is controlled by a “destruction” complex composed of axin, adenomatous polyposis coli (APC), glycogen synthase kinase-3β (GSK-3β), and so on [26]. The β-catenin could be phosphorylated by GSK-3β, leading to its degradation. In response to Wnt, the GSK-3β will be inhibited, resulting in the accumulation of nonphosphorylated β-catenin, which could avoid degradation and translocate into the nucleus. In the nucleus,
Figure 5: Honokiol downregulated the Wnt/β-catenin signaling and its downstream in SAS SP cells. (a) Honokiol decreased the protein levels of β-catenin and TCF-4 as well as their downstream c-Myc and Cyclin D1, accompanied with the increase of GSK-3α/β in SP cells. The expression of stem cell marker CD44 was also decreased. Cells were treated with honokiol for 48h and the cell lysates were analyzed with Western blot using β-actin as loading control. (b) TOPFLASH TCF reporter assay in honokiol-treated SP cells. Cells were transfected with pTOPFLASH luciferase reporter constructs as described in Section 2 and treated with honokiol for 48 h, lysed, and analyzed using the Dual Luciferase Assay System kit. Results are expressed in relative luciferase units (RLU), normalized to transfection efficiency as described in Section 2. Each bar represents the mean of three independent transfections ± S.E. (c) Honokiol downregulated the EMT markers Slug and Snail proteins as well as Cyclin D1 while suppressed the β-catenin in SP cells. Cells were treated with honokiol for 48 h and then analyzed by Western blot as described in (a).

β-catenin forms a complex with T cell transcription factor (TCF) and other cofactors to activate the transcription of Wnt signaling target genes such as c-Myc and Cyclin D1 [26, 32]. In accordance with these events of Wnt signaling, honokiol decreased β-catenin and TCF-4 of SP cells accompanied with the elevated GSK-3α/β and downregulated c-Myc and Cyclin D1 protein levels. The elevated GSK-3α/β might increase the formation of β-catenin/axin/GSK-3β destruction complex, a known precursor to β-catenin degradation. Based on our data, we deduce that the effects of honokiol on CSC elimination are highly related to Wnt signaling inhibition.

CD44, one of the well-known CSC markers, is initially used for breast CSC identification. Although CD44 alone is not sufficient for precisely isolating CSC in head and neck cancer cells, the CD44 expressing cells appear to have elevated tumorigenicity [10]. Consistently, our results showed the SAS SP cells expressed much higher level of CD44 than that of the Non-SP cells. Similar results were also observed...
in another OSCC cells by Zhang et al. [1]. As CD44 is also a target gene of Wnt signaling [33], the decreased CD44 in honokiol-treated SAS SP cells might also attribute to the inhibition of Wnt signaling cascade. It is known that β-catenin could mediate epithelial to mesenchymal transition (EMT) [34], which plays a crucial role in cancer invasion and metastasis. The EMT markers such as Snail and Slug are also the target genes of β-catenin [35, 36]. Thus, the suppression of Snail and Slug in honokiol-treated SP cells might also result from the inhibition of Wnt/β-catenin signaling pathway. On the other hand, Mani et al. showed that EMT could generate cells with properties of stem cells [37]. Following this finding, extensive studies had demonstrated the link between EMT and CSC phenotype [38, 39]. Therefore, the suppressing effects of honokiol on the above EMT markers might also coincide with its effects against the stemness of CSC.

A number of nonsteroidal anti-inflammatory drugs (NSAIDs), such as aspirin, celecoxib, and sulindac, and natural compounds like epigallocatechin-3-gallate (EGCG), resveratrol, quercetin, curcumin, and so forth had been identified as inhibitors and/or modulators of Wnt/β-catenin signaling pathway [26]. Many of them such as EGCG, resveratrol, and curcumin were shown to have potential in CSC elimination [30]. Here, we demonstrated the substantial effects of honokiol on Wnt/β-catenin signaling inhibition and apoptosis induction in oral CSCs. As the biology of CSC is comprehensive and contains a considerable crosstalk in signaling pathways, combining honokiol with other CSC-eliminating agents listed above might provide better therapeutic effects. Further future studies to investigate these combination effects on CSC elimination are warranted.

5. Conclusions

Honokiol eliminated the CSC-like SP cells in SAS human oral squamous cell carcinoma cells. The underlying mechanisms were associated with apoptosis induction and the inhibition of Wnt/β-catenin cascade and related EMT markers. As CSC is a very important target for cancer therapy, our results further demonstrate the anticancer properties of honokiol and point out its potential application in CSC targeted therapy of oral cancer.

Authors’ Contributions

Chih-Jung Yao and Gi-Ming Lai contributed equally to this paper.

Conflict of Interests

The authors declare no conflict of interests.

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