Abstract. The function of sodium cantharidinate on inducing hepatocellular carcinoma cell apoptosis was investigated for the first time. Sodium cantharidinate inhibits HepG2 cell growth mainly by LC3 autophagy pathway. MTT results show that sodium cantharidinate effectively inhibits the proliferation of HepG2 cells in a dose- and time-dependent manner and induce cell apoptosis by caspase-3 activity. The further western blotting and FACS detection show that sodium cantharidinate initiates HepG2 cell autophagy program by LC3 pathway. Autophagy-specific inhibitor 3-MA reduce sodium cantharidinate-induced caspase-3 activity and HepG2 cell apoptosis. Silence of the LC3 gene in HepG2 cell lines also reduce sodium cantharidinate-induced cell apoptosis. Collectively, our data indicate that sodium cantharidinate induces HepG2 cell apoptosis through LC3 autophagy pathway. Sodium cantharidinate has potential for development as a new drug for treatment of human HCC.

Introduction

The incidence of hepatocellular carcinoma (HCC) is becoming the second leading cause of cancer-related death worldwide, which accounting for approximately 800,000 deaths every year (1). Hepatic resection and liver transplantation have progressed in surgical procedures for HCC. The improved outcomes are limited because of the frequent recurrence, even after liver transplantation (2-5). Thus, it is urgent to develop novel approaches for hepatocarcinoma prevention and treatment. At present, chemotherapy is also a focus for tumor treatment (6). Sorafenib, the molecular targeting agent, was reported to improve survival rates and outcomes in patients with non-resectable or early stage HCC (7,8). However, sorafenib is the only approved molecular targeted treatment for advanced HCC. Other targeted agents are under investigation. Trials comparing new agents in combination with sorafenib are ongoing. Combinations of systemic targeted therapies with local treatments are being evaluated for further improvement in HCC patient outcomes (9-11).

In recent years, increased data concerning the traditional Chinese medicine with a remarkable activity on the influence with tumor cell death pathway could guide tumor treatment decisions and clinical management (12). Cantharidin, also together with its acid form cantharidinate, was first extracted from Chinese blister beetle, have been used in traditional Chinese medicine for many years (13,14). Sodium cantharidinate has powerful antitumor activity proved in clinical practices in recent years (15). The compound directly inhibits multiple malignant tumors, including myeloma, oral buccal carcinoma, leukemia, gastric cancer, Colo205 CRC, and has low toxic/adverse effects so far (16). In recent years, researchers have confirmed through in vitro experiments that sodium cantharidinate and its derivatives directly kill liver cancer cell lines (17).

Autophagy is the natural, destructive cellular mechanism that degrades damaged proteins and cytoplasm components in lysosomes and thus maintains cellular homeostasis and supplies substrates for energy generation. It is a critical pathway for homeostasis, development and other pathophysiological processes (18). Autophagy plays an important role in the healthy and diseased liver (19,20). Autophagy is considered as an important cellular metabolic process (21). Its function on cell fate is double-edged, which can promote cell survival, therefore may also promote cell death via different mechanisms (22). Autophagy plays different roles depending on the drug, cell type or time of drug action, and the mechanism is not fully understood (23,24). Therefore, the study of the dual role of autophagy may provide new clues for tumor treatment.
In the present study, we investigated whether sodium cantharidinate induces the HepG2 cell line apoptosis and whether it depends on the autophagy pathway.

**Materials and methods**

**Reagents.** Sodium cantharidinate, Hoechst 33258, MTT and RNase were purchased from Sigma (St. Louis, MO, USA). Z-DEVD-FMK (HY-12466, Caspase-3 inhibitor) was purchased from MedChem Express (Monmouth Junction, NJ, USA). Propidium iodide and Annexin V-FITC Apoptosis Detection kit was purchased from BD Biosciences (San Jose, CA, USA). LC3 siRNA reagent kit was purchased from Cell Signaling Technology (Danvers, MA, USA). Dulbecco’s modified Eagle’s medium (DMEM), t trypsin, fetal bovine serum (FBS), PBS, penicillin and streptomycin were obtained from Gibco BRL Life Technologies (Grand Island, NY, USA).

**Preparation of sodium cantharidinate.** Sodium cantharidinate was dissolved in PBS (pH 7.2) to prepare a stock solution at a concentration of 1.0 mM which was stored at -20°C. DMEM complete medium was added to dilute the sodium cantharidinate to the appropriate concentrations prior to use.

**HepG2 cell culture and treatment.** HepG2 cells were routinely cultured in DMEM complete medium which contains 50 U/ml antibiotics and 10% fetal bovine serum (FBS) under the conditions of 5% CO₂ at 37°C in cell incubator (HERAcell 150; Thermo Fisher Scientific, Waltham, MA, USA). Following trypsinization to passage the cells in 75 T flask 3-5 days, the cells were counted and reseed in 96-well plate in DMEM complete medium without or with sodium cantharidinate for MTT array or apoptosis detection.

**The effect of sodium cantharidinate on HepG2 proliferation.** The inhibitory effect of sodium cantharidinate on the proliferation of HepG2 cells were detected via MTT assays. All experiment steps were performed following the instructions of the kit. Briefly, the cells were seeded on 96-well plates at a density of 5x10⁴/ml at a volume of 200 µl per well. All groups without or with sodium cantharidinate (0, 2.0, 5.0, 12.5 µM) were incubated 6-24 h. MTT (1.0 mg/ml) was added to each well, and the cells were incubated for 4 h. The MTT solution was then aspirated, and 100 µl DMSO was added. The 96-well plates were read using a microplate spectrophotometer (Synergy H1, BioTek, Winoozki, VT, USA) at 540 nm. The experiments were repeated in triplicate. The inhibition percentage was calculated as (1 - the value in experimental group / the value in the control group) x100%.

**FCM for cell apoptosis.** Annexin V-FITC and PI double staining flow cytometry analyses were employed. The HepG2 cells were plated in 96-well plates containing 200 µl medium at a density of 5x10⁴ cells/well. The induction of apoptosis in the HepG2 cells were examined without or with sodium cantharidinate (5.0 µM). After culture, the cells were collected in 1.5 ml centrifuge tubes, washed three times with cold PBS and binding buffer, and then stained with Annexin V-FITC and PI (Annexin V-FITC Apoptosis Detection kit; BD Biosciences) for apoptosis detection. Briefly, HepG2 cells in centrifuge tubes were re-suspended in binding buffer. Then, 5 µl of FITC-Annexin V was added to the tubes, which were incubated for 10 min followed by the addition of 5 µl PI. The samples were then incubated with PI for another 15 min and immediately analyzed using a flow cytometer (FACScan; BD Biosciences) with the Flowjo FACS analysis software. The cells in the different portions represented the different cell states as follows: the late-apoptotic cells were present in the upper right portion, the viable cells were present in the lower left portion, and the early apoptotic cells were present in the lower right portion.

**Western blotting.** HepG2 cell lysates were separated by SDS-PAGE under non-reducing conditions on a 10% polyacrylamide gel. The proteins were then transferred onto PVDF membranes (GE Healthcare Life Sciences, Piscataway, NJ, USA) by electroblotting. The membranes were blocked with blocking buffer overnight at 4°C and then incubated with the caspase-3, cleaved caspase-3, LC3-I and LC3-II antibodies for 1.5 h at room temperature. The membranes were then washed by washing buffer six times and incubated with HRP-conjugated secondary antibodies for 1 h. After washing, protein bands were visualized using an enhanced chemiluminescent system (Thermo Fisher Scientific). The primary antibodies used (caspase-3, cleaved caspase-3, LC3-I, LC3-II and β-actin) were all obtained from Cell Signaling Technology.

**Indirect immunofluorescence staining and confocal laser microscopy.** Hoechst 33258 (Sigma) were used to assessed apoptotic nuclear changes. After treatment with 5.0 µM sodium cantharidinate for 0, 6, 12, and 24 h, cells were fixed with 4% paraformaldehyde, stained with Hoechst 33258 (2 µg/ml) for 30 min, washed in PBS, and examined using Olympus FV1000 confocal laser microscopy to reveal cell chromatin condensation.Briefly, HepG2 cells were cultured on coverslips overnight, then treated with 5.0 µM sodium cantharidinate for 6 h and rinsed with PBS at least three times. Cells were fixed for 20 min with 4% paraformaldehyde after incubation, then permeabilized with 0.1% Triton X-100 for 5 min, finally blocked with bovine serum albumin. Then incubated with primary antibodies against LC3 (1:50 dilution) overnight at 4°C, then in FITC/Rhodamine Red-conjugated secondary antibodies (1:400 dilution) (all antibodies, Santa Cruz Biotechnology, CA, USA) by electroblotting. The membranes were blocked with blocking buffer overnight at 4˚C and then incubated with the caspase-3, cleaved caspase-3, LC3-I and LC3-II antibodies for 1.5 h at room temperature. The membranes were then washed by washing buffer six times and incubated with HRP-conjugated secondary antibodies for 1 h. After washing, protein bands were visualized using an enhanced chemiluminescent system (Thermo Fisher Scientific). The primary antibodies used (caspase-3, cleaved caspase-3, LC3-I, LC3-II and β-actin) were all obtained from Cell Signaling Technology.

**Statistical analysis.** All data were analyzed and assessed for significance using the Pearson omnibus normality test. All data are presented as the mean ± the standard error of the mean. Mean values were compared using paired t-tests (two groups) followed by the Bonferroni correction for multiple comparison tests. p-values <0.05 were considered to indicate a statistically significant result. All statistical tests were performed with prism software (GraphPad, San Diego, CA, USA).

**Results**

**Sodium cantharidinate induces apoptosis in HepG2 cells by caspase-3 activity.** HepG2 cells were treated with different
doses of sodium cantharidinate for different time intervals, and MTT results showed that sodium cantharidinate effectively inhibited the proliferation of HepG2 cells in a dose- and time-dependent manner (Fig. 1A). We then selected 5.0 µM sodium cantharidinate for treatment of HepG2 cells at different time intervals, and the apoptotic effector caspase-3 was detected by western blotting. The results showed that the expression of cleaved caspase-3 was increased at all three time points (4-fold change, p<0.01, n=3), (D) Cells were stained with Hoechst 33258. Cell morphology was observed by laser scanning confocal microscopy. (E and F) Flow cytometric analysis of apoptosis in HepG2 cells treated with sodium cantharidinate. The cells were exposed to either control solution (0.1% DMSO in medium) or sodium cantharidinate at 5.0 µM and incubated for 6-24 h. (G) A caspase-3 inhibitor, Z-DEVD-FMK (100 µM) was added to the well, or not. Cells were treated by sodium cantharidinate for 6 h, then apoptosis was determined by FACS. I-Cas3, caspase-3 inhibitor; SC, sodium cantharidinate. The experiments were repeated at least three times. The data are expressed as the means ± SD of three experiments (**p<0.01 vs. control).

Figure 1. Sodium cantharidinate induces apoptosis of HepG2 cells. (A) HepG2 cells were treated with varying doses of sodium cantharidinate for 6-24 h. Cell viability was determined by MTT assays. (B) Western blot analysis for the expression of caspase-3 and cleaved caspase-3 protein in HepG2 cells treated with 5.0 µM sodium cantharidinate. (C) Quantitation of cleaved caspase-3 protein levels (**p<0.01, n=3). (D) Cells were stained with Hoechst 33258. Cell morphology was observed by laser scanning confocal microscopy. (E and F) Flow cytometric analysis of apoptosis in HepG2 cells treated with sodium cantharidinate. The cells were exposed to either control solution (0.1% DMSO in medium) or sodium cantharidinate at 5.0 µM and incubated for 6-24 h. (G) A caspase-3 inhibitor, Z-DEVD-FMK (100 µM) was added to the well, or not. Cells were treated by sodium cantharidinate for 6 h, then apoptosis was determined by FACS. I-Cas3, caspase-3 inhibitor; SC, sodium cantharidinate. The experiments were repeated at least three times. The data are expressed as the means ± SD of three experiments (**p<0.01 vs. control).
Figure 2. LC3 protein expression in HepG2 cells treated with sodium cantharidinate. (A and B) Western blot analysis and quantitation of the ratio of LC3-II to LC3-I for the expression of LC3 in HepG2 cells treated with 5.0 µM sodium cantharidinate for 6 h. (**) p<0.01, n=3). (C) HepG2 cells were stained with Hoechst 33258 and LC3-FITC. Cell morphology was observed by laser scanning confocal microscopy. (D and E) Flow cytometric analysis of LC3 expression in HepG2 cells treated with sodium cantharidinate. The HepG2 cells were exposed to either control solution (0.1% DMSO in medium) or sodium cantharidinate at 5.0 µM and incubated for 6 h. The data are expressed as the means ± SD of three experiments (**p<0.001 vs. control).

Figure 3. Inhibition of autophagy reduces sodium cantharidinate-induced cell apoptosis. (A) Western blot analysis for the protein expression of LC3-II to LC3-I for the expression of LC3 in HepG2 cells treated with 5.0 µM sodium cantharidinate combined with 3-MA for 6 h. (B) Quantitation of the ratio of LC3-II to LC3-I. Data are presented as means ± SD, compared with the control group (***p<0.001, n=3). (C) HepG2 cells were stained with Hoechst 33258 and LC3-FITC. Cell morphology was observed by laser scanning confocal microscopy. (D) Flow cytometric analysis of apoptosis in HepG2 cells treated with sodium cantharidinate combined with 3-MA. The data are expressed as the means ± SD of three experiments (***p<0.001 vs. control).
The results showed that the protein expression ratio of LC3-II and LC3-I was significantly increased by sodium cantharidinate treatment for 6-24 h (Fig. 2A and B). Furthermore, indirect immunofluorescence showed that LC3 had translocated to the cytoplasm, forming punctate aggregates, and the fluorescence intensity of LC3 was also enhanced (Fig. 2C), suggesting that sodium cantharidinate induced autophagy in HepG2 cells. The expression of LC3 expressed in HepG2 cells was analyzed by flow cytometry. The results are shown in Fig. 2D and E. The LC3 expression level was much higher in HepG2 cells treated with sodium cantharidinate than controls (MFI: 2508±165 vs. 1458±89, p<0.001). These results showed that sodium cantharidinate induced HepG2 cell autophagy through LC3 pathway.

**Inhibition of autophagy reduces sodium cantharidinate-induced cell apoptosis.** Previous research used 3-MA to inhibit autophagy and prove that autophagy is involved in the growth inhibition of hepatoma cells (26). Therefore, we combined 5 mM 3-MA and 5 µM sodium cantharidinate treatment in HepG2 cells for 6 h, and detected the protein expression of LC3-II and LC3-I by western blotting. Sodium cantharidinate combined with 3-MA resulted in a reduction of protein expression ratio of LC3-II and LC3-I compared with sodium cantharidinate alone (3-fold change, p<0.01, Fig. 3A and B). Indirect immunofluorescence demonstrated that LC3 was distributed in both the cytoplasm and the nucleus, and the fluorescence intensity was significantly reduced (Fig. 3C), showing that 3-MA inhibited sodium cantharidinate-induced autophagy effectively. Furthermore, Annexin V-FITC and PI double staining assay was also performed to confirm the LC3 inhibitor influences the cytotoxicity of sodium cantharidinate on HepG2 cells (Fig. 3D). The results showed that compared with the control group, the numbers of early and late apoptotic cells decreased significantly when 3-MA was combined with sodium cantharidinate. The proportion of early and late apoptotic cells in the sodium cantharidinate treatment group reached 37.2%, but it decreased to 22.2% when 3-MA was added (p<0.01).

**Silence of LC3 inhibits autophagy to reduce sodium cantharidinate-induced cell apoptosis.** We applied RNAi technology to inhibit LC3 expression on HepG2 cells. After the LC3 siRNA treatment of HepG2 cells for 24 h, we observed the expression rate of green fluorescent protein (GFP) to be
were significantly decreased detected by western blotting. At expression. After the LC3 siRNA treatment in HepG2 cells for sodium cantharidinate-induced apoptosis of HepG2 cells were compared to the control cells (p<0.001, Fig. 4G and H). These results showed that LC3 autophagy pathway played an important role in the sodium cantharidinate induced HepG2 cell apoptosis.

Discussion

The incidence of hepatocellular carcinoma is becoming the second leading cause of cancer-related death worldwide accounting for approximately 800,000 deaths every year. Hepatic resection and liver transplantation have progressed in surgical procedures for HCC. However, the outcomes have improved only slightly because of the frequent recurrence, even after liver transplantation. The pathogenesis on HCC remains unclear, but the genetic mutations of normal cells affected by environmental deterioration or other risk factors became a generally accepted carcinogenic factor (27).

Sodium cantharidinate kills liver cancer cell lines directly, which provided the favorable theoretical basis for the application of treatment of primary liver cancer (17). The present study demonstrated that sodium cantharidinate was able to inhibit the proliferation of HepG2 cells within the ranges of 2.0-12.5 µM and 6-24 h. Sodium cantharidinate enhanced the apoptotic effector of caspase-3 activity and induced cell death. Nucleus stained with Hoechst 33258 and Annexin V-FITC and PI double staining was consistent with MTT results. Caspase-3 activation could be initiated by many upstream signal-regulated molecules (28-30). Previous studies suggested that drugs could promote autophagy in human cancer cell lines, prompting speculation that apoptosis may be involved in the antitumor effect (26). Some research also demonstrated that oxidative stress can induce autophagy then inhibit the proliferation of liver cancer cells (26). In this study, we found that LC3 punctate aggregates and nucleation appeared in HepG2 cells after sodium cantharidinate treatment of the LC3 silenced HepG2 cells and late apoptotic cells decreased significantly when sodium cantharidinate treatment. The results showed that LC3 autophagy pathway played an important role in the sodium cantharidinate induced HepG2 cell apoptosis.

Acknowledgements

This study was supported in part by grants from the Jilin Provincial Natural Science Foundation of China (no. 20140520014JH), the 4th Young Scientist Fund of Jilin University (no. 2013068), the National Major Scientific, the Technological Special Project for 'Significant New Drugs Development' (no. 2014ZX09303030), the Interdisciplinary Chemistry and Medicine Foundation of Jilin University (JDYYJCHX004) and the National Natural Science Foundation of China (no. 31470418, to Y.H.).

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