Research Article

Evaluation of *Cinnamomum osmophloeum* Kanehira Extracts on Tyrosinase Suppressor, Wound Repair Promoter, and Antioxidant

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1. Introduction

Skin, made up of three layer cells including, epidermis, dermis, and hypodermis, is the largest vertebrates organ in the human body. Human skin is commonly exposed to oxidative stresses from solar ultraviolet (UV) radiation and free radicals as well as its induced cellular reactive oxygen species (ROS) [1, 2], which are the common reasons for tumor genesis or skin aging. To protect skin from UV radiation, skin operates complex defense system including skin thickening, pigment synthesis, and a network of nonenzymatic and enzymatic antioxidative mechanisms [2]. In addition to a significant responsibility, in the prevention of human skin UV-caused damage, which increases melanocytes transfer of melanosomes to keratinocytes, melanin determines skin color [3]. Hyperpigmentation is commonly cared with therapeutic drugs or cosmetics of pigment-reducing or skin-whiten abilities. During the melanin synthesis processes, tyrosinase is classified to be the rate-limiting oxidase at first two steps [4]. It catalyzes the pigments production such as...
eumelanin and phenomelanin. Two types of pigments production were reported, including the L-tyrosine hydroxylation to 3,4-dihydroxy-L-phenylalanine (L-DOPA) and then the L-DOPA oxidation to dopaquinone (a biochemical precursor to pigments) [5]. In the active site for tyrosinase, two copper ions are essential to catalyze colorful pigments or melanin by oxidative stress. To antagonize tyrosinase activity can reduce the syndrome of hyperpigmentation and dermatological disorders.

Skin as the first immune defense line of human plays a noteworthy role in avoiding various biological, chemical, mechanical, and physical damages [1, 2]. Chronic or acute severe injuries on the skin, such as abrasions, burns, leg ulcers, or lesions, in consequence considerable losses of dermal tissues pose huge challenges to the therapeutical processes. Keratinocytes in epidermis and fibroblasts in dermis are the first stop for body protection against external stimulus or for the skin wound healing [6]. In terms of wound healing, wound closure is known to be initiated by fibroblast migration from its margins. Based on the migratory force, resistance from the regenerated tissue may lead to fibroblast differentiation [7], which is featured by the local expression profiles of skin cells, such as several growth factors and the extracellular matrix. Skin wound healing is a cutting edge study for many medicine fields [8].

Smoking factors, salted food, or environmental toxicants bring about various oxidative stresses to human being [9]. The level of excessive free radicals produces a high oxidative stress which is a negative effect against the normal skin and results in aging or some diseases. Through biochemical processes, the intracellular physiological oxidants are engendered from nonenzymatic systems such as those involving enzymatic catalysis, transition metals, various oxidases that transformed them into the reactive nitrogen species, or reactive oxygen species [1]. If antioxidants are invigorated, they can significantly prevent or reduce the oxidative pressure damages [10]. There are several important components constructed to cellular membrane lipids from the phospholipids, membrane proteins, polyunsaturated fatty acids, cholesterol, and nucleic acids [11]. Excessive free radicals and ROS cause oxidative pressure injury on lipids, proteins, and DNA, and the damage eventually induced cellular damage, aging, neural disorders, diabetes, atherosclerosis, inflammatory, cancer, and cardiovascular disease, especially unwanted pigment accumulation [12].

2. Materials and Methods

2.1. Reagents and Materials. All the reagents were purchased from Sigma Chemical (St. Louis, MO), including dimethyl sulfoxide (DMSO), 1,1-diphenyl-2-picrylhydrazyl (DPPH), 3-(4,5-dimetylthiazol-2-yl)-2,5-diphényl, tetrazolium bromide (MTT), 3-tert-butyl-4-hydroxyanisole (BHA), ethylenediaminetetraacetic acid (EDTA), FeCl₃, FeCl₂·4H₂O, kojic acid, L-tyrosine, mushroom tyrosinase, potassium ferricyanide [K₃Fe(CN)₆], trichloroacetic acid and vitamin C, and other highest purity chemical buffers and reagents. Cell culture reagents were purchased from GIBCO BRL (Gaithersburg, MD), including fetal bovine serum (FBS) and Dulbecco’s modified Eagle medium (DMEM).

2.2. Extraction of C. osmophloeum Kanehira Leaves. The plant specimen was authenticated by Ladies Biotech Co., LTD, where voucher specimens were kept. Dry leaves of C. osmophloeum Kanehira (0.6 kg) were sliced and soaked in 3 L ethyl alcohol for one day before further three ethyl alcohol extractions. After filtration, the extracts were evaporated to final weight of 8.49 g.

2.3. B16-F10 Melanoma Cell Cultures. The melanoma B16-F10 cells (BCRC 60031 in ATCC) were maintained at 37°C under 5% CO₂ atmosphere by feeding the medium (10 mM
HEPES, 13.4 mg/mL DMEM, 100 μg/mL streptomycin sulfate, 143 U/mL benzylpenicillin potassium, and 24 mM NaHCO₃, pH 7.1) with 10% FBS [17].

2.4. B16-F10 Cell Viability. MTT assay was used to evaluate the effects of cell viabilities for the treatments of C. osmophloeum Kanehira extracts [17]. Briefly, cells (6 × 10⁵ cells/well) were plated in 96-well plates for overnight. Cells were treated with either vehicle (DMSO) or indicated concentrations of each sample for 24 h. Subsequently, 0.5 mg/mL MTT in 100 μL of fresh medium was used to replace the medium and the reaction was performed in a 37 °C cell culture incubator for 2 h. The generating crystals were dissolved in 100 μL of DMSO with smooth shaking for 10 min in darkness. Finally, the absorbance (A) value of this reaction was detected at 595 nm by multiplate reader (UV-vis, BioTek, Winooski, VT). Cell viability (%) was formulated as follows:

\[
\text{Cell viability} (%) = 100 \times \frac{OD_{\text{sample}}}{OD_{\text{control}}},
\]

2.5. B16-F10 Cellular Tyrosinase Activity. The tyrosinase activity was dependent on the dopachrome formation rate as previously described [17]. Melanoma B16-F10 cells (10⁵ cells/well) were plated in a 12-well plate with 1,000 μL of medium, and they were treated with indicated concentrations of extracts for 48 h. After PBS washing, B16-F10 cells lysed with 1% triton X-100/PBS and 50 μL of 2 mM L-tyrosine were added. Standing for 3 h at 37 °C in darkness, its absorbance at 490 nm was examined spectrophotometrically, where the tyrosinase activity evaluation formula was similar to (2).

2.6. B16-F10 Cellular Melanin Contents. The cellular melanin contents were measured with minor modifications as previously described [17]. Briefly, B16-F10 melanoma cells (2.5 × 10⁵ cells/well/1500 μL of medium) were plated in 6-well plates with extracts and incubated for 48 h. After dissolving in 10% DMSO with 50 μL of 2.0 M NaOH for 1 h at 90 °C, cell lysates were centrifuged at 10,000 × g for 10 min to collect the supernatants for melanin determination using spectrophotometer at 475 nm with similar formula to (2).

2.7. Mushroom Tyrosinase Activity. The mushroom tyrosinase activity was measured with minor modifications as previously described [18, 19]. The various concentrations of extracts were added: 2 μL with 68 μL of 50 mM phosphate buffer (pH 6.8), 10 μL of 0.5 units/mL of mushroom tyrosinase, and 10 μL of the mixture. The absorbance of the mushroom tyrosinase inhibition assay at 490 nm was determined at 5 min per interval until 30 minutes with a 96-well plate spectrophotometer, where kojic acid was regarded as a positive control. Mushroom tyrosinase activity (%) was formulated as follows:

\[
\text{Mushroom tyrosinase activity} (%) = 100 - \left[100 \times \frac{(A - B) - (C - D)}{(A - B)}\right],
\]

where \(A\) is the OD value under no sample; \(B\) is the OD value under no sample and tyrosinase; \(C\) is the OD value under sample; and \(D\) is the OD value under sample but no tyrosinase.

2.8. Animal Experiments. Six-week-old male Wistar rats are used. These rats are kept on standard rat chow and water ad libitum for 1 week before challenging. The animal studies were performed under authorization from the Animal Use Committee of Kaohsiung Medical University. The experimental rats were housed on a 12/12-hour light-dark cycle with the air conditioner and adequate supply of food and water. Twelve rats were grouped into two sets, that is, petroleum jelly and C. osmophloeum Kanehira extracts (experimental groups). The wound healing of skin was measured with minor modifications as previously described [20]. After rats were anesthetized, its dorsal hair was shaved and the wounds of 1 cm in diameter were generated. After a back skin excising, the wounds of all experimental rats were quickly covered with the petroleum jelly (8% DMSO, w/v) or 0.8 mg extracts, where the petroleum jelly was used as a reference.

2.9. Measurement of the Wound Area. Digital camera (Coolpix P6000, Nikon, Japan) was used to record the progression of skin wound after 0, 1, 3, and 5 days with protocol parameters (aperture: F/7.2, shutter speed: 1/60). The wound dressing was not removed under healing period unless the substance was easy to detach manually. The area of each skin wound was determined by SPOT software (Diagnostic Instruments, Inc., Sterling Heights, MI, USA). Some random Sani-Chips were visualized on wound sites; however, wound size measurements were not interfered. The wound healing index was formulated as follows:

\[
\frac{\text{Wound area of day } N}{\text{Wound area of day } 0} \times 100\%.
\]

2.10. Determination Antioxidation Ability by DPPH * Radical Scavenging. The principle of antioxidation determination is based on the color change of DPPH to light yellow if free radicals are scavenged [21]. The more the light color rendered, the higher the antioxidant capacity from the component. Suitable concentration doses of C. osmophloeum Kanehira extracts were added to 1 μL and with 99 μL of DPPH solution. When DPPH reacted with antioxidants or vitamin C (positive control), it changed to reduced form and led to a lower absorbance at 517 nm. The scavenging activity (%) of DPPH radical was formulated as follows:

\[
\text{Scavenging activity} (%) = 100 \times \frac{(OD_{\text{control}} - OD_{\text{sample}})}{OD_{\text{control}}}
\]

2.11. Metal Chelating Activity. Metal chelating activity was measured with slight modifications as previously described [5]. Briefly, C. osmophloeum Kanehira extracts dissolved in DMSO were mixed with a reagent containing 10 μL of 2 mM FeCl₂·4H₂O. To initiate the reaction with the addition of
20 μL of 5 mM ferrozine, the solution was vigorously shaken and then it was stood for 10 min at room temperature. EDTA was regarded to be a positive control. Its absorbance at 562 nm was calculated as the chelating activity (%) and its method was alike to (4).

2.12. Reducing Power. The reducing power of C. osmophloeum Kanehira extracts was determined according to the previous method [21]. The extracts were incubated with 85 μL of 67 mM phosphate buffer (pH 6.8) and 2.5 μL of 20% K$_3$Fe(CN)$_6$ at 50°C for 20 min. After the addition of 160 μL of trichloroacetic acid (10%), it was centrifuged at 3,000 × g for 10 min to collect the supernatant (75 μL) for reacting with 2% FeCl$_3$ (25 μL). BHA was regarded to be a positive control. Finally, its absorbance at 700 nm was measured by spectrophotometer.

2.13. Statistics. Data are indicated as a mean and standard deviation in triplicate at least. Its difference significance was evaluated by Student’s t-test.

3. Results and Discussion

3.1. B16-F10 Cytotoxicity of C. osmophloeum Kanehira Extracts. Melanoma B16-F10 cells were cultured in the indicated doses of tested extracts (10, 25, 50, 100, and 200 μg/mL). In Figure 2, the cell viability was determined by MTT assay. The proliferation of B16-F10 cell was inhibited by extracts in a dose-responsive manner ranging from 10 to 200 μg/mL. When the mouse melanoma cells were incubated in a higher assay surrounding 100 μg/mL, the viabilities of extract-treated B16-F10 cells were more than 50% at 48 h treatment, suggesting that extracts had discernable cytotoxic effect on mouse melanoma cells.

3.2. C. osmophloeum Kanehira Extracts on B16-F10 Cellular Tyrosinase Activity and Melanin Content. We further investigated the in situ cellular tyrosinase and melanin suppressions of extracts. The melanin generation mechanisms contain the L-tyrosine hydroxylation and the L-DOPA oxidation to its corresponding dopaquinone to form pigment by additional multiple biosynthesis steps through the enzymatic tyrosinase. We verified that the extracts had the tyrosinase-inhibiting ability and melanin content effectiveness in mouse melanoma cell, B16-F10. In Figure 3(a), the extracts had revealed superior obvious suppressions even at a moderate quantity concentration to both tyrosinase activity and melanin content. Additionally, Figure 3(b) shows that melanin contents and tyrosinase activities were highly correlated under the same dose-responsive manner upon C. osmophloeum Kanehira treatments. Both tyrosinase activity and melanin content decreased in a similar dose-dependent tendency, when we increased dosages of extracts, indicating that the inhibition of cellular tyrosinase activity might induce the epidermal melanin reduction. But interestingly, extracts at the concentration 100 μg/mL, the cell viability remained 52% (Figure 2), and the tyrosinase activity was 18% lower than cell viability (Figure 3(a)). Melanin content did not show evident reduction. On the contrary, with highest concentration of C. osmophloeum Kanehira extracts at 50 μg/mL, the melanin content was decreased for low cell viability. As the tendency of tyrosinase activity was lower than cell viability, the melanin content was a bit higher than cell viability.

3.3. Measurement of C. osmophloeum Kanehira Extracts on Mushroom Tyrosinase Activity. We previously reported that UV exposure may induce the oxidative stress which is prone to be skin darkening and ROS generation for tumor progression [22,23]. For the prevention of skin darkening and the hyperpigmentation, we evaluated the inhibitory effects of C. osmophloeum Kanehira extracts using in vitro mushroom tyrosinase inhibitory assay. The inhibition effectiveness of C. osmophloeum Kanehira extracts demonstrates moderate suppression to the activity of mushroom tyrosinase at 200 μM (Figure 4). Accordingly, these compounds have the potential use for supplements in industry of cosmetics and pharmaceuticals.

3.4. Evaluation of the In Vivo Wound Size Assay. In order for C. osmophloeum Kanehira extracts to be utilized as a dermal agent, they should exhibit minimal toxicity towards normal tissues. The wound-healing performance of the C. osmophloeum Kanehira extracts was measured by an animal model in terms of the full thickness wound assay and monitored by image analysis of excision wound area. We discovered that the mice displayed no obvious evidences of gross toxicity during the course of the treatment period. The body weight of the mice was also monitored at an interval of every other day over the course of this study. The results showed that the body weight of mice in the treatment and the control groups was not significantly different over the duration of the experiment (data not shown). Additionally, the mean values of the heart, liver, and kidney weights after sacrifice between these two groups of mice were not significantly changed.

![Figure 2](image_url)
Figure 3: The inhibitory effects of various concentrations of *C. osmophloeum* extracts on B16-F10 cells. (a) The tyrosinase activity and (b) the melanin content of B16-F10 cells were incubated with indicated concentrations of *C. osmophloeum* extracts. Data: the mean value ± SD (triplicate values for three independent experiments); ∗ < 0.01, ∗∗ < 0.001.

Figure 4: The inhibitory effects of various concentrations of *C. osmophloeum* extracts and kojic acid on mushroom tyrosinase for 5 minutes. Data: the mean value ± SD (triplicate values in three independent experiments); ∗ < 0.01, ∗∗ < 0.001.

The wound areas of both petroleum jelly (8% DMSO, w/v) and 0.8 mg extract treatment groups were decreased in a time-dependent manner (Figure 5). Wound areas of extract-treatment groups at days 0, 1, 3, and 5 were 100 ± 0.1%, 86.7 ± 5.8%, 63.0 ± 6.1%, and 42.7 ± 6.4%, respectively, which were smaller than those of petroleum jelly group (100 ± 0.4%, 96.7 ± 5.8%, 83.3 ± 11.5%, and 73.0 ± 11.3%) (Figure 5(a)). In the beginning of the wound creation, the experimental group displayed smaller area than that of petroleum jelly group which showed a constant repair trend after 3 days. The *C. osmophloeum* Kanehira-treated wound healing displayed over 30% and 50% wound area closure after 3 and 5 days, respectively (Figure 5(b)). Therefore, the repairing ability and the wound shrinking ratio of the extract experimental group were higher and more effective.

3.5. Antioxidative Properties of *C. osmophloeum* Kanehira Extracts. Accumulating evidence shows that free radical increases can lead to skin melanin overexpression and speed up the variety of oxidations of lipids in manufactured food.
Table 1: Measurement of C. osmophloeum extracts on antioxidant experiments. The data were expressed as a mean value in three independent experiments.

| C. osmophloeum extract \(\mu g/mL\) | DPPH scavenging (%) | Metal chelating ability (%) | Reducing power (OD 700) |
|-----------------------------------|----------------------|----------------------------|------------------------|
| 1                                 | ≤10.0                | ≤10.0                      | 0.90 ± 0.00            |
| 10                                | ≤10.0                | ≤10.0                      | 0.10 ± 0.00            |
| 50                                | ≤10.01               | ≤10.0                      | 0.18 ± 0.02            |
| 100                               | 13.23 ± 0.01         | ≤10.0                      | 0.27 ± 0.03            |
| 250                               | 38.97 ± 0.02         | ≤10.0                      | 0.48 ± 0.02            |

Vitamin C\(^a\) 80.82 ± 0.00 — —
EDTA\(^b\) — 80.76 ± 0.01 —
BHA\(^c\) — — 0.56 ± 0.03

\(^a\) Vitamin C was used as a positive control on DPPH assay at 100 \(\mu\)M.
\(^b\) EDTA was used as a positive control on metal chelating ability at 100 \(\mu\)M.
\(^c\) BHA was used as a positive control on reducing power at 100 \(\mu\)M.

and reducing power ability except metal chelating. In the future, we will analyze purification from C. osmophloeum Kanehira extracts to find an effective antioxidant and tyrosinase inhibitor.

### Conflict of Interests

The authors had no conflict of interests.

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