Antiproliferative activity of gambogic acid isolated from *Garcinia hanburyi* in Hep3B and Huh7 cancer cells

PARRY NGAN HON LEE and WING SHING HO

School of Life Sciences, The Chinese University of Hong Kong, Hong Kong, SAR, P.R. China

Received November 8, 2012; Accepted December 28, 2012

DOI: 10.3892/or.2013.2291

**Abstract.** The anticancer activities of gambogic acid (GA) on two hepatocellular carcinoma cells with either p53 deletion (Hep3B) or p53 mutation (Huh7) were investigated in the present study. GA inhibited the growth of Hep3B and Huh7 through similar apoptotic pathways. After treatment of Hep3B and Huh7 with GA for 24 h, the IC\textsubscript{50} was determined for both cell lines at 1.8 and 2.2 µM, respectively. The results showed that both cancer cells underwent morphological changes and DNA fragmentation. GA induced apoptosis in the two cell lines through caspases-3/7, -8 and -9 in the mitochondrial pathway. The results suggest that both the caspases in the extrinsic death receptor pathway and the mitochondrial-dependent pathway are involved in the GA-induced cell apoptosis. The inhibitory effects of GA on Hep3B and Huh7 are independent of p53-associated pathway.

**Introduction**

Liver cancer remains one of the leading causes of cancer-related mortality. Herbal medicine is a valid source of new therapeutic agents since an immense chemical diversity is found in different herb medicines. Some of the active principles from herbal medicines have been isolated and characterized for cancer drug development. Herbal medicine offers promise with its complementary role in the cancer treatment with cancer drugs. Gambogic acid (GA) is one of the naturally occurring compounds present in a brownish-to-orange resin called gamboge, which is derived from *Garcinia hanburyi* (Fig. 1). *Garcinia hanburyi* has a long history of medicinal use in Southeast Asia, and it is used as a folk medicine and coloring agent (1). An improved separation method for the determination of twelve xanthones in gamboges from *Garcinia hanburyi* enabled researchers to purify GA from *Garcinia hanburyi* (2). Previous studies have reported that GA has potent antitumor activity via induction of reactive oxygen species accumulation which consequently led to apoptosis of SMMC-7721 cells (3). GA was reported to covalently modify I\kappaB kinase \beta subunit to inhibit lipopolysaccharide-induced activation of NF-\kappaB in macrophages (4). Other studies showed GA mediated the control of nucleophosmin and nucleoporins in the programmed cell death of Jurkat cells (5). GA caused microtubule depolymerization and phosphorylation of c-Jun N-terminal kinase-1 leading to cell cycle arrest in MCF-7 cells (6). The induction of apoptosis in cancer cells is vital in cancer treatment. The present study explored the antitumor activity of GA in Hep3B and Huh7 human liver cancer cell lines.

**Materials and methods**

**Chemicals and reagents.** Trypsin-EDTA (1X), Dulbecco's minimal essential medium (DMEM), Roswell Park Memorial Institute (RPMI)-1640 medium, penicillin-streptomycin (PS) antibiotic mixture (100X) and qualified fetal bovine serum (FBS) were purchased from Invitrogen. Chemicals and reagents were of analytical grade purchased from Sigma Chemicals, St. Louis, MO, USA. Dimethylsulfoxide (DMSO) was purchased from Fisher Scientific, USA. Bromophenol Blue, Agarose, Tris-Base, Boric Acid, NaCl and SDS powder were ordered from USB, Cleveland, OH, USA. Ethanol was purchased from BDH.

**Cell culture and treatment.** Hep3B, Huh7 and WRL68 were purchased from ATCC and cultured according to their protocols. GA was a generous gift from the Chinese Medicine Laboratory, Hong Kong Jockey Club Institute of Chinese Medicine, and was isolated by the established method (2). GA was prepared by dissolving 4 mg of dry GA into 1 ml of DMSO. Cancer cells were seeded in a 96-well plate at the density of 5.0x10\textsuperscript{4} cells/well and treated with various concentrations of GA for 24 h. Methylthiazoletetrazolium (MTT) solution (5 mg/ml) was added to the assay mixture and incubated for 4 h. The culture media was removed prior to addition of DMSO. Each experiment with GA treatment was repeated three times. By comparing the absorbance of the wells of cells treated with different concentrations of GA with the control, the viability of cells after GA treatment was calculated. The concentration of GA that reduced the cell viability of 50% (IC\textsubscript{50}) was recorded.

**DNA extraction.** The 100 bp DNA ladder was from Fermentas and the Cell Death Detection ELISA kit was purchased from...
Roche Applied Science. The cell samples were processed according to the manufacturer's protocols. The supernatant was subsequently removed and the DNA pellet was allowed to air dry for 15 to 20 min. TE buffer (20-50 µl) containing 0.2 mg/ml of RNase A was added to the DNA pellet. The samples were incubated at 37°C for 90 min to completely dissolve the sample. The DNA solution (2 µl) was added to 998 µl TE buffer and the concentration was measured using UV spectrophotometry (DU 650; Beckman-Coulter) with OD 

**DNA agarose gel electrophoresis.** Ten microliters of the dissolved DNA samples were mixed with 2 µl 6X DNA loading dye. The mixtures were loaded on the 2% agarose gel and run at 75V for 1 h. The DNA bands were examined under UV illuminator and the gel was photographed for documentation.

**DAPI stain.** Changes in cell morphology during apoptosis were examined by fluorescence microscopy of DAPI-stained cells. The monolayer of cells was washed in PBS and fixed with 4% paraformaldehyde for 30 min at room temperature. The fixed cells were permeabilized with 0.2% Triton X-100 in PBS three times and incubated with 1 µg/ml of DAPI for 30 min. The cells were washed with PBS three times. The apoptotic nuclei (intensely stained, fragmented nuclei and condensed chromatin) were examined under x400 magnification using a fluorescent microscope with a 340/380 nm excitation filter for at least two investigations.

**Cell death detection with ELISA.** ELISA was carried out according to the manufacturer's protocol (Invitrogen). The cells were incubated with different concentrations of GA in a 96-well plate for 24 h prior to centrifugation at 200 x g for 10 min. The lysis buffer (200 µl) was added to lyse the cell pellet. The plate was kept at room temperature for 30 min prior to centrifugation at 200 x g for 10 min. The supernatant (20 µl) was placed in the streptavidin-coated plate for analysis. The immunoreagent (80 µl) was added to the well and covered by aluminum foil with shaking at 300 rpm for 2 h. The solution in each well was removed and the well was rinsed with 250 µl incubation buffer three times. ABTS solution of 100 µl was added into the wells prior to shaking in the plate shaker at 250 rpm for color development. After 10-20 min, 100 µl ABTS Stop Solution was added to stop the reaction and the well was measured at 405 nm. Absorbance of wells was recorded and the enrichment factor, which shows the extent of apoptosis in the GA-treated sample-blank was calculated using the following expression:

\[
\text{Enrichment factor} = \frac{\text{Absorbance of the GA-treated sample-blank}}{\text{Absorbance of GA-treated cells}}
\]

**Western blotting.** The SDS-PAGE was completed when the dye front (blue in color) reached the bottom of the gel. The gel was removed and immersed into transfer buffer. The Whatman 3 MM paper (6 pieces/gel) and PVDF membrane were cut into the same dimension as the resolving gel. The PVDF membrane was washed with 100% methanol for 1 min and was immersed into the transfer buffer. The Whatman papers were soaked into the transfer buffer. A gel sandwich was assembled with the PVDF membrane and the resolving gel layered between two stacks of Whatman papers in the semi-dry Trans-Blot electroblotter (Bio-Rad). Proteins were transferred to the membrane at constant voltage at 10 V for 1 h. The membrane was collected and rinsed briefly with TBST buffer. Then, the membrane was immersed into blocking solution with primary antibody for 1 h at 4°C with continuous agitation. The primary antibody used was mouse monoclonal anti-p53 (1:1,000), anti-BID (1:1,000), anti-Bcl-2 (1:1,000), anti-Bax (1:500), anti-cytochrome c (1:2,000), anti-caspase-8 (1:1,000), anti-PARP (1:1,000), anti-transferrin receptor (1:1,000) and anti-GAPDH (1:5,000). The unbound primary antibody was removed by washing with TBST for 20 min, thrice. The membrane was immersed into secondary antibody (goat anti-mouse HRP/mouse anti-rabbit HRP) for 1 h at room temperature with slow agitation. The excess antibody was removed by washing with TBST three
times (20 min). Millipore Immobilon Western HRP kit (0.5 ml of each reagent) was applied to the membrane for 3 to 5 min. After removing excess reagent, protein bands on the membrane were visualized and recorded on Fuji Super RX film (Fujifilm). Intensity of the bands was measured using ImageJ program.

**Immunoprecipitation.** Cells were seeded into 100 mm dishes. After 24 h, different concentrations of GA (2, 1, 0.5 µM) were added before incubation for another 24 h. The cells were collected and lysed in CHAPS buffer. The concentration of the lysate was measured and diluted 500 µg protein/500 µl buffer. The anti-Bax 6A7 monoclonal antibody (2 µg) was added to the diluted lysate and the mixture was incubated overnight at 4°C. After the incubation, 25 µl of protein A-agarose gel bead was added and the lysate was incubated for 3 h. The gel beads were washed three times using the CHAPS buffer and collected by centrifugation at 14,000 x g for 10 min. The sample loading dye (50 µl) was added to the beads prior to boiling for 10 min, followed by a centrifugation of 14,000 x g at 4°C. The supernatant was subjected to western blot analysis using mouse anti-Bax antibody.

**Caspase cascade study of GA-induced cancer cells.** Apo-One™ caspase-3/7 assay kit was purchased from Promega. Caspase inhibitor z-IETD-fmk and z-LEDH-fmk were from Calbiochem. Trypan blue was purchased from Invitrogen. Hemocytometer was obtained from Sigma Chemicals. Caspase-3/7 activity was examined using Apo-One™ Caspase-3/7 Assay kit (Promega). Cells (Hep3B or Huh7) were seeded in a 96-well plate in the medium for 24 h followed by GA treatment of the cell lines. Subsequently, z-DEVD-rhodamine 110 from the assay kit was added into the well. After 2 h of incubation at room temperature, the fluorescence intensity of the wells was measured by a fluorescence plate reader at excitation 492 nm and emission 535 nm.

**Results**

**GA induces apoptosis in hepatocellular cells.** The effects of GA on the viability of Hep3B and Huh7 were investigated using MTT assay. GA effectively reduced the viability of cancer cells after incubation for 24 h. The IC₅₀ of GA for Hep3B was 1.8 and 2.2 µM for Huh7 (Figs. 2 and 3). After treatment of cells with GA, cell shrinkage and budding were recorded. DNA fragmentation and condensation were visualized by the DAPI staining investigation using fluorescence microscopy (Figs. 4 and 5). DNA fragmentation was detected after GA treatment in HCC cell lines (Fig. 5). The amount of cytosolic DNA fragment was determined with cell ELISA kit (Fig. 6). The results showed that the amount of DNA in the cytosol significantly increased after addition of GA. Apoptosis is characterized by the occurrence of cell shrinkage and membrane blebbing (7). In addition, chromatin condensation and DNA cleavage were visualized by the DAPI staining investigation using fluorescence microscopy (Figs. 4 and 5). DNA fragmentation was detected after GA treatment in HCC cell lines (Fig. 5). The amount of cytosolic DNA fragment was determined with cell ELISA kit (Fig. 6). The results showed that the amount of DNA in the cytosol significantly increased after addition of GA. Apoptosis is characterized by the occurrence of cell shrinkage and membrane blebbing (7). In addition, chromatin condensation and DNA cleavage were visualized by the DAPI staining investigation using fluorescence microscopy (Figs. 4 and 5). DNA fragmentation was detected after GA treatment in HCC cell lines (Fig. 5). The amount of cytosolic DNA fragment was determined with cell ELISA kit (Fig. 6). The results showed that the amount of DNA in the cytosol significantly increased after addition of GA. Apoptosis is characterized by the occurrence of cell shrinkage and membrane blebbing (7).
tested cells was collected and the amount of released DNA was detected by the ELISA kit. The results showed that following GA treatment, the levels of cytoplasmic DNA in HCC cells was markedly elevated (Fig. 7). These results provide evidence that GA mediated cell death through apoptosis.

Caspase cascade studies in GA-induced apoptosis. To study caspase activity in the GA-induced apoptosis, Apo-One™ Caspase-3/7 Assay was used. An increase of caspase-3/7 activity was found in the two HCC cells after GA treatment (Fig. 7). The activation of caspase-8 in the death receptor pathway and caspase-9 in the mitochondrial pathway were determined using western blot analysis (Figs. 8-10). The results indicate that the cleavage of the active form of caspases-8 and -9 after GA-treatment occurred in a time- and dose-dependent manner (Fig. 11).

Caspase-8 activation in GA-treated cells leads to Bid cleavage. The involvement of Bid and its relationship with caspase-8 activation in GA-induced apoptosis were investigated. Full
length Bid (24 kDa) was found to be cleaved in a time- and dose-dependent manner upon GA treatment (Fig. 12). A cell fractionation experiment was performed to investigate the level of tBid in the mitochondria. Fig. 13 indicates an accumulation of tBid in the mitochondria in the HCC cell line.

GA induces Bax conformational changes and cytochrome c release. It has been reported that Bax plays an important role in Bid-mediated apoptosis through conformational change and mediation of signaling process (11,12). Fig. 14 shows that the expression level of Bax was constant in GA-induced apoptosis. In order to further examine the role of Bax in GA-induced
apoptosis, immunoprecipitation (IP) of Bax using Bax (6A7) monoclonal antibody was performed. This antibody recognizes the Bax protein with conformational change but not the native form (11). It was reported that the conformational change of Bax was triggered by the truncated form of Bid (tBid) (13). This tBid is subsequently translocated to the mitochondria with Bax. The translocation of tBid to mitochondria was shown in western blot analysis (Fig. 15). Cytochrome c, which is originally present in the mitochondria, was released.

Discussion

In the present study we provided insights into the effects of GA on the inhibition of two different types of liver cancer cells. Present chemotherapy based on the treatment of liver cancer with drugs is not yet satisfactory. Specific anticancer agents against different types of liver cancer need to be developed. We demonstrated that GA induced apoptosis via the mitochondrial pathway in two types of liver cancer cells. p53 protein is a crucial factor in cellular stress responses and acts as an essential tumor suppressor (14). Upon activation, p53 controls the signaling process associated with the cell cycle based on the severity of the DNA damage. Thus, it can inhibit cell cycle progression or induce apoptosis. More than 50% of human tumors have been reported to have p53 mutations which affect p53 function. We found that GA induced caspase-associated apoptotic pathway which is important for induction of caspase-regulated apoptosis. We detected caspase protein in both cancer cells. The two HCC cell lines differ in the expression pattern of Bax and tBid to the outer membrane followed by the release of cytochrome c is a possible prerequisite for the mitochondrial apoptotic pathway (13,18). GA induces apoptosis in p53-deficient cancer cells. GA exerts inhibitory effects on Hep3B and Huh7 cell lines through similar mode of action. Based on these results, it seems unlikely that GA-induced apoptosis is due to the p53 status. This study showed the anticancer activity of GA is mediated via both the caspases in the extrinsic death receptor pathway and the mitochondria-dependent pathway. The present study demonstrated that deletion or deficiency of p53 does not affect cell cycle. The present results indicate that GA positively regulates the cancer cell apoptosis. Therefore, GA activity in liver cancer cells may represent one of the molecular mechanisms involved in anticancer agent-induced apoptosis. The low concentration of GA towards cancer cells is one of the fundamental criteria for efficient drug development and targeting. The results suggest that GA can be developed as an anticancer drug for p53-deficient cancer cells. The present study on GA may provide a promising therapeutic strategy for different types of liver cancer.
Acknowledgements

This study was in part supported by grant no. 6903292 kindly provided by Luck Tissue MFY Ltd. We thank Mr. Matt Cheung for his technical assistance.

References