

Delicaflavone Reverses Cisplatin Resistance via Endoplasmic Reticulum Stress Signaling Pathway in Non-Small Cell Lung Cancer Cells

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Yuxia Sui Department of Pharmacy, Fujian Provincial Hospital, No. 134 Dong Street, Fuzhou 350001, People's Republic of China Tel/Fax +86 15959090652 Email suiyuxia123@163.com **Background:** The incidence and mortality of lung cancer continue to increase around the world; in 2018, new lung cancer cases accounted for 11.6% of all cancer cases, and lung cancer deaths accounted for 18.4% of cancer deaths. Cisplatin (DDP) is a first-line chemotherapy drug for lung cancer; however, DDP resistance can lead to a poor prognosis in patients with lung cancer. Therefore, reversing DDP resistance is a treatment goal.

Materials and Methods: Cell counting kit-8 (CCK8) assays, wound healing analyses, Transwell assays, in vitro tumor xenografts, and flow cytometry were used to detect the proliferation, migration, invasion, and apoptosis of multidrug resistant A549/DDP and PC9/DDP cells, respectively. Western blot was performed to detect protein levels of cleaved caspase-3, CHOP, and GRP78.

Results: Delicaflavone inhibited DDP resistance of lung cancer cells and decreased proliferation in a dose- and time-dependent manner. It also decreased migration and invasion and enhanced apoptosis. Western blots showed that delicaflavone overcame DDP resistance by increasing the expression of GRP78 and CHOP and the apoptosis-related protein cleaved caspase-3.

Conclusion: Delicaflavone can reverse DDP resistance in A549/DDP and PC9/DDP cells by inhibiting cell proliferation and migration and enhancing apoptosis and cleaved caspase-3 levels by increasing the expression of CHOP and GRP78 protein via the endoplasmic reticular stress pathway. It could be a useful therapeutic adjunct to treat DDP-resistant lung cancer.

Keywords: delicaflavone, Cisplatin; DDP, lung cancer, endoplasmic reticulum stress

Introduction

Lung cancer incidence and mortality continue to increase worldwide; in 2018, new lung cancer cases accounted for 11.6% of all cancer cases, and lung cancer deaths accounted for 18.4% of all cancer deaths. Because early clinical manifestations of lung cancer are not typical, early detection and treatment are limited. Chemotherapy is the primary treatment for middle- and late-stage lung cancer. Cisplatin (DDP) is a chemotherapeutic drug widely used in the treatment of lung cancer, but drug resistance of tumors had affected its application. Exploring ways to reverse DDP drug resistance will improve the treatment of lung cancer.

Delicaflavone is a biflavonoid compound existing in the Selaginella doederleinii Hieron and other herbals or plants. It has good antitumor activity and excellent safety profiles. It is worth further research and development. Previous studies showed that Delicaflavone could induced apoptosis, autophagy, cell cycle arrest and inhibits PI3K/AKT/mTOR, Ras/ MEK/Erk, Akt/mTOR/p70S6K signaling pathways and MAPK signaling cascades in colorectal cancer cell, lung cancer and cervical cancer HeLa cells. 7-9 However, it has not been reported whether delicaflavone can reverse DDP resistance in the human lung adenocarcinoma multidrug resistant A549/DDP and PC9/DDP cell line. The aims of this study is to investigate DDP resistance reversal by delicaflavone in A549/DDP and PC9/DDP cells and explore its possible molecular mechanism.

Our results confirmed that delicaflavone could attenuate DDP resistance in A549/DDP and PC9/DDP cells by inhibiting cell proliferation and migration and enhancing apoptosis and cleaved caspase-3 levels by increasing the expression of CHOP and GRP78 protein via the endoplasmic reticular stress pathway. It could be a useful therapeutic adjunct to treat DDP-resistant lung cancer.

Materials and Methods

Cell Culture

A549, A549/DDP, PC9, and PC9/DDP were purchased from the Cell Collection Committee of the Chinese Academy of Sciences (Shanghai, China) and the cells were cultured with RPMll640+10% FBS at 37°C, 5%CO₂.

CCK8 Analysis

5×10³/well A549, A549/DDP, PC9, and PC9/DDP cells were inoculated into 96 well plates for 12 h, then, the different concentrations of delicaflavone (5, 10, 20, 40 and 80 µM) or DDP (2, 4, 8, 16 and 32 µM) were added. In the meantime, A549/DDP cells were cultured with delicaflavone (20 µM) or DDP (16 µM) or delicaflavone (20 µM)+DDP (16 µM). The control group was added with an equal volume of PBS, and each group was repeated 3 times. 22 h after, incubation with 10µL CCK-8 (Sigma-Aldrich, USA) solution for 2 h, The OD value at the wavelength of 450 nm was measured by microplate assay (EnSpire 2300, PerkinElmer, USA), and the cell growth curve was measured according to the OD value.

Wound Healing

A549/DDP and PC9/DDP cells density were adjusted to make 2×10⁶/mL cell suspension absord 0.2 mL and inoculated into a 6-well plate. Twenty-four hours later, delicaflavone (20 μ M) or DDP (16 μ M) or delicaflavone (20 μ M) +DDP (16 µM) were added into culture bottle for 24 h. The

control group was added with an equal volume of PBS, and each group was repeated 3 times. We observed the width of the scratch under the inverted microscope and take photos.

Transwell Assay

A549/DDP, PC9/DDP cells cultured with were Delicaflavone (20 µM) or DDP (16 μM) Delicaflavone (20 µM)+DDP (16 µM) for 24 h. The control group was added with an equal volume of PBS, and each group was repeated 3 times. 4×10⁴/well A549/DDP, PC9/DDP cells were inoculated onto the upper chamber with 2% FBS and 500 mL medium containing 10% FBS was added to the lower chamber. Incubated for 24 h, removal of the upper chamber of cells, the cells were fixed with 4% paraformaldehyde, stain with crystal violet, washed, dried and counted.

In vitro Tumor Xenografts

A549/DDP, PC9/DDP cells density were adjusted to make 2×10⁷/mL cell suspension absord 0.2 mL and injected into the back of mice. 10d later, delicaflavone (20 µM) or DDP (16 μM) or delicaflavone (20 μM)+DDP (16 μM) peritoneal injection twice a week. The control group was added with an equal volume of normal saline. Observed continuously for 3 weeks and the tumor size was measured by scaler every week, and each group was repeated 3 times. The laboratory animals were approved by the medical laboratory animal ethics committee of Fujian Medical University. Instructive notions with respect to caring for laboratory animals (which is released by the Ministry of Science and Technology of the People's Republic of China in September 30th, 2006) were followed for the welfare of the animals.

Flow Cytometry Assay

A549/DDP, PC9/DDP cells were cultured with delicaflavone (20 μM) or DDP (16 μM) or delicaflavone (20 μM) +DDP (16 µM) for 24 h. The control group was added with an equal volume of PBS, and each group was repeated 3 times. A549/DDP, PC9/DDP cells were digested with trypsin without ED-TA and cell suspension was prepared, then 800r/min centrifugation for 10min at 4° C. 5µL Annexin V-FITC and 5µL propidium iodide (PI) (Life technologies, USA) were added to mix. The apoptosis rate was detected by flow cytometry (BD, USA).

Western Blot Assay

Extraction of total protein and BCA quantitative method was used to detect the protein concentration (Thermo Scientific,

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USA). Then the antibody was added to incubate, cleaved-caspase-3, CHOP, GRP78, β-actin (Cell Signaling Technology, USA), ECL reagent (Amersham Biosciences) chemiluminescence, development and Image J was used for analysis, and each group was repeated 3 times.

Statistical Analysis

SPSS 20.0 statistical software was used to analyze test data. The measurement data is expressed as mean \pm standard ($\ddot{y}x\pm s$), T-test was used for comparison between the two groups, and ANOVA was used for comparison between multiple groups. P<0.05 was considered statistically significant (*P < 0.05 represents compared with control group. $^{\#}P$ < 0.05 represents compared with the Delicaflavone-alone group or DDP-alone group.)

Results

Delicaflavone United with DDP Regulated the Cell Viability of Lung Cancer DDP Resistant Cells

In this study, we firstly examined the cell viability of A549, PC9, A549/DDP, and PC9/DDP by CCK8 assay. The result showed that the cell proliferation inhibitory rate of DDP on A549 and PC9 cells was higher than that of A549/DDP and PC9/DDP cells in a dose-dependent manner (Figure 1A and B), the half maximal inhibitory concentration (IC50) of A549 was 16 μ M and of A549/DDP cells was 48 μ M by DDP. A549/ DDP exhibited more significant DDP resistance than A549. In order to investigate the cell proliferation inhibitory effect of delicaflavone on A549/DDP and PC9/DDP cells, CCK8 assay showed delicaflavone can inhibited A549/DDP and PC9/DDP cells proliferation in a dose-dependent manner and the nonlethal concentration of A549/DDP and PC9/DDP were 20 µM delicaflavone (Figure 1C and D). Further, the results revealed that 20 µM delicaflavone combined with 16 µM DDP had a high cooperative proliferation inhibition rate (Figure 1E and F). The cell viability of A549/DDP and PC9/DDP cells were detected by CCK8 assay at the concentrations of delicaflavone $(20 \mu M)$ or DDP $(16 \mu M)$ or delicaflavone $(20 \mu M)$ +DDP $(16 \mu M)$ μM) for 24h, 48h, 72 h. The results indicated that compare with the delicaflavone united with DDP group, the cell viability of A549/DDP and PC9/DDP cells were significantly lower than that of the delicaflavone-alone group or DDP-alone group (Figure 1G and H), revealing that delicaflavone united with DDP can significantly inhibit the cell viability.

Delicaflavone United with DDP Regulated Lung Cancer DDP Resistant Cells' Invasion, Migration

In order to evaluated whether delicaflavone affects the migration and invasion effect of A549/DDP and PC9/DDP cells. Wound healing showed compared with the control group, the cell migration rate of A549/DDP and PC9/DDP cells were decreased in delicaflavone united with DDP group, delicaflavone-alone group and DDP-alone group, compare with the delicaflavone united with DDP group, the migration ability of A549/DDP and PC9/DDP cells were significantly reduced than that of the delicaflavone-alone group or DDP-alone group (Figure 2A–D).

Transwell assay showed compared with the control group, the migration number of cells were decreased in delicaflavone united with DDP group, compared with the delicaflavone-alone group or DDP-alone group, the migration number of cell were decreased was significantly reduced in the delicaflavone united with DDP group (Figure 2E–H). In vitro tumor Xenografts, the results of tumor volume was similar reduced in the delicaflavone united with DDP group (Figure 2I–L). Dates revealed that delicaflavone reversed A549/DDP cells invasion, migration in vivo and in vitro experiments.

Delicaflavone United with DDP Regulated Lung Cancer DDP Resistant Cells Apoptosis and the Expression of Apoptosis-Related Proteins

To assess whether delicaflavone could affect A549/DDP and PC9/DDP cells apoptosis. Flow cytometry assay was used to evaluate A549/DDP and PC9/DDP cells apoptosis by delicaflavone (20 μ M) or DDP (16 μ M) or delicaflavone (20 μ M)+DDP (16 μ M) for 24 h. Flow cytometry assay showed compared with the control group, the apoptosis rate of A549/DDP and PC9/DDP cells were increased in delicaflavone united with DDP group, compared with the delicaflavone-alone group or DDP-alone group. The apoptosis rate was increased in delicaflavone united with DDP group, compared with the delicaflavone-alone group (Figure 3A–D). Subsequently, Western blot revealed that protein expression level of cleaved-caspase-3 was a similar result in A549/DDP and PC9/DDP cells (Figure 3E–H).

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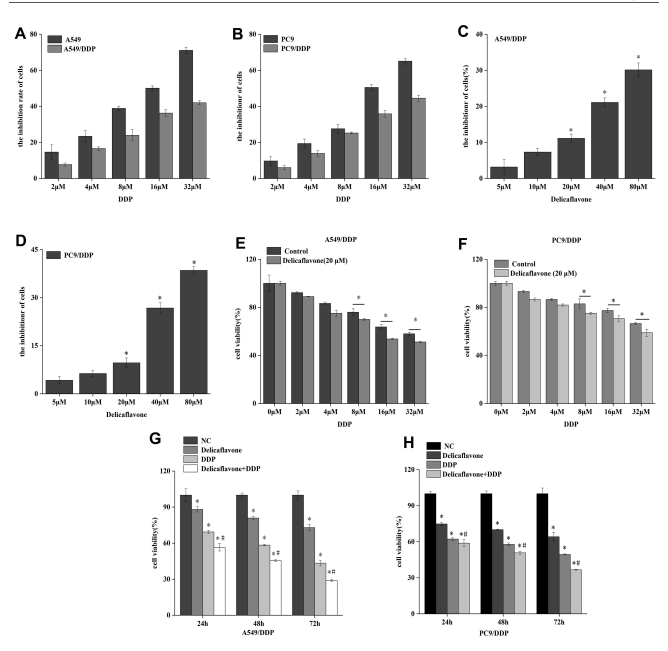


Figure 1 Delicaflavone united with DDP could reduced the cell viability of lung cancer DDP resistant cells. (A) the cell viability of A549 and A549/DDP cells were detected with the concentrations of DDP for 24h. (B) the cell viability of PC9 and PC9/DDP cells were detected with the concentrations of DDP for 24h. (C) the cell viability of A549 and A549/DDP cells were detected with the concentrations of Delicaflavone for 24 h. (D) the cell viability of PC9 and PC9/DDP cells were detected with the concentrations of Delicaflavone for 24 h. (E) A549/DDP cells were treated with 20 µM Delicaflavone combined with indicated concentrations of DDP for 24 h. (F) PC9/DDP cells were treated with 20 µM Delicaflavone combined with indicated concentrations of DDP for 24 h. (G) the cell viability of A549/DDP cells were detected by CCK8 at the concentrations of Delicaflavone (20 µM) or DDP (16 µM) or Delicaflavone (20 µM)+DDP (16 µM) for 24h, 48h, 72 h. (H) the cell viability of PC9/DDP cells were detected by CCK8 at the concentrations of Delicaflavone (20 µM) or DDP (16 µM) or Delicaflavone (20 µM)+DDP (16 µM) for 24h, 48h, 72 h. Data represent mean ± SD. *P < 0.05. $^{\#}P < 0.05$ represents compared with the Delicaflavone alone group or DDP alone group.

Delicaflavone United with DDP Regulated Lung Cancer DDP Resistant Cells Invasion, Migration, Apoptosis Through **Endoplasmic Reticulum Stress Signaling Pathway**

To explore whether Delicaflavone affect A549/DDP and PC9/DDP cells invasion, migration, apoptosis by ER stress signaling pathway. The protein expression of ER stress-related protein CHOP and GRP78 were detected by Western blot. The protein expression of GRP78 and CHOP were increased in DDP group compared with the control group, and also increased in delicaflavone united with DDP group, compared with the Delicaflavone-alone group or DDP-alone group (Figure 4A-D).

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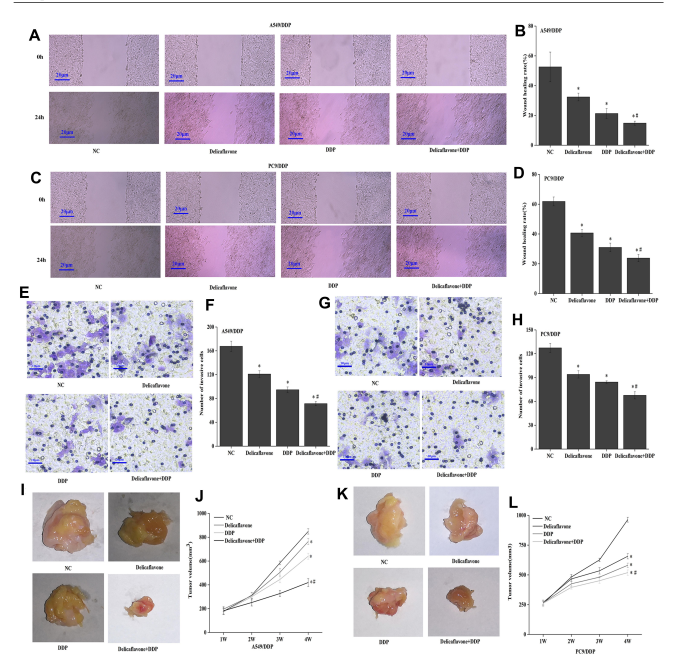


Figure 2 Delicaflavone united with DDP regulated lung cancer DDP resistant cells invasion, migration. (**A–D**) Wound healing showed the migration ability of the Delicaflavone united with DDP group was significantly reduced than that of the Delicaflavone alone group or DDP alone group. (**E–H**) Transwell assay showed the migration number of cell were decreased was significantly reduced in the Delicaflavone united with DDP group. (**I–L**) In vitro tumor Xenografts, the results of tumor volume was reduced in the Delicaflavone united with DDP group. Data represent mean \pm SD. *P < 0.05 represents compared with the Delicaflavone alone group or DDP alone group.

Discussion

Lung cancer has the characteristics of high morbidity and mortality, and non-small cell lung cancer accounts for 85% of lung cancer. ¹⁰ Chemotherapy is a strategy for the treatment of lung cancer. DDP was a common chemotherapy drug for cancers and plays an anti-tumor role by inducing apoptosis, autophagy and necrosis. ^{11–13} However, the drug resistance of DDP in tumor was limited the clinical

application. The resistance of lung cancer cells to DDP is a complex process, involving multiple factors and genes, including the mechanisms of the accumulation and deactivation of intracellular drugs, DNA damage repair, autophagy, and apoptosis. 14–17

Recently more and more research has focused on the mechanism of Delicaflavone in tumors. It exerts antitumor activity by inducing apoptosis and autophagy and

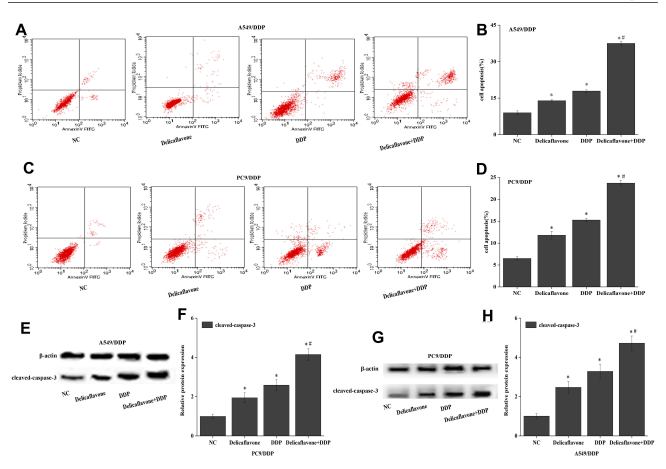


Figure 3 Delicaflavone united with DDP regulated lung cancer DDP resistant cells apoptosis and the expression of apoptosis related proteins. (A-D) Flow cytometry showed that NC, Delicaflavone, DDP, Delicaflavone united with DDP could regulated lung cancer DDP resistant cells apoptosis. (E-H) The protein expression of cleavedcaspase-3 was detected by Western blot. *P < 0.05 represents compared with control group. #P < 0.05 represents compared with the Delicaflavone-alone group or DDPalone group.

inhibiting proliferation in in vitro tumor xenografts of colorectal and cervical cancer cells.⁷⁻⁹ However, the research focus on reversing drug resistance was lacked. In this study, Dates revealed that the cell viability of A549/ DDP and PC9/DDP in Delicaflavone united with the DDP group was significantly lower than that of the delicaflavone-alone group or DDP-alone group. Wound healing, Transwell assay, and in vitro tumor xenografts also showed that delicaflavone decreased migration and invasion in vivo and in vitro experiments.

Endoplasmic reticulum (ER) stress is an important cellular protective mechanism. In some tumors, ER stress activates a series of compensatory mechanisms, and discovered that Hypoxia, oxidative stress and drugs can induce ER stress and cleaved-caspase-3 can promotes apoptosis by promotion the expression of GRP78 and CHOP protein in some tumors. 18-21 Recent research suggests that ER stress may be related molecular mechanism of tumor drug resistance. 22-25 In our study, Delicaflavone could enhance A549/DDP and PC9/DDP cells apoptosis by Flow cytometry assay, Western blots showed that delicaflavone overcame DDP resistance by increasing the expression of GRP78 and CHOP and the apoptosisrelated protein cleaved caspase-3.

Our results proved that Delicaflavone could attenuate DDP resistance in A549/DDP and PC9/DDP cells by inhibiting cell proliferation, migration, and enhancing apoptosis and cleaved-caspase-3 protein by increasing the expression of GRP78 and CHOP via the ER stress pathway. This study provided a theoretical foundation for the combination of Delicaflavone and chemotherapy drugs in the treatment of lung cancer.

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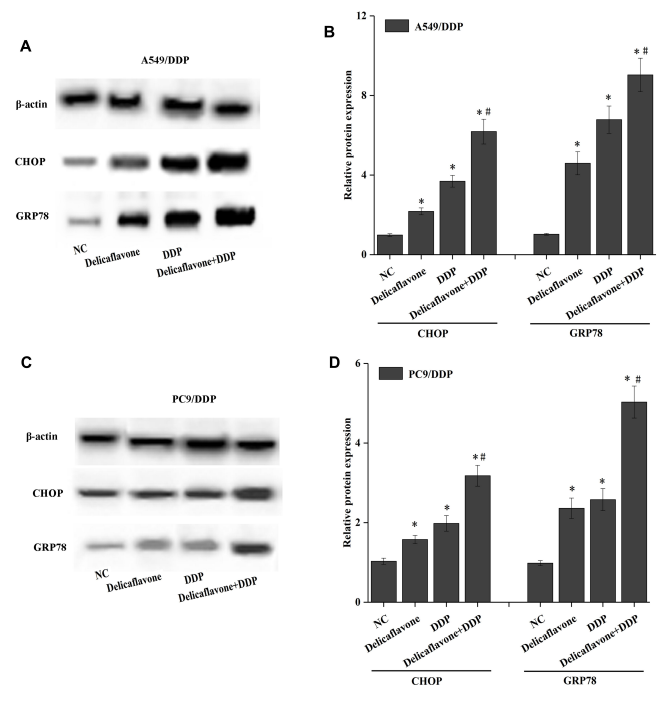


Figure 4 Delicaflavone united with DDP regulated lung cancer DDP resistant cells invasion, migration, apoptosis through ER stress signaling pathway. (A-D) The protein expression of ER stress-related protein CHOP and GRP78 were detected by Western blot. *P < 0.05 represents compared with control group. *P < 0.05 represents compared with the Delicaflavone-alone group or DDP-alone group.

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Disclosure

The authors report no conflicts of interest for this work.

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