INTRODUCTION

*Panax ginseng* (P. ginseng) C.A. Meyer (Araliaceae) is a valuable herb in East Asia that has also gained popularity in the West due to its pharmacological properties for various diseases including atherosclerosis, liver dysfunction, cerebrovascular disease, hypertension, and post-menopausal disorder [1]. In addition, total crude extract from ginseng included in dermatological formulations was reported to have several beneficial effects on human and animal skin [2-4]. Other studies report that ginsenoside Rb1 [5] and Rb2 [3] promote burn wound healing, and topical applications of ginsenoside compound K might prevent or improve skin deteriorations such as xerosis and wrinkles, partly ascribed to age-dependent decreases in human skin [6]. Furthermore, research has shown the protection of red ginseng extract against acute UVB-irradiated skin aging, such as in-
creases in skin thickness and pigmentation and reductions in skin elasticity [7].

Cultivated ginseng is systematically farmed on open land and harvested after a 5 to 6 year cultivation period. Many are aware of the value of ginseng roots, and several studies have been conducted to elucidate the unexplained healing activities of ginseng. Since most studies have focused on ginseng roots, scientists show less interest in ginseng leaves. Yet, it is possible to harvest ginseng leaves every year unlike ginseng roots. In addition, groups of scientists have found that ginseng leaves are rich in polysaccharides, polyphenol, flavonoids, and ginsenosides [8-10]. Shi et al. [11] reported that the content of ginsenoside Re in *P. ginseng* leaves and root-hairs is about five times higher than that of Rg1, but Re content is lower than that of Rg1 in the root. They also suggested that *P. ginseng* leaf may be more beneficial than other plant parts. Rg1, Re, and Rb1 are the three main ginsenosides in extracts of *P. ginseng* roots. Ginseng leaf is known to have six major ginsenosides, Rb1, Rb2, Re, Rd, Re, and Rg1 [12]. It is generally believed that dammarane saponins in ginseng leaf play major roles in its antioxidant activity in animal models [13]. In addition, the polyphenols and flavonoids in ginseng leaf exhibit very potent antioxidant activities [9]. Recent studies have employed new technologies to maximize the herbal benefits of ginseng, and have focused on the development of new ginseng health products [14]. Powders and extracts from ginseng roots have been used to make cosmetics [15]. However, products incorporating ginseng roots are typically too costly because ginseng is not only expensive, but also difficult to cultivate. The objective of this study was to evaluate the potential use of ginseng leaf as a cosmetic material. In this research, we employed enzymatic treated ginseng leaf by utilizing Ultraflo L, which has side activities such as cellulose, β-xylanase and α-arabinofuranosidase to improve ginsenoside recovery. We also studied the bioavailability and skin safety of the enzymatic treated ginseng leaf as a cosmetic material.

**MATERIALS AND METHODS**

**Chemical material and plant material**

Standard ginsenosides including compound K, Rh2, Rk1, Rg5, Rg3, F2, Rd, Rh2, Rc, Rb1, Rg2(s), Rg2(r), Rh1, Rf, Re, and Rg1 were purchased from Ambo Institute (Seoul, Korea). Folin Ciocalteu’s phenol reagent, 2,2-diphenyl-2-picrylhydrazyl hydrate (DPPH), and 2,2'-azino-bis(3-ethylbenothiazoline-6-sulfonic acid) diammonium salt (ABTS) were purchased from Sigma Chemical Co. (St. Louis, MO, USA). The Ultraflo L was purchased from Novozymes A/S (Bagsvaerd, Denmark). Ultraflo L is a b-glucanase (endo-1,3(4))-), derived from *Humicola insolens*, and has side activities such as cellulose and b-xylanase. This enzyme apparently contains significant levels of one or more a-arabinofuranosidase side activities [16]. And then, Ultraflo L standardized to contain 45 FBG/g (fungal beta glucanase units). Cultivated leaves of Korean *P. ginseng* were collected from a good agricultural practice cultivation garden in Ganghwa-do, which is the major ginseng producing area of Korea. The Korean ginseng leaf was freeze-dried, made into a fine powder, and stored at -70°C until further use.

**Preparation of Korean ginseng leaf extraction**

The Korean ginseng leaf powder (3 g) was rehydrated with distilled water (DW, 50 mL) and 0.5 wt% Ultraflo L was added at 40°C for 12 hours to hydrolyze cellulose and pectin in the ginseng leaf [16-18]. The hydrolyzed mixture was extracted twice with 150 mL of ethanol under reflux in a water bath at 90°C for 2 hours. The extract was then centrifuge at 10,000×g for 30 minutes. The supernatant was evaporated and then lyophilized. The non-enzymatic treated ginseng leaf (3 g/50 mL DW) was incubated without enzyme at 40°C for 12 hours and then extracted.

**Determination of chemical composition and ginsenoside content of ginseng leaf extracts**

The amount of total carbohydrate was measured by the phenol-sulfuric acid method [19] using glucose as the respective standard. Uronic acid was estimated by the 3-phenylphenol method [20] using galactronic acid as a standard. The total polyphenol and total flavonoid contents of the extracts were determined by employing Folin-Ciocalteu and p-dimethylamino-cinnamaldehyde reagents, respectively, using protocols reported elsewhere [21]. The total phenolic compound and flavonoid contents were expressed as gallic acid equivalents and catechin equivalents, respectively.

The levels of 16 major ginsenosides were analyzed using an HPLC-based technique developed by Lee et al. [22]. The utilized HPLC system was a Varian Prostar 200 HPLC system (Varian Inc., Palo Alto, CA, USA) equipped with a quaternary solvent delivery system, an auto-sampler, and UV detector, measuring at 203 nm. The column was a IMtakt Cadenza CD-C18 column (4.6×75 mM; IMtakt Corporation, Kyoto, Japan). The peak identifications were based on retention times and
comparisons with injected standard samples.

**Antioxidant activities of ginseng leaf extracts**

Antioxidant activity was measured by DPPH radical scavenging activity and ABTS radical scavenging activity. The DPPH radical scavenging activity was measured by the method described by Quang et al. [23]. ABTS radical scavenging activity was determined as described by Wang and Xiong [24]. All tests were performed in triplicate. The antioxidant activities of the test samples were expressed as the median effective concentration for radical-scavenging activity (IC₅₀), i.e., the amount of tested extract required for a 50% decrease in absorbance of DPPH radicals or ABTS radicals.

**Determination of ginsenoside content of ginseng leaf extracts**

The levels of 16 major ginsenosides were analyzed using an HPLC-based technique developed by Lee et al. [22]. The utilized HPLC system was a Varian Prostar 200 HPLC system equipped with a quaternary solvent delivery system, an auto-sampler, and UV detector, measuring at 203 nm. The column was an IMtakt Cadenza CD-C18 column. The peak identifications were based on retention times and comparisons with injected standard samples.

**Cosmetic formulations**

The formulations that were investigated were based on essence types, as described in Table 1. The formulations included supplemented basic ingredients contained in base essence (BsE), 0.04% adenosine contained in base essence (AdE), 2% arbutin contained in base essence (ArE), and 10% Ultraflo L treated ginseng leaf extract (UTGL) contained in base essence.

**Table 1. Composition of four cosmetic formulations**

<table>
<thead>
<tr>
<th>Formulations (%)</th>
<th>BsE</th>
<th>AdE</th>
<th>ArE</th>
<th>UTGL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deionized water</td>
<td>61.5</td>
<td>62.5</td>
<td>59.5</td>
<td>51.5</td>
</tr>
<tr>
<td>Carborner (1%)</td>
<td>30</td>
<td>30</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>Hyaluronic acid</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Beta glucan</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Triethanol amine</td>
<td>0.3</td>
<td>0.3</td>
<td>0.3</td>
<td>0.3</td>
</tr>
<tr>
<td>Methyl paraban</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
</tr>
<tr>
<td>DS-49</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
</tr>
<tr>
<td>Active ingredient</td>
<td>-</td>
<td>0.04</td>
<td>2</td>
<td>10</td>
</tr>
<tr>
<td>Total</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
</tbody>
</table>

BsE, base essence type cosmetic formulations; AdE, 0.04% adenosine essence type cosmetic formulation (BsE+adenosine); ArE, 2% arbutin essence type cosmetic formulation (BsE+arbutin); UTGL, 10% Ultraflo L treated ginseng leaf (BsE+UTGL extracts); DS-49, disodium-2,2’-dihydroxy-4,4’-dimethoxybenzophenone sulphonic acid.

**Topical treatment and UVB irradiation**

UVB irradiation assays were performed on HRS/J strain male hairless mice weighing 25 to 30 g according to the method of Marquele-Oliveira et al. [25], with some modifications. The experimental protocol was reviewed and approved by the Korea University Animal Care Committee (KUACUC-2010-106).

The skin of the mice was irradiated by UVB irradiation using an ultraviolet lamp, a model TL/12RS 40 W Philips. This particular source emits in the range of 270 to 400 nm with an output peak at 313 nm, and is equipped with UVB and UV detectors. A 100 mg amount of sample was applied 3 times a day for 7 days on the upper side of the animals and then the animals were irradiated after 3 days. During irradiation, the back was irradiated at 5.2075×10⁻² W/cm² at a distance of 40 cm as measured by a 1,700 radiometer (International Light Inc., Japan). The animals were fed a standard diet and allowed to drink water ad libitum. They were housed in cages with a 12-hour light and 12-hour dark cycle. The animals were irradiated within their cages. The mice were sorted into a UVB group group (UVB+no essence) as a negative control group, BsE group (UVB+BsE), AdE group (UVB+AdE), ArE group (UVB+ArE), UTGL group (UVB+ArE), and a No UVB group (No UVB+No essence) as a positive control group.

**Evaluation of photoprotective effects**

Skin wrinkling in each group was evaluated by blinded investigators using a grading scale (Table 2) based on the experimental model proposed by Bissett et al. [26].

The animals were killed by decapitation 6 hours after the last UVB exposure on 7 days and the full thickness of the dorsal skin was removed. The skin was washed in 0.15 M NaCl solution, blotted onto filter paper, weighed,

**Table 2. Grading scale of mouse skin wrinkling**

<table>
<thead>
<tr>
<th>Grade</th>
<th>Description of skin</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Numerous fine striations covering back and flanks of body. Fine striations run length of body (head-to-tail direction) and appear and disappear with motion</td>
</tr>
<tr>
<td>1</td>
<td>All fine striations on back along spine are gone. A few shallow coarse wrinkles across back (run perpendicular to head-to-tail direction) which appear and disappear with motion</td>
</tr>
<tr>
<td>2</td>
<td>All fine striations gone. Some coarse wrinkles across back (run perpendicular to head-to-tail direction) which are permanent</td>
</tr>
<tr>
<td>3</td>
<td>All fine striations gone. Several deep coarse wrinkles across back (run perpendicular to head-to-tail direction) which are permanent</td>
</tr>
<tr>
<td>4</td>
<td>Deep-wrinkled, leathery skin with some flesh-coloured lesions</td>
</tr>
</tbody>
</table>
Skin safety of Ultraflo L treated ginseng leaf

Skin safety was evaluated according to the guideline for toxicity of functional cosmetics (KFDA Notification 2008-75) and the experimental protocol was reviewed and approved by the Korea University Animal Care Committee (KUACUC-2010-106). Hartley strain guinea pigs weighing 300 to 400 g were obtained from Nara biotech (Seoul, Korea). They were individually housed in stainless cages.

The primary irritation test was conducted according to the method of Draize et al. [30]. Each animal was clipped free of hair with an electric clipper and shaved with an electric razor 1 day before testing. A 0.1 mL amount of 10% UTGL (vehicle: saline) was spread on cotton lint (1.5 × 1.5 cm) and then lints were applied to two sites (one site intact, and the other abraded with a sterilized syringe needle). The application sites were covered and wrapped with an elastic adhesive bandage (3M Transpore®; 3M Health Care, St. Paul, MN, USA). Approximately 24, 48, and 72 hours after application, the animals were examined for signs of irritation. The skin reactions were evaluated in accordance with the following Draize method [30]: (1) Erythema and eschar formation: Score 0, no erythema; Score 1, very slight erythema; Score 2, well-defined erythema; Score 3, moderate to severe erythema; Score 4, severe erythema and slight eschar formation, and (2) Edema formation: Score 0, no edema; Score 1, very slight edema; Score 2, slight edema; Score 3, moderate edema; Score 4, severe edema.

The primary irritation index (PII), the sum of the scored reactions (both for erythema/eschar and edema formation) for all animals at 72 hour divided by animal number, was calculated following test completion. Body weights were measured on the day before administration and at test termination for all animals.

Skin sensitization was evaluated according to the maximization test method, as described by Magnusson and Kligman [31]. The first stage of the induction was performed as follows. On 0 day (D-0), the first day of the first stage of induction, (i) 0.1 mL of injection solvent+Freund’s complete adjuvant emulsion (1:1); (ii) 0.1 mL sample; and (iii) 0.1 mL sample+Freund’s complete adjuvant emulsion (1:1) were injected intra-dermally at the four corners of the clipped and shaved induction area. After 48 hours (D-2), these patches were removed, and the test sites were gently sponged to remove any remaining sample residue. On day D-6, 0.1 mL of 10% sodium lauryl sulfate (Sigma Chemical Co.) in petrolatum was applied to this area. In the second stage of induction, day D-7, 0.1 mL of each tested material was applied to the area and patched for 48 hours. On day D-22, the animals, whose back hair had been clipped and shaved on day 21, were exposed to a challenge dose. A 0.1 mL amount of sample was applied osseclusively to a challenge area and patched in a similar way as for the inductions. After 24 hours, the patches were removed, and the test sites were gently sponged to remove any remaining sample residue. Skin reactions were evaluated visually 24 and 48 hours after patch removal according to a well-defined sensitization scoring system [32]. Skin reaction: Score 0, no visible changes; Score 1, discrete or patchy erythema; Score 2, moderate and confluent erythema; Score 3, intense erythema and swelling.

Statistical analysis

All expressed values are the means of triplicate determinations. All statistical analyses were performed using the SPSS ver. 12.0 (SPSS Inc., Chicago, IL, USA). The statistical significance of differences was determined using unpaired Student’s t-tests and one-way ANOVA, evaluating significant differences at p-values<0.05. All data are at the 5% significance level and are reported as means±SD.

RESULTS

Chemical composition and ginsenoside content of ginseng leaf extract

In a preliminary study, ginseng leaf was treated with 8 enzymes (Celluclast 1.5 L, Cytolase PCL5, Econase CE, Optidex L-400, Pectinex 5XL, Rapidase TF, Ultraflo L, and Viscozyme L) to improve its yield of ginsenosides. The Ultraflo L treated ginseng leaf extract had
the highest ginsenoside content (data not shown). The chemical composition and ginsenoside compositions of non-enzymatic treated ginseng leaf (NEGL) and UTGL were compared (Table 3 and Fig. 1). Total sugar, uronic acid, and polyphenol, flavonoid contents were higher in UTGL than NEGL. The total ginsenoside contents of NEGL and UTGL in ginseng leaf extract were 271 mg/g and 406 mg/g, respectively. In addition, the level of metabolite ginsenosides (sum of Rg2, Rg3, Rg5, RK1, compound K, Rh1, Rh2, and F2) was higher in UTGL (93.1 mg/g) than in NEGL (62.4 mg/g). The increased rate of ginsenoside types in UTGL compared to NEGL was generally 140% to 157%.

**Antioxidant activities**

As shown in Table 3, total polyphenol was higher in UTGL (33.68 µg/mg) than NEGL (23.22 µg/mg). The antioxidant properties of polyphenols and flavonoids are due to their redox properties and the ability to chelate metals and quench singlet oxygen [33]. The radical scavenging activities are summarized in Fig. 2. UTGL exhibited relatively higher DPPH (IC$_{50}$ 2.8 mg) and ABTS (IC$_{50}$ 1.6 mg/mL) radical scavenging activities than UTGL (4.8 mg/mL and 2.2 mg/mL, respectively).

**In vivo study to evaluate photoprotective effects**

The UTGL only showed increased yields without conversion of ginsenosides. Therefore, we evaluated the photoprotective effects of an essence type formulation containing 10% UTGL (total ginsenoside concentration was 4.06%). We used 0.04% AdE and 2% ArE contained in base essence as positive controls. UV exposure, particularly UVB, causes the generation of free radicals and related reactive oxygen species (ROS), such as H$_2$O$_2$. ROS generated as a consequence

Table 3. Chemical composition of ginseng leaf

<table>
<thead>
<tr>
<th></th>
<th>NEGL</th>
<th>UTGL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total sugar (mg/mg)</td>
<td>0.47±0.04</td>
<td>0.67±0.06</td>
</tr>
<tr>
<td>Uronic acid (µg/mg)</td>
<td>36.21±1.70</td>
<td>51.23±0.29</td>
</tr>
<tr>
<td>Polyphenol (µg/mg)</td>
<td>23.22±0.44</td>
<td>33.68±0.13</td>
</tr>
<tr>
<td>Flavonoid (µg/mg)</td>
<td>0.10±0.00</td>
<td>0.16±0.00</td>
</tr>
</tbody>
</table>

Values are presented as mean±SD (n=3). NEGL, non-enzymatic treated ginseng leaf; UTGL, Ultraflo L treated ginseng leaf.

**Fig. 1.** Gensenoside content of ginseng leaf. The levels of 16 major ginsenosides were analyzed using an HPLC system, IMtakt Cadenza CD-C18 column and UV detector at 203 nm. The peak identifications were based on retention times and comparisons with injected standard samples. Metabolites were the sum of Rg2, Rg3, Rg5, RK1, compound K, Rh1, Rh2, and F2. Each contents that represents mean±SD (n=3). NEGL, non-enzymatic treated ginseng leaf; UTGL, Ultraflo L treated ginseng leaf.

**Fig. 2.** Antioxidant activities of ginseng leaf. Antioxidant activity was measured by 2,2-diphenyl-2-picrylhydrazyl hydrate (DPPH) radical scavenging activity and 2,2’-azino-bis(3-ethylbenzothiazoline-6-sulfonic acid) diammonium salt (ABTS) radical scavenging activity. All tests were performed in triplicate. The antioxidant activities of the test samples were expressed as the median effective concentration for radical-scavenging activity (EC$_{50}$), i.e., the amount of tested extract required for a 50% decrease in absorbance of DPPH radicals or ABTS radicals. Each amount represents means±SD (n=3). NEGL, non-enzymatic treated ginseng leaf; UTGL, Ultraflo L treated ginseng leaf.
UVB exposure produce oxidative stress, intracellular GSH depletion, and increased products of lipid peroxidation. This may be the reason for wrinkle formation in photoaged skin [34,35]. As shown in Fig. 3, the base essence group (BsE) had reduced hydrogen peroxide formation and a lower visual wrinkling grade compared to the no cream group since the BsE formulation contained some available compound. However, treatments of active ingredients such as adenosin or arbutin were significantly more effective than BsE. The UTGL group showed normalized hydrogen peroxide and lipid peroxidation induced-UVB exposure that was similar to the non exposure group. Finally, in terms of visual wrinkling grade, the UTGL group showed a similar tendency to the non-UVB treated control group with anti-wrinkles effects against UVB exposure.

Primary irritation and skin sensitization testing of UTGL

The skin safety of cosmetic ingredients is more important than their function. Therefore, we evaluated the safety of UTGL using primary irritation and skin sensitization tests. Table 4 shows the results of the primary skin irritation test. The guinea pigs had 10% UTGL dermally attached for 24 hours. The UTGL did not induce any adverse reactions such as erythema and edema on intact skin sites; however, some guinea pigs treated with UTGL on abraded skin sites showed very slight erythema. The PII score of UTGL was 0.05 and it was classified as a practically non-irritating material (PII, 0

![Fig. 3. Effects of cream type formations with added active ingredients on hydrogen peroxide level (A), reduced glutathione level (B), lipid peroxidase level (C), and visual wrinkling grade (D) in UVB-irradiated hairless mice. Different letters at each measurement indicate significant differences among groups (p<0.05). Each bar represents mean±SD (n=5). BsE, base essence; AdE, BsE+adenosine; ArE, BsE+arbutin; UTGL, BsE+Ultraflo L treated Korean ginseng leaf extracts; GSH, reduced glutathione; TBARS, thiobarbituric acid reactive substance.](image-url)
In the skin sensitization test, the guinea pigs were sensitized with intradermal injections of 10% UTGL for 48 hours. After 1 week, they were treated on the injection site with the same concentration and then challenged with a 1% solution 2 weeks later, and observed for 24, 48, and 72 hours.

Table 5 shows the skin sensitization test results of the guinea pigs. In the negative control group (saline), the positive rate after evocation was 0% and the mean evaluation score was 0. In the UTGL group, the positive rate 72 hours after evocation was 40%, and the mean evaluation score was 0.4. In the positive control dinitrochlorobenzene group, the positive rate was 100% and the mean evaluation score was 2.2.

**DISCUSSION**

*P. ginseng* has been shown to possess a wide range of biological and pharmacological activities and is used as a general tonic in traditional oriental medicine to increase vitality, health, and longevity, especially in older
persons. Previous research has reported that ginseng has several beneficial effects on human and animal skin [3-7]. Kim et al. [36] reported that topical applications of total ginseng saponins (10 pg or 100 ng/mouse) and ginsenoside Rb1 (100 fg, 10 pg, or 1 ng/mouse) isolated from red ginseng significantly inhibited increases in skin thickness and wrinkle formation and reduced skin elasticity induced by long-term UVB irradiation. Furthermore, Lee et al. [37] reported that the topical administration of a cream base containing red ginseng (percentage of ginseng saponins was about 3.3%) for 22 weeks resulted in significantly decreased wrinkling scores, minimal epidermal hyperplasia, slightly increased dermal cellularity, and a lack of cyst proliferation by UV-irradiation. Thus, ginseng root powder and extracts have been used to make cosmetics [15]. However, products containing ginseng root ingredients are too costly because ginseng is not only expensive, but also difficult to cultivate. Ginseng should be harvested after a 5 to 6 year cultivation period. However, it is possible to harvest ginseng leaves every year, unlike ginseng roots.

Traditionally, ginseng leaf has been consumed mostly in the form of tea. Ginseng leaf is rich in polysaccharides, polyphenols, and flavonoids [8-10], and contents of some ginsenosides are higher than in the roots [11]. However, these compounds are tightly linked with carbohydrates such cellulose, pectin, or β-glucan. We prepared enzymatic treated ginseng leaf using Ultraflo L to improve ginsenoside recovery. Ultraflo L is a β-glucanase (endo-1,3(4)-) and it has side activities such as cellulose, β-xylanase, and α-arabinofuranosidase. The observed increase in polyphenol content in UTGL was accompanied by increases in flavonoid content, antioxidant activities, and ginsenoside contents as compared to NEGL.

The ginsenoside Rb2 has been reported to improve wound healing, stimulate epidermal cell proliferation, and enhance the expression of protein factors related to cell proliferation, namely, epidermal growth factor and its receptor fibronectin, and its receptor keratin 5/14, as well as collagenase I [3]. In the present study, the ginsenoside Rb1 and Rb2 contents of the ginseng leaf treated with Ultraflo L were 9.50 mg/g and 20.30 mg/g, and in the non-enzymatic treated ginseng leaf they were 6.86 mg/g and 14.5 mg/g, respectively.

UV exposure causes oxidative stress, intracellular GSH depletion, increased products of lipid peroxidation, and ultimately wrinkle formation of the skin [34,35]. The ginseng leaf treated with Ultraflo L (147.9 μm/g and 10.8 μm/g) normalized hydrogen peroxide and lipid peroxidation induced-UVB exposure compared with no cream (716.2 μm/g and 43.59 μm/g). We assume that these effects were due to its antioxidant activity through polyphenols and flavonoids [33,38]. In this study, the polyphenol and flavonoid contents of the ginseng leaf treated with Ultraflo L were 33.7 μg/g and 0.16 μg/g, and in the non-enzymatic treated ginseng leaf they were 23.2 μg/g and 0.1 μg/g, respectively. UTGL exhibited relatively higher DPPH and ABTS (IC50, 2.8 mg and 1.6 mg, respectively) radical scavenging activities than UTGL (4.8 mg and 2.2 mg, respectively). Many natural substances such as carotenoids, tocopherols, and polyphenols are able to act as antioxidants and are widely contained in foods and plants. Flavonoids and other polyphenols have the ability to scavenge free radicals, and therefore, delay lipid oxidation [38].

Ginseng may cause a very serious skin reaction called Stevens-Johnson syndrome (SJS). SJS is a life-threatening condition affecting the skin in which cell death causes the epidermis to separate from the dermis. The syndrome is thought to be a hypersensitivity complex affecting the skin and the mucous membranes [39]. In our results, the 10% ginseng leaf treated with Ultraflo L did not induce any adverse reactions such as erythema and edema on intact skin sites, but some guinea pigs treated with UTGL on abraded skin sites showed very slight erythema. UTGL also showed skin sensitization by using guinea pig maximization testing. It is difficult to determine the reasons between dosage and self-side effects. But 10% is an extremely high dosage of active ingredient in a cosmetic.

In this study, ginseng leaf treated with Ultraflo L had increased contents of polyphenols and ginsenosides compared to non-enzymatic treated ginseng leaf. The ginseng leaf treated with Ultraflo L had anti-wrinkles effects against photo-damage. However, 10% ginseng leaf treated with Ultraflo L showed adverse skin safety effects in skin sensitization tests. Therefore, it is considered that ginseng leaves treated with Ultraflo L retain an optimal usage range. From these results, in terms of cost and source availability, ginseng leaves treated with Ultraflo L have advantages over untreated ginseng leaves and have potential as a cosmetic ingredient.

ACKNOWLEDGEMENTS

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REFERENCES


