Xanthoangelol, a Major Chalcone Constituent of Angelica keiskei, Induces Apoptosis in Neuroblastoma and Leukemia Cells

Keiichi Tabata,a Kou Motani,a Noriya Takayanagi,a Reiko Nishimura,a Satoru Asami,a Yumiko Kimura,a Motohiko Ukiya,c Daisuke Hasegawa,c Toshihiro Akihisa,c and Takashi Suzuki*a,b,d

a Clinical Pharmacy; b Chemical Analysis Center, College of Pharmacy, Nihon University; 7–7–1 Narashinodai, Funabashi, Chiba 274–8555, Japan; c Department of Materials and Applied Chemistry, College of Science and Technology, Nihon University; 1–8 Kanda Surugadai, Chiyoda-ku, Tokyo 101–8301, Japan; and d Department of Pediatrics, School of Medicine, Nihon University; 30–1 Oyaguchikami-cho, Itabashi-ku, Tokyo 173–0032, Japan.

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Xanthoangelol, a major chalcone constituent of the stem exudates of Angelica keiskei, was evaluated for cell toxicity and apoptosis-inducing activity in human neuroblastoma (IMR-32) and leukemia (Jurkat) cells. Xanthoangelol concentration-dependently reduced the survival rates of both cell lines as revealed by the trypan blue exclusion test. Early apoptosis induced by 4 h incubation with xanthoangelol was detected using flow cytometry after double-staining with annexin V and propidium iodide (PI). Western blot analysis showed that xanthoangelol markedly reduced the level of procaspase-3 and increased the level of cleaved caspase-3, but Bax and Bcl-2 proteins were not affected. These results suggest that xanthoangelol induces apoptotic cell death by activation of caspase-3 in neuroblastoma and leukemia cells through a mechanism that does not involve Bax/Bcl-2 signal transduction. Therefore, xanthoangelol may be applicable as an effective drug for treatment of neuroblastoma and leukemia.

Key words xanthoangelol; apoptosis; neuroblastoma; leukemia

Neuroblastoma is the most common solid pediatric tumor and remarkable for its clinical heterogeneity. Despite recent advances in chemotherapy, the prognosis of advanced neuroblastoma is still very poor. However, some favorable types of neuroblastoma, especially in infants under 1 year of age, are known to regress spontaneously or mature even if widespread metastases to bone marrow, skin and/or liver (special stage: stage IVS) are present. Apoptosis is known to occur in normal development of nervous systems, and neuroblastoma is generated from neural crest cells when the apoptotic systems do not carry out. Delayed implementation of the normal apoptotic pathway has been proposed as an explanation for the spontaneous regression of favorable neuroblastoma.1) It is reported that resistance to apoptosis plays a contributory role in the mechanism of the aggressive behavior shown by advanced neuroblastoma.2) Acute lymphocytic leukemia, like advanced neuroblastoma, is also a pediatric disease that is difficult to treat, especially in older children or those with a high amount of leukemic cells in the peripheral blood.

Angelica keiskei has been used traditionally in Japan as a diuretic, laxative, analeptic and galactagogue, and an A. keiskei extract was previously reported to affect metabolic activity3) and vasoconstriction5) in rats. Moreover, A. keiskei and a major chalcone constituent of this plant, xanthoangelol, reportedly have inhibitory effects against tumor promoter activity6,7) and metastasis.8) Xanthoangelol possesses a chalcone structure, and some compounds related to calmoterin activity6,7) and metastasis.8) Xanthoangelol markedly reduced the level of precursor caspase-3 and increased the level of cleaved caspase-3, but Bax and Bcl-2 proteins were not affected. These results suggest that xanthoangelol induces apoptotic cell death by activation of caspase-3 in neuroblastoma and leukemia cells through a mechanism that does not involve Bax/Bcl-2 signal transduction. Therefore, xanthoangelol may be applicable as an effective drug for treatment of neuroblastoma and leukemia.

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MATERIALS AND METHODS

Materials Xanthoangelol (3′-C-geranyl-2′,4,4′-trihydroxychalcone) was isolated from the stem exudate of A. keiskei6) and dissolved in dimethyl sulfoxide (DMSO) (final concentration 0.2%). IMR-32 and Jurkat were maintained in RPMI-1640 medium (Invitrogen) supplemented with 100 U/ml penicillin, 100 µg/ml streptomycin and 10% fetal bovine serum (FBS) (Invitrogen). The cells were maintained at 37 °C/5% CO2 in a humid environment.

Trypan Blue Exclusion Assay The cells (1×10⁶) were plated into a 60-mm dish and maintained for 24 h. Xanthoangelol (final concentrations 10⁻⁶, 10⁻⁵, 10⁻⁴ M) and vehicle were applied for 48 h. For the IMR-32 cell protocol, the cells were stripped using 0.05% trypsin-EDTA solution after washing them in phosphate-buffered saline (−). They were then washed in RPMI-1640 medium (with 10% FBS) and counted with a phase-contrast microscope immediately after addition of an equal volume of 1% trypan blue solution. Cell viability was calculated as the ratio of surviving cells to total cells.

3-[4,5-Dimethylthiazol-2-yl]-2,5-diphenyltetrazolium Bromide (MTT) Assay IMR-32 cells (1×10⁴/well) were spread onto a 96-well culture plate with phenol red-free RPMI 1640 medium (with 10% FBS) and maintained for 24 h. Then xanthoangelol (final concentration 10⁻⁸—10⁻⁴ M) and vehicle were applied for 48 h. After addition of 0.5% MTT solution as a 1/10 volume of medium in the well, incubation was continued for a further 4 h at 37 °C/5% CO2. An equal volume of stop solution (0.04 N HCl in isopropanol) to that of the culture medium was then added to each well and the absorbance at 570 nm (peak) and 630 nm (bottom) was
measured after thorough pipetting to disperse the generated blue formazan.

**Flow Cytometry** Apoptosis was detected using a Vi
tart™ apoptosis assay kit #3 (Molecular Probes). Cells (1×10^6 cells/dish) were exposed to xanthoangelol (10^{-5}–10^{-6} M) or the vehicle for 4 h. To prepare the cell sample for flow cytometry, cells were washed with annexin-binding buffer and stained with annexin V-fluorescein isothiocyanate (FITC) and propidium iodide (PI) for 15 min. The cell samples were analyzed by a FC500 flow cytometer (Beckman Coulter) using the FL1 and FL4 range for annexin V FITC and PI, respectively.

**Immunoblotting** Cells were collected and lysed with the lysis buffer (20 mM Tris–HCl pH 8.0, 137 mM NaCl, 1% Nonidet P-40, 10% glycerol, 1 mM phenylmethylsulfonylfluoride, protease inhibitor cocktail I (1 : 200; Sigma), phosphatase inhibitor cocktail II (1 : 100; Sigma), and 1 mM dithiothreitol). Protein concentration was determined using the Bradford method. Cell lysates containing 20 μg of total protein were loaded onto 12.5% or 15/25% SDS–polyacrylamide gels with Tris/glycine running buffer and transferred to polyvinylidene difluoride (PVDF) membranes (Amersham Bioscience). Each membrane was blocked with blocking buffer (5% skim milk, 137 mM NaCl and 20 mM Tris–HCl, pH 7.6) for 1 h at room temperature and incubated with the primary antibody (anti-caspase-3 1 : 3000, anti-Bax (2D2) 1 : 2000 and anti-Bcl-2 (Bcl-2-100) 1 : 1000 (Sigma), diluted in 5% bovine serum albumin) at 4 °C overnight. After washing with Tris–buffered saline containing 0.1% Tween-20, the membrane was incubated with a secondary antibody conjugated with horseradish peroxidase (1 : 10000, Sigma), diluted in 5% skim milk milk) at room temperature for 1 h. The signal was detected using an enhanced chemiluminescence Western blotting detection system (Amersham Bioscience). The density of the band was analyzed by NIH image-J software.

**RESULTS**

**Reduction of Neuroblastoma and Leukemia Cells Survival by Xanthoangelol** In the trypan blue exclusion study, 48 h exposure of IMR-32 and Jurkat cells to xanthoangelol markedly reduced their viability in a concentration-dependent manner (Figs. 1A, B). The highest concentration (10^{-4} M) of xanthoangelol showed potent cytotoxicity against both cell lines. IMR-32 cells were more sensitive than Jurkat cells to xanthoangelol at intermediate concentration. Both the MTT and trypan blue assays demonstrated concentration-dependent cytotoxicity of xanthoangelol, with an EC_{50} of 2.49×10^{-9} M for IMR-32 cells (Fig. 1C).

**Detection of Early Apoptosis** To clarify whether or not the xanthoangelol-induced cell death involved apoptosis, early apoptosis was examined using flow cytometry after annexin V and PI double staining. The distribution of stained cells is shown in Fig. 2A (control; vehicle only) and B (treated with 3×10^{-5} M xanthoangelol) in IMR-32 cells. As shown in Fig. 2C, the ratio of apoptotic IMR-32 cells was significantly increased at a xanthoangelol concentration of 3×10^{-5} M, and higher concentrations induced cell death with increased cell permeability. The low survival level even in the control cells (shown in Fig. 2C) seems to be caused by the cell membrane damage in the detaching process of adherent neuroblastoma cells from culture dishes. In Jurkat cells, xanthoangelol induced concentration-dependent early apoptosis followed by secondary cell death (Fig. 2D). In both cell lines, xanthoangelol in the concentration of 10^{-5} M, which 48 h exposure significantly reduced the cell viability in trypan blue assay, did not affect the parameter of apoptosis for 4 h application.

**Activation of Caspase-3** To clarify the mechanism by which xanthoangelol induces apoptotic cell death, caspase-3 activity was evaluated by immunoblotting. In both cell lines, the bands of caspase-3 (32 kDa) were gradually diminished by xanthoangelol (1×10^{-4} M) in a time-dependent manner. Especially in IMR-32 cells, activated caspase-3 (19 kDa) was detected in parallel with a reduction of 32-kDa caspase-3 (Fig. 3). These data revealed that xanthoangelol-induced cell death occurs through activation of caspase-3.

**Effects on Bax and Bcl-2** Expression of the proapoptotic protein Bax and the anti-apoptotic mitochondrial protein Bcl-2 was examined by immunoblotting analysis to clar-
ify the partial mechanism of xanthoangelol-induced apoptosis. Xanthoangelol (10^{-5}—10^{-4} M) slightly reduced the level of Bax and increased the level of Bcl-2 in IMR-32 cells (Fig. 4A). Bax was detected in Jurkat cells, but the expression level was very low compared to IMR-32 cells (these data were obtained from same gel). And xanthoangelol had no effect on the expression of Bax and Bcl-2 protein at the concentration of 10^{-4} M (Fig. 4B). The Bax/Bcl-2 ratio is one of the indices of the intrinsic mechanism of apoptosis in mitochondria. Xanthoangelol reduced this ratio in IMR-32 cells, it seems that xanthoangelol-induced apoptosis does not involve Bax/Bcl-2 signal transduction (Fig. 4C).

DISCUSSION

Our study investigated the cytotoxic activity of xanthoangelol, a major constituent of *A. keiskei*, which is a Japanese traditional health food, and its contribution to apoptosis in human neuroblastoma and leukemia cells.

The root of *A. keiskei* has been traditionally used as a diuretic, laxative, analeptic and galactagogic. Recently, scientific data on the active ingredients of *A. keiskei* have been accumulating, and some of them appear to have antitumor activity.6,8,11) Xanthoangelol has been isolated from *A. keiskei* extract as a major effective compound, which shows anti-tumor-promoting activity in vitro6) and anti-tumor growth and antimetastatic activity in vivo.8) Kimura and Baba have reported that the mechanism of this antitumor activity of xanthoangelol in Lewis lung carcinoma (LLC) cells involves inhibition of DNA synthesis within a concentration range of 10^{-5}—10^{-4} M.8) In this study, the same dose of xanthoangelol induced apoptosis with caspase-3 activation. Thus, apoptosis-inducing activity may contribute in part to the antitumor effect of xanthoangelol.

Proapoptic Bax and antiapoptic Bcl-2 are representative members of the Bcl-2 protein family. Expression of Bcl-2 is associated with an unfavorable histology of neuroblastoma.12) Up-regulation of Bax is required in order for apoptotic death to be induced in neuroblastoma cells by many drugs, such as cisplatin and ginsenoside Rh2.13,14) Hence, we started the immunoblotting experiment to expect that Bax/Bcl-2 ratio was increased by xanthoangelol. But, in our study, xanthoangelol reduced the Bax/Bcl-2 ratio in IMR-32 cells, a finding that is

**Fig. 2. Early Apoptosis and Secondary Necrosis Detected by Flow Cytometry Using Annexin V-FITC and PI Staining**

Xanthoangelol (10^{-5}—10^{-4} M) was applied for 4 h. The distribution of cells stained by annexin V FITC (FL1 Log) and PI (FL4 Log) in IMR-32 control cells (A) and cells treated with xanthoangelol (3×10^{-5} M) (B). The letters a, b, c and d in the graph represent alive, early apoptosis, secondary necrosis, and cells stained by PI alone, respectively. Concentration-dependent change in the percentage of each area was shown in C (IMR-32) and D (Jurkat). ∗p<0.05, ∗∗p<0.01 vs. control compared by one-way ANOVA followed by Bonferroni’s post-hoc test.

**Fig. 3. Activation of Caspase-3 by Xanthoangelol**

Caspase-3 (32 kDa) and cleaved caspase-3 (19 kDa) were detected by immunoblotting using anti-caspase-3 antibody in IMR-32 (A) and Jurkat (B) cells. Xanthoangelol (1×10^{-4} M) was applied for 0—24 h. The density of the bands for IMR-32 (shown in A) and Jurkat (shown in B) were analyzed and are shown in C and D respectively.
and like them, xanthoangelol also possesses potent apoptosis-inducing activity. Moreover, unlike cisplatin, continuous administration of an effective dose of xanthoangelol has been shown not to reduce the body, liver or lung weight of mice in vivo.\textsuperscript{20,21} Xanthoangelol-induced DNA synthesis inhibition was observed in LLC cells, but not in human umbilical vein endothelial cells (HUVECs).\textsuperscript{22} \textit{A. keiskei} has been used as a health food in Japan, and its dried yellow sap contains more than 10% xanthoangelol. Therefore, xanthoangelol has the potential to be a valuable therapeutic drug for neuroblastoma without producing adverse effects.

In summary, we have revealed that xanthoangelol induces caspase-3-dependent apoptotic cell death, which does not directly involve the Bax/Bcl-2 pathway. The apoptosis-inducing activity of xanthoangelol has not been reported previously. These results suggest that xanthoangelol may become a candidate as an antitumor agent.

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**REFERENCES**