Hair-Growth-Promoting Effects of Fermented Red Ginseng Marc and Traditional Polyherb Formula in C57BL/6 Mice

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Abstract: An abnormal hair-growth cycle induces hair loss, which affects psychological distress and impairs life quality. Red ginseng marc (RGM) is usually discarded as a byproduct after extracting red ginseng, but several studies have shown that the RGM still has bioactive components including ginsenosides. Therefore, the hair-growth effects of fermented RGM (fRGM) and traditional polyherb formula (PH) were examined in C57BL/6 mice. The dorsal hairs of mice were depilated, and they were topically treated with fRGM or PH at 400, 200 and 100 mg/kg or the combination of all middle doses (combi) once a day for two weeks. The hair-covering regions were significantly increased with higher doses of fRGM and PH and in combi groups, compared with the control treated with distilled water. Hair length, thickness and weight also increased in the treatment groups. In particular, the fRGM and PH increased the anagen-phased hair follicles, the follicular diameters and the dermal thickness. Immunostains for Ki67 showed the anagen-phased cell division in the treatment groups. The beneficial effects were greater in the high doses of fRGM and PH and the combi groups. These suggest hair-growth-promoting effects of fRGM, PH and the combination by enhancing the hair-growth cycle.

Keywords: hair growth; growth cycle; red ginseng marc; polyherb; hair length; hair thickness; hair weight; proliferation; anagen

1. Introduction
Scalp hair has an important psychosocial function related to human appearances, as well as functions in insulation and protection. The hair normally sheds up to 100 a day [1], and the hair follicle constantly undergoes the specific growth cycle characterized by anagen (growth phase), catagen (regression phase) and telogen (resting phase) [2]. The hair-growth cycle is regulated by growth factors, cytokines and hormones; however, the altered growth cycle can cause hair loss. The most common hair losses related to the abnormal growth cycle are androgenetic alopecia and chronic alopecia areata [3,4]. The other factors include heredity, aging, medication and psychological stress [4]. Hair loss is not considered a life-threatening condition; however, it causes psychological distress and impairment of life quality [4,5]. In this context, the treatment is a major concern for the cosmetologists and dermatologists. Until now, finasteride (as a 5α-reductase type II inhibitor) and minoxidil (as an antihypertensive vasodilator) have been approved by the Food and Drug Administration (FDA) of the United States for hair growth [6]. Although the
drugs promote the anagen phase, they have some side effects, in particular in finasteride (i.e., reduced libido, erectile dysfunction and gynecomastia), and the effectiveness varies greatly among individuals [7]. Therefore, there is an urgent need to develop treatments that prevent hair loss and promote hair growth.

Ginseng (*Panax ginseng* C.A. Meyer) has long been used as a traditional herbal medicine in Eastern Asia. Red ginseng is a steamed root of the ginseng that is harvested after six years, and the extracts have shown various pharmacological properties such as antioxidant, anti-stress, antitumor, antimutagenic, anti-diabetic and immunomodulatory activities [8,9]. The red ginseng has been reported to promote hair growth in animal models by preventing apoptosis of hair follicle cells and inhibiting 5-α reductase [10,11]. It also increases the hair follicle density and size in the patients with androgenetic alopecia [12]. The functional main ingredients are saponins (ginsenosides), polysaccharides and phenolic compounds. In particular, ginsenosides increase epithelial cell genesis/development and endothelial progenitor cell migration/proliferation [13], and they also involve promoting hair growth through anti-apoptotic effects [14,15]. Furthermore, some components composing traditional polyherb formulas used for treating hair loss have been proved to have promoting effects on hair growth, even though the combination effects remain to be clarified: *Cornu cervi pantotrichum* facilitates hair growth through upregulation of fibroblast growth factor (FGF)-7 [16]; *Eclipta prostrata* stimulates the anagen phase through regulation of FGF-7 and -5 [17]; *Dictamnus dasycarpus* Turczaininov, *Polygonum multiflorum* Thunberg and *Lycopus lucidus* Turczaininov are often used in combination for the herbal formulas in alopecia [18]. In addition, *Eclipta prostrata* and *Polygonum multiflorum* Thunberg have significant inhibitory activities against 5-α reductase [19].

Red ginseng marc (RGM) is a byproduct obtained after extracting the red ginseng, which is usually discarded as waste. However, RGM or fermented RGM has shown antioxidant, immunomodulatory and anticancer effects [20,21]. It is believed that the bioactive components including ginsenosides and polysaccharides contained in RGM are still effective despite small amounts [20,22]. Moreover, the fermentation process of RGM enhances the beneficial effects, especially strong antioxidant and anti-inflammatory effects [23]. Effective microorganism (EM) is a complex of microorganisms including photosynthetic bacteria, lactic acid bacteria and yeast, and various natural products fermented with EM enhance the antioxidant, anti-inflammatory, anticancer, anti-stress and neuroprotective effects [24–26]. In the present study, the extracts of RGM fermented with EM contained ginsenosides including Rb1, Rb2, Rb3, Rc, Rd, Re, Rf, Rg2, Rg3 and Rh1. Among them, Rb1 and Rd have been reported to prevent hair loss and stimulate proliferation of hair follicle cells [15,27]. Therefore, to find out whether the fermented type of RGM (which is generally considered useless), traditional polyherb formula (which is traditionally used for treating hair loss, but there is a lack of scientific evidence) can actually enhance hair growth, and whether the combination can have the synergic effects, we examined the promoting effects in hair-removed C57BL/6 mice.

## 2. Materials and Methods

### 2.1. Preparation of Fermented RGM and Traditional Polyherb Formula

Fermented RGM was provided by Punggi Ginseng Farming Corp. (Yeongju, Korea). Briefly, after red ginseng was extracted by boiling with distilled water (DW) for 3 h at 80 °C, the remaining RGM was dried and grounded. The powder-form of RGM was fermented with EM containing *Lactobacillus casei*, *Saccharomyces cerevisiae* and *Rhodobacter capsulatus* (Ever Miracle™, Ever Miracle Co., Ltd., Jeonju, Korea) in DW at 45 °C for 15 days. Then, it was extracted by boiling three times in DW for 3 h at 80 °C. The yield efficiency was 40.5%. The extracts of fermented RGM (fRGM) were lyophilized. The components of fRGM were analyzed based on a standard of the Korean FDA: For ginsenosides, the powder of fRGM was dissolved in 80% ethanol and analyzed using an Agilent 1100 high-performance liquid chromatography (HPLC) with Agilent C18 column under the standard
curves for Rg1, Re, Rf, Rg2, Rh1, Rb1, Rc, Rb2, Rb3, Rd, Rg3 and Rh2. For soluble sugars and polysaccharides, the powder was incubated in 80% ethanol at 80 °C for 30 min and centrifuged at 4000×g for 20 min. The supernatant was mixed with 2% anthrone–ethyl acetate and 98% H2SO4 for sugars. The precipitates were homogenized in 1 mol/L HCl for the polysaccharides and then neutralized with 10% NaOH, followed by centrifuging at 4000×g for 10 min. The supernatants were measured at 630 nm under the standard curve using D-glucose. For carbohydrates, the powder of fRGM was incubated with 1 mol/L NaOH in anhydrous ethanol at 80 °C for 10 min and centrifuged at 4000×g for 10 min. The supernatants were incubated with 0.09 mol/L NaOH, 1 mol/L acetic acid and 10% iodine solution for 10 min, and the absorbance was measured at 580 nm. The contents for Pb, As, Hg, Sb and Cd were analyzed using an HPLC system under their standard curves. The potential toxic substances were examined by the Korea Conformity Laboratories (Seoul, Korea) as a certified material test report. All medicinal herbs composing traditional polyherb formula (PH) were purchased from Omniherb pharmacy (Jecheon, Korea) and are listed in Table 1. The fRGM and PH were dissolved in DW and stored at 4 °C until use.

Table 1. Composition of traditional polyherb formula used here.

<table>
<thead>
<tr>
<th>Herbs</th>
<th>Scientific Name</th>
<th>Amounts</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pilose Antler</td>
<td>Cornu cervi pantotrichum</td>
<td>20%</td>
</tr>
<tr>
<td>Old Antler</td>
<td>Cornu cervi</td>
<td>60%</td>
</tr>
<tr>
<td>Ecliptae Herba</td>
<td>Eclipta prostrata L.</td>
<td>1%</td>
</tr>
<tr>
<td>Dictamnii Radicis Cortex</td>
<td>Dictamnus dasycarpus Turczaininov</td>
<td>2%</td>
</tr>
<tr>
<td>Polygoni Multiflori Radix</td>
<td>Polygonum multiflorum Thunberg</td>
<td>8%</td>
</tr>
<tr>
<td>Lycopi Herba</td>
<td>Lycopus lucidus Turczaininov</td>
<td>8%</td>
</tr>
<tr>
<td>Red ginseng</td>
<td>Panax ginseng C. A. Meyer</td>
<td>1%</td>
</tr>
</tbody>
</table>

All of the medicinal herbs were purchased from Omniherb Co, Ltd. (Jecheon, Korea). The amounts of herbs were expressed as percentages contained in a polyherb formula.

2.2. Animal

All experiments were conducted according to the national regulations of the usage and welfare of laboratory animals and approved by the Daegu Haany University—Institutional Animal Care and Use Committee (Gyeongsan, Korea, approval number: DHU2015-019). Six-week-old male C57BL/6 mice were purchased from OrientBio (Seungnam, Korea). The mice were housed in a temperature- (20–25 °C) and humidity- (40–45%) controlled room, with a light/dark cycle of 12/12 h. Feed and water were supplied ad libitum. Animals were acclimatized for 2 weeks (Figure 1).

**Figure 1.** A scheme of study design. Hair-removed model was topically treated with fermented red ginseng marc (fRGM) or traditional polyherb formula (PH) at 400, 200 and 100 mg/kg, or the combination at 200 mg/kg of fRGM and PH in each (combi). The results were compared with those of the control group treated with distilled water (DW).
2.3. Hair Removal and Treatments

The dorsal skin hairs of mice were shaved completely using an electric clipper and a hair waxing cream containing 80% thioglycolic acid (NiClean™, Ildong Pharmaceutical Co., Seoul, Korea) for inducing anagen synchronization, as described elsewhere [28–30]. The next day, a total of 48 mice was divided into eight groups (n = 6 per group) based on their body weights; a control group with DW as a vehicle (DW), six treatment groups with fRGM (fRGM400, fRGM200 and fRGM100) or PH (PH400, PH200 and PH100) at three different doses of 400, 200 and 100 mg/kg in DW. One group was for a combination group with fRGM and PH at 200 mg/kg in each (combi). Treatments were topically applied on the dorsal-hair-removed area of approximately 3 × 5 cm² in a volume of 200 μL per mouse, once a day for 2 weeks.

2.4. Macroscopic and Microscopic Assessment

The dorsal skin image was obtained using a digital camera, and hair growth was assessed as darkening of the dorsal skin, as described previously [28,29]. After treatments for 2 weeks, mice were euthanized using CO₂ gas, and the dorsal skin area was sampled. The half of the samples with hairs were weighed, and the hairs were carefully plucked. Among the hairs, ten hairs per a mouse were collected, and hair length and thickness were assessed under microscopy [31]. They were analyzed using a computer-based image analysis program (iSolution FL ver 9.1, IMT i-solution Inc., Vancouver, BC, Canada). In addition, hair weight was calculated by subtracting the weight of hair-removed skin samples from that of the samples with hairs. Then, the other samples close to the neck were subjected to histopathological analyses.

2.5. Histopathology

The skin sample was fixed in 10% neutral buffered formalin and trimmed crossly and longitudinally based on the sagittal axis of a mouse body. The paraffin-embedded samples were serially sectioned at a thickness of 3 μm. The sections were stained with hematoxylin and eosin (H&E) or immunostained [32]. In H&E stain, histomorphometric analyses were examined for dermal thickness (μm/skin) and numbers of hair follicles (follicles/mm²) in the cross sections, and hair shafts and follicular sizes with hair root sheath (μm/follicle) in the longitudinal sections, using an image analysis program (iSolution FL ver 9.1, IMT i-solution Inc.), as described previously [28,30]. The analyses were performed at least in three histological fields in each section by a histopathologist blinded to the groups.

2.6. Immunohistochemistry

The serial sections were de-paraffinized and conducted for citrate buffer antigen retrieval pretreatment, as described previously [32]. Briefly, the sections were heated in 10 mM citrate buffers (pH 6.0) at 95–100 °C for 20 min. They were fixed in methanol for 30 min and then blocked with normal horse serum solution (1:100, Vector Laboratories, Burlingame, CA, USA) for 1 h. The sections were incubated with a mouse anti-Ki-67 antibody (ab6526, 1:200, Abcam, Cambridge, UK) overnight at 4 °C, and then with Alexa Fluor 488-conjugated goat anti-mouse IgG (1:2000, Molecular Probes, Eugene, OR, USA) for 1 h, followed by mounting with Vectashield™ containing 4,6-diamidino-2-phenylindole (Vector Laboratories). The immunostains were performed in a humidity chamber, and sections were rinsed in 0.01 M PBS three times between each step. The Ki-67 antibody-omitted stains were used as a negative control. Hair follicle cells occupied by over 30% of immunofluorescence were regarded as positive cells. The immunostained cells were expressed as a percentage of total follicle cells more than 100 in the cross and longitudinal sections to the craniocaudal axis of mice. It was assessed by a histopathologist blinded to the groups.
2.7. Statistical Analyses

All data were expressed as means ± standard error (SEM) of six sample sizes. Variance homogeneity was examined using the Levene test. Since the test indicated no significant deviations, the data were analyzed by one-way analysis of variance (ANOVA), followed by Tukey post hoc test. The kinetic changes on hair growth were examined by two-way ANOVA with main factors for the group and the day measured, and the day was treated as repeated measurements. A $p < 0.05$ was considered significant.

3. Results

3.1. Components of fRGM

The fRGM contained 0.2 g/mL carbohydrates and 0.1 g/mL saccharides; however, fats including saturated and trans-fat and cholesterol were not detected. It had crude ginsenosides of 1.71 mg/g and polyphenols of 9.82 ± 0.03 mg/g. The contained ginsenosides were Rb1 (0.6 mg/kg), Rb2 (0.9 mg/kg), Rb3 (1.8 mg/kg), Rc (1.9 mg/kg), Rd (3.5 mg/kg), Re (3.1 mg/kg), Rf (3.9 mg/kg), Rg2 (12.1 mg/kg), Rg3 (9.6 mg/kg) and Rh1 (17.9 mg/kg). The potential toxic substances, such as Pb, As, Hg, Sb and Cd, were few under detection.

3.2. Effects on Hair Growth Region

The hair was observed a little in all groups until day 6 post-treatment and had markedly grown from day 9 post-treatment regardless of treatments. Although hair growth until day 14 post-treatment showed individual differences even in the same group, it tended to be faster in the treatment groups of fRGM, PH and combi than the DW control (Figure 2). Two-way ANOVA for the kinetic hair growth showed no significances for the groups. There were significances for the days measured ($F = 374.1; p < 0.01$, Figure 3), representing just daily hair growth. There were no significant interactions between the groups and the days. However, the total hair growth from the initial (day 0) to the last treatment (day 15) showed differences among groups ($F = 4.2; p < 0.01$). The post hoc versus the DW control revealed significant increases by 1.5-, 1.4-, 1.5- and 1.5-folds in the fRGM400, fRGM200, PH400 and combi groups, respectively ($p < 0.05$).

Figure 2. Kinetic changes in hair-covering regions. The representative images for hair-regrown regions in three of six mice per group are shown on days 0, 6, 9, 12 and 14 after treatments.
Figure 3. Effects on hair growth. The kinetic changes in hair growth (a) and the total hair growth from the initial to the last treatments (b) were assessed. Values are expressed means ± SEM of 6 sample sizes. ** $p < 0.01$ and * $p < 0.05$ versus the DW group.

3.3. Effects on Hair Length, Thickness and Weight

Hair length and thickness were increased in all treatment groups comparing to the DW (Figure 4). There were significant differences among the groups for hair length ($F = 15.9; p < 0.01$) and thickness ($F = 5.2; p < 0.01$). Hair length was significantly increased by 1.3-fold in the fRGM400, fRGM200 and combi groups and by 1.2-fold in the fRGM100 and PH groups ($p < 0.01$). Hair thickness was increased by 1.4-fold in the fRGM400 and combi and 1.3-fold in the other treatment groups ($p < 0.05$). There were also significant differences among the groups for hair weight ($F = 3.7; p < 0.01$). The post hoc tests versus the DW group revealed significant increases by 2.4-, 2.2-, 2.3-, 2.1 and 2.3-fold in the fRGM400, fRGM200, PH400, PH200 and combi groups, respectively ($p < 0.05$). The weights were also increased by 2.0- and 1.7-fold in the fRGM100 and PH100, respectively; however, they were not significant.
Figure 4. Effects on hair quality. After treatments for 2 weeks, ten hairs per mouse and the dorsal skin tissues were sampled. The hairs were analyzed for the length (a) and thickness (b). Hair weight was measured by subtracting weights of hair-removed samples from those of the samples with hairs and normalized by sample areas (c). Values are expressed as mean ± SEM of 6 sample sizes. ** $p < 0.01$ and * $p < 0.05$ versus the DW group.

3.4. Effects on Development of Anagen-Phased Hair Follicle

In both cross and longitudinal sections stained with H&E stain, the dermal thickness was increased in the treatment groups of fRGM, PH and combi than the DW control. Number of the follicles, especially in follicles located at the lower subcutis, and the follicular diameters were also observed as increased in the treatment groups (Figure 5). One-way ANOVA showed significant differences among the groups for the dermal thickness ($F = 3.4; p < 0.01$) and the follicular sizes ($F = 4.1; p < 0.01$). For both data, the post hoc tests versus the DW group revealed significant increases by 1.2-fold in the fRGM400, fRGM200, PH400, PH200 and combi groups ($p < 0.05$).
Figure 5. Effects on hair-growth cycle. Tissue sections of the dorsal skin samples were stained in hematoxylin and eosin, and the representative images are shown in (a). In the cross sections, the dotted boxes were high-magnified in the lower. Scale bars indicate 200 μm. Thickness of the dermis (DE) and diameters of hair follicles in the subcutis (SC) were measured in the cross and longitudinal sections, respectively (b,c). Values are expressed as means ± SEM of 6 sample sizes. ** $p < 0.01$ and * $p < 0.05$ versus the DW group.

The anagen-phase-like follicles with complete root sheath surrounding the bulb were evidently observed in the subcutis of the fRGM, PH and combi groups. One-way ANOVA showed significant differences among the groups for the number of hair follicles in the subcutis ($F = 5.8; p < 0.01$) and the total hair follicles ($F = 4.3; p < 0.01$), however there was no differences in the number in the dermis (Table 2). The ratio of the follicles in the subcutis to the total follicles was also significantly different among the groups ($F = 2.3; p < 0.05$). The post hoc tests versus the DW group revealed significant increases in hair follicles of the subcutis in the fRGM400, fRGM200, PH400, PH200, PH100 and combi groups ($p < 0.05$). The total number of follicles was also significantly increased in the fRGM400, fRGM200, PH400, PH200 and combi groups compared with the DW control ($p < 0.05$). The ratio of the follicles in the subcutis to the total follicle was increased in the PH400 ($p = 0.08$), PH200 ($p = 0.06$) and combi group ($p = 0.07$); however, it failed to approach the significance.
Table 2. Histomorphometric analysis for hair follicles.

<table>
<thead>
<tr>
<th>Groups</th>
<th>Numbers of Hair Follicle/mm² (% in Total)</th>
<th>Subcutis</th>
<th>Dermis</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>DW</td>
<td>26.00 ± 1.68 (52.90 ± 1.39)</td>
<td>23.47 ± 2.31 (47.10 ± 1.39)</td>
<td>49.47 ± 3.80 (100)</td>
<td></td>
</tr>
<tr>
<td>fRGM400</td>
<td>38.30 ± 1.93 ** (58.98 ± 0.98)</td>
<td>26.54 ± 1.03 (41.02 ± 0.98)</td>
<td>64.85 ± 2.69 * (100)</td>
<td></td>
</tr>
<tr>
<td>fRGM200</td>
<td>37.34 ± 1.71 ** (59.27 ± 2.13)</td>
<td>25.68 ± 1.59 (40.73 ± 2.13)</td>
<td>63.02 ± 1.91 * (100)</td>
<td></td>
</tr>
<tr>
<td>fRGM100</td>
<td>33.73 ± 2.80 (56.77 ± 2.04)</td>
<td>25.32 ± 1.33 (43.23 ± 2.04)</td>
<td>59.05 ± 3.45 (100)</td>
<td></td>
</tr>
<tr>
<td>PH400</td>
<td>39.88 ± 2.07 ** (59.60 ± 1.83)</td>
<td>26.98 ± 1.45 (40.40 ± 1.83)</td>
<td>66.86 ± 2.53 ** (100)</td>
<td></td>
</tr>
<tr>
<td>PH200</td>
<td>38.14 ± 1.81 ** (59.93 ± 0.87)</td>
<td>25.41 ± 0.77 (40.07 ± 0.87)</td>
<td>63.55 ± 2.35 * (100)</td>
<td></td>
</tr>
<tr>
<td>PH100</td>
<td>35.88 ± 2.32 * (58.98 ± 1.62)</td>
<td>24.82 ± 1.33 (41.02 ± 1.62)</td>
<td>60.70 ± 3.15 (100)</td>
<td></td>
</tr>
<tr>
<td>combi</td>
<td>41.26 ± 1.05 ** (59.83 ± 1.33)</td>
<td>27.97 ± 2.06 (40.18 ± 1.33)</td>
<td>69.23 ± 2.88 ** (100)</td>
<td></td>
</tr>
</tbody>
</table>

The hair follicles were assessed in hematoxylin–eosin stain of the cross sections to the cranio-caudal axis of mice. Values are expressed as mean ± SEM of 6 sample sizes. ** p < 0.01 and * p < 0.05 versus the DW group.

3.5. Proliferating Effects on Hair Follicle

The immunostain for Ki-67 as a proliferating marker was few in the DW control in the cross and longitudinal sections. However, a few were observed in the bulb of hair follicles in the subcutis of the treatment groups (Figure 6). Although the immunostained hair follicles were approximately 2–3%, they were observed more in the treatment groups than the DW group. Actually, ANOVA showed significant differences among the groups in the cross (F = 4.3; p < 0.01) and longitudinal sections (F = 3.1; p < 0.05). The immunostained follicles were increased by 2.0-fold in the fRGM400, fRGM200, PH400 and fRGM200 groups, and 2.3-fold in the combi groups in the cross sections, compared with the DW group (p < 0.05). They were also increased by 2.6-fold in the fRGM400 and PH400, and 2.8-fold in the combi in the longitudinal sections (p < 0.05).

Figure 6. Effects on cellular proliferation. The serial sections were immunostained for Ki-67 as a maker for proliferation (green) and counterstained with 4,6-diamidino-2-phenylindole (blue) (a). Scale bars indicate 100 μm. The immunostained
cells were assessed in the cross (b) and longitudinal sections (c) and expressed as a percentage of total follicle cells. Values are mean ± SEM of 6 sample sizes. ** $p < 0.01$ and * $p < 0.05$ versus the DW group.

4. Discussion

Hair growth was rapidly increased from day 9 after hair removal and waxing, similar with other studies [28–30]. The kinetic hair growth on each day was not significantly different among the groups, probably due to small sample sizes or limitation of the measurements to detect the subtle changes on each day. In addition, the individual differences shown in the same group might be involved in the incomplete anagen synchronization, which may result in a failure to approach the significant differences. However, the higher doses of fRGM and PH increased the final grown hair regions on day 0 to 15 post-treatment. Hair length and thickness increased for all doses of fRGM and PH, and hair weight increased for the treatment groups except the low-dose groups. The combination treatments also showed beneficial effects on hair growth and hair quality, even though the combination dose was half the high doses of fRGM and PH. Hair weighing and microscopic measurements of hair length and width are generally used in clinical studies to evaluate the effects of drugs or cosmetic molecules as a hair tonic [33]. Further, clinical diagnosis for hair loss needs scalp biopsy to observe the follicular quantity and diameter and other morphometric data for abnormal hair cell growth. The current histopathological analyses revealed that components of fRGM and PH contribute to increasing follicular numbers, diameters and proliferation, in particular in the hair follicles located at the deep subcutis. These suggest that the fRGM and PH may promote hair-growth cycle, probably by enhancing anagen-phased cell division, as well as functions as a hair tonic.

The length and thickness of a hair shaft depend on the duration of the hair follicular cycle, in particular duration of anagen phase [34]. Indeed, androgenetic alopecia produces thinning of scalp hair, and the miniaturization of hair follicles involves shortened anagen phase and reduced sizes of the hair matrix [3]. In this context, enhancing the induction of anagen or prolonging the duration can be a therapeutic strategy for hair loss. The fRGM, PH and the combination increased the hair follicles closed to the deep layer of subcutis, suggesting promoting effects on development of the follicles in anagen III to VI or catagen stages [34]. It might result in the increased total hair follicles in the treatments of the higher doses of fRGM, PH and the combination. The increases in anagen-phased follicles might be due to induction of an earlier anagen phase or transition of hair growth from telogen to anagen [35,36]. The transformation from telogen- to anagen-phased hair follicles undergoes through rapid proliferation of follicular keratinocytes and elongation and thickening of the hair shaft [37]. Here, considering that the fRGM and PH and the combination increased Ki-67 positive cells in the bulb of the subcutis, the anagen-like hair follicles could be regenerated by rapid proliferation of the matrix keratinocytes, producing increased follicular sizes for a new hair fiber. It suggests that the components contained in the fRGM and PH may contribute to induction of anagen-phased follicles and improve the altered hair-growth cycles or destroyed hair follicles in hair loss [2,38].

Hair growth is regulated by various hormones, cytokines and growth factors. For example, androgens reduce the anagen phase, increasing the other phased follicles and delaying the telogen to anagen transitions [39]. Basic FGF promotes proliferation of the papilla cells, contributing to elongate the hair shaft [40], and hepatocyte growth factor and vascular endothelial growth factor (VEGF) stimulate growth of the hair follicle and hair shaft [41,42]. The growth factors and others including epidermal growth factor, keratinocyte growth factor (KGF), insulin-like growth factor-I and transforming growth factor have shown mitogenic and motogenic effects on keratinocyte [43]. The traditional poly- herb formula used here is composed of seven herbs, and the components may contribute to enhancing the hair-growth cycle. Cornu cervi pantotrichum and Eclipta prostrata facilitate hair growth through upregulation of FGF-7 (KGF) and FGF-5 [16,17]. In red ginseng, Rb1 and Rd stimulate proliferation of hair follicle cells [15,27], and other phenolic compounds
and vitamins give specific support to the maintenance of hair follicle health [44]. In particular, vitamin C is known to improve blood vessel formation, increasing blood flow in the scalp by stimulating VEGF synthesis [45]. There is a report showing promoting potentials of Rb1 on hair growth through activating extracellular signal-regulated kinase and AKT signaling pathways [10]. The mitogen-activated pathway plays an important role in cell growth and proliferation [46,47]. Further, *Eclipta prostrata, Polygonum multiflorum Thunberg* and red ginseng have been reported to inhibit 5-α reductase [10,11,19]. The paracrine factors have been assumed to stimulate or regulate hair growth and the proliferation of hair matrix cells, and further studies need to clarify the exact mechanisms.

As increases in the concern on the well-being in life, the demands to improve one’s appearance instigate the consumption of functional products for hair health. Further, alopecia is a widespread condition affecting both sexes: 50% over 50 years of age in males and about 50% in females [6]. The clinical signs are usually milder in females than males; however, hair loss creates emotional stress in both. Most of the hair-loss-related disorders need daily controls, which increases interests in functional shampoo products in the market because of an easy cure. Although a variety of medicated shampoos are commercially available, the therapeutic benefits are ambiguous. Further, thousands of natural products including traditional polyherbs are used as hair tonics, based on traditional experiences rather than the scientific effectiveness. Several herbs have shown beneficial effects on hair loss as a complementary and alternative medicine [48,49], while effectiveness of many other herbs needs to be clarified. Here, topical treatments of the fRGM and PH for 2 weeks showed promoting effects on hair growth, probably by enhancing the induction of anagen or prolonging the duration. Interestingly, the fRGM contained a very low concentration of ginsenosides compared with those of the red ginseng [50]; however, it still had the functional effects on hair growth as an additive component rather than the byproduct. Given that the fRGM contained few toxic compounds, and all treatments had little dermatological side effects, it is expected to use fRGM in combination with the effective traditional polyherb formula for the synergistic effects.

5. Conclusions

In the present study, the hair-depilated mice model showed individual differences even in hair growth in the same treatment group, probably due to the incomplete anagen synchronization. Although there were no significant differences in hair-covering regions on a specific day during the treatments, the fRGM and PH significantly increased the regions that were finally regrown on days 0 to 14 and also increased hair length, thickness and weight. In addition, follicular numbers, thickness and proliferation were observed in deep layer of the subcutis, mainly composed of anagen-phased follicles. The beneficial effects were greater in the high doses of fRGM and PH and the combi groups. It suggests that the fRGM and PH may promote hair growth via anagen-phased cell divisions, along with functions as a hair tonic. Future study needs to clarify the promoting effects on the growth cycle in proper models and the relevant mechanisms. These provide useful information for guiding the use of fRGMs and PH in patients with hair loss.


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**References**


