Skin Whitening Effects of Angelica koreana and Cnidium monnieri Extracts

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Abstract The objective of this study is to estimate the inhibition of skin melanin formation by extract of Angelica koreana and Cnidium monnieri and the possibility of functional cosmetic materials through anti-irritation and stability test. The extract used in this experiment is White-AK™ and the INCI name is Osthole. The main component of White-AK™ was identified as coumarin and EC50 value was 2.7ppm by mouse melanoma B 16 cell test. White-AK™ showed inhibitory effects 100 times lower concentration than arbutin. The main mechanism for skin whitening effect thought to be inhibition of tyrosinase-related gene expression. The basic essence formulation of White-AK™ 5% solution applied to the skin showed the effect of relieving skin irritation. White-AK™ in an opaque container, under UV conditions for 4 weeks, and showed close to 100% recovery and 97% recovery under 50 oC for 4 weeks. Therefore, it is thought that White-AK™ which helps skin whitening, relieving skin irritation and stable in UV condition is able to be used as the functional component in the cosmetic formulation.

Key Words : Angelica koreana, Cnidium monnieri, tyrosinase, skin whitening, cosmetics

요약 본 연구의 목적은 강활과 벌사상자 추출물에 의한 피부 멜라닌 형성 저해와 안정성과 자극성 시험을 통하여 기능성 화장품의 사용 물질 가능성을 평가하고자 하였다. 본 실험에 사용된 추출물은 White-AK™이고 INCI 명칭은 Osthole이다. White-AK™의 주성분은 쿠마린이고 EC50 값은 마우스 멜라노마 B16 세포 시험에서 2.7ppm이었다. White-AK™는 알부틴보다 100배 낮은 농도에서도 저해 효과를 보였다. 미백 효과의 주 기작은 티로시나제-관련 유전자 발현을 저해하는 것으로 생각되었다. White-AK™ 5% 용액을 함유한 기본적인 에센스 처방을 피부에 적용 시 피부자극을 경감시켰다. White-AK™은 불투명한 용기에서 4주간 자외선 하에서 100%의 회수율을 보였고 50 oC에서 4주간 97%의 회수율을 보였다. 따라서 피부 미백효과와 자극성 경감효과가 있고 자외선 조건하에서 안정한 White-AK™은 화장품 처방에서 기능성 원료로 사용될 수 있을 것으로 사료된다.

주제어 : 강활, 벌사상자, 티로시나제, 피부 미백, 화장품

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1. Introduction

In recent years, the number of Asian women aspiring for a white skin complexion has increased dramatically. This is due partly to the discovery of many effective skin whitening agents, particularly those derived from natural sources such as arbutin, kojic acid, and licorice extract. Implementation of a wide range of consumer desire for skin color is possible due to the development of computer graphic and algorithm [1][2].

Promotion of these substances by the cosmetic industry in various media has played a critical role in the widespread popularity of skin whitening products among women, especially in the Far East and Southeast Asia. Also, it is traditional perception of the women in this region that a lighter skin provides them the appearance of grace, nobility and youthfulness, as evidenced in many Asian poems and songs, which often describe the beauty of a lady in association with the fairness of her skin.

The skin color is mainly determined by the content of an epidermal pigment called melanin. Its major function is to provide protection against ultraviolet (UV) radiation. However, excessive melanin production is not desirable in many people since it may cause a darker or uneven skin color. The initial process of melanin production (melanogenesis) is controlled by tyrosinase, which is an enzyme catalyzing the hydroxylation of tyrosine, the precursor of melanin, into dihydroxyphenylalanine (DOPA) and other intermediates. Thus, inhibition of tyrosinase activity or its production can prevent melanogenesis [3]. Several plant extracts were found to exhibit strong anti-tyrosinase activity in vitro such as those from Areca catechu L. [4], Artocarpus incisus [5], Broussonetia spp. (paper mulberry root bark extract) [6], Glycyrrhiza glabra (licorice extract) [7], Prunus spp. [8], and Rheum officinale [9]. Many of these extracts have been tested in vivo and commercially developed as skin whitening agents in cosmetic preparations such as kojic acid, licorice extract and Morus alba extract [10]. However, they are quite expensive and some of the users may develop skin hypersensitivity upon application especially when using high concentrations.

Artocarpus lakoocha Roxb is a tropical tree about 20-30 meters high commonly found throughout south and southeast Asia. The main constituent in the heartwood extract is 2,4,3',5'-tetrahydroxystilbene or oxyresveratrol [11].

Recently, Sritularak et al. [12] have screened a large number of plants for their in vitro antityrosinase activity and found that the heartwood extract of A. lakoocha exhibited the highest activity. Purification of the extract yielded two active components, namely, oxyresveratrol and resveratrol. The main objective of this study was to evaluate the whitening effect of extracts of Angelica koreana, Cnidium monnieri. The results were subsequently compared with tyrosinase inhibitor commonly used in skin whitening products (arbutin). Furthermore, the stability of the extract in cosmetic vehicles was assessed for the cosmetic application.

2. Materials and Methods

2.1 Chemicals

All tissue culture medium and components were purchased from GIBCO BRL (Long Island, NY USA). MTT, sodium phosphate, HCL and charcoal were from Sigma–Aldrich (St. Louis, MO, USA). Protease inhibitor cocktail was from Boehringer–Mannheim (Indianapolis, IN, USA).

Angelica koreana and Cnidium monnieri were purchased from the oriental medicine market located in Seoul, South Korea. Each of plants was sliced and weighed. 100g of powder from each plant was extracted with 500 mL of ethanol: water (80:20, v/v) at...
room temperature for 7 days. After filtration, this extract was evaporated to dryness under vacuum, and then completely dried by lyophilization. Each extract was fractionated with different solvents (ether, hexane, chloroform, ethylacetate, butanol, and water), and was used as the sample in this study.

The sample (White-AK™) used in this experiment is powder form and the main component was identified as coumarin as shown in [Fig. 1].

[Fig. 1] The chemical structure of coumarin.

2.2 Cells and culture

B16 melanoma cells were purchased from Korean Cell line bank and cultured in DMEM (Dulbecco’s modified eagle’s medium, sigma, D-2902, St. Louis, MO 63178 USA) supplemented with penicillin (100 U/mL), streptomycin (100 U/mL), and 10 % fetal bovine serum (Fetal bovine serum, Gibco, 26140-079, Invitrogen Co.) at 37 °C in an incubator flushed continuously with 5 % CO₂[13].

2.3 Cell treatment

B16 Melanoma cells were seeded 6-well plate at a density of 2.5x10⁴ cells per well and allowed to attach for 24 hr. After then, triplicate cultures were fed with fresh medium containing various concentrations of compound. After 48 hr, the medium was replaced with the same, fresh test medium. After further 48 hr, cells are harvested with 0.5 mL of 0.25% trypsin/EDTA. After dislodging the cells with occasional agitation, 2mL of medium were immediately added to inactivate the trypsin, and 100 mL aliquot were seeded into 96-well plate for MTT assay, as described below. The remainder cell suspensions were centrifuged for 5 min at 2,500g, washed with PBS and then solubilized in 200 mL of extraction buffer (1% Nonidet P-40, 0.01% SDS, 0.1 M Tris–HCl pH 7.2 and protease inhibitor cocktail). Extracts were solubilized at 4 °C for at least 1 hr and then assays were conducted for each sample, in triplicate.

2.4 MTT assay

The test materials were added and incubated in 5% CO₂ incubator at 37°C for 12hrs by the same procedure of NR assay and 0.01 ml of MTT at the conc, of 5mg/ml in PBS was added into well and incubated for 4hrs, and then the residual contents were removed by turn over. To dissolve formazan crystal, isopropanol containing 0.04N HCl was added and stirred for 20min and it was read by ELISA reader in the dual ranges of 570nm and 630nm[4].

2.5 Inhibition of melanization

B-16 melanoma cells were cultured in DMEM supplemented with 10% FBS in humidified incubator at 37 °C under 5% CO₂ in 6well plate at density of 2x10⁴ cells/well. After cells were attached, medium was replaced with DMEM containing 10% FBS, 0.2 mM α-MSH, 2mM theophylline after samples addition. After 4days, trypsin was added and cells were collected by centrifugation. Then cell pellets were dried and dissolved in 1N NaOH. Melanin synthesis inhibition rates were measured at 490 nm using ELISA reader[7].

2.6 Western Blotting

Cells were treated as described above. At the end of each treatment period, cells were washed in PBS and were lyzed in extraction buffer containing 1 % Nonidet P-40, 0.01 % SDS, and the protease inhibitor cocktail. Protein contents were determined with a BCA assay kit (Pierce, Rockford, IL) and equal amounts of each protein extract (10 mg per lane) were resolved using 8
% SDS polyarylamide gel (Koma Biotech, Korea) and transblotted onto nitrocellulose membranes (Amersham, Piscataway, NJ) and the membranes were blocked with 5% nonfat milk in TBS buffer. Following the blocking, the membranes were incubated with a PEP7 (anti-tyrosinase) a PEP1 (anti-TRP1), or a PEP8 (anti-TRP2) (each at a 1:1000 dilution). The membranes were then incubated with HRP-conjugated anti-rabbit IgG at a dilution of 1:2000. Immunoreactive bands were detected with enhanced chemiluminescence using an ECL kit (Amersham, Piscataway, NJ) according to the manufacturer’s instruction[14].

2.7 Skin irritation test

5% of White–AK™ in dipropylene glycol was formulated into control essence and cream vehicles. The final concentration of White–AK™ was 0.2 wt %. Test materials were applied to the upper outer arm of subjects for up to 4 h, following the specific testing format of the test developed for chemical irritation prediction. A positive control was tested in each study. Liquid test materials were applied neat, in a 0.2 ml volume to 25 mm Hill Top Chambers (Hill Top Research, Inc., Cincinnati, OH). The duration of exposure to the test materials increased progressively from 30 min through 1, 2, 3, and 4 h. The sites were graded at 24, 48, and 72 h after patch removal using a 4-point (0–3+) grading scale of increasing irritation as follows[15]:

0: No reaction.
+: Weak positive reaction, mild erythema across most of the treatment site.
++: Moderate positive reaction: distinct erythema across entire treatment site, possibly spreading beyond treatment site.
++++: Strong positive reaction: spreading erythema with edema.

2.8 Statistics

All data were expressed as means ±S.D. The statistical significance for the comet assay was evaluated with student’s t-test and for the clinical study with one-way ANOVA. P<0.05 was considered significant.

3. Results and Discussion

3.1 MTT assay

When selecting whitening agents, one of the important points is that they should have minimal effect on melanocyte cell proliferation. Thus, the proliferation of the cells treated with White–AK™ was evaluated by MTT assay. As shown in [Fig. 2], White–AK™ showed little inhibitory effect on cell proliferation at the tested concentration. This result suggests that White–AK™ can be a safe candidate of whitening agent.

![Fig. 2] White–AK™ did not show any inhibitory effect on cell proliferation. Fibroblast cells were treated with or without White–AK™. After 4 days, cells were harvested. Values are the averages of three determinations ±S.D.

3.2 Skin Irritation Test

We tested primary irritation test of White–AK™ in cosmetic vehicles. The primary skin irritation was normally the main problem of natural extract in application on the skin. We tested several concentration
of White–AKTM When cosmetic essence and cream with White–AKTM was tropically applied, there was no detectable irritation as shown in <Table 1>. Thus White–AKTM could be used as an anti-irritation agent.

(Table 1) Primary skin irritation score of White–AKTM.

<table>
<thead>
<tr>
<th>Vehicles</th>
<th>Concentration of White–AKTM</th>
<th>Irritation Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Essence Type A</td>
<td>-</td>
<td>0.16 ± 0.01</td>
</tr>
<tr>
<td>Essence Type A</td>
<td>0.1 wt%</td>
<td>0 ± 0.01</td>
</tr>
<tr>
<td>Essence Type A</td>
<td>0.3 wt%</td>
<td>0 ± 0.01</td>
</tr>
<tr>
<td>Essence Type B</td>
<td>-</td>
<td>0.02 ± 0.01</td>
</tr>
<tr>
<td>Essence Type B</td>
<td>0.2 wt%</td>
<td>0.01 ± 0.01</td>
</tr>
<tr>
<td>Cream Type</td>
<td>-</td>
<td>0 ± 0.01</td>
</tr>
<tr>
<td>Cream Type</td>
<td>0.2 wt%</td>
<td>0 ± 0.01</td>
</tr>
</tbody>
</table>

3.3 Effect of White–AKTM on pigmentation of B16 melanoma cell

The quantization of the melanin was analyzed through ELISA to examine the White–AKTM for inhibitory activity against melanin synthesis. The result is given [Fig. 3]. The White–AKTM show excellent whitening effects on the B16 melanoma cells. The calculated value of EC50 was 2.7ppm.

3.4 Effect of White–AKTM on expression of tyrosinase

To explore the mechanism responsible for the decreased pigmentation, we examined changes in the proteins levels of three important melanogenic enzymes (tyrosinase, TRP-1, β-actin) using western blotting. The results were shown in Figure 4. When compared with untreated control, 5 and 10ppm White–AKTM inhibited protein expression dramatically without any effect on TRP-1 and β-actin. These results suggest that whitening potency of White–AKTM results mainly from down regulation of expression of tyrosinase.

3.5. Stability of White–AKTM

[Fig. 5] and [Fig. 6] demonstrated the stability of 5 % of White–AKTM dipropylene glycol solution in transparent and opaque bottle during 1 month storage at under outdoor ultraviolet (UV) condition or indoor at 50°C.

[Fig. 4] Western blotting image of specific detection of tyrosinase, TRP-1, β-actin.

[Fig. 5] Recovery percent of 5 % of White–AKTM in DPG solution under outdoor ultraviolet(UV) condition: ●; in transparent bottle, ■ in opaque bottle.

[Fig. 6] Recovery percent of 5 % of White–AKTM in DPG solution at 50°C.
4. Conclusions

The results of this study clearly demonstrated that the extract of Angelica koreana, Cnidium monnieri, was able to reduce melanin formation in B-16 melanoma cell. The major component of extract was coumarin. Comparing to arbutin commonly used in whitening products like kojic acid and licolice extract, White-AK™ showed quite effective efficacy on skin whitening. The mechanism of whitening effect is thought to be down regulation of expression of tyrosinase. From the primary skin irritation test, the extract appears to have a very promising potential for use as a safe, effective and economical whitening agent in the cosmetic industry. More studies were carried out to determine the optimal concentration, stability as well as safety and efficency of White-AK™.

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