Physiological and Whitening Effects of *Morus alba* Extracts

**Kyungmee Gug†**

**Abstract**

Mulberry extracts can be incorporated into skin-whitening products. The compound attributed to lighten the skin is arbutin, a form of hydroquinone that inhibits melanin release by suppressing the tyrosinase enzyme. For the cosmetic applications, the physiological effects of mulberry (*Morus alba*) extracts were investigated. The water soluble fraction of mulberry contains higher amount of protein (16.28~4.47%) in contrast to fat (1.55~1.41%). In addition, the fraction abundantly contains succinic acid (972.4-275.8 mg/g) and phosphoric acid (1,628.4-121.9 mg/g) in different parts of mulberry. The free radical scavenging ability in water soluble fraction was found to display remarkable effects in comparison with methanol and ethyl acetate fraction. The ethyl acetate-soluble of root and leaf showed remarkable tyrosinase inhibition activity by IC₅₀ (µg/ml). The anticancer activity of methanol fraction obtained from mulberry using human cancer cell lines showed growth inhibition effect (270.14 mg/ml in Calu-6 cells, 295.29 mg/ml in HCT-116, and 332.29 mg/ml in MCF-7 cells, respectively). Based on the results, *Morus alba* extracts include cosmetic ingredients with antioxidizing and whitening properties.

**Key words**: *Morus alba*, Organic Acid, Antioxidant, Tyrosinase, MTT, Anticancer Activity, Whitening Property

**1. Introduction**

The Interests of the middle-aged in healthy skin have been increased under the influence of social trend of pursuing healthy beautiful appearance. In particular, various methods including chemical, physical, and biological means are being researched in order to improve the aging skin. Active oxygens are always generated during our respiration process to obtain necessary energy for keeping our health. The active oxygens are very harmful for our skin cells, leading to the pigmentation, wrinkles, and elasticity decrease on skin[1,2]. As the results, active oxygens suppress the metabolism of our skins, results in the increased formation of melanin pigments which is the defense mechanism for protecting our skins. Although melanin produced in the skin protects skin from UV light, too much melamines leave dark brown stains[3]. Such stain formation on skin is not good for beauty culture. It was reported from the in-vivo studies that second metabolic products originated from plants suppress the formation of free radical species and active oxygens, preventing oxidative cell loss[4,5]. Kojic acid and arbutin currently used for the inhibitor of tyrosinase show the unwanted side effects such as cytotoxicity and mutation[6]. Thus, the development of novel safe materials with high whitening activity[7-10].

Mulberry is widely distributed thru east asian countries including Japan, China (in particular, Manchu area), Mongolia. Mulberry leaves are widely used for silk-farming, leaves, fruits, branches, root barks of mulberry are also used for producing Chinese medicine to lower blood sugar (by γ-GABA) and blood pressure (by DNJ/deoxino-jirimicin)[11,13]. Mulberry contains inorganics (such as Ca, Fe, K, Zn, Mg), vitamins (A, B, C), and vegetable fibers[12]. The cultivation of mulberry is relatively easy and simple thru the well-established mechanized farming technology. Thus, mulberry is the plant resources of global competitive power.

Here we report a study on the development of possible natural antioxidant and functional cosmetics by measuring the antioxidant and whitening effect of leaves, branches, root barks, and fruits of mulberry.
2. Materials and Methods

2.1. Preparation of Extract Samples
The leaves, branches, roots, fruits of mulberry used for the research were given by Dongeunara, Inc.. The mulberry materials were washed with distilled water and then dried in the shaded area. The mulberry materials were extracted with water. 10% (g/l) water extracts were subject to fractionation with methanol, hexane, ethyl acetate solvents, in order, to give extracts. The fractionated extracts were subject to centrifugation with a speed of 12,000 rpm for 30 min and were then filtered thru Whatman No. 4 filter paper. The resulting solutions were condensed by a rotary evaporator (EYELA N-N series) and finally freeze-dried to give powder samples.

2.2. Analyses of General Components
The analysis of general components of the samples were performed according to AOAC method\textsuperscript{[14]}. The ash content for rough ash was determined when no weight change was observed after heating at 550°C. The protein content for rough protein was determined after complete decomposition using sulfuric acid and decomposition promoter at heating condition. The volatile compounds were collected by using distillation setup and nitrogen content then was measured. The nitrogen content was converted to protein content by using protein coefficient. Rough lipid was extracted using a Soxhlet setup (Soxtex 2050 Auto Extraction Unit, Foss Tecater., Hillerod, Denmark) to give lipid. The amount of lipid was weighted to determine rough lipid content. Rough fiber content was measured by using the modified Henneberg-Stohman method. Vitamin C was extracted with 5% metaphosphoric acid and then reacted with DNP (2,4-dinitrophenyl hydrazine) to be subject to colorimetric method (measuring absorbance at 520 nm).

2.3. Analysis of Organic Acids
Organic acids were analyzed according to Lee method\textsuperscript{[15]}. The 3 gram sample was dissolved in 100 mL of distilled water and was subject to centrifugation at 4°C and 15,000 rpm. The above layered solution was filtered thru a Whatman No. 2 filter paper. The filtered solution was refiltered thru a 0.45 µm membrane filter. The refiltered solution was purified using Sep-pak C\textsubscript{18} cartridge and was finally analyzed by ion chromatography. For the ion chromatography a column [ICE-AS6(9×250 nm)], 0.4 mM heptfluorobutryric acid as a mobile phase, 5 mM tetrabutylammonium hydroxide as a regenerator, an Electro conductivity detector, and n-ICE micromembrane as a blocking equipment were used at a moving speed of 1 mL/min. Organic acid standards include acetic acid, butyric acid, citric acid, formic acid, lactic acid, malic acid, succinic acid, phosphoric acid, propionic acid, pyrogallic acid, tartaric acid (Sigma Chemical Co., St. Louis, MO, USA).

2.4. Measurement of Electron-donating Ability
The electron donating ability of the extracts was determined by adapting Chu method\textsuperscript{[16]}. The 50 µl of 2×10\textsuperscript{-4} M DPPH (1,1-diphenyl-2-picylhydrazyl, Sigma Chemical Co., St. Louis, MO, USA) solution was added to each 100 µl samples, was well mixed for 10 sec, and then let it be alone at ambient temperature for 10 min. The absorbance was measured at 517 nm using a UV spectrophotometer. The radical elimination ability of DPPH was obtained by expressing the percent between a sample and a control.

Free radical scavenging activity (%) =

\[
1 - \frac{\text{Absorbance of control at 517 nm}}{\text{Absorbance of sample at 517 nm}} \times 100
\]

2.5. Measurement of Tyrosinase Suppressing Activity
The suppressing effect of Tyrosinase was measured by dopachrome method\textsuperscript{[17]}. 150 µl of mushroom tyrosinase (110 U/ml, Sigma Chemical Co., St. Louis, MO, USA), 225 µl (2.5 µM) of L-tyrosine, 225 µl of 0.175 M phosphate buffer (pH 6.8), and 300 µl ethanol solution or a sample solution (1 mg/ml) were well mixed. The solution was incubated at 37°C for 10 min. Finally, DOPA chrome produced at 475 nm was measured. The suppressing degree of Tyrosinase was measured by using the following equation.

Tyrosinase inhibition (%) = \(\frac{A - B}{A} \times 100\)

A is the absorbance of control reaction solution and B is the absorbance of reaction solution after adding a sample. In order to know the suppressing ability of tyrosinase the half concentration (to decrease initial tyrosinase activity to its half; IC\textsubscript{50}) and arbutin whitening
material were obtained by using a standard curve and then compared.

### 2.6. Anticancer Effect and Cell Toxicity Analysis

The cell lines used for this research include Calu-6 (Human pulmonary carcinoma), MCF-7 (Human breast adenocarcinoma), and HCT-116 (Human colon carcinoma). 10% FBS (fetal bovine serum, Gibco, Rocville, MD, USA) added RPMI 1640 medium was inoculated at 37°C and then incubated at a 5% CO₂ incubator. MTT assay was carried out to evaluate the cytotoxicity of mulberry methanol extract on cancer cell line[18]. A 0.1 mL (105 cells/mL) of cell suspension solution was inoculated at each well of 96 well plate, incubated at a CO₂ incubator at 37°C for 24 hrs. 0.1 mL of 0.3 mg/mL concentration of extract was added, cubated for 48 hrs, and upper layered solution was removed. A 5 mg/mL concentration of MTT solution (10 [µL]) was added to PBS (pH 7.4) and 90 [µL] medium was then added. The mixture was injected to each well and then reacted for 3 hrs in the dark. The upper layer was removed. DMSO solvent (100 [µL]) was added and untouched for 10 min at ambient temperature. The absorbance was measured at 550 nm by using ELISA reader (ELx800, Bio-Tek, USA).

### 3. Results and Discussion

#### 3.1. Content of Ingredients with Part of Mulberry

The general component contents of rough ash, protein, lipid, fiber and vitamin C in the leaves, roots, branches, and fruits are in the range of 3.09~8.94% for rough ash, 4.47~16.28% for rough protein, 1.41~1.55% for rough lipid, and 3.52~43.21% for rough fiber.

<table>
<thead>
<tr>
<th>Composition</th>
<th>Leaf</th>
<th>Root</th>
<th>Branch</th>
<th>Fruit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crude ash (%)</td>
<td>8.94±0.11</td>
<td>3.55±0.15</td>
<td>3.09±0.17</td>
<td>3.82±0.17</td>
</tr>
<tr>
<td>Crude protein (%)</td>
<td>16.28±0.02</td>
<td>11.87±0.15</td>
<td>4.47±0.28</td>
<td>6.78±0.70</td>
</tr>
<tr>
<td>Crude fat (%)</td>
<td>1.44±0.08</td>
<td>1.55±0.06</td>
<td>1.51±0.05</td>
<td>1.41±0.05</td>
</tr>
<tr>
<td>Crude fiber (%)</td>
<td>10.78±1.40</td>
<td>20.78±0.91</td>
<td>43.21±0.68</td>
<td>3.52±0.34</td>
</tr>
<tr>
<td>Vitamin C (mg/100 g)</td>
<td>tr²)</td>
<td>tr</td>
<td>tr</td>
<td>18 mg</td>
</tr>
</tbody>
</table>

³Mean ± SD with different superscripts within a row is significantly different (p<0.05) by Duncan's multiple range tests
²tr represents trace.

### 3.2. Analyses of Organic Acids

Organic acids are the source of sour taste of fruit juice and important material for determining sugar and fragrance of fruits[19]. The organic acids contained in mulberry were given in Table 2. Among 11 standard

<table>
<thead>
<tr>
<th>Organic acids</th>
<th>Contents¹</th>
<th>Leaf</th>
<th>Root</th>
<th>Branch</th>
<th>Fruit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Citric</td>
<td>7.1</td>
<td>394.2</td>
<td>39.2</td>
<td>3,529.4</td>
<td></td>
</tr>
<tr>
<td>Phosphoric</td>
<td>460.6</td>
<td>121.9</td>
<td>230.2</td>
<td>1,628.4</td>
<td></td>
</tr>
<tr>
<td>Succinic</td>
<td>972.4</td>
<td>623.6</td>
<td>275.8</td>
<td>494.6</td>
<td></td>
</tr>
<tr>
<td>Acetic</td>
<td>88.0</td>
<td>68.7</td>
<td>58.7</td>
<td>26.9</td>
<td></td>
</tr>
<tr>
<td>Lactic</td>
<td>55.4</td>
<td>24.5</td>
<td>130.4</td>
<td>18.4</td>
<td></td>
</tr>
<tr>
<td>Tartaric</td>
<td>84.0</td>
<td>12.4</td>
<td>44.9</td>
<td>95.1</td>
<td></td>
</tr>
<tr>
<td>Malic</td>
<td>33.8</td>
<td>56.0</td>
<td>11.1</td>
<td>44.3</td>
<td></td>
</tr>
<tr>
<td>Formic</td>
<td>10.9</td>
<td>6.8</td>
<td>16.9</td>
<td>6.3</td>
<td></td>
</tr>
<tr>
<td>Pyroglutamic</td>
<td>23.6</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Butyric</td>
<td>13.2</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Propionic</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>1725.4</td>
<td>1333.7</td>
<td>807.2</td>
<td>5843.4</td>
<td></td>
</tr>
</tbody>
</table>

¹Mean ± SD with different superscripts within a row is significantly different (p<0.05) by Duncan's multiple range tests
²Not detected

Although vitamin C was found to be 18 mg per gram in the fruit, vitamin C was not found in leaves, roots, and branches of mulberry. Although the content of rough ash and protein was found to be greatest in leaves, the content of rough lipid was found to be similar in the all the regions of mulberry as seen in Table 1. The content of rough fiber in the branches was 43.21% and was greater by 12 times in the fruits.
organic acids used for analysis, 10 organic acids (citric acid, phosphoric acid, succinic acid, acetic acid, lactic acid, tartaric acid, malic acid, formic acid, pyroglutamic acid, and butyric acid) were found at all the regions of mulberry. However, propionic acid was not detected at all the regions of mulberry.

In the leaves of mulberry total organic acid content was 1,725.4 mg/100 g and the content of succinic acid was greatest (972.4 mg/100 g). The content in the leaves then decrease in the order: phosphoric acid > acetic acid > tartaric acid > lactic acid > malic acid > butyric acid > formic acid > citric acid. In the roots of mulberry total organic acid content was 1,333.7 mg/100 g and the content of succinic acid was greatest (634.6 mg/100 g). Next, the content in the leaves decrease in the order: citric acid > phosphoric acid > acetic acid > malic acid > pyroglutamic acid > lactic acid > tartaric acid > formic acid. In the branches of mulberry total organic acid content was 807.2 mg/100 g and was smaller than the other regions of mulberry. The content of succinic acid was greatest (275.8 mg/100 g).

The content in the branches then decrease in the order: phosphoric acid > lactic acid > acetic acid > tartaric acid > citric acid > formic acid > malic acid. In the fruits of mulberry total organic acid content was 5,843.4 mg/100 g and the content of citric acid was greatest (3,529.4 mg/100 g). The content in the leaves then decrease in the order: phosphoric acid > succinic acid > tartaric acid > malic acid > acetic acid > lactic acid > formic acid. Among 10 organic acids citric acid, phosphoric acid, and succinic acid were found to be high content in all regions of mulberry. In the fruits of mulberry the content of citric acid was greater by 497 times than in the leaves. Citric acid is used as a fragrance additive for soft drinks and as a stabilizer for organic compounds. Phosphoric acid is used for additives for sour taste and fruit fragrance\[20]. However, Butyric acid was detected only in the leaves and pyroglutamic acid was found only in the roots.

3.3. Measurement of Electron-donating Ability

DPPH is a stable radical species and is used for

![Fig. 1. Antioxidant activities of extracts and their fractions Morus alba part. DPPH radical scavenging activities of leaves, root, branch and fruit.](image)
detecting the antioxidizing activity of antioxidants[21]. The antioxidizing activities of extracts (with hot water, methanol, hexane, ethyl acetate) in each region of mulberry were summarized in Fig. 1. The electron donating ability increased with increasing the extract concentration. The antioxidizing activity of ethyl acetate extract was greatest.

3.4. Tyrosinase Suppressing Effect

Tyrosine biosynthesis suppresser can suppress the synthesis of melanin polymer in the skin and can be used for the evaluation of whitening agent[25]. In the point of attaining 50% of the initial tyrosinase activity the fractionation concentration (IC_{50}) of mulberry with extracting solvent are given in Fig. 2. As seen in the experimental results, the greatest tyrosinase suppressing activity was obtained in the case of ethyl acetate fractionation.

3.5. Anticancer Effect

The growth suppressing effect on the cells (Calu-6, MCF-7, HCT-116) treated with methanol extract of mulberry was given in Fig. 3. For IC_{50}, the cell growth activity of Calu-6 cell was greatest to be 270.14 mg/ml. The cell growth activities of MCF-7 cell and HCT-116 were found to be 295.29 mg/ml and 332.29 mg/ml, respectively. Therefore, the extracts of mulberry (extracted with methanol) could be used as additives for drinks or tea. The cytotoxicity on NIH/3T3 cell was about 80% growth activity at 0.5 mg/ml (high concentration)[27].

Thus, the extracts of mulberry (extracted with methanol) could be used as additives for cosmetics.

4. Conclusion

In order to search chemicals which can be applied for cosmetic industry we investigated the extracts of Mulberry in terms of antioxidizing and whitening effects depending on the part and condition of extraction. From the hydrothermal extraction process of Mulberry 10 organic acids were detected along with several other components including proteins, cellulose, fats. Among the organic acids the content of citric acid, phosphoric acid, and succinic acid was higher than the other organic acids.

The compound attributed to lighten the skin is arbutin, a form of hydroquinone that inhibits melanin release by suppressing the tyrosinase enzyme. For the cosmetic applications, the physiological effects of mulberry (Morus alba) extracts were investigated. The water soluble fraction of mulberry contains higher amount of protein (16.28–4.47%) in contrast to fat (1.55–1.41%). In addition, the fraction abundantly contains succinic acid (972.4–275.8 mg/g) and phosphoric acid (1,628.4–121.9 mg/g) in different parts of mulberry. The free radical scavenging ability in water soluble fraction was found to display remarkable effects in comparison with methanol and ethyl acetate fraction. The ethyl acetate-soluble fractions of root and leaf showed remarkable...
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Tyrosinase inhibition activity by IC50 (µg/ml). The anti-cancer activity of methanol fraction extracted from mulberry using human cancer cell lines showed growth inhibition effect (270.14 µg/ml in Calu-6 cells, 295.29 µg/ml in HCT-116, and 332.29 µg/ml in MCF-7 cells, respectively). From the results above, *Morus alba* extracts include cosmetic ingredients with antioxidizing and whitening properties.

Acknowledgments

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References


