Cell cytotoxicity and anti-glycation activity of taxifolin-rich extract from Japanese larch, *Larix kaempferi*

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**A B S T R A C T**

The larches, the *Larix* genus of plants are known as a natural source of taxifolin (dihydroquercetin), and extracts of its taxifolin rich xylem are used in dietary supplements to maintain health. In the present study, to assess biological activities of a methanol extract of the Japanese larch, *Larix kaempferi* (LK-ME), the effects of LK-ME on cell viability, inflammatory cytokine expression, and glycation were investigated. The effects of taxifolin which is known to be a main compound of LK-ME, and its related flavonoids, quercetin and luteolin were also examined. The results show that taxifolin exhibits lower growth inhibition activity and lesser induction activity of inflammatory cytokines in a human monocyte derived cell line, THP-1 cells, while in vitro anti-glycation activities of taxifolin were inhibiting at comparable levels to those of quercetin and luteolin. The growth inhibition and the cytokine induction activities, and the anti-glycation effects of LK-ME are assumed to have properties similar to taxifolin. The results of high performance liquid chromatography (HPLC) analysis indicated that taxifolin was detected as the main peak of LK-ME at the absorbance of 280 nm, and the concentration of taxifolin was measured as 3.12 mg/mL. The actual concentration of taxifolin in LK-ME is lower than the concentration estimated from the IC50 values calculated by the results of glycation assays, suggesting that other compounds contained in LK-ME are involved in the anti-glycation activity.

1. Introduction

Japanese larch, *Larix kaempferi* is a deciduous needle leaved tree and afforested in Hokkaido and Nagano, Japan on a large scale. Among other *Larix* genus plants, *Larix sibirica* and *Larix gmelinii* are known to contain abundant taxifolin (dihydroquercetin), a flavonoid, in the xylem, and taxifolin rich extracts of these larch xylems are used in a dietary supplement [1, 2, 3]. Taxifolin is known to be an anti-oxidative agent [4], and beneficial effects of taxifolin have been reported. Previous reports showed possible beneficial effects of taxifolin using animal models, including improvement of microcirculation [5], hepatoprotective effects [6], anti-viral activity [7], and prevention of diabetic nephropathy [8] as well as cardiomyopathy [9]. Further, in vitro studies demonstrated that taxifolin exhibits anti-bacterial [10], anti-fungal [11], and anti-parasitic [12] effects, and taxifolin inhibits acetylcholinesterase and carbonic anhydrase isoenzymes [13]. Taxifolin also inhibits oligomer formation of amyloid β proteins in mice, and is thought to be effective to prevent Alzheimer related diseases [14]. Similar to *L. sibirica* and *L. gmelinii*, the xylem of *L. kaempferi* is also known to contain much taxifolin [15]. At present *L. kaempferi* extracts have not been used in dietary supplements, and the effects of *L. kaempferi* extracts on the health of humans or animals are not well known.

Advanced glycation end products (AGEs) are produced by non-enzymatic reactions (Maillard reaction) of sugars and proteins. Excess energy intake, especially over-consumption of hydrocarbons increases blood sugar levels followed by induced glycation reactions. Elevated blood AGE levels are found in patients with diabetes mellitus [16], and it is thought to be involved in the disease onset of diabetic complications, such as in diabetic neuropathy [17], diabetic retinopathy and diabetic nephropathy [18]. In addition, accumulation of AGEs is known to progress with aging [16, 19]. It has been reported that accumulation of AGEs in the frontal lobe is found in Alzheimer dementia patients [20]. Overall, inhibition of production and accumulation of AGEs are thought to be important to prevent age related diseases.

In the present study, to evaluate potential of *L. kaempferi* extract to use for a supplement to maintain health of humans and animals, the effects of...
**L. kaempferi** methanol extract (LK-ME) on cell viability, induction of inflammatory cytokine mRNAs, and inhibition activity of glycation were investigated. These effects were also examined on taxifolin, a main compound of LK-ME, and compared with other taxifolin related flavonoids, quercetin and luteolin.

2. Results

2.1. Quantification of the taxifolin concentration in LK-ME

A methanol extract of *L. kaempferi* saw dust was used as LK-ME in this study. In this extraction condition, taxifolin is thought to be effectively extracted from the saw dust of *L. kaempferi*, and to substantiate this, and assess the quality of the extract, the concentration of taxifolin in LK-ME was quantified using high performance liquid chromatography (HPLC). As shown in Fig. 1A, taxifolin in LK-ME was detected as the main peak. The purity of the standard taxifolin was calculated as 95.4% (Fig. 1B). The concentration of taxifolin in LK-ME was calculated as 3.12 mg/ml by a comparison with the peak area and the purity of taxifolin standard solution (20 μg/ml).

2.2. The effects of LK-ME on THP-1 cells

To investigate the effects of LK-ME on immune cells, the effects of LK-ME on the cell viability of a monocyte derived cell-line, THP-1 cells [21], were examined. Effects of taxifolin, a main component in the xylem of *L. kaempferi*, and the taxifolin related compounds, quercetin and luteolin on the cell viability were also examined. The chemical structures of taxifolin, quercetin, and luteolin are shown in Fig. 2A. The results show that LK-ME inhibits growth of THP-1 cells in a dose dependent manner (Fig. 1B). The inhibition by taxifolin on the growth of THP-1 cells was weaker than those of quercetin and luteolin (Fig. 1C-E). The growth inhibition activity of a 100-fold dilution of LK-ME was slightly higher than that of 300 μM Taxifolin.

Next, the immune stimulation activity of LK-ME against THP-1 cells was investigated using real-time RT-PCR analysis. Here, THP-1 cells were stimulated with LK-ME, taxifolin, quercetin, or luteolin, and the mRNA expressions of inflammatory cytokines, interleukin-8 (IL-8), and tumor necrosis factor-α (TNF-α), were monitored. As shown in Fig. 3, the results indicate that the expressions of IL-8 and TNF-α mRNAs are significantly increased after the stimulation with LK-ME, as well as with taxifolin, quercetin, and luteolin. The induction activities of IL-8 and TNF-α mRNAs were different for these compounds, and these mRNAs were more effectively induced after stimulation with luteolin and quercetin than that of taxifolin.

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**Fig. 1.** HPLC analysis of the taxifolin concentration in LK-ME. LK-ME (A) and purified taxifolin (B, 20 μg/ml) were loaded onto an octa decyl silyl column (YMC-Pack Pro C18, YMC, Kyoto, Japan), and eluted by a linear gradient of phosphate buffer and acetonitrile. The eluted compounds including taxifolin were detected spectrophotometrically at column wavelength of 280 nm. Taxifolin peak is indicated with arrow head.
2.3. Anti-glycation activity of LK-ME

To investigate the anti-glycation activity of LK-ME, glucose, fructose, and glyceraldehyde were reacted with albumin, collagen, and elastin, respectively. Production of glycated proteins were determined by measurement of the fluorescence intensity. The anti-glycation activities of taxifolin, quercetin, and luteolin were also examined. As shown in Fig. 4A-C, LK-ME inhibited glycation in a dose dependent manner. Quercetin is known to be an anti-glycation agent [22]. Similar to quercetin (Fig. 4G-I), taxifolin (Fig. 4D-F) and luteolin (Fig. 4J and K) also exhibited anti-glycation activities in a dose dependent manner. The half maximal inhibitory concentration (IC50) calculated from these data are shown in Table 1. In the albumin-glucose reaction, luteolin exhibits the strongest anti-glycation activity. Taxifolin was more efficiently inhibiting glycation in the collagen-fructose reaction than quercetin and luteolin, and in the elastin-glyceraldehyde reaction more effectively than quercetin.

3. Discussion

In present study, the effects of LK-ME on cell growth, induction of inflammatory cytokines, and glycation were investigated. Overall, the biological activities of LK-ME investigated in this study were similar to those of its main compound, taxifolin. The inhibition activity to cell growth and the induction activities of IL-8 and TNF-α mRNAs were more strongly induced after stimulation with quercetin and luteolin when compared with that of taxifolin (Fig. 3). The TNF-α and IL-8 are known to be inflammatory cytokines induced by various stressors [23, 24, 25]. Based on this, the stress induced after stimulated with the flavonoids is thought to be involved in the induction of TNF-α and IL-8, and the stress induction after treatment with taxifolin is assumed to be less pronounced than that of the other flavonoids examined in this study, quercetin and luteolin.

The results of the HPLC analysis showed that taxifolin was detected as the major compound which absorbs at 280 nm in LK-ME used in this study (Fig. 1). There are several minor peaks in the chromatogram chart of LK-ME. A previous report showed that a metanol extract of L. kaempferi contains dihydrokaempferol, naringenin, 4-Hydroxybenzaldehyde, p-Coumaryl aldehyde as minor compounds [15]. Therefore, the minor peaks found in the HPLC chart are thought to be including these compounds.

The results of the cell viability analysis indicate that the growth inhibition activity of a 1,000-fold dilution of LK-ME is equivalent to 30–300 μM of taxifolin (Fig. 2B and C). This gives the taxifolin concentration in LK-ME, estimated by the growth inhibition activity, as 9.1–91.3 mg/ml. The IL-8 and TNF-α mRNA induction activities of a 100-fold dilution of LK-ME extract are comparable to 300 μM of taxifolin (Fig. 3). The taxifolin concentration in LK-ME is estimated as <9.1 mg/ml by the mRNA induction activities. Further, based on the IC50 values indicated in Table 1, anti-glycation activities of LK-ME against albumin-glucose, collagen-fructose, and elastin-glyceraldehyde were equivalent to 6.70, 15.51, and 9.12 mg/ml taxifolin, respectively. The actual concentration of taxifolin in LK-ME was calculated as 3.12 mg/ml by the HPLC analysis, and these concentrations of taxifolin in LK-ME estimated by the experimental results are higher than the concentrated quantities determined by the HPLC analysis. The difference between the estimated concentrations
between the experimental results and the actual concentrations suggest that other compounds contained in LK-ME are involved in the biological activities investigated in this study.

Japanese larch, *L. kaempferi* is thought to be a good source of taxifolin, and as shown in this study, taxifolin is simply extracted from the saw dust of the xylem of *L. kaempferi*. Although, further investigations including toxicity tests are required before it becomes possible to use *L. kaempferi* extract as a dietary supplement, the results shown in this study suggest *L. kaempferi* extract to be of promise for a supplement to maintain health of humans and also animals.

4. Materials and methods

4.1. Preparation of *L. kaempferi* extract (LK-ME)

LK-ME used in this study was prepared by methanol extraction from saw dust of *L. kaempferi*. Saw dust of *L. kaempferi* was obtained from the Forestry cooperative of Shimokawa town, Hokkaido, Japan. The saw dust (7.43 g) was extracted with 40 ml of methanol overnight at room temperature. The debris was removed by centrifugation, and the supernatant was filtrated with a 0.45 μm filter. Then, 1 ml of the extract was dried, resolved into 150 μl of dimethyl sulfoxide (DMSO, for assays using cultured cells) or ethanol (for glycation assay), and used in this study.

Fig. 3. The expressions of *IL-8* and *TNF-α* mRNAs after stimulation with LK-ME. THP-1 cells were stimulated with LK-ME (A, E), taxifolin (B, F), quercetin (C, G), and luteolin (D, H) at the concentrations indicated in the figure. After 6 h of stimulation, the cells were harvested, and total RNA isolated from the cells subjected to real-time RT-PCR analysis using specific primer sets for *IL-8* (A–E) and *TNF-α* (E–H) mRNA. The data are indicated as relative expression values compared with the expression in unstimulated control cells after normalization with the GAPDH mRNA expression. Error bars indicate standard deviations calculated from three independent experiments, and the asterisks (*) indicate that the difference is statistically significant (p < 0.01) and larger than two-fold, compared with that of the control.

4.2. Quantitation of taxifolin concentrations in LK-ME

The concentration of taxifolin in LK-ME was quantified using high performance liquid chromatography (HPLC). The HPLC system (Shimadzu Corporation, Kyoto, Japan) consist of a Model LC-20AD high pressure pump, a Model CTO-20AC column oven, a Model SIL-20AC total-volume injection-type auto-sampler, and a Model SPD-20A variable wavelength UV–Vis detector. Samples were separated using YMC-Pack Pro C18 (internal diameter: 3.0 mm, length: 150 mm, YMC, Kyoto, Japan) at 40 °C and the mobile phase consisted of 10 mM phosphoric acid (A) and acetonitrile (B) at 0.5 ml/min flow rates. Purified taxifolin (20 μg/ml) was used as the standard compound, and the concentration of taxifolin in LK-ME was calculated by the peak area of absorbance units at 280 nm compared with the standard. The HPLC analysis was performed by the Biodynamic Plant Institute, Sapporo, Hokkaido, Japan.
4.3. Cell culture and monitoring of cell viability

A human monocyte-derived cell line, THP-1 cells (ATCC TIB-202) [21] were grown and maintained in RPMI 1640 medium supplemented with 10% fetal bovine serum, 100 U/ml penicillin, 100 mg/ml streptomycin (Life Technologies, Carlsbad, CA, USA). The cells were grown at 37 °C in 5% CO2 in a humidified incubator. The cell viability was monitored using a Cell Counting Kit-8 (Dojindo, Kumamoto, Japan) in

Fig. 4. Effect of LK-ME on glycation. Human albumin and glucose (left column), bovine collagen and fructose (middle column), and bovine elastin and glyceraldehyde (right column) were incubated with the indicated concentrations of LK-ME (A–C), taxifolin (D–F), quercetin (G–I), and luteolin (J, K). At the indicated time points, the concentrations of advanced glycation end products (AGEs) was measured by monitoring the fluorescence intensity (excitation: 365 nm, emission: 410–460 nm) using a multimode microplate reader. Error bars indicate standard deviations (n = 3).
Table 1
The half maximal inhibitory concentration (IC50) of LK-ME against production of AGEs.

|                | LK-ME | Taxifolin | Quercetin | Luteolin |
|----------------|-------|-----------|-----------|----------|
| Albumin-Glucose | ×474.97 dil. | 46.34 μM | 47.28 μM | 19.55 μM |
|                | 2.11 μl/ml | 14.10 μg/ml | 14.29 μg/ml | 5.60 μg/ml |
| Collagen-Fructose | ×3214.56 dil. | 15.92μM | 48.76 μM | 41.42 μM |
|                | 0.31 μl/ml | 4.83 μg/ml | 14.74 μg/ml | 11.86 μg/ml |
| Elastin         | ×102.14 dil. | 293.32 μM | 597.30 μM | n.d.     |
| Glyceraldehyde  | 9.80 μl/ml | 89.24 μg/ml | 180.53 μg/ml | n.d.     |

These IC50 values were calculated from the results of the glycation assay (Fig. 3).

n.d.: no data.

4.4. Monitoring the production of advanced glycation end products (AGEs)

Human serum albumin (Fraction V; Nacalai tesque, Kyoto, Japan), elastin derived from bovine neck ligament (MP Biomedicals, Irvine, CA, USA), and collagen derived from bovine skin (Type I, acid soluble; Nippi, Tokyo, Japan) were purchased as commercially available products. To solubilize elastin, 100 mg of bovine elastin was heated in 1ml of 0.1N NaOH at 99 °C, and the supernatant recovered. After repeating this step twice, the remaining pellets were autoclaved in 1 ml of 0.1N NaOH, and the supernatant was collected. This step was also repeated twice, and then the collected supernatant containing solubilized elastin was neutralized using 1N HCl, and sterilized using a 0.22 μm filter. The protein concentration was measured using a Pierce BCA Protein Assay Kit (Thermo Fisher Scientific Waltham, MA, USA) according to the manufacturer instructions, and used in this study.

The production of AGEs was measured by monitoring the increment in fluorescence intensity. Human albumin (8 mg/ml) and glucose (0.2 M), bovine collagen (0.3 mg/ml) and fructose (0.2 M), or bovine elastin (0.5 mg/ml) and glyceraldehyde (0.05 M) were reacted in 0.05 M phosphate buffer (pH 7.4) with a series of concentrations of LK-ME, taxifolin, quercetin, or luteolin at 60 °C. After the incubation, the fluorescence intensity (excitation: 365 nm emission: 410–460 nm) was measured using a multimode microplate reader (GloMax Multi Detection System; Promega, Madison, WI, USA).

The half maximal inhibitory concentration (IC50) values were calculated using the IC50 calculator, a web based program developed by AAT Bioquest, Sunnyvale, CA, USA (https://www.aatbio.com/tools/ic50-calculator).

4.5. Real-time reverse transcriptase polymerase chain reaction (RT-PCR) analysis

The total RNA was extracted from the cultured cells using TRIzol reagent (Invitrogen, Carlsbad, CA, USA). The isolated total RNA was treated with DNase1 (Takara, Otsu, Shiga, Japan) and then subjected to oligo-dT- and random-primed reverse transcription using ReverTra Ace (Toyobo, Osaka, Japan). Real-time PCR was performed using Thunderbird SYBR qPCR Mix (Toyobo), and the PCR reactions and analysis of the mRNA expressions were performed using the CFX96 Real-Time PCR Detection System (Bio-Rad, Hercules, CA, USA). The procedures were performed according to the manufacturer protocols. The following listed specific primer sets for IL-8, TNF-α, and glyceraldehyde-3-phosphate dehydrogenase (GAPDH) mRNAs were used in this study. TNF-α sense primer 5'-GCCAGGACCTCTCTCTAACGACG-3'; TNF-α antisense primer 5'-ATGGCTACAGGCTTGTCACT-3'; IL-8 sense primer 5'-GTCGAGTTTGGCCAAGAGT-3'; IL-8 antisense primer 5'-CCTGACAGCCAGATTTCTTT-3'; GAPDH sense primer 5'-TTCTTTGGCTGCAGCCCCG-3'; and GAPDH antisense primer 5'-GTTGACCGAGGGCCCAATCAGG-3'.

4.6. Statistical analysis

To determine statistically significant differences between data pairs, a two-tailed unpaired Student’s t-test was performed in this study.

Declarations

Author contribution statement

Daisuke Muramatsu: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

Hirofumi Uchiyama: Conceived and designed the experiments; Performed the experiments.

Hiromi Kida: Contributed reagents, materials, analysis tools or data; Wrote the paper.

Atsushi Iwai: Conceived and designed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

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Additional information

No additional information is available for this paper.

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