**Lycii cortex radicis** extract inhibits glioma tumor growth *in vitro* and *in vivo* through downregulation of the Akt/ERK pathway

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Abstract. *Lycii cortex radicis* (LCR) is a traditional Korean medicinal herb. The present study was undertaken to examine the effect of an LCR extract on glioma cell growth and to determine its molecular mechanism in U87MG human glioma cells. The LCR extract resulted in apoptotic cell death in a dose- and time-dependent manner. The LCR extract-induced cell death was associated with generation of reactive oxygen species (ROS). Western blot analysis showed that the LCR extract caused downregulation of Akt and ERK. The LCR-induced cell death was prevented by transfection with the constitutively active forms of Akt and MEK. Oral administration of LCR extracts in subcutaneous U87MG xenograft models reduced glioma tumor volume. Taken together, these findings suggest that the LCR extract results in human glioma cell death through mechanisms involving ROS generation, downregulation of Akt and ERK, and caspase activation *in vitro* and reduces glioma tumor growth *in vivo*. These data suggest that the LCR extract may serve as a potential therapeutic agent for malignant human glioblastomas.

Introduction

Glioblastoma is the most common and highly aggressive primary brain tumor and is characterized by marked angiogenesis and extensive tumor cell invasion into the normal brain parenchyma (1,2). Glioblastoma is a rapidly growing, highly infiltrative tumor making complete surgical removal impossible. After diagnosis of glioblastoma multiforme, the median survival time of 9-12 months has remained unchanged despite aggressive treatment including surgical removal of the tumor, radiotherapy, and chemotherapy (3,4). Since the localization of tumors in the brain limits drug availability (5), the use of novel therapeutic approaches may be required.

Natural products derived from plants have recently received much attention as potential chemopreventive and chemotherapeutic agents. Among them great attention has been given to naturally occurring polyphenolic compounds such as flavonoids. These compounds may be potential candidates for cancer prevention and treatment in preclinical models and clinical trials (6-8). Considering that the use of synthetic agents in long-term chemopreventive strategies is associated with toxicity problems and leads to development of multidrug resistance, flavonoids may be a very promising group of compounds exerting the chemopreventive and chemotherapeutic effects.

*Eucalyptus* *cortex radicis* (LCR) has been extensively used as a traditional Korean medicinal herb for centuries due to its biological activities such as cooling blood and bringing down fever (9). Previous studies have shown that the LCR extract improves insulin resistance and lipid metabolism in obese-diabetic rats (10). Since it has been known that LCR contains a variety of physiologically active compounds such as apigenin, luteolin, kaempferol, quercetin, oleandric acid, and urosolic acid (11,12) which may have anticancer activities (13-15), LCR may exert an inhibitory effect on tumor growth. However, little information is available regarding the effect of LCR on cancer cell growth.

The present study was undertaken to investigate whether the LCR extract affects cell growth and to characterize its molecular mechanism in U87MG human glioma cells. The results of the present study show that the LCR extract inhibits glioma tumor growth through apoptotic cell death mediated by down-regulation of the Akt/ERK pathway *in vitro* and *in vivo*.

Materials and methods

Reagents. N-acetylcycteine (NAC), Trolox, catalase, Hoechst 33258, 3-,[4,5-dimethylthiazol-2-yl]-2,5-diphenyterotiazolium bromide (MTT), and propidium iodide were purchased from Sigma-Aldrich Chemical (St. Louis, MO, USA). Tween-20, U0126 and LY984002 were purchased from Calbiochem (San Diego, CA, USA). 2',7'-Dichlorofluorescein diacetate (DCFH-DA) was obtained from Molecular Probes (Eugene, OR, USA). Antibodies were obtained from Cell Signaling Technology,
Inc. (Beverly, MA, USA). All other chemicals were of the highest commercial grade available.

Preparation of the LCR extract. The LCR extract was prepared and authenticated by Dr Sun Dong Park (College of Oriental Medicine, Dongguk University, Kyung Ju, Korea). A voucher specimen (no. DGBS019) has been deposited in the Herbarium of Oriental Medicine, Dongguk University. The crushed LCR (1,000 g) was extracted 3 times each time with 3 volumes of methyl alcohol at 60˚C for 24 h. The extract was filtered and evaporated under a reduced pressure using a rotary evaporator to yield 72.97 g (yield 7.30%).

Cell culture. U87MG cells were obtained from the American Type Culture Collection (Rockville, MD, USA) and maintained by serial passages in 75-cm² culture flasks (Costar, Cambridge, MA, USA). The cells were grown in Dulbecco's modified Eagle's medium (DMEM, Gibco-BRL, Invitrogen, Carlsbad, CA, USA) containing 10% heat-inactivated fetal bovine serum (HyClone, Logan, UT, USA) at 37˚C in humidified 95% air/5% CO² incubator. When the cultures reached confluence, subculture was prepared using a 0.02% EDTA-0.05% trypsin solution. The cells were grown on tissue culture plates and used 1-2 days after plating when a confluent monolayer culture was achieved. Cells were treated with LCR extract in serum-free medium.

Measurement of cell viability and cell death. Cell viability was evaluated using an MTT assay (16). After washing the cells, the culture medium containing 0.5 mg/ml of MTT was added to each well. The cells were incubated for 2 h at 37˚C, the supernatant was removed and the formed formazan crystals in viable cells were solubilized with 0.11 ml of dimethyl sulfoxide. A 0.1 ml aliquot of each sample was then transfected to 96-well plates and the absorbance of each well was measured at 550 nm with an ELISA Reader (FLUOstar OPTIMA, BMG Labtech, Offenburg, Germany). Data were expressed as a percentage of control measured in the absence of LCR extract. Unless otherwise stated, the cells were exposed to 10 mg/ml LCR extract for 24 h. Test reagents were added to the medium 30 min before exposure to the LCR extract.

Cell death was estimated by the trypan blue exclusion assay. The cells were harvested using 0.025% trypsin and incubated with 4% trypan blue solution. The number of viable and non-viable cells was counted using a hemocytometer under light microscopy. Cells failing to exclude the dye were considered non-viable.

Measurement of apoptosis. Cells were grown in 6-well plates and were treated as indicated. Then, attached and floating cells were pooled, pelleted by centrifugation, washed in PBS, and fixed with cold 70% ethanol containing 0.5% Tween-20 at 4˚C overnight. Cells were washed and resuspended in 1.0 ml of propidium iodide solution containing 100 µg of RNase A/ml and 50 µg propidium iodide/ml and incubated for 30 min at 37˚C. Apoptotic cells were assayed using a FACSort Becton-Dickinson flow cytometer at 488 nm and data were analyzed with the CellQuest software. Cells with sub-G₁ propidium iodide incorporation were considered as apoptotic. The percentage of apoptotic cells was calculated as the ratio of events on sub-G₁ to events from the whole population.

Measurement of reactive oxygen species (ROS). The intracellular generation of ROS was measured using DCFH-DA. The nonfluorescent ester penetrates into the cells and is hydrolyzed to DCFH by the cellular esterases. The probe (DCFH) is rapidly oxidized to the highly fluorescent compound 2’7’-dichlorofluorescein (DCF) in the presence of cellular peroxidase and ROS, such as hydrogen peroxide or fatty acid peroxides. Cells cultured in 24-well plate were preincubated in the culture medium with 30 µM DCFH-DA for 1 h at 37˚C. After the preincubation, the cells were exposed to 10 mg/ml LCR extract for various durations. Changes in DCF fluorescence was assayed using the FACSort Becton-Dickinson Flow Cytometer (Becton-Dickinson Bioscience, San Jose, CA, USA) and data were analyzed with the CellQuest software.

Western blot analysis. Cells were harvested at various times after treatment of the LCR extract and disrupted in lysis buffer (1% Triton X-100, 1 mM EGTA, 1 mM EDTA, 10 mM Tris-HCl, pH 7.4). Cell debris was removed by centrifugation at 10,000 x g for 10 min at 4˚C. The resulting supernatants were resolved on a 12% SDS-PAGE under denatured reducing conditions and transferred to nitrocellulose membranes. The membranes were blocked with 5% non-fat dried milk at room temperature for 30 min and incubated with primary antibodies. The membranes were washed and incubated with horseradish peroxidase-conjugated secondary antibody. The signal was visualized using enhanced chemiluminescence (Amersham, Buckinghamshire, UK).

In vivo tumor growth assay. U87MG cells (2x10⁶) were injected subcutaneously into the right hind leg of 4-week male Balb/c nude mouse. After injection, animals were fed with 50 or 100 mg/kg LCR extract (n=5) in 50 µl saline or with 50 µl saline (vehicle, n=5) by oral gavage daily. After 7 weeks, the tumors were excised and tumor volume was calculated using the equation: tumor volume (mm³) = (length x width²) x π/6. Tumors were fixed in formalin, embedded in paraffin, and sectioned by standard methods for immunohistochemical analyses.

Immunohistochemistry. Paraftin-embedded tumor sections were deparaffinized and dehydrated through a series of graded alcohols. Sections were blocked with 8% BSA in PBS and incubated overnight at 4˚C with rabbit anti-Ki-67 (1:500 dilution) antibody. Sections were washed and incubated with appropriate fluorescein (FITC) secondary antibodies at room temperature for 1 h. After washing, counterstaining was carried out with Hoechst 33258 for 15 min. Sections were viewed under a fluorescence microscope (Leica, Wetzlar, Germany). The positive cells were counted and presented as the average of the four highest areas.

Terminal deoxynucleotidyl transferase-mediated dUTP nick end labeling (TUNEL) assay. The TUNEL assay was performed using an In Situ Cell Death Detection kit (Roche, Indianapolis, IN, USA) according to the manufacturer’s instructions and pictures were captured on an inverted fluorescence microscope. Briefly, slides were deparaffinized and treated with permeabilization solution (0.1% Triton X-100 + 0.1% sodium citrate) at 4˚C for 2 min to enhance the staining. For detection of single- and double-stranded DNA breaks,
slides were incubated for 1 h at 37°C with TUNEL reaction mixture (enzyme solution + labeling solution). Slides were stained with Hoechst 33258 for 15 min. TUNEL-positive cells were counted and expressed as the percentage of total cells.

Statistical analysis. The data are expressed as mean ± SEM and the difference between the two groups was evaluated using the Student’s t-test. Multiple group comparison was performed using one-way analysis of variance followed by the Tukey post-hoc test. A probability level of 0.05 was used to establish significance.

Results

Effect of LCR extract on cell viability and cell death. To determine the effect of the LCR extract on cell viability, cells were exposed to 1-30 mg/ml for 24 and 48 h. The LCR extract caused loss of cell viability in a dose- and time-dependent manner as evidenced by a decrease in MTT reduction (Fig. 1A). After addition of 10 mg/ml LCR extract for 24 and 48 h, the cell viability decreased up to ~55 and 25% of control, respectively. To ascertain whether reduction in cell viability was attributed to cell death, a trypan blue exclusion assay was performed. As shown in Fig. 1B, the LCR extract induced cell death in a dose- and time-dependent manner, similarly to those estimated by the MTT assay, suggesting that the reduction in cell viability by the LCR extract was mainly due to induction of cell death.

To determine whether the LCR extract-induced cell death was attributed to apoptosis, cells were exposed to 10 mg/ml LCR extract for 24 h, and flow cytometric analysis was performed for cells exposed to LCR extract. The sub-G1 peak (M1 gate) that represents a population of cells with reduced DNA staining, probably due to DNA fragmentation, increased from 2.92% in the control to 41.11% in the LCR extract-treated cells (Fig. 1C). The LCR extract induced glioma cell apoptosis in a time-dependent manner (Fig. 1D). These data suggest that the LCR extract-induced cell death was largely attributed to induction of apoptosis.

Role of ROS in LCR extract-induced cell death. To determine whether the LCR extract induces ROS generation in human
glioma cells, cells were exposed to the LCR extract and changes in DCF fluorescence were measured by flow cytometry. ROS generation increased in cells exposed to 10 mg/ml LCR extract for 12 h as assayed by increased DCFH-DA oxidation (Fig. 2A). To ascertain if ROS generation by the LCR extract is associated with ROS generation, the effect of antioxidants on the downregulation of these kinases was examined. As shown in Fig. 2E, the LCR extract-induced inhibition of Akt and ERK phosphorylation was blocked by NAC and catalase. These results suggest that ROS generation acts upstream for downregulation of Akt and ERK in cells exposed to the LCR extract.

Effect of the LCR extract on glioma tumor growth in vivo. To determine the antineoplastic effect of the LCR extract in vivo, U87MG cells were injected subcutaneously into Balb/c nude mouse. After injection, the LCR extract (50 or 100 mg/kg) or vehicle (saline) were administered by oral gavage daily. All mice were developed subcutaneous tumors with volume of~688 mm$^3$ after 7 weeks. LCR extract administration caused reduction of tumor volume in a dose-dependent fashion (Fig. 4A). There were no differences in body weight between the two groups (data not shown).

To assess whether the LCR extract inhibits tumor proliferation in vivo, Ki-67 expression was evaluated. Treatment of LCR extract decreased the proliferative rate of tumor cells in a dose-dependent manner (Fig. 4B), supporting the results obtained in vitro. We also obtained evidence that the LCR extract significantly increased apoptosis (Fig. 4C).

To determine whether ERK and Akt are downregulated by the LCR extract in vivo, phosphorylation of ERK and Akt was examined in tumor tissues of animals treated with 50 mg/kg LCR extract. Similarly to in vitro data, treatment of the LCR extract inhibited activation of ERK and Akt (Fig. 4D).

Discussion

Inhibition of cancer cell growth through induction of differentiation and apoptosis may be an attractive approach to human cancer therapy. It has been reported that natural products derived from plants exert anticancer effects via induction of apoptosis, although the molecular mechanisms by which they induce apoptosis have not been yet clarified (20,21). Recently,
we observed that mulberry fruit (Moris fructus) extract, a traditional Korean medicinal herb, induces human glioma cell death through the ROS-dependent mitochondrial pathway and inhibits glioma tumor growth (22). In the present study, we examined the effect of the LCR extract on glioma cell growth. The LCR extract has been extensively used as a traditional Korean medicinal herb for centuries (9). Although the LCR extract has been shown to improve insulin resistance and lipid metabolism in obese-diabetic rats (10), the effect of LCR on glioma cell growth has not been explored.

The present study demonstrated that the LCR extract caused loss of cell viability in a dose-and time-dependent manner and its effect was attributed to apoptotic cell death (Fig. 1).

Flavonoids, the major component of natural products derived from plants, behave as an antioxidant (20) or a pro-oxidant generating ROS (23-25). We previously observed that flavonoids such as silibinin and kaempferol induce cell death through ROS generation (14,15). Similarly, the LCR extract in the present study increased ROS generation and the LCR extract-induced cell death was prevented by antioxidants (Fig. 2), indicating that ROS production plays an important role in the LCR extract-induced glioma cell death.

The Akt signaling pathway mediates glioma cell survival and growth (26). The aberrant activation of Akt signaling has been identified as crucial to the malignant features of glioblastoma multiforme such as rapid tumor growth, invasiveness, resistance to cytotoxic treatments, and massive neovascularization (27-29). Therefore, regulation of the Akt signaling pathway may be a promising target for the clinical management of patients with glioma. Indeed, Akt was highly activated in U87MG human glioma cells and the Akt activa-
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Inhibition was inhibited by the LCR extract (Fig. 3A) and the LCR extract-induced cell death was also prevented by transfection with caAkt (Fig. 3D). These data imply that downregulation of Akt plays an important role in the LCR extract-induced glioma cell death.

ERK is activated by a variety of extracellular signals including mitogens contributes to the proliferative responses in cells, and is considered to be an essential common element of mitogenic signaling (17,30). Its constitutive expression causes cell transformation and plays a putative role in the carcinogenesis process and drug resistance (31-33). However, the effect of flavonoids on ERK activation is controversial. Activation of ERK is inhibited by flavonoids in vascular smooth muscle cells (34), human epidermal carcinoma cells (35), and neuronal cells (36), whereas it is increased following flavonoid treatment in lung cancer cells (37). Previously, we observed that ERK is activated by silibinin (15) and inhibited by kaempferol (14) in human glioma cells. These studies suggest that the effect of flavonoids on ERK activation may be dependent on cell types and flavonoid structures. These data may support reports that flavonoid-induced inhibition of phosphatidylinositol 3-kinase and protein kinase C is dependent on flavonoid structures (38).

In the present study, the LCR extract caused an inhibition of ERK phosphorylation (Fig. 3A) and the LCR extract-induced cell death was prevented by transfection of caMEK (Fig. 3D). The data imply that downregulation of the ERK signaling pathway plays an important role in the LCR extract-induced glioma cell death.

ROS induce the activation of multiple signaling pathways including Akt and ERK, leading to cell survival or cell death (39). On the other hand, the ability of ROS to trigger downregulation of the ERK and Akt pathway has also been reported.

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References


