Hair Growth Promoting Activity of *Nothopanax scutellarium* Merr. Leaves

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ABSTRACT

Objective: The aims of this study were to know the safety using Hen's eggs-chorioallantoic membrane (HET-CAM) test method and hair growth promoting activity of the ethyl acetate fraction of *Nothopanax scutellarium* leaves. Methods: Safety test was measured by scoring and categorizing irritation on HET-CAM. Meanwhile, activity test was conducted by applying the hair tonic of the ethylacetate fraction on the back of the rabbits, and the length of hair was measured in the 1st, 2nd, and 3rd week. In the 3rd week, the hair growth was weighed and hair diameter also was measured using scanning electron microscope (SEM). Results: The result showed that 0.2 gram of ethyl acetate fraction of *N. scutellarium* leaves have mild irritation effect, whereas the formulation with 0.5% and 1% of fraction increased hair growth and hair diameter. Conclusion: The ethyl acetate fraction of *N. scutellarium* have mild iritan effect, and the hair tonic demonstrated hair growth promoting activity.

Key words: Hair Tonic, Hair Growth Activity, *Nothopanax scutellarium*, Safety, HET-CAM.

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INTRODUCTION

Hair loss and thinning are common problems that found in term of clinical dermatology. There are a number of products of hair tonic that claimed to increase hair growth, either using natural products or synthetic compounds. Some studies also reported the prospect of the natural product for hair growth promoting. There has been no report on its main active ingredient responsible for the hair growth activity. In the current work, cedrol as a major constituent from *P. orientalis* was evaluated for its potential on hair growth in vivo, different concentration of cedrol (10, 20 and 30 mg/mL) promoted hair growth in a dose-dependent manner. *Nothopanax scutellarium* Merr from Araliace family is the tropical plant. The leaves were traditionally known to have hair growth promotion activity.

Based on previous research, ethanolic extract of the leaves was proven to have hair growth activity. However, the results of studies showed that hair growth activity of the ethanolic extract was lower than Minoxidil. Another study, the combination of ethanolic extract of the leaves with ethanol extract of tea leaves, also resulted in lower hair growth activity than positive control. Some studies reported that flavonoid demonstrated hair growth promoting activity. To increase hair growth activity of leaves extracts from *N. scutellarium*, further fractionation process was needed using the suitable solvent to increase the level of flavonoids, which required in increasing hair growth. Ethyl acetate was chosen as a solvent for a fractionation process of *N. scutellarium* leaves.

This study presented the safety test of the ethyl acetate fraction of *N. scutellarium* leaves using HET-CAM (Hen's Egg Test - chorioallantoic membrane) method. Then, the hair tonic formula of the fraction on hair growth activity was performed in rabbits.

MATERIALS AND METHODS

Materials

The leaves were obtained from the Indonesian Spice and Medicinal Crops Research Institute (ISMCRI), Bogor, and authenticated in Botany Herbarium Research Institute, Cibinong, West Java. The voucher specimen was deposited in Herbarium of Pharmacognosy, Faculty of Pharmacy, Universitas Indonesia. Solvents (ethanol, ethyl acetate, n-hexane), nipagin, nipasol, sodium metabisulfite, propylene glycol, chorioallantoic membrane (CAM), and minoxidil as the positive control were purchased from local suppliers.

Extraction and Fractionation

The leaves powder (4 kg) were macerated with ethanol (40L). Solvents were filtered and concentrated using rotary vacuum evaporator at 50°C. The ethanolic extract was dispersed in water and partitioned with n-hexane and ethyl acetate, subsequently. The ethyl acetate phase was concentrated using a rotary vacuum evaporator at a temperature of 50°C to give a viscous fraction of ethyl acetate. Total flavonoid of the extract was determined by AlCl₃ according to Kabir et al, 2016 with slight modification.

Safety test by HET-CAM method

The HET-CAM bioassy was performed to evaluate the level of irritation on mucous membrane based on slight modification of Steiling method. Briefly, the fraction (0.2 g) solution was applied to chorioallantoic membrane (CAM) and left for 20 seconds until the sample was spread smooth. Then, the membrane was evaluated within 5 minutes to notice any symptoms of hemorrhage, lysis, and/or coagulation. The evaluation determined based on the result of scores and category of irritation (Table 1). The results of irritation scores can be calculated using the following equation:

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where:
Hemorrhage time = time (in seconds) of the first appearance of blood hemorrhages.
Lysis time = time (in seconds) of the first appearance of vessel lysis.
Coagulation time = time (in seconds) of the first appearance of protein coagulation.

**Formulation of hair tonic**

Formulation of hair tonic prepared the variation of concentration of ethyl acetate fraction the leaves. The calculation of the percentage composition of each hair tonic can be seen in Table 2. Nipagin and nipasol were dissolved in ethanol, while sodium metabisulfite was dissolved in distilled water. Both solutions were mixed, and ethyl acetate fraction of *N. scutellarium* leaves was added into solution. Propylene glycol was added until the desired volume. All material were stirred until homogenously mixed.

**Evaluation of hair tonic**

Evaluation of hair tonic preparations was conducted by observing the color and odor during storage. Furthermore, examination of pH by using a pH meter and hair tonic stability test at low temperature (4°C±2°C), room temperature (25°C±2°C), and high temperature (40°C±2°C) for 12 weeks by observing organoleptic appearance and pH every 2 weeks.15,16

**Hair growth activity**

White male rabbits of New Zealand strain (4-5 months, with weights ranging from 2-4 kg) were obtained from Indonesia Research Institute of Animal Production (IRIAP). The study was approved by the Ethical Committee, Faculty of Medicine, Universitas Indonesia (No.683/H2.F1/ETIK/2012).

Four white male rabbits were used to examine hair growth activity. The hair on the back of the rabbit was shaved and divided into 6 areas with size of 4 x 4 cm; 3 areas on the left side, and another 3 areas on the right side with 2 cm distance of each area. Table 3 showed the treatment of samples. Hair tonic was applied in the shaved areas and observed for 3 weeks. During the experiment the hair growth was observed at week 1, 2, and 3, the 10 longest rabbit hair removed in each test area and measured using calipers.17 At the end of the application hair weight of rabbits were determined by pulling hair that grows on the test area and then weighed. The hair diameter was observed at week 3 by using SEM (Scanning Microscope Electron).

**Statistical analysis**

Data were reported as mean ± standard deviation. Statistical analyses were performed using Student’s t-test for the significance of the results (P < 0.05).

**RESULTS**

**Fractionation**

The result of fractionation that used in the preparation of hair tonic was ethyl acetate fraction. Characteristics of ethyl acetate fraction are shown in Table 4. The fraction contained total flavonoid 4.79%.

**Safety test by HET-CAM method**

Table showed the results of HET-CAM test. The irritation score of the test fraction is 4.54, indicated mild irritation on chorioallantoic membranes (Table 5).
Evaluation of hair tonic

On this study, the stability test conducted by accelerated stability test. The results of organoleptic consisted of observations color and odor of hair tonic of the ethyl acetate fraction stored at low temperature (4±2°C), room temperature (25°±2°C), and high temperature (40°±2°C) looks stable. Results of hair tonic pH measurement of the fraction of ethyl acetate did not change significantly at low temperatures (4±2°C), room temperature (25°±2°C) and high temperature (40°±2°C) for 12 weeks of storage.

Effectivity of hair growth

The results showed that there were significant differences among groups. After observation for 3 weeks, the resulting hair growth on the groups of formula II containing fraction of test of 0.5% and formula III containing the fraction of test of 1%; has faster hair growth activity (Figure 1).

Hair length

The result of measurement of hair length of rabbit at week 1 shows the average on the groups of normal control, negative control, the formula I, formula II, formula III, and positive control respectively are 4.72 mm, 4.80 mm, 8.39 mm, 10.48 mm, 12.37 mm, and 9.16 mm. Furthermore, the results of the statistical analysis showed that the average hair length rabbit of each treatment group there were significant differences (p<0.05).

At week 2, the average of hair length rabbit on the groups of normal control, negative control, the formula I, formula II, formula III, and positive control respectively are 8.10 mm, 8.13 mm, 12.09 mm, 14.29 mm, 18.24 mm, and 13.78 mm. The result of statistical calculation show the average hair length rabbit of each treatment group there were significant differences (p<0.05). At week 3, the average of hair length rabbit on the groups of normal control, negative control, the formula I, formula II, formula III, and positive control respectively are 12.33 mm, 12.35 mm, 16.25 mm, 18.36 mm, 28.25 mm, and 17.24 mm. The result of statistical calculation shows the average hair length rabbit of each treatment group there were significant differences (p<0.05). Based on results of calculation of average hair length rabbits show formula II and formula III has faster activity of hair growth than the positive control (Figure 1). The result of statistical calculation shows the average hair length rabbit of each treatment group there were significant differences (p<0.05).

Hair weight

The result of average of hair weight at normal control group, the negative control, formula I, formula II, formula III, and positive control each is 56.02 mg/cm², 55.86 mg/cm², 70.28 mg/cm², 73.02 mg/cm², 84.25 mg/cm², and 71.18 mg/cm². The result of average hair weight indicated that test area which is given formula II and formula III containing ethyl acetate fraction of N. scutellarium leaves 0.5% and 1% had a greater weight than positive control (Figure 2). There was significant difference in each treatment (p<0.05).

Hair diameter

The results of average rabbit’s hair diameter show that the formula II and formula III containing of 0.5% and 1% ethyl acetate fraction of Nothopanax scutellarium leaves, respectively, has a larger diameter than the positive control (Figure 3). In addition to the observation of hair diameter, morphology hair of rabbit also was observed by SEM. The results show that the morphology hair of rabbit in the groups of formula II and formula III has composed regularly of hair cuticle and overlap, which consists of layers of flat keratin (Figure 4).

DISCUSSION

In this study, fractionation is conducted to increase the levels of flavonoids of Nothopanax scutellarium leaves, which believed to have potential activity of hair loss and thinning treatment. Flavonoid has been reported to have some pharmacological activities, such as antioxidant, hepatoprotective and anticancer. Based on several studies, a compound that plays a role in hair growth activity are proanthocyanidin compounds and procyanidin. One of the mechanisms hair growth promoting through Endothelial nitric oxide synthase (eNOS). Daidzein, genistein, isorhamnetin, kaempferol, quercetin, naringenin, and pelargonidin inhibited iNOS protein and mRNA expression and also nitric oxide (NO) production. Myricetin, quercetin, baicalein and fisetin demonstrated hair growth activity via inhibition of the type I 5α-reductase.
Evaluation of irritation level was performed by using the HET-CAM method determined by the score and category irritation by comparing the changes in the chorioallantoic membrane between the negative control, positive control and fractions control. Results showed that there was a difference between the positive control (1%SLS) with test fractions containing ethyl acetate fraction of N. scutellarium leaves (Table 5). In some studies, testing by the HET-CAM method has been done especially for plants that can be developed into cosmetic products, such as plants of Lansium domesticum and Phyllanthus niruri. Both of these plants has efficacy as an antioxidant and antityronase. Based on the previous study, at a concentration of 2.5% either extract of Lansium domesticum or extract of Phyllanthus niruri cause moderate irritation of chorioallantoic membranes.23

In this study, we used the ethyl acetate fraction as an active compound of the hair tonic formulation. This formulation was evaluated by observing the color, odor, pH, and stability of the hair tonic stored at low temperature (4±2°C), room temperature (25°±2°C), and high temperature (40°±2°C) for 12 weeks with a method of accelerated stability. The results showed hair tonic stable during storage. A product which stable in accelerated stability tests, that the that is stable in storage at room temperature for one year.24 Meanwhile, result of observations showed a hair tonic pH has a pH in the range of physiological pH "acid mantle" the skin or also called pH balance, which ranges from 4.5 to 6.5, and still in the scalp pH range, i.e., between 4, 0 to 5.8.25,26

Treatment of hair loss and thinning from natural product was widely needed. Proanthocyanidin from grape seeds stimulated the proliferation of hair follicle cells of mice in vitro and stimulate hair growth cycle of the telogen phase to anagen phase in vivo. Root extract of Sophora flavescent has been reported to stimulate hair growth.5-27

In this study, formula II and III, each containing 0.5% and 1% of ethyl acetate fraction have faster hair growth than positive control (Figure 1). The mechanism of hair growth from the ethyl acetate fraction is still unclear, but the activity of hair growth resulting from the ethyl acetate fraction N. scutellarium leaves is suggested due to flavonoids. Based on several studies, flavonoids contribute to the activity of hair growth by improving blood circulation to nourish hair follicles, as well as improve blood circulation to nourish hair follicles that can increase hair growth.25,29 Other studies also shown that flavonoids can shorten the telogen phase and can prolong the anagen phase. Based on research, minoxidil can prolong the anagen phase by extending the dermal papilla cell survival in the hair follicles by increasing proliferation and anti-apoptotic effects that can stimulate hair growth in the anagen phase.28 The role of flavonoids on the activity of hair growth is caused by several factors hair growth, such as insulin-like growth factors-1 (IGF-1), vascular endothelial growth factors (VEGF), keratinocyte growth factors (KGF), and hepatocyte growth factors (HGF), these factors have the effect of stimulating hair growth.5-27

In this study, the hair tonic on the formula III has a larger diameter than the positive control and has hair cuticle composed regularly and overlap (Figure 4). Some research suggests that healthy hair is observed with SEM (Scanning Electron Microscope) can be seen clearly has the hair cuticle smooth edges, patterns of cuticle layer to are arranged neatly to the inside of the hair seemed protected.25,29 Meanwhile, damaged hair has cuticle edges were chipped or loose. If the hair cuticle is almost completely peeled off, will make cortex layer is exposed, for example, the branched hair ends and hair fragile.25

**CONCLUSION**

Based on the results of this research concluded that the ethyl acetate fraction of Nothopanax scutellarium leaves could mild irritate the membranes chorioallantois and hair tonic preparations with the concentra-

**Table 5: The result of average score and category of irritation**

<table>
<thead>
<tr>
<th>Sample</th>
<th>Average of Irritation score</th>
<th>Category of irritation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Negative control: 0.9%NaCl solution</td>
<td>0</td>
<td>Nonirritant or practically no irritation</td>
</tr>
<tr>
<td>Positive control: 1%SLS</td>
<td>10.23</td>
<td>Strong or severe irritation</td>
</tr>
<tr>
<td>Fraction of test: ethyl acetate fraction of N. scutellarium leaves</td>
<td>4.54</td>
<td>Weak or slight irritation</td>
</tr>
</tbody>
</table>

**Figure 3:** The average of rabbit hair diameter (µm) at the end of experiment.

**Figure 4:** Scanning Microscope Electrone magnification: x2000. (a) normal control; (b) negative control; (c) formula I; (d) formula II; (e) formula III; (f) positive control.
tion of 0.5% and 1% have hair growth activity the faster and can enlarge the diameter of the hair compared to the positive control.

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CONFLICT OF INTEREST

There is no conflict of interest.

REFERENCES