Research Article

Potential Therapeutic Role of Z-Isochaihulactone in Lung Cancer through Induction of Apoptosis via Notch Signaling

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Lung cancer is one of the leading causes of cancer death in worldwide and required for novel therapeutic strategy. Our previous research demonstrated that the crude acetone extract of Bupleurum scorzonerifolium (BS-AE) and its component isochaihulactone induce antiproliferative and apoptotic effects on the lung adenocarcinoma cell line. Structural analysis has identified isochaihulactone as a lignan, with a chiral center and two racemic forms (Z-isochaihulactone and E-isochaihulactone). In this study, Z-isochaihulactone displayed significantly higher tumor cytotoxicity than E-isochaihulactone in A549 cells. The notch signaling pathway plays a pivotal role in determination of cell fate during development, while in lung cancer, it might have oncogenic or tumor-suppressive controversial functions. We showed that Z-isochaihulactone induced morphological changes in the A549 cells, inhibited cell growth, and arrested the cell cycle at the G2/M phase. It also induced upregulation of the active form of Notch1 (notch intracellular domain, NICD), which further induced p21 and c-Myc expression in time- and dose-dependent manners. Administrations of Z-isochaihulactone in nude mice can significantly inhibit tumor growth due to enhancement of NICD expression confirmed by immunohistochemical analysis. Taken together, our results supported that Z-isochaihulactone can efficiently inhibit tumorigenicity and be a potential compound for therapy.

1. Background

Cancers present a serious clinical problem, causing significant social effects and a considerable economic burden on the healthcare system. Surgery is the only potentially curative treatment; however, complete resection is often impossible because of widespread disease [1, 2]. Other conventional cancer treatments, such as chemotherapy and external beam radiation, are largely ineffective against carcinoids [3]. The development of novel therapeutic approaches for advanced carcinoid tumor disease is therefore of considerable urgency. Previous studies have reported that ectopic expression of the notch receptor Notch1 in carcinoid and other neuroendocrine tumor cells resulted in decreased production of neuroendocrine tumor markers and suppression of cancer cell growth [4–8]. The authors concluded that activation of Notch1 signaling was an attractive approach for the development of new treatments for carcinoids and other neuroendocrine tumors. Previous studies have also shown that notch signaling plays an oncogenic role in a number of tumors, such as T-ALL [9] and mouse mammary tumors [10], by inhibiting apoptosis and inducing proliferation [11, 12]. However, recent investigations have shown that depending on the cell context, notch signaling might also
display opposing functions in tumorigenesis. In SCLC, expression of activated Notch1 or Notch2 upregulated the expression of p21* and p27*kip1, leading to cell cycle arrest [13]. A direct transcriptional target of aberrant Notch1 signaling, c-Myc, plays an indispensable role in Notch1C-induced murine mammary tumorigenesis [14]. Chen et al. further reported that the Notch1 protein is downregulated in NSCLC cell lines, and that the expression of the constitutively active Notch1 in adenocarcinoma of the lung (ACL) cells caused cell death. Under hypoxic conditions, Notch1 is markedly upregulated, which might be essential for cell survival in ACL, a type of NSCLC [15]. These results indicate that oxygen concentration determines the biological effects of Notch1 signaling in ACL. Zheng et al. have reported that overexpression of activated Notch1 (NICD) inhibited the growth of the lung adenocarcinoma A549 cells in vitro through induction of cell cycle arrest and suppressed A549 tumor growth in nude mice [16]. Notch signaling might therefore have tumor-suppressive function in human lung adenocarcinoma cells [17]. In mammals, at least 2 families of bHLH proteins are induced following Notch activation: the HES (hairy enhancer of split) family and the HRT (hairy-related transcription factor; also known as HEY, HESR, HERP, or CHF) family. Both the HES and HRT families function as transcriptional repressors [12]. Besides, they have other downstream targets of NICD, such as p21, c-Myc, NF-kB, and cyclin D1 [12, 14, 18, 19], and the expression of these genes via NICD also has to depend on the cell context. Valproic acid (VPA) is a branched chain fatty acid used in the treatment of patients with epilepsy and other neuropsychiatric disorders [20]. Valproic acid is a well-established histone deacetylase (HDAC) inhibitor and currently under evaluation in various cancer clinical trials. Stockhausen and colleagues recently described the ability of VPA to increase Notch1 protein levels in neuroblastoma cells [21]. Studies have also described the activation of Notch1 signaling and induction of cell apoptosis by VPA in several tumor cell lines [22, 23]. In the present study, VPA was used as a positive control.

Nan-Chai-Hu (Chai Hu of the South), the root of *Bupleurum scorzonerifolium*, is a Chinese herb used in the treatment of influenza, fever, malaria, cancer, and menstrual disorders in China, Japan, and other countries of Asia. It is also a major ingredient of traditional Chinese medicine formulations such as Sho-Saiko-To-Tang (T'J9 in Japan) and Bu-Zhong-Yi-Qi-Tang. Partitioning of the acetone fraction extract of Nan-Chai-Hu enabled the identification of a novel lignan, isochaihulactone (K8), which displays antitumor activity against A549 cells in vitro and in vivo [24–26]. Previous research has also identified the target gene of isochaihulactone, its mediation by NAG-1 upregulation, and tumor apoptotic effects [27]. Structural analysis of isochaihulactone has identified it as lignan, with a chiral center and two racemic forms (the Z-isochaihulactone and E-isochaihulactone). The present study’s analyzes showed that Z-isochaihulactone displays significantly higher tumor cytotoxicity than E-isochaihulactone in A549 cells in vitro, with Z-isochaihulactone treatment resulting in dose-dependent inhibition of carcinoid cell growth. Western blot and flow cytometric analyses demonstrated that cell cycle arrest at the G2 phase mediated this growth inhibition. In cervical cancer cells, overexpression of the active form of Notch1 inhibited the proliferation of tumor cells [28]. Constitutive activation of Notch1 also inhibited the growth of human lung adenocarcinoma A549 cells through induction of cell cycle arrest and apoptosis [16]. The present study describes Z-isochaihulactone-induced inhibition of carcinoid tumor growth in vivo in a xenograft model and suggests that the activation of Notch1 signaling mediates these effects. These data form the basis for a clinical trial on Z-isochaihulactone in the treatment of patients with advanced carcinoid cancer, a disease for which few effective therapies currently exist.

2. Materials and Methods

2.1. Chemistry. The synthesis of anhydropodorhizol 1 was previously described in our previous report [26], while the synthesis of isochaihulactone 2, to the best of our knowledge, has never been reported. The target compounds 1 and 2 were prepared using the synthetic strategy shown in Scheme 1. Compound 3 was readily obtained via Stobbe condensation of piperonal and diethyl succinate. Subsequent hydrogenation of 3 afforded 4 in excellent yield, and the chemoselective reduction of the potassium salts of 4 with CaCl2/NaBH4 in ethanol gave the desired lactone 5. Aldol condensation of 5 with 3,4,5-trimethoxybenzaldehyde gave the alcohol 6 which was treated with either acetic anhydride or methanesulfonyl chloride to afford the corresponding acetate or mesylate 7. Base promoted elimination of the formed acetate afforded 1 exclusively, while elimination of the mesylate intermediate afforded 1 and 2 in 1:3 ratio.

2.2. Chemicals and Reagents. Z-isochaihulactone was dissolved in DMSO to a concentration of 50 mM and stored in –20°C as a master stock solution. VPA (2-propylpentanoic acid), dimethyl sulfoxide (DMSO), 3-(4,5-dimethyl thiazol-2-yl)-2,5-diphenyl tetrazolium bromide (MTT), propidium iodide (PI), and Hoechst 33342 were purchased from Sigma Chemical Co. (St. Louis, MO, USA). RPMI 1640 medium, fetal bovine serum (FBS), penicillin, trypsin/EDTA, and Nu PAGE Bis-Tris Electrophoresis System (precast polyacrylamide mini-gel) were purchased from Invitrogen (Carlsbad, CA, USA). Cyclin B1, cdc25C, cdc2, phosphor-Asp330 caspase-9, caspase-3, PARP, and β-actin antibodies were purchased from Cell signaling Technology. Polyvinylidene-fluoride (PVDF) membranes, BSA Protein Assay Kit, and Western blot chemiluminescence reagent were purchased from Amersham Biosciences (Arlington Heights, IL). [siRNA against Notch-1 and nonspecific siRNA (Santa Cruz Biotechnology, sc-44226 and sc-37007)].

2.3. Cell Lines and Cell Culture. A549 human lung adenocarcinoma cells were obtained from American Type Culture Collection (Manassas, VA) and propagated in culture dishes at the desired densities in RPMI 1640 medium.
Supplemented with 10% fetal bovine serum and 1% penicillin/streptomycin. The cells were incubated at 37°C in a humidified atmosphere containing 5% CO₂.

2.4. Growth Inhibition Assay. The viability of the cells after treatment with various chemicals was evaluated using the MTT assay performed in triplicate. Briefly, the cancer cells (3 × 10³) were incubated in 96-well plates containing 200 μL of the culture medium. Cells were permitted to adhere for 12–18 h then washed with phosphate-buffered saline (PBS). Solutions were always prepared fresh by dissolving 0.2% DMSO or drugs in culture medium and then were added to A549 cells. After 24–72 h of exposure, the drug-containing medium was removed, washed with PBS, and replaced by fresh medium. The cells in each well were then incubated in culture medium with 500 μg/mL MTT for 1 h. After the medium was removed, 100 μL of DMSO was added to each well. Absorbance at 570 nm of the maximum was detected by a Power Wave X Microplate ELISA Reader (Bio-Tek Instruments, Winooski, VT). The absorbance for DMSO-treated cells was considered as 100%. The results were determined by three independent experiments.

2.5. Hoechst 33342 Staining. After a 48-h treatment of the cells with K8 (2.5 to 10 μM), Hoechst 33248 staining was performed. A549 human lung adenocarcinoma cells were stained with Hoechst 33248 dye to evaluate apoptosis. The cells were fixed with 4% paraformaldehyde at room temperature and stained with Hoechst 33342 working solution (5 μmol/L) for 30 min, then washed with PBS. Fluorescence was visualized using a fluorescent microscope (Zeiss) under 400x magnification.

2.6. Cell Cycle Analysis. The cell cycle was determined by flow cytometry with DNA staining to reveal the total amount of DNA. Approximately 5 × 10³ cells were incubated in various concentrations of K8 and VPA for 48 h. Cells were harvested by treating the cells with trypsin/EDTA. The cells were collected, washed with PBS, fixed with cold 70% ethanol overnight, and then stained with a solution containing 20 μg/mL PI and 0.1% Triton X-100 for 1 h in the dark. The cells will then pass through FACScan flow cytometer (equipped with a 488-nmargon laser) to measure the DNA content. The data was obtained and analyzed with Cell Quest 3.0.1 (Becton Dickinson, Franklin Lakes, NJ) and ModFitLT V2.0 software.

2.7. Western Blot Analysis. Approximately 5 × 10⁶ cells were cultured in 100 mm² dishes and then incubated in various concentrations of K8 for 48 h. The cells were lysed on ice with 150 mL of lysis buffer (50 mmol/L Tris-HCl, pH 7.5, 0.5 mol/L NaCl, 5 mmol/L MgCl₂, 0.5% nonidet P-40, 1 mmol/L phenylmethylsulfonyl fluoride, 1 mg/mL pepstatin, and 50 mg/mL leupeptin) and centrifuged at...
13000×g at 4°C for 15 min. The protein concentrations in the supernatants were quantified using a BSA Protein Assay Kit. Electrophoresis was performed on a NuPAGE Bis-Tris Electrophoresis System using 50 mg of reduced protein extract per lane. Resolved proteins were then transferred to polyvinylidene fluoride (PVDF) membranes. Filters were blocked with 5% nonfat milk overnight and probed with appropriate dilution of primary antibodies for 2 h at room temperature. Membranes were washed three times with 0.1% Tween 100 and incubated with HRP-conjugated secondary antibody for 1 h at room temperature. All proteins were detected using Western Lightning Chemiluminescence Reagent Plus and quantified using a densitometer.

2.8. In Vitro Transfection. To create the notch-expressing A549 stable cell line, A549 cells were stably transfected with an expression vector containing the activated Notch1 coding region in the sense orientation (pEF.hICN1.CMV.GFP) using GeneJammer reagent. After 48 h, the gene expression was measured by fluorescence microscopy and then assessed for overexpression of notch by Western blot analysis.

2.9. Detection of Apoptosis. The viability of the cells after treatment with various chemicals was evaluated using an MTT assay performed in triplicate. Briefly, the cancer cells (5×10^5) were incubated in 96-well plates containing 200 μL of serum-containing medium. Cells were permitted to adhere for 1 h, then washed with phosphate-buffered saline (PBS). Solutions were always prepared fresh by dissolving 0.2% DMSO or drugs in culture medium and added to A549 cells. For inhibitor treatment experiments, cells were treated with 5 mM isochaihulactone and preincubated for 1 h with γ-secretase inhibitor L-685. After 3, 6, 12, and 24 h of exposure, the drug-containing medium was removed, washed with PBS, and replaced by fresh medium. The apoptosis was analyzed according to the method described by Van Engeland et al. to detect the integrity of cellular membrane and the externalization of phosphatidylserine [29]. In brief, approximately 1×10^6 cells were grown in 10 mm diameter plates. The cells were incubated in various concentrations of K8 for the indicated time and then labeled with FITC Annexin V and PI prior to harvested. After labeling, the cells were washed with binding buffer and harvesting by scraping. Cells were resuspended in binding buffer at a concentration of 2×10^5 cells/mL before analysis by flow cytometry (FACScan). The data was analyzed on WinMDI V2.8 software. The percentage of cells undergoing apoptosis was determined by three independent experiments.

2.10. Antitumor Activity In Vivo. Xenograft Mice as a Model system to Study Cytotoxicity Effect of Isochaihulactone in vivo. the implantation of cancer cells was carried out similarly as previous reports. Male congenital athymic BALB/c nude (nu/nu) mice were purchased from National Sciences Council (Taipei, Taiwan), and all procedures were performed in compliance with the standard operating procedures of the Laboratory Animal Center of Ilan University (Ilan, Taiwan).

All experiments were carried out using 6–8-week-old mice weighing 18–22 g. The animals were sc. implanted with 1×10^7 A549 cells into the dorsal subcutaneous tissue. When the tumor reached 80–120 mm^3 in volume, animals were divided randomly into control and test groups consisting of 4 mice per group (day 1). Daily subcutaneous (sc.) administration of Z-K8, dissolved in a vehicle of 10% DMSO in vitamin K (v/v), was performed from days 1 to 5 far from the inoculated tumor sites (>1.5 cm). The control group was treated with vehicle only. The mice were weighed three times a week up to day 37 to monitor the effects, and at the same time, the tumor volume was determined by measurement of the length (L) and width (W) of the tumor. The tumor volume at day n (TVn) was calculated as TV (mm^3) = (L×W^2)/2. The relative tumor volume at day n (RTVn) versus day 1 was expressed according to the following formula: RTVn = TVn/TV1. Tumor regression (T/C (%)) in treated versus control mice was calculated using the following formula: T/C (%) = (mean RTV of treated group)/(mean RTV of control group) × 100. Xenograft tumors as well as other vital organs of treated and control mice were harvested and fixed in 4% formalin, embedded in paraffin, and cut in 4-mm sections for histologic study.

2.11. Immunohistochemical Staining. All tumor tissues (sc. A549 tumors with or without isochaihulactone treatment) were fixed in 4% formalin at 4°C for 16 h and then embedded in paraffin. Paraffin sections were deparaffinized in xylene and rehydrated through a graded series of ethanol solutions power, 1% bovine serum albumin in phosphate-buffered saline for 60 min at room temperature followed by an overnight incubation with a 100-fold dilution of Notch1 or caspase-3 rabbit polyclonal antibody (Cell Signaling Technology) in blocking solution. Subsequently, the immune complexes were visualized using horseradish peroxidase-conjugated anti-goat IgG secondary antibodies (1:1000 dilution; Santa Cruz Biotechnology Inc.) and the LSAB2 system (Dako North America Inc., Carpinteria, CA), respectively, and then incubated for 10 min with 0.5 mg/mL diaminobenzidine and 0.03% (v/v) H2O2 in PBS. Finally, sections were counterstained with hematoxylin, mounted, observed under a light microscope at magnifications of 400x, and photographed.

2.12. Statistical Analysis. The data was shown as mean with standard deviation. The statistical difference was analyzed using the Student’s t-test for normal distributed values and by nonparametric Mann-Whitney U test for values of unnormal distribution. Values of P < 0.05 were considered significant.

3. Results and Discussions

3.1. Z-Isochaihulactone Inhibited Cell Growth in Human Lung Adenocarcinoma. Isochaihulactone is a lignan with a chiral center and two racemic forms (Z-isochaihulactone and E-isochaihulactone), as identified using NMR analysis (Figure 1). Treatment of human lung cancer A549 cells with
isochaihulactone, its Z-form or its E-form, in concentrations ranging from 2.5 μM to 20 μM, for 48 h, revealed that Z-isochaihulactone displayed significantly higher cytotoxicity (IC₅₀ = 5.0 μM) than E-isochaihulactone (IC₅₀ > 20 μM) in A549 cells (P < 0.05), and equal cytotoxicity to isochaihulactone (IC₅₀ = 5 μM) (Table 1). As shown in Table 1, the antitumor effects of Z-isochaihulactone (IC₅₀ = 4.5 μM) on Taxol-resistant A549T-12 cells indicated that the A549T-12 cell line is more sensitive to Z-isochaihulactone than the A549 cell line. Following treatment of A549 cells with different concentrations of Z-isochaihulactone (from 1.25 μM to 20 μM) for 24, 48, and 72 h, MTT assay was used to determine cell viability (Figure 2(a)). Z-isochaihulactone-treated cells exhibited reduced cell viability in time- and dose-dependent manners. The A549 cells were treated with the IC₅₀ of isochaihulactone (determined as 5 μM), or with different concentrations of Z-isochaihulactone (from 2.5 μM to 10 μM), for 48 h, then stained with Hoechst 33342 for 30 min before observation of the nuclear condensation (Figure 2(b)). The association between Z-isochaihulactone-induced growth inhibition and apoptosis was then evaluated. As shown in Figure 2(c), activation of caspase-9 and caspase-3 occurred in a dose-dependent manner following Z-isochaihulactone treatment. Z-isochaihulactone-induced apoptosis thus mediated through a caspase-dependent pathway. Caspase-9 activation, cleavage of caspase-3, and PARP all occurred in a time-dependent manner following Z-isochaihulactone treatment (Figure 2(d)).

3.2. Z-Isochaihulactone-Induced A549 Cell Cycle Arrest in the G2/M Phase. To elucidate Z-isochaihulactone’s mode of action, the present study evaluated its effects on cell cycle progression. Flow cytometric analysis showed that Z-isochaihulactone treatment induced the accumulation of cells in the G2/M phase in a dose-dependent manner (Figure 2(e)). This suggested that Z-isochaihulactone might cause G2/M phase arrest, similar to paclitaxel and vincristine. Z-isochaihulactone also regulated the expression of the G2/M regulatory proteins, including cyclin B1 and survivin, in time- and concentration-dependent manners (Figure 2(f)).
**Table 1: Growth inhibition of K8, E-K8, or Z-K8 against human tumor cell lines.**

| Cell line | Origin                                      | IC<sub>50</sub> (μM) |
|-----------|---------------------------------------------|----------------------|
|           |                                             | K8       | E-K8    | Z-K8    |
| AGS       | Gastric carcinoma                           | ND       | ND      | 2.5     |
| J5        | Hepatocarcinoma                             | 21.5     | ND      | 2.5     |
| LNCaP     | Androgen-dependent, prostate adenocarcinoma | 4.3      | >20     | 1       |
| PC3       | Androgen-independent, prostate adenocarcinoma | >20      | >20     | 4.5     |
| A549      | Lung adenocarcinoma                         | 18.5     | >20     | 5       |
| H460      | Lung adenocarcinoma                         | 20.0     | >20     | 10.0    |
| H1299     | Lung adenocarcinoma                         | 20.5     | >20     | 12.5    |
| A549-T12  | Lung adenocarcinoma                         | 10.0     | >20     | 4.5     |
| OEC-M1    | Oral squamous cell carcinoma                | ND       | >1      | 1       |

IC<sub>50</sub>: drug concentration that inhibits cell division by 50% after 48 h.
ND: not determined, or concentration &gt;100 μM.

**Z-Isocaihulactone Activated Notch Signaling Association Gene Expression in A549 Cells.** Z-isocaihulactone increased Notch1 protein expression in concentration- and time-dependent manners (Figure 3(a)). To investigate the transfer of activated Notch1 (NICD) from the cytoplasm to the nucleus, cells were treated with Z-isocaihulactone in the presence of the γ-secretase inhibitor L-685. The nuclear and cytoplasmic proteins were then collected before separation and analysis of NICD expression. Data showed that a higher Z-isocaihulactone concentration was associated with higher NICD expression in the nucleus (Figure 3(a)). The A549 cells also overexpressed the notch association proteins (the downstream proteins of Notch1, such as c-Myc, NF-κB, and HES-1) following Z-isocaihulactone treatment (Figures 3(b) and 3(c)).

**3.3. Involvement of NICD Activation in Z-Isocaihulactone-Induced Inhibition of A549 Cell Growth.** To evaluate the effects of Z-isocaihulactone treatment on Notch1 expression, A549 cells were stably transfected with an expression vector containing the activated Notch1 coding region in the sense orientation (pEEhICN1.CMV.GFP); the transfection rate was then measured using fluorescence microscopy (Figure 3(d)). To evaluate Notch1 involvement in Z-isocaihulactone-induced apoptosis in A549 cells, cells were treated with Z-isocaihulactone and then measured using fluorescence microscopy. The data showed that Z-isocaihulactone-induced apoptosis was in a dose-dependent manner. Further analysis revealed that L-685 had no effects on the activation of Notch1 (Figure 3(e)).

**3.4. Z-Isocaihulactone Exhibits In Vivo Anticancer Activity.** Subcutaneous (sc.) injection of approximately 1 x 10<sup>7</sup> A549 cells into the backs of nude mice established human lung cancer xenografts to evaluate the antitumor activity of Z-isocaihulactone in vivo. After the tumors had reached approximately 80 mm<sup>3</sup> to 100 mm<sup>3</sup> in size, mice were randomized into vehicle control and treatment groups (5 animals each) and provided a daily sc. injection of DMSO (control group) or 15 mg/kg and 30 mg/kg of Z-isocaihulactone (treatment groups) for 5 days. Results showed that Z-isocaihulactone inhibited tumor growth at low doses and induced tumor regression at higher doses (Figure 3(f)). Further histological examination revealed the NICD expression in vivo (Figure 3(g)). After sacrifice of the animals and lysis of the A549 tumor tissues, Western blot analysis measured the protein levels. The data showed that Z-isocaihulactone increased Notch1 expression and induced activation of caspase-9, caspase-3, and PARP in a dose-dependent manner. Z-isocaihulactone-induced apoptosis in vivo is thus mediated through a caspase-dependent pathway (Figures 3(h) and 3(i)).

**3.5. Discussion.** The group’s previous research demonstrated that isocaihulactone induced tubulin depolymerization and inhibited normal spindle formation in NSCLC cells, resulting in mitotic arrest and cell death [14, 23]. Isochaihulactone contains a γ-butyrolactone-centered heterocyclic group, forming a Z or E configuration, and has a chiral center for each isomer forming (+) and (−) enantiomers. The present study aimed to elucidate the biological effects of two pure isomers of isocaihulactone on human lung cancer A549 cells in vitro and in vivo. It is the first to identify that Z-isocaihulactone displays significantly higher tumor cytotoxicity (IC<sub>50</sub> = 0.96 ± 0.02 μM) than E-isocaihulactone (IC<sub>50</sub> = 6.5 ± 0.02 μM) in A549 cells.

Chen et al. reported that overexpression of NICD inhibited the growth of the lung adenocarcinoma A549 cells in vitro by inducing cell cycle arrest and suppressed A549 tumor growth in nude mice [15]. These findings suggested that notch signaling might have tumor-suppressive function in human lung adenocarcinoma cells. In the present study, treatment of A549 tumor cells with Z-isocaihulactone, followed by Western blot analysis, elucidated the effects of Z-isocaihulactone on NICD expression. As shown in Figure 3(e), Z-isocaihulactone increased NICD protein expression in time- and dose-dependent manners. The group’s previous study confirmed the antitumor effects of isocaihulactone [23]. The present study extends on this research by evaluating inhibition of tumor growth based on NICD protein expression in a nude mice xenograft animal.
**Figure 2: Continued.**
**Figure 2**: Z-isochaihulactone caused growth inhibition by inducing G2/M phase arrest of human lung adenocarcinoma cells *in vitro* and induces apoptosis. (a) Growth inhibition effect was assessed in human lung cancer cell lines A549. The cells were treated with various concentrations ranging from 1.25 to 20 μM Z-isochaihulactone for various times (24, 48, and 72 h) as indicated. Growth inhibition effect (IC₅₀) was determined by MTT assay. (b) Cells were treated with 10 μM Z-isochaihulactone for 48 h then stained with Hoechst 33342 for 30 min. The chromatin morphology of A549 cells was measured by fluorescence microscopy (magnification 400x). (c) The Western blot analysis was done and as described in Section 2. Cells were probed with Z-isochaihulactone with various concentrations ranging from 1.25–20 μM for 24 h. Cells were probed with Cleaved caspase-9, caspase-9, Cleaved caspase-3, caspase-3, Cleaved PARP, and PARP antibodies. Expression of β-actin was used as an internal control. (d) Cells were treated with 5 μM Z-isochaihulactone for the indicated time (6–48 h) and analyzed by Western blot analysis. Cells were probed with Cleaved caspase-9, caspase-9, Cleaved caspase-3, caspase-3, Cleaved PARP, and PARP antibodies. Expression of β-actin was used as an internal control. (e) The cell cycle analysis was done as described in Section 2. The cell cycle was assessed in A549 cells. Cells were treated with various concentrations from 2.5 to 10 μM Z-isochaihulactone and VPA 10 mM for 48 h. Cells were collected, fixed in alcohol, then stained with propidium iodide (PI), and analyzed by flow cytometry. (f) Cells were treated with various concentrations of Z-isochaihulactone from 1.25 to 20 μM and VPA 10 mM for 48 h. The cell cycle association proteins were determined by Western blot analysis. The Western blot analysis was done and as described in Section 2. Cells were probed with cyclin B1 and phosphor-cdc2 (Thr161) antibodies. Expression of β-actin was used as an internal control.

Although the tumor volume of mice treated with Z-isochaihulactone was not regressed as those treated with racemic compound [26], Z-isochaihulactone did have potential in tumor inhibition, when compared with the control group (Figure 4(a)). Whether E- and Z-isochaihulactone have synergistic effect in tumor inhibition, it needs to be further investigated. The obtained data indicate that the expression of Notch1 is an essential factor in the antitumor activity of Z-isochaihulactone. Experiments using the γ-secretase inhibitor L-685 supported the notion that NICD expression induces apoptosis-promoting effects by silencing NICD expression (Figure 3(e)). Results indicated that the upregulation of NICD by Z-isochaihulactone is sequenced by upregulation of c-Myc and HES-1 protein expression (Figures 3(b) and 3(c)). The effects of Z-isochaihulactone on upregulation of NICD expression suggest that this tumor suppressor protein mediates the antitumor effects of Z-isochaihulactone. The c-Myc and HES-1 appear to be key downstream targets of NICD.

Results from analyses of Notch1 receptor expression in A549 cells following treatment with Z-isochaihulactone showed that Z-isochaihulactone induced morphological changes in A549 cells (Figure 2(b)), inhibited A549 cell growth (Figure 2(a)), and arrested the cell cycle at the G2/M phase (Figure 2(e)). The tumor suppressor protein p53 plays a role in the molecular response to DNA damage. Acting as a DNA-binding transcription factor, it regulates specific target genes to arrest the cell cycle and initiate apoptosis. Following DNA damage, the cyclin-dependent kinase inhibitor p21 is expressed in a p53-dependent or p53-independent manner [30]. The p21 might assist with maintenance of G2 cell cycle arrest by inactivating the cyclin B1/cdc2 complex, disrupting the interaction between the proliferating cell nuclear antigen and cdc25C [31]. In the present study, treatment with Z-isochaihulactone-induced increased p53 and p21 expression in A549 cells (Figure 2(f)). It is well known that the cdc25-mediated activation of the cyclin B1/cdc2 complex triggers the transition from the G2 phase to mitosis, and that dephosphorylation of Thr15 by the phosphatase cdc25 induces cyclin B1/cdc2 activation [32, 33]. The present study's findings indicate that Z-isochaihulactone mediated A549 cell arrest at the G2/M phase (Figure 2(e)), with cell arrest accompanied by decreased cyclin B1 expression. Z-isochaihulactone had no influence on survivin expression. Results therefore suggest that Z-isochaihulactone treatment causes inappropriate accumulation of G2/M regulators leading to apoptosis.

Z-isochaihulactone induces apoptosis in A549 cells following mitotic arrest. The mechanism by which microtubule-damaging agents induce apoptosis is not well understood. The activation of aspartate-specific cysteine protease (caspase) represents a crucial step in the induction of drug-induced apoptosis, with cleavage of poly (ADP-ribose) polymerase (PARP) by caspase-3 considered to be one of the hallmarks of apoptosis [34]. In the present study, Z-isochaihulactone induced cleavage of caspase-9 and...
caspase-3 and activation of PARP (Figure 2(c)). Evaluation of the association between Z-isochaihulactone-induced cell death and Bcl-2 phosphorylation indicated that Z-isochaihulactone treatment activated caspase-9 and caspase-3 (Figure 2(c)). Z-isochaihulactone-induced cell death is thus mediated through a caspase-dependent pathway. Z-isochaihulactone induced caspase-9 activation, Bcl-2 phosphorylation, and cleavage of caspase-3 and PARP in a time-dependent manner (Figure 2(d)).

4. Conclusions

In summary, Z-isochaihulactone is a novel lignan compound which causes inappropriate expression of cyclin B1/cdc2 kinase G2/M regulatory proteins and initiates the apoptotic cascade. The present study’s findings indicate that Z-isochaihulactone regulates Notch1 expression and demonstrate that Notch1 has antitumorigenic and proapoptotic activities in vitro and in vivo. Furthermore, mice treated with Z-isochaihulactone had no obvious body weight alternation and histopathological change in liver; H&E staining analysis (data not shown) suggested that Z-isochaihulactone had no critical adverse effects. Although more detailed analyses are needed to clarify the mechanisms of the antitumor effects of Z-isochaihulactone, these results should encourage further investigation of a potential role for Z-isochaihulactone in future as a potential novel clinical anticancer drug.

Conflict of Interests

The authors declare that they have no conflict of interests.

Authors’ Contribution

D. Chan and Y. S. Chen contributed equally to this work.
Figure 4: Z-isochaihulactone induced in vivo growth inhibition of A549 xenografts in nude mice. Inhibition of tumor growth from A549 cells for upregulation of Notch1 protein. Nude mice injected with approximately $1 \times 10^7$ A549 cells into the dorsal subcutaneous tissue. (a) When the tumor volumes reached 80–120 mm$^3$, A549 tumor-bearing mice were administered sc. with vehicle control (●), 15 mg/kg Z-isochaihulactone (■), and 30 mg/kg Z-isochaihulactone (▲) on days 1–5 for 5 days. These figures show A549 cells relative tumor volume of control and therapeutic groups. (b) Immunohistochemical staining was analyzed in A549 tumor tissues (at day 6 after initiation of drug treatment). Representative photographs of sections of the control group and Z-K8-treated group. A549 tumors, immunohistochemically stained with NICD and Cleaved caspase-3 rabbit monoclonal antibody, and NICD and Cleaved caspase-3 positive cells were stained brown (black arrowheads) (×400). [Scale bars, 100 μm. n = 4 different experiments.] Means ± SD. $b P < 0.05$, $c P < 0.01$ versus the control. (d) Western blot was analyzed in A549 tumor tissues (at day 34 after initiation of drug treatment). Tumor lysate was probed with Cleaved caspase-9, caspase-9, Cleaved caspase-3, caspase-3, Cleaved PARP, PARP, and NICD antibodies. Expression of β-actin was used as an internal control.

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