The efficacy of a pseudo-ceramide and eucalyptus extract containing lotion on dry scalp skin

Yutaka Takagi1
Xuanhe Ning1
Akihiko Takahashi1
Chiaki Ishizaki2
Takahiro Nishizaka1
Kazuhiro Kaizu1
Masahiro Miyaki1
Hotaka Yamamuro1
Makoto Kawashima3

1Skin Care Product Research, Kao Corporation, Tokyo, Japan; 2Department of Skin Research Center, Inforward Corporation, Tokyo, Japan; 3Department of Dermatology, Tokyo Women’s Medical University, Tokyo, Japan

Introduction

Ceramides are major components of intercellular lipids in the stratum corneum1 and play a crucial role in the barrier function2 and moisture-retention ability (MRA) of the skin.3 Skin with decreased stratum corneum functions, such as seen in patients with atopic dermatitis,4,5 has low ceramide levels in the stratum corneum.6–7 The efficacy of topically applied ceramides, pseudo-ceramides, or ceramide precursors on atopic dermatitis has been reported.8–13

Pruritic inflammatory skin with a reduced cutaneous barrier function and MRA is also observed on the scalps of patients with atopic dermatitis, despite the conditions of high humidity caused by thick hair, and sweat and sebaceous glands. Dry and scaly eruptions on the scalp are also observed in diseases such as seborrheic dermatitis and pityriasis capitis.14 Harding et al reported that scalps with dandruff have low ceramide levels.15 Thus, an increase in ceramide levels may improve scalp conditions, not only in patients with atopic dermatitis but also in patients with seborrheic dermatitis and/or pityriasis capitis.

Imokawa et al synthesized a pseudo-ceramide, cetyl PG hydroxyethyl palmitamide (Figure 1), to compensate for natural ceramides8 and found that the topical application
of pseudo-ceramide on the skin of patients with atopic dermatitis and on subjects with sensitive skin enhanced their stratum corneum functions.\textsuperscript{10,12} Furthermore, we reported that a eucalyptus extract is effective in increasing ceramide levels in the epidermis.\textsuperscript{16}

In this study, we evaluated the efficacy of a lotion containing pseudo-ceramide and a eucalyptus extract on dry and scaly scalp conditions of patients with atopic dermatitis, seborrheic dermatitis, or pityriasis capitis. Environmental factors are known to affect the functions of the stratum corneum.\textsuperscript{17} Since the scalp is often covered with thick hair and is protected from exposure to the outer environment, we performed this evaluation during the winter season when the meteorological influence is lowest.

**Methods**

**Subjects**

This evaluation was performed as a single dermatologist-controlled baseline comparison study and adhered to the tenets of the Declaration of Helsinki. All clinical studies were reviewed and approved by the Review Board of the Shinkohkai Medical Corporation (Tokyo, Japan), and written informed consent was obtained from each subject before the study.

Subjects with dry skin symptoms on their scalp were prescreened from candidates with atopic dermatitis, seborrheic dermatitis, or pityriasis capitis by visual observation and by medical interviews conducted by the dermatologists. Subjects who were receiving medical treatment on the scalp and/or were using anti-dandruff or anti-itchiness materials on the scalp were excluded from the study. They were excluded prior to the screening.

Of the 67 Japanese females who were pre-screened, 33 subjects aged 22–55 years (mean=40.5±9.9 years) were enrolled in this study. Nineteen subjects with atopic dermatitis (mild: 12 subjects; moderate: 7 subjects, classified by the grading system of Rajka & Langeland),\textsuperscript{18} 6 subjects with seborrheic dermatitis, and 9 subjects with pityriasis capitis were enrolled. Two subjects with moderate atopic dermatitis were taking an oral medical treatment as noted below, and none of the subjects were receiving topical medical treatment on the scalp.

**Materials**

All test materials used in this study were provided by Kao Corporation (Tokyo, Japan). The test scalp lotion was a fragrance-free, color-free, and weakly acidic, and contained synthetic pseudo-ceramide (cetyl PG hydroxyethyl palmitamide)\textsuperscript{8} and eucalyptus leaf extract.\textsuperscript{16} The other ingredients of the test lotion included: water (aqua), butylene glycol, betaine, glycerin, polyethylene glycol, *Thujopsis dolabrata* branch extract, L-arginine, sodium methyl stearoyl taurate, and methylparaben.

The test lotion was applied once a day after each patient used a shampoo. Changes of shampoo, conditioner, hair styling products, and/or hair care products during this study were prohibited. Hair perms or coloration were also prohibited.

**Evaluations**

Prior to evaluating the efficacy of the scalp lotion, the severity of skin symptoms on each part of the scalp was assessed as follows: The scalp was divided into 23 areas (5 areas on the front and top of the head, 5 areas on the right side of the head, 5 areas on the left side of the head, and 8 areas on the back of the head) as shown schematically in Figure 2. Following acclimation in a climate-controlled room at 20°C±2°C and 40%±5% relative humidity for 20 minutes, 2 well-trained specialists performed assessments of scaling and erythema on each area of the scalp. Scaling and erythema were rated on a scale of 1–4 (1=none, 2=mild, 3=moderate, and 4=severe). Each mean value (0.5) was also obtained as a score. To
compare the severity of scalp skin symptoms between the areas of the head, the average score of each region was used. The skin condition of each area was recorded using a Skin-Watcher® SW-40 (COREFRONT Corp., Tokyo, Japan).

The efficacy of the lotion was assessed from evaluations at the end of weeks 2 and 4 compared with the start of the study (week 0). All assessments were performed at the instrumental analyzing area after removal of the hair with scissors and acclimation in the climate-controlled room at 20°C±2°C and 40%±5% relative humidity for 20 minutes. The dermatologist performed assessments of dryness, scaling, erythema, itchiness, and scabbing on the whole scalp, and rated those symptoms on a scale of 1–5 (1=none, 2=slight, 3=mild, 4=moderate, and 5=severe).

Instrumental analysis
The instrumental measurements were performed on a 10×10 mm area on the left side of the head (70 mm above the ear) of each patient, where the hair was removed by scissors. MRA and transepidermal water loss (TEWL) were measured using a Corneometer® (Courage + Khazaka electronic GmbH, Cologne, Germany) and a Vapometer® (Delfin Technologies Ltd, Kuopio, Finland), respectively. The average values of 3 MRA measurements and 2 TEWL measurements were used for analysis.

Subjective evaluation by the test subjects
Subjective evaluation of the itchiness of the scalp was obtained from each patient using the visual analog scale (VAS). The VAS consisted of a 100-mm horizontal line marked 0 (no itch) to 100 (maximum intensity of itch). Patients marked their itch on this line by themselves.19 The quality of life (QoL) was analyzed using the Skindex-16®.20

Judgment of safety
At the week 4 evaluation, the dermatologist judged the safety and usefulness of the test material. The safety was judged according to a diary kept by each subject and by inquiry on each examination day. In case of interruption of the trial, the suspected cause and reason were described. Immediately after an interruption, an assertion of safety was performed.

Statistical analysis
All data were statistically analyzed using the one-way repeated measures analysis of variance test or the Wilcoxon signed-rank test using StatView 5.0 (SAS Institute Inc., Cary, NC, USA).

Results
This study was performed in Tokyo, Japan, from January through February in 2015. The average temperature was 5.8°C±2.0°C and the relative humidity was 55.1%±17.7%. There were relatively small changes in temperature and relative humidity during this period even though there were daily variations (Figure 3).

In the initial evaluation (week 0), the back of the head had the most severe scaling and erythema, whereas there was less on the side of the head, and least on the front and top of the head (Figure 4). These differences in the severity of skin symptoms based on the head regions were similar to other kinds of skin diseases (data not shown).

Table 1 indicates the changes in severity of skin symptoms following usage of the scalp lotion. There were no statistically significant differences in any of those skin symptoms between the patients with atopic dermatitis and those with the other 2 diseases at week 0. At the week 4 evaluation, dryness, scaling, erythema, and itchiness were significantly improved in all 34 subjects. Highly significant improvements in scaling, erythema, and itchiness were observed even at week 2. Scabbing was found only on 5 subjects at the baseline (week 0) with a score of 2, which decreased to 4 subjects at week 4 (no significance, data not shown). The efficacy of the scalp lotion was found to be similar for patients with atopic dermatitis and those with seborrheic dermatitis or pityriasis capitis. Figure 5 shows representative photographs demonstrating the
Figure 4 Differences in the severity of scaling (A) and erythema (B) at week 0 in the scalp region.

Notes: Each column indicates the number of subjects with the range of average scores. *p<0.05 and **p<0.01, Wilcoxon signed-rank test.

Table 1 Dryness, scaling, erythema, and itchiness on the scalp of subjects with AD or SD/PC at weeks 0, 2, and 4

<table>
<thead>
<tr>
<th>Score</th>
<th>Total</th>
<th>AD</th>
<th>SD/PC</th>
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<tbody>
<tr>
<td></td>
<td>W0</td>
<td>W2</td>
<td>W4</td>
</tr>
<tr>
<td>Dryness</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 (none)</td>
<td>3</td>
<td>4</td>
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</tr>
<tr>
<td>2 (slight)</td>
<td>11</td>
<td>16</td>
<td>17</td>
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<tr>
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<td>4 (moderate)</td>
<td>5</td>
<td>7</td>
<td>5</td>
</tr>
<tr>
<td>5 (severe)</td>
<td>3</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
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<td>**</td>
<td>ns</td>
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<tr>
<td>Scaling</td>
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<td>**</td>
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<tr>
<td>p-value</td>
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</table>

Notes: *p<0.05, **p<0.01, and “ns” indicates no significant difference from week 0. Obtained using the Wilcoxon signed-rank test.

Abbreviations: AD, atopic dermatitis; SD, seborrheic dermatitis; PC, pityriasis capitis.
The efficacy of lotion on dry scalp skin

Clinical improvement of the symptoms on the different areas of the head. In patients with atopic dermatitis (Figure 5A) and in those with seborrheic dermatitis (Figure 5B), scaling and erythema on all areas of the scalp were diminished at the end of the study.

Itchiness, measured using the VAS, improved significantly at weeks 2 and 4 (Figure 6), and the MRA gradually and significantly increased at weeks 2 and 4 (Figure 7A). The average TEWL value of all subjects at week 0 was consistent with the TEWL value of healthy scalp skin (6.6±1.6 g/m²·hour)²¹ and there were no significant changes during the test period (Figure 7B).

No adverse events due to treatment with the test material were recorded in the diary kept by each subject throughout this evaluation. From this result and the inquiry at each visual evaluation at weeks 2 and 4, this scalp lotion was judged as “safe” for all subjects.

Analysis by the Skindex-16 revealed significant improvements in symptoms, emotions, functioning, and globally at week 4 (Figure 8).

Discussion
The scalps of patients with seborrheic dermatitis or pityriasis capitis have lower ceramide levels in their stratum corneum.¹⁵,²² This decrease in ceramide induces a lower MRA and cutaneous barrier function, causing skin diseases such as atopic dermatitis.³,⁵ We have confirmed that the topical application of pseudo-ceramide (cetyl PG hydroxyethyl palmitamide) improves the decreased stratum corneum function of facial and body skin of patients with atopic dermatitis.¹⁰,¹² Furthermore, we found that eucalyptus extract is effective in

Figure 5 Typical clinical features at weeks 0 and 4.
Notes: (A) A 23-year-old female subject with mild atopic dermatitis. (B) A 51-year-old female subject with seborrheic dermatitis.

Figure 6 Itchiness evaluated by VAS at weeks 0, 2, and 4. 0 mm indicates no itch and 100 mm represents a maximum intensity of itch.
Note: **p<0.01, a significant difference from week 0, obtained using a one-way repeated measures analysis of variance test.
Abbreviation: VAS, visual analog scale.
increasing ceramide levels in the epidermis. In this study, we clarified the efficacy of the topical application of a lotion containing pseudo-ceramide and eucalyptus extract on the dry and scaly scalps of patients with atopic dermatitis, seborrheic dermatitis, or pityriasis capitis, as shown by the decrease of scaliness, erythema, and itchiness.

In this evaluation, all subjects constantly had scalp symptoms prior to evaluation. Furthermore, there were only slight changes in the climate environment, such as the daily temperature and humidity, around weeks 0, 2, and 4, even though the daily relative humidity did change. Additionally, changes of shampoo, conditioner, hair styling products and hair care products, and hair perms or coloration were prohibited during this study. Thus, we conclude that any changes in scalp conditions are caused by usage of the scalp lotion. Because the subjects used various shampoos, conditioners, and hair styling products each day, the selection of a placebo scalp lotion was quite difficult. For these reasons, this evaluation was performed as a single dermatologist-controlled baseline comparison study.

Because of the presence of pseudo-ceramides in the lotion, the total ceramide level in the scalp could not be analyzed in this study. However, we have previously confirmed that a topically applied eucalyptus extract increases the ceramide level of the stratum corneum in the scalp. Furthermore, the pseudo-ceramide remained in the stratum corneum for 16 hours after topical application (data not shown). Since the efficacy of topically applied ceramides, pseudo-ceramides, or ceramide precursors on atopic dermatitis has been reported, it is surmised that both increased ceramide and applied pseudo-ceramides might improve the condition of the stratum corneum.

Cleansing with high-performance detergent is also a major factor that induces skin irritation by extracting natural moisturizing factors and intercellular lipids. However, in this study, the subjects did not change their shampoos, conditioners, or hair styling products. Thus, this study clarified the efficacy of the scalp lotion to increase ceramide levels to improve scalp conditions, as it does in other areas of the body for patients with atopic dermatitis.

Two subjects with moderate atopic dermatitis had an oral medical treatment to take care of their body skin symptoms. Both of them ingested an antihistamine medicine and they had kept taking this medicine before the start of this evaluation. The dermatologist judged that this medication would have no effect on the stratum corneum condition and unless
changing this treatment, this oral medical treatment may have little or no effect on result. Thus, they kept taking that same medication.

There were no statistically significant changes in TEWL during the 4-week test period. The subjects in the study had TEWL value similar to that of a healthy scalp (6.6±1.6 g/m²·hour),29 so changes of the TEWL value, if any, were too small to be detected.

There was no control group who did not use a scalp lotion but there were only small changes in temperature and relative humidity during this period (average temperature and relative humidity at the week 4 evaluation was almost same the week 0 evaluation). Thus, the environment might have less effect on the improvement of scalp conditions.

There are several studies of the placebo effects on itch.26 Because there were no placebo treatments, it is presumed that the decrease in itchiness might be caused by placebo effects, but the improvement of other skin symptoms, such as dryness, scaling, erythema, and the increase in MRA support the improvement of itchiness by this test scalp lotion.

Analysis by the Skindex-16 revealed significant improvements in symptoms, emotions, functioning, and globally at week 4. Functioning was also improved but the score at week 0 was low and thus the decrease was not significantly different. Similar to the improvements of itchiness, some proportion of the benefit may be due to a placebo effect, but improvements of dryness, scaling, and erythema, and also the increase in MRA support the improvements of QoL.

Limitations

There are some limitations to this study that need to be addressed in future studies. 1) The study assessed only females and whether similar effects occur on males with these skin diseases needs to be assessed. 2) The study did not include a control lotion without pseudoceramide nor eucalyptus extracts, so lotions with either of those actives or none should be performed to clarify which active(s) are effective. 3) The study covered only 4 weeks of evaluations, so whether a longer-term study would identify continued beneficial effects, or potentially adverse effects, should be elucidated.

Conclusion

From these data, the efficacy of the daily applications of a scalp lotion containing pseudo-ceramide and a eucalyptus extract to increase the MRA and improve the condition of the scalp was shown and the QoL of the patients was improved through the use of this lotion.

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Author contributions

All authors contributed toward data analysis, drafting and revising the paper and agree to be accountable for all aspects of the work.

Disclosure

Akihiko Takahashi, Xuanhe Ning, Takahiro Nishizaka, Kazuhiro Kaizu, Masahiro Miyaki, Yutaka Takagi, and Hotaka Yamamuro are employees of the Kao Corporation. Chiaki Ishizaki is an employee of Infoward Inc. Makotokawashima received an honorarium from Kao Corporation for supervising this study. The authors report no other conflicts of interest in this work.

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