Interaction Analysis of Glycyrrhizin on Licorice Extract-Induced Apoptosis of Human Leukemia Cells by Knockout Extract

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Abstract
Licorice (Glycyrrhiza spp.) is the most frequently prescribed herb in traditional Chinese medicine (TCM) and Japanese Kampo medicine [1,2]. In Western countries, licorice has been widely used as a flavoring and sweetening agent in foods, beverages, candies, tobacco, and dietary supplements. Licorice has various pharmaceutical activities including anti-inflammatory, anti-ulcer, anti-cancer, anti-virus, anti-allergy, and hepatoprotective activities [3-5]. Numerous phytochemical investigations reported that licorice contains at least 470 constituents including triterpenes, saponins, flavonoids, isoflavonoids, chalcones, polysaccharides, sugars, and amino acids [6]. Glycyrrhizin (GC; Figure 1) is a major bioactive triterpene glycoside in licorice and elicits anti-inflammatory, anti-ulcer, anti-tumor, anti-allergy, and hepatoprotective effects [4,5,7]. In contrast, various licorice constituents, such as flavonoid glycosides, and their aglycones induce apoptosis and act as anti-oxidative, anti-microbial, superoxide scavenging, and anti-tumor agents [4,5]. However, due to technical difficulties in preparing the GC-free licorice extract (LE), the potential function of GC in LE and the interaction between GC and other components of LE are not well understood.

Our laboratory has established several monoclonal antibodies (MAb) against naturally occurring bioactive compounds [6-11]. In our previous work, we prepared an anti-GC MAb for competitive enzyme-linked immunosorbent assay (ELISA) and Western blotting to identify and quantify GC [12]. Also we reported one-step purification of antigen molecule by immunoaffinity column conjugated with MAb. In these previous studies, we selected high specific MAb against forskolin [13] and ginsenoside Rb1 [14] for the separation of their antigens, and wide cross-reactive MAb against solasodine glycosides for the separation of all solasodine glycosides [15]. Furthermore, we reported a one-step GC isolation procedure using immunoaffinity columns conjugated with anti-GC MAb to prepare GC-knockout (KO) extract [16,17]. Recently, several groups also reported the preparation of KO extracts using different approaches such as preparative HPLC and TLC [18-20]. In this study, we investigated the potential role of GC in LE-induced apoptosis using GC-KO extract. GC-KO extract is useful tool to understand the synergistic interaction between GC and other compounds contained in LE. KO extracts provide a novel approach in pharmacological investigations that clarify the functions of bioactive compounds and the interactions between target compounds and other constituents of medicinal and functional food plant extracts.

Keywords: Licorice; Glycyrrhizin; Knockout extract; Immunoaffinity column; Apoptosis; HL-60

Introduction
Licorice (Glycyrrhiza spp.) is one of the oldest and most frequently prescribed herbs in traditional Chinese medicine (TCM) and Japanese Kampo medicine [1,2]. In Western countries, licorice has been widely used as a flavoring and sweetening agent in foods, beverages, candies, tobacco, and dietary supplements. Licorice has various pharmaceutical activities including anti-inflammatory, anti-ulcer, anti-cancer, anti-virus, anti-allergy, and hepatoprotective activities [3-5]. Numerous phytochemical investigations reported that licorice contains at least 470 constituents including triterpenes, saponins, flavonoids, isoflavonoids, chalcones, polysaccharides, sugars, and amino acids [6]. Glycyrrhizin (GC; Figure 1) is a major bioactive triterpene glycoside in licorice and elicits anti-inflammatory, anti-ulcer, anti-tumor, anti-allergy, and hepatoprotective effects [4,5,7]. In contrast, various licorice constituents, such as flavonoid glycosides, and their aglycones induce apoptosis and act as anti-oxidative, anti-microbial, superoxide scavenging, and anti-tumor agents [4,5]. However, due to technical difficulties in preparing the GC-free licorice extract (LE), the potential function of GC in LE and the interaction between GC and other components of LE are not well understood.

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Materials and Methods

Chemicals
Licorice root powder was purchased from Uchida Wakanyaku Corporation (Tokyo, Japan), and LE was prepared according to our previously published procedure [14]. Antibodies against poly (ADP-ribose) polymerase (PARP), caspase-3, and β-actin were purchased from Cell Signaling Technology (Beverly, MA, USA). Fetal bovine serum (FBS) was purchased from Gibco (Gaithersburg, MD, USA). All other chemicals were purchased from Wako Pure Chemical Industries (Osaka, Japan).

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Preparation of GC-KO extract using immunoaffinity columns conjugated with anti-GC MAb

Anti-GC MAb immunoaffinity columns and GC-KO extracts were prepared as described previously [16,17]. In brief, purified anti-GC MAb in coupling buffer (Bio-Rad Affi-gel HX coupling buffer) was dialyzed and oxidized with NaO2. Glycerol was then added to the reaction mixture and dialyzed. Affi-Gel HX gel (Bio-Rad) was added to the above reaction mixture, and the resulting hydrazone gel was packed into a plastic column. Twelve milligrams of LE in loading buffer (5% MeOH) was applied to the column, and the loading buffer was continuously circulated overnight at 4°C to enhance binding efficiency. After washing the column with loading buffer, the binding fraction was eluted with elution buffer (20 mM phosphate buffer containing 30% MeOH). Finally, each fraction was deionized and the solvent was lyophilized. The concentrations of GC in LE and GC-KO fractions were determined using ELISA. Fractions were then subjected to thin layer chromatography (TLC) with n-ButOH: H2O:CH3COOH (7:2:1) as the developing solvent, and stained with H2SO4 or analyzed by Eastern blot analysis.

Eastern blot analysis

GC, LE, and GC-KO extracts were applied to a polyethersulfone (PES) membrane and developed with acetonitrile:H2O:formic acid (45:55:2). The developed PES membrane was dried and immersed in NaO2 (10 mg/ml) for 1 h. After washing with water, 50 mM carbonate buffer solution (pH 9.6) containing 1% BSA was added and stirred for another 3 h. The PES membrane was washed and stained using standard protocols of Eastern blot analysis with anti-GC MAb [12,21].

HPLC fingerprints of LE and GC-KO extract

The standard compounds, i.e., GC, liquiritin, liquiritigenin and isoliquiritigenin, were dissolved in MeOH as 2 mg/ml stock solutions and stored at 4°C until use. The mixture of standard compounds was prepared by mixing equal volumes of stock solutions. A lyophilized LE and GC-KO extract were dissolved in MeOH as 30 mg/ml stock solution. HPLC analysis was performed using a TOSOH 8020 (Tokyo, Japan) equipped with a TSKgel ODS-100V (5 µm, 250×4.6 mm) and a UV-8020 detector (Tosoh, Tokyo, Japan). 20 µl of the mixture of standard compounds or 10 µl of GC-KO extract were injected into the column and eluted at room temperature with a constant flow rate of 1.0 ml/min. Acetonitrile (solvent A) and 0.15% acetic acid-H2O (solution B) were used in the mobile phase. Analysis was performed as follows: 0-20 min, hold 20% A; 20-60 min, linear gradient from 20% A to 100% A; 60-70 min, linear gradient from 100% A to 20% A. The detection wavelength was set to 254 nm.

Cell culture and treatment

Human leukemia HL-60 cell line and murine macrophage RAW264.7 were obtained from RIKEN Bio Resource Center Cell Bank, and cultured at 37°C under 5% CO2 in RPMI 1640 medium or Dulbecco’s modified Eagle’s medium (DMEM) containing 10% FBS, respectively. LE, GC, and GC-KO extracts were dissolved in dimethyl sulfoxide (DMSO) and diluted in medium at the final concentration of 0.2% DMSO (v/v). Control cells were treated with the same concentration of DMSO (0.2%, v/v).

Cell viability

Cell viability was determined by the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay, as described previously [22,23]. HL-60 cells (1×106 cells/well) in 96-well plates were treated with LE, GC, or GC-KO extract for 24 h or 48 h. At the end of treatment, MTT solutions were added to each well, and the cells were incubated for another 4 h. The precipitated MTT-formazan was dissolved with 0.04 N HCl-isopropanol, and formazan absorbance was measured at 595 nm using a microplate reader (Immuno Mini NJ-2300, Nihon InterMed, Tokyo, Japan).

Nuclear morphology assay

HL-60 cells (1×106 cells) were plated in 6-cm dishes, and then treated with LE for 48 h. At the end of treatment, cells were harvested, washed with PBS, and fixed with 1% glacial acetic acid for 30 min. After washing with PBS, cells were stained with Hoechst 33258 for 10 min and washed again with PBS. Nuclear morphology was then observed using a fluorescence microscope (Eclipse E600, Nikon, Tokyo, Japan).

DNA fragmentation analysis

DNA fragmentation was examined using a previously described method [22,23]. HL-60 cells (1×106 cells) were treated with LE for various periods. Treated cells were harvested and resuspended in lysis buffer (50 mM Tris-HCl, pH 8.0, 10 mM EDTA, and 0.5% SDS) with 0.2 mg/ml RNase A and incubated at 56°C for 30 min. Proteinase K was then added and incubated for another 3 h. DNA was separated on a 2% agarose gel and visualized under UV illumination after staining with ethidium bromide.

Western blot analysis

HL-60 cells were plated in 3×6 cm dishes (1×106 cells/dish), and then treated with LE, GC-KO extract, or GC for various periods. Harvested cells were lysed, and the supernatants were boiled for 5 min. Protein concentration was determined using a dye-binding protein assay kit (Bio-Rad) according to the manufacturer’s instructions. Equal quantities of protein were subjected to SDS-PAGE and electrophoretically transferred to PVDF membranes. After blotting, the membrane was incubated with specific primary antibody overnight at 4°C, and further incubated for 1 h with HRP-conjugated secondary antibody. Bound antibodies were detected using the ECL plus Western Blotting Detection System with a LAS4000mini (GE Healthcare). β-Actin was used as the internal control of whole cell lysate.
Analysis of cellular GC accumulation

HL-60 cells (1×10⁶ cells) were plated in 6-cm dish, and then treated with GC for various periods. After incubation, cells were washed twice with PBS and resuspended in 200 µl of PBS. Cells were sonicated and protein concentration was determined using a dye-binding protein assay kit (Bio-Rad). GC in cell protein was measured using competitive ELISA with anti-GC MAb, as described previously [12,17].

Immunocytochemistry

RAW264 cells (1×10⁵ cells) in chamber slide were treated with GC for 6 h. The treated cells were fixed for 10 min in 4% paraformaldehyde in PBS on ice. After wash with PBS, the cells were treated with 0.2% Triton X-100 for 2 min for membrane permeabilization. Then the cells were blocked with 1% BSA for 1 h and incubated with anti-GC MAb overnight at 4°C. The cells were rinsed and then incubated with FITC-labeled anti-mouse IgG for 1 h. Images of cellular immunofluorescence were examined under a confocal laser scanning microscope, LSM5Pascal (Carl Zeiss).

Statistical analyses

All data were derived from at least three independent experiments. Differences between groups were tested using Student's t-test and P values less than 0.01 were considered significant.

Results and Discussion

Preparation and characterization of the GC-KO extract

In our previous study, we demonstrated the preparation of anti-GC MAb [12]. Figure 1 shows the chemical structure of GC, glycyrrhetic acid-3-O-glucuronide, and glycyrrhetic acid. Cross reactivity of the anti-GC MAb with glycyrrhetic acid-3-O-glucuronide and glycyrrhetic acid was 0.585% and 1.865%, respectively. Cross reactivity with other related compounds (deoxycholic acid, ursolic acid, and oleanolic acid) was less than 0.005%. Using the GC-specific MAb, we prepared anti-GC immunoaffinity columns for the preparation of the GC-KO extract [17]. Twelve milligrams of LE, which contains 1275 µg of GC, was dissolved in loading buffer and applied to the anti-GC immunoaffinity column. After circulation of the loading buffer to enhance the binding efficiency of GC, the unbound fraction was collected. Subsequently, the column was washed completely, and the bound fraction was removed using the elution buffer. After deionization and lyophilization of each fraction, the concentration of GC was determined by ELISA using anti-GC MAb. The unbound fraction contained 3.50 µg of GC (0.27% of the loaded GC), whereas the bound fraction contained 1269.26 µg of GC (99.55% of the loaded GC), suggesting that the anti-GC immunoaffinity column functioned and then eliminated 99.55% of GC from LE.

In order to confirm this evidence, we next performed Eastern blot analysis, the high-sensitivity on-membrane quantitative staining using anti-GC MAb [12,21]. Figure 2A clearly showed the absence of GC in the unbound fraction (lane 3). Furthermore, TLC analysis demonstrated high concentration of GC exists together with the other components in LE (lane 2; Figure 2B). However, the spot of GC in unbound fraction specifically disappeared (lane 3; Figure 2B).

In order to make clear the more detailed characterization of GC-KO extract, we analyzed the HPLC fingerprint of LE and GC-KO extract because we have indicated that the HPLC fingerprint of the leaf extract of Eriobotrya japonica, which strongly suppressed the production of prostaglandin E₂ and nitric oxide in lipopolysaccharide-treated macrophages [24], showed the typical profile of the triterpene glycyrrhizin on Licorice Extract-Induced Apoptosis of Human Leukemia Cells by Knockout Extract. Nat Prod Chem Res 1: 105. doi:10.4172/2329-6836.1000105
constituents, and four major triterpenes including corosolic acid, ursolic acid, maslinic acid and oleanolic acid were highlighted relatively [25]. As shown in Figure 3, GC-KO extract contained three licorice flavonoids, such as liquiritin, liquiritigenin, and isoliquiritigenin, in the same pattern as LE. However, the peak corresponding to GC was not significantly suppress cell viability, it was expected that the GC-KO extract would suppress cell proliferation to the same degree as GC-KO extract.

Taken together, these results clearly indicate the effective elimination of GC from LE by the anti-GC immunoaffinity column, and suggest that the unbound fraction was the GC-KO extract.

**LE-Induced apoptosis of HL-60 cells**

We examined the effects of LE on cell viability of HL-60 cells using the MTT assay. Cells were treated with 25-200 µg/mL LE for 24 h and 48 h. As shown in figure 4A, LE significantly inhibited cell proliferation in a dose and time-dependent manner with IC50 values of 189.7 µg/mL and 130.2 µg/mL after 24 h and 48 h, respectively. To determine whether LE-induced cell death is mediated by apoptosis, nuclear morphological features of apoptotic cells were examined. Staining of the cells with Hoechst 33258 showed that treatment with 200 µg/mL LE for 48 h resulted in chromatin condensation (Figure 4B). For further confirmation, DNA fragmentation was investigated (Figure 4C). Moreover, Western blot analysis demonstrated that LE induced PARP cleavage and activated caspase-3; out of which both are hallmarks of apoptosis (Figure 4D). Taken together, these findings indicate that LE induced apoptosis in HL-60 cells.

**Treatment with GC alone fails to suppress cell proliferation**

At 200 µg/mL, cell proliferation was strongly suppressed by LE, and typical apoptotic features were observed. ELISA using the anti-GC MAb indicated that 200 µg LE contained 21.2 ± 1.24 µg of GC. In order to determine whether GC alone can induce cell death, we treated cells with GC at 2.5-40 µg/mL for 48 h. As shown in figure 5A GC had no significant effect on cell viability, suggesting that at these concentrations GC alone does not directly induce cell death. Next, we investigated whether or not GC uptakes into cells to exert unknown potential function. Cells were treated with 21.2 µg/mL GC for 1-12 h, and then GC concentrations in cell lysates were determined. ELISA experiments using the anti-GC MAb indicated that GC was indeed taken up by the cells (Figure 5B). Furthermore, in order to confirm the uptake of GC in the different cell line, we examined it using a murine macrophage RAW264 cells stimulated with lipopolysaccharide (LPS), which is inflammation model, resulted that clear incorporation of GC into cells was observed as shown in (Figure 6). These results indicated that, although GC alone cannot induce the cell death, GC can accumulate in cells.

**GC synergistically induced cell death with the other constituents of LE**

To further explore the role of GC in LE-induced apoptosis, we treated cells with LE (200 µg/mL), GC alone (21.2 µg/mL), GC-KO extract (178.8 µg/mL), or the combination of GC (21.2 µg/mL) and GC-KO extract (178.8 µg/mL). Treatment with LE inhibited cell viability (inhibition ratio (IR) = 83.0%) (Figure 7A). Because GC alone did not significantly suppress cell viability, it was expected that the GC-KO extract would suppress cell proliferation to the same degree as LE. However, the inhibitory effect of GC-KO extract was attenuated compared with LE (IR=24.8%), and the combination of GC-KO extract and GC significantly rescued this inhibitory effect (IR=43.5%). Similarly, Western blotting experiments demonstrated that caspase-3 activation by the GC-KO extract was weaker than that of LE, and addition of GC
This newly developed system may open the wide applications in the field of natural medicines like that an inhibitor and/or an activator in crude medicine can be found out easily using KO extracts.

The health benefits of medicinal and functional food plants have recently attracted increasing interest. However, the plant extracts are complex mixtures of active phytochemicals, which act synergistically or additively on specific and/or multiple molecular and cellular targets. Therefore, KO extracts are critical tools for determining the potential functions of target constituents from crude plant extracts using in vitro and in vivo assays.

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