Original article

In-vitro antioxidative, antiinflammatory properties of *Aurea helianthus* leaf extract a Korean traditional medicinal plant

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

The leaf of *Aurea helianthus* (*A. helianthus Jinhuaikui*) is popularly used in China traditional medicine, however, scientific evidence on its antioxidant properties rarely studied. In this study, biological activities of *A. helianthus* leave’s 80% ethanol extract (AHL) were investigated. The measured total polyphenol and flavonoid content of AHL was 184.24 ± 5.01 mg GAE/g and 102.53 ± 0.98 mg NAR/g. AHL showed the highest *α*-diphenyl-β-picrylhydrazyl (DPPH) and 2,2'-azino-bis-3-ethylbenzothiazoline-6-sulfonic acid (ABTS) radical scavenging activities of 98.30 ± 0.18% at 1000 μg/mL. DPPH and ABTS radical scavenging activities significantly increased in a AHL concentration-dependent manner. AHL treatment significantly suppressed the generation of pro-inflammatory mediators, including nitric oxide (NO), in lipopolysaccharide (LPS)-stimulated RAW 264.7 macrophages. AHL demonstrated strong anti-inflammatory activity that reduced NO production in LPS-stimulated RAW 264.7 cells. To test the potential protective effect of AHL, the antioxidant capacity, on the cell growth, viability of a human hepatoma cell (HepG2) and Raw 264.7 cell were investigated. AHL also enhanced cytoxicity on the proliferation of HepG2 cells and was capable of inhibiting 56% against LPS at 400 μg/mL. The results of this study the potential of AHL as an excellent antioxidant substance for inhibiting inflammatory mediators. Therefore, AHL may be used as a therapeutic approach to various inflammatory diseases.

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1. Introduction

Antioxidant medicinal plants, including phenolic and flavonoid, are considered beneficial because of their protective actions in diseases as cancer. Phenol and flavonoids have been showed a wide range of biological activities (Bravol, 1998), including anticarcinogenic actions. Most of the beneficial health effects of flavonoids are attributed to their antioxidant and chelating abilities. Over product ion of reactive oxygen species (ROS) has shown to have detrimental effects on human health leading to cell/tissue damage and degenerative disorders such as inflammation, cardiovascular and neurogenic diseases, cancer, and aging – related disorders. ROS are capable of oxidizing cellular proteins, nucleic acids, and lipids. Lipid peroxidation is a free-radical mediated propagation of oxidative insult to polysaturated fatty acids involving several types of free radicals, and termination occurs through enzymatic means or by free radical scavenging by antioxidants (Korkina and Afans’ev, 1997). ROS, including hydroxyl radicals, superoxide anions, singlet oxygen, and hydrogen peroxyde, are generated as by-products of cellular metabolism (Crack and Taylor, 2005). If oxidative stress going on, oxidative damage to biomolecules increase and results in diverse biological damage such as mutagenesis and cell death (Ernak and Davies, 2002). Many reports suggest that ROS are principal mediators of apoptosis (Simbula et al., 2007). Antioxidants are added to food to slow the rate of oxidation and, if used properly, they can extend the length life of the food. ROS are produced by mitochondrial electron transfer processes and cytochrome P450 systems in hepatocytes (Masella et al., 2005). Human hepatoma cell line (HepG2) is quite suitable for cytotoxicity evaluation due.
to the quality and stability of its enzymic and metabolic background (Osseni et al., 2000). Many biological, chemical, and physical agents can generate inflammation with increased danger of human cancers (Oshshima et al., 2003). Inflammation is an important physiological defense process by infection, or exposure to endotoxins such as lipopolysaccharid (LPS) (Medzhitov, 2008). LPS is derived from gram-negative bacteria, is a strong inducer of pro-inflammatory cytokines and immune responses in animals (Choi et al., 2015). It has been involved in non-pathogenic aspects of bacterial ecology, and bacteriophage sensitivity. Overproduction of NO in response to iNOS during the inflammation process is an important target in drug development for inflammation disease (Wilson et al., 1996). NO has harmful acts with tissue injury, septic shock and apoptosis (Kubes and McCafferty, 2000). So, many studies are currently going to develop inhibitors from medicinal plants to prevent or cure chronic inflammatory conditions for minimal side effects (Lampiasi and Montana, 2016).

*A. helianthus* is a short-lived perennial herb in *Abelmoschus* and the tropics but an annual in cooler climates. Its main effects are anti-pyretic, anti-inflammatory, analgesic and it has the functions of blood lipid regulation, tumor cells inhibition, immune regulation, and anti-oxidation and so on (Lu and Jia, 2015). The young leaves and stems may be eaten raw, steamed, boiled, stir-fried or added to soups. As the leaves cook quickly, add them last to steamed vegetable or stir-fry. The leaves contain mucilage, which can give a slightly slimy feel in the mouth. *A. helianthus* is a very nutritious vegetable; the leaves are high in vitamins A, C, iron, and 2% protein by dry weight. However the importance of this plant is that it is one of the world’s most nutritious leafy vegetables because of its high protein content. A paste of the bark is used to treat wounds. In Nepal, the root juice is applied to sprains and the flower juice is used to treat protein content. A paste of the bark is used to treat wounds. In Nepal, the root juice is applied to sprains and the flower juice is used to treat protein content.

2. Materials and methods

*A. helianthus* leaves were collected from market, 2015. 05., China. For analysis, each 2 g of *A. helianthus* leaves was extracted with 80% EtOH, (3 x 250 mL) by reflux and evaporated in vacuo. Evaporation was conducted using an evaporator system under reflux in vacuo from BUCHI, Meierseggrasse, Flawil, Switzerland. UV/VIS spectrometer was performed by Synergy H1 Hybrid Reader purchased from BioTek Instruments Winooski, VT, USA. Human liver (HepG2) and macrophage (Raw264.7) cells were purchased ATCC (Rockville, MD), and stored at 4 °C until used.

2.1. Reagents

All reagents were purchased from Sigma Aldrich, St. Louis, MO, USA, except indicated. Dulbecco’s modified eagle’s medium (DMEM) and fetal bovine serum (FBS) were purchased Gibco, NY, USA. Penicillin/streptomycin and trypsin/EDTA were obtained Welgene, Gyeongsan-si, Gyeongsangbuk-do, South Korea. The reagents 1,1-diphenyl-1-picrylhydrazyl (DPPH), 2,2-azinibis-3-ethyl benzothiazoline-6-sulfonic acid (ABTS), α-tocopherol, 2,4,6-tris(2-pyridyl)-s-triazine (TPTZ), iron (III) chloride hexahydrate, gallic acid, folin and ciocalteu’s phenol reagent, 3-(4,5-dimethyl thiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) and LPS were purchased from Sigma-Aldrich (St. Louis, MO, USA). Acetic acid was purchased from Junsei (Tokyo, Japan).

2.2. Determination of total phenol content

The concentration of total phenols in plant extracts was estimated by Folin-Denis’s phenol method (Vernon et al., 1999). 100 µl of the samples with 400 µl of water were mixed with 50 µl of a 2 N folin-ciocalteu reagent. After 3 min, the Na2CO3 solution was added. The absorbance was determined after 1 h at 20 °C, versus a zero-absorbance reagent blank. The blue color produced was measured at 725 nm using a UV/VIS spectrometer. The concentration of total phenolic compounds in extracts was calculated by comparison with a standard curve similarly prepared with 0–1 mg/mL garlic acid. Total phenolic content values were determined using an equation that was obtained from the standard curve of garlic acid graph (r² = 0.992). The total phenolic content of the sample was expressed as garlic acid equivalents which reflected the phenolic content as the amount of garlic acid (mg/g) of extracts.

2.3. Determination of total flavonoid content

The total flavonoid content was determined using the method (Chae, 2002). 1 mL of glycol in methanol was mixed with the 100 µl of the samples and 100 µl 1 N NaOH. Absorption readings at 420 nm using a UV/VIS spectrometer were taken after 1 h against a blank sample consisting of a 100 µl samples with 1.1 mL water. The total flavonoid content was determined using a curve of naringin (r² = 0.979) as the standard. Total flavonoid content is expressed as mg of naringin equivalents CAE/g of extract.

2.4. DPPH scavenging assay

This assay was carried out as described by Brand-Williams et al. (1995) with some modifications. 40 µL of various dilutions of the test materials (pure antioxidants or plant extracts) were mixed with 160 µl of a 0.25 mM DPPH solution. After 30 min, the absorbance at 517 nm of maximum absorbance of DPPH, was recorded as A_{sample}, using a UV/VIS spectrophotometer. A blank experiment was also carried out applying the same procedure to a solution without the test material and the absorbance was recorded as A_{blank}. Ascorbic acid was used as a positive control. The radical scavenging activity of each solution was then calculated as percent according to the following equation:

\[
\text{Radical scavenging activity (\%)} = \frac{(A_{\text{blank}} - A_{\text{sample}})}{A_{\text{blank}}} \times 100
\]

where A_{blank} is the absorbance of the control reaction (containing all reagents except the test sample), and A_{sample} is the absorbance of the test sample. Tests were carried out in triplicate. The experiment was performed in triplicate.

2.5. ABTS scavenging assay

Radical cation scavenging capacity of extract was examined against ABTS generated by the method (Re et al., 1999). ABTS was dissolved in water to a 7 mM concentration. ABTS radical cation was produced by reacting ABTS stock solution with 2.45 mM potassium peroxodisulfate and allowing the mixture to stand in the dark at room temperature for 12–16 h before use. The ABTS+ reaction mixture contained 180 µl of ABTS+ and 20 µl of antioxidant testing sample with an absorbance at 734 nm, recorded as A_{sample}. The radical scavenging activity of each solution was then calculated as percent according to the following equation:

\[
\text{Radical scavenging activity (\%)} = \frac{(A_{\text{blank}} - A_{\text{sample}})}{A_{\text{blank}}} \times 100
\]

Ascorbic acid was used as a positive control.

2.6. Reducing power assay

To determine the reducing capacity of extracts, use the method of Yen and Chen (1995). Extracts (tested from 125 to 1000 g/mL) 250 µL were added in 200 mM sodium phosphate buffer 250 µL. For a 500 µL sample solution, 250 µL of 1% potassium ferricyanide were added and incubation was carried out for 20 min at 50 °C.
After added 250 µL of 10% trichloroacetic acid (TCA) to sample solution, centrifuge at 3000 rpm. Then, 500 µL supernatants of sample solutions added 100 µL of 0.1% ferric chloride (FeCl₃). After incubation at room temperature for 10 min, optical density was determined using a microplate reader set at 700 nm.

2.7. Cell culture

HepG2 and Raw 264.7 cells were cultured in DMEM medium supplemented with 10% FBS and 1% penicillin/streptomycin. Cells were maintained in a humidified atmosphere at 37 °C, 5% CO₂. Cells were subcultured at 70–80% confluence.

2.8. Cell viability

The cell viability was measured using MTT assay (Angius and Floris, 2015). HepG2 cells were seeded in 96 well plates at 2 × 10⁵ cell/well. After overnight growth, cells were treated with AHL for 1 h, followed in the presence or absence of H₂O₂ (20 mM) for the next 24 h. And Raw 264.7 cells were seeded in 96 well plates at 2 × 10⁵ cell/well. After overnight growth, cells were treated with AHL for 1 h, followed in the presence or absence of LPS (1 mg/mL) for the next 24 h MTT solution (5 mg/mL) was added and the cells were further cultured for 2 h. After the incubation, the formazan crystals in each well were dissolved in 125 µL of DMSO for 15 min. The absorbance at 570 nm was measured with microplate reader.

2.9. Intercellular reactive oxygen species (ROS) generation

Intercellular ROS generation was measured by DCF-DA (2’,7’-dichlorodihydrofluorescein diacetate) with temperature maintained at 37 °C (Wang and Joseph, 1999). After 24 h cultured HepG2 cells with extract in black 96 well plate, treated with 20 µM DCF-DA in serum free medium (SFM) for 50 min and treated with 200 µM H₂O₂ in Krebs-Henseleit buffer (KHB) for 30 min and 1 h. The fluorescence at the excitation filter at 485 nm and the emission filter at 535 nm was measured by a microplate reader.

2.10. Nitro oxide (NO) assay

Nitrate concentration as an index for NO synthesis was determined using greiss reagent as the manual directs (Kyung et al., 2012). After 24 h cultured Raw 264.7 cells treated extracts with or without LPS, supernatants 50 µL and greiss reagent 50 µL were added to 96 well plates to each well. After incubation at room temperature for 10 min, optical density was determined using a microplate reader set at 540 nm. The nitric oxide content was determined using a curve of sodium nitrite (r² = 0.998) as the standard.

2.11. Statistical analysis

All statistical analyses were performed using Graphpad prism 6 software. Results were compared by one-way analysis of variance (ANOVA). A P-value of <.05 was considered to be significant.

3. Results and discussion

Flavonoids and polyphenolics in general depending on their glycoside, isoprenoids and aliphatic ethers content can acquire almost and polarity. The content of extractable polyphenolics in solvent extracts of AHL, determined from regression equation of calibration curve y = (x + 0.0099)/0.0039, were expressed as naringin equivalent (GAE). Polyphenols were present in A. helianthus (186.24 ± 5.01). Flavonoids have been reported to indicate a wide range of biological effects containing antibacterial, antiviral, anti-inflammatory, and anti-allergic among others (Cao and Prior, 1999). The content of extractable flavonoids in solvent extracts of AHL, determined from regression equation of calibration curve y = (x + 0.0099)/0.0039, were expressed as naringin equivalent (Table 1). It is important that flavonoid content of plants may diverse in plant species and the morphology or plant part used. No other studies have shown that flavonoids are the important bioactivity compounds in A. helianthus and are responsible for their antioxidant activity. The flavonoids present in AHL may contribute to the antioxidant activity (radical scavenging activity) in HepG2 cells and Raw 264.7 cells.

The DPPH free radical scavenging capacity of AHL is showed in Table 2. AHL showed the highest DPPH scavenging percentage 60.35% at 1000 mg/mL. Thanks to polyphenolic compounds, the most plants have antioxidant effects and more effective antioxidants in vitro than ascorbate (Vitamin C) (Shahidi, 2000). The correlation between the scavenging activity and the polyphenol content of AHL were highly significant scavenging capacity against DPPH free radical. So it can be assumed that the polyphenols in AHL is major contributors to its DPPH scavenging capacity, hence its antioxidant activity.

Table 3 shows reducing power based on their ability to reduce ferric (Fe³⁺) to ferrous (Fe²⁺) ion through the donation of an electron, with the resulting Fe²⁺ formation monitored spectrophotometrically at 700 nm (Hassas-Roudsari et al., 2009). Overall, result indicates that AHL had the strongest reducing power of the antioxidants activity at any given concentration. AHL results from the present investigation (125–1000 μg/mL) exhibited higher reducing power due to the presence of higher concentration. So, absorbance increased dependent on concentration.

As shown in Fig. 1 cell viability was maintained more than 99% after treatment of AHL at concentration of 2 mg/mL. Therefore, our AHL not showed cell toxicity even the concentration of 1 mg/mL in

| Table 1 | Extract yield, total phenol and flavonoid contents in ethanol extract of A. helianthus LEAVES. |
|---------|-----------------------------------------------|
| Sample  | Extract yield (%) | Total phenol (mg GAE/g)* | Total flavonoid (mg NAR/g)# |
| A. helianthus LEAVES. | | | |
| 23.3 | 186.24 ± 5.01 | 102.53 ± 0.98 |

All values are mean ± SD of triplicates analysis. SD: Standard deviations, GAE: gallic acid, NAR: naringin, AHL: Extract of Aurea helianthus leaves.

* Total phenol content analyzed as GAE equivalents.
# Total flavonoid content analyzed as NAR equivalents.

| Table 2 | DPPH and ABTS scavenging activity of ethanol extract of A. helianthus LEAVES. |
|---------|-------------------------------------|
| Sample  | Assay | Concentration (µg/mL) | RSA (%) |
| A. helianthus LEAVES | | | |
| 100 | ABTS | 19.85 ± 0.33 |
| 250 | 38.32 ± 0.09 |
| 500 | 65.39 ± 0.56 |
| 750 | 86.94 ± 0.48 |
| 1000 | 98.30 ± 0.18 |
| Ascorbic acid | | | |
| 1000 | DPPH | 7.92 ± 0.11 |
| 250 | 14.92 ± 0.11 |
| 500 | 34.06 ± 0.28 |
| 750 | 51.13 ± 0.11 |
| 1000 | 60.35 ± 0.29. |
| Ascorbic acid | | | |
| 1000 | | 99.35 ± 0.39 |

All values are mean ± SD of triplicates analysis. SD: Radical scavenging activity, DPPH: 1,1-diphenyl-2-pycrylhydrazyl, ABTS: 2’-azino-bis(3-ethylbenzothiazoline-6-sulphonic acid), 1: Extract of Aurea helianthus leaves.
H₂O₂ treated HepG2 cells. The cytotoxicity can be measured by assessing cellular damage (Tsuchiya, 2010). Cytotoxic effect of AHL on the proliferation of human liver hepatoma cells (Hep G2) was assessed with the MTT assay. Results showed the AHL capacity to inhibit the proliferation of the cancer cells in a concentration dependent manner (Fig. 1A). At 5 mg/mL, the AHL was capable of scavenging 40% of cancerous cells. So it can be assumed that the cytotoxic effect of AHL on the proliferation of the HepG2 cancer cells is due to their antioxidant capacity which interfere with important biological processes in cell development. Polyphenols owe their antioxidant properties to their ability to pair the unpaired electron of the free radical.

Cell viability in HepG2 cells treated with AHL for 24 h. All values are means ± SD from three independent experiments. (⁎P < .01, ⁎⁎⁎P < .0001; significant effect between the control without H₂O₂ and control with H₂O₂, *P < .01, **P < .001, ***P < 0.0001; significant effect between the control with H₂O₂ and treated extract with H₂O₂) AHL: Extract of A. helianthus leaves.

We studied AHL cytotoxic effect on cell growth and viability in RAW 264.7 macrophage cells with MTT assay. Macrophage cells are important immune cells in the host defense against infections and cancer cells and they activated when they encounter antigens like as pathogens or cancer cells (Medzhitov and Janeway, 1997). RAW 264.7 cells were used to assess the effect of AHL on inflammatory mediator synthesis. AHL did not affect normal cell growth and various concentrations of AHL (0, 50, 10, 200, 400, 800 and 1000 μg/mL) were applied to RAW 264.7 macrophages for 24 h (Fig. 3).

Cell viability of AHL in Raw 264.7 cells treated with or without LPS (1 μg/mL) for 24 h. All values are means ± SD from three independent experiments. (⁎P < 0.1, ⁎⁎P < .05, ⁎⁎⁎P < .001; significant effect between the control with LPS and treated extract with LPS.) AHL: Extract of A. helianthus leaves.

Macrophage cells produce several pro-inflammatory mediators interleukin-1β, tumor necrosis factor-α, or NO (Beutler, 1999). The process of acute inflammation is initiated by resident immune cells already present in the involved tissue, mainly resident macrophages, dendritic cells, histiocytes, Kupffer cells and mast cells. We first investigated the inhibitory influence of A. helianthus and assessed the anti-inflammatory effect in LPS-stimulated RAW 264.7 cells as an in vitro model. Production of NO was detected following the treatment with various concentrations of AHL. AHL showed dose-dependent inhibition of LPS-induced NO production at 200 μg/mL (44%). In contrast, pretreatment of cells with AHL led to a significant decrease in LPS-induced NO production (Fig. 4).

NO production of AHL in Raw 264.7 cells treated with or without LPS (1 μg/mL) for 24 h. All values are means ± SD from three independent experiments. (⁎P < .05, ⁎⁎P < .01, ⁎⁎⁎P < .001; significant effect between the control and treated extract) AHL: Extract of A. helianthus leaves.

We performed DCF-DA assay for measuring ROS production in HepG2 cells. H₂O₂-induced ROS generation was increased up to 1.4-fold when compared with control group (Fig. 2). In contrast AHL effectively reduced intracellular ROS production to more than 20 μM of H₂O₂ treated cells.

Intercellular ROS generation were measured by DCF-DA treated with AHL for 30 min and 1 h with H₂O₂ (20 μM) in HepG2 cells. All values are means ± SD from three independent experiments. H₂O₂ treated for 30 min, H₂O₂ treated for 1 h (⁎⁎⁎P < .0001; significant effect between the control without H₂O₂ and control with H₂O₂, *P < .01, **P < .001, ***P < .0001; significant effect between the control with H₂O₂ and treated extract with H₂O₂). AHL: Extract of A. helianthus leaves.

Table 3
Reducing power of AHL A. helianthus leaves.

| Sample | Concentration (μg/mL) | Absorbance (700 nm) |
|--------|-----------------------|---------------------|
| AHL    | 125                   | 0.12 ± 0.01         |
|        | 250                   | 0.18 ± 0.03         |
|        | 500                   | 0.27 ± 0.02         |
|        | 1000                  | 0.47 ± 0.03         |

All values are mean ± SD of triplicates analysis. SD: Standard deviations, AHL: Extract of Aurea helianthus leaves.
independent experiments. (**P < .01; significant effect between the control without LPS and control with LPS. "P < .1, "**P < .001; significant effect between the control with LPS and treated extract with LPS.) AHL: Extract of A. helianthus leaves.

Previous research did not demonstrated the presence of antioxidants in different from the A. helianthus, which are being used in China as traditional and modern medicinal herbs and supplements. The yield, total polyphenolic and flavonoid content, and the antioxidant properties in vitro of AHL extract has not been reported. These results remain important as the first step in screening the antioxidant activity of A. helianthus.

4. Conclusion

In vitro antioxidant activity represents only a status in the basic assessment of A. helianthus pharmacological activity. Further evaluation of specific flavonoid present in the leaf is ensured. Evaluation of the antioxidative properties of AHL in vivo is needed to ensure functionality and long term safety.

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Conflict of interest

The authors have declared that no competing interests exist.

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