A Novel Cream Formulation Containing Nicotinamide 4%, Arbutin 3%, Bisabolol 1%, and Retinaldehyde 0.05% for Treatment of Epidermal Melasma

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Epidermal melasma is a common hyperpigmentation disorder that can be challenging to treat. Although current treatment options for melasma are limited, topical skin-lightening preparations have widely been used as alternatives to hydroquinone. In this prospective, single-arm, open-label study, treatment of epidermal melasma with a novel cream formulation containing nicotinamide 4%, arbutin 3%, bisabolol 1%, and retinaldehyde 0.05% was associated with reductions in Melasma Area and Severity Index (MASI) scores as well as total melasma surface area as measured by medical imaging software. Treatment outcomes including tolerance and safety profiles as well as patient satisfaction and product appreciation showed this novel cosmetic compound may be valuable in the treatment of epidermal melasma.


Epidermal melasma is a common hyperpigmentation disorder characterized by the appearance of abnormal melanin deposits in different layers of the skin. Melasma can be difficult to treat and often recurs due to the migration of new melanocytes from hair follicles to the skin’s surface. A novel cream formulation containing nicotinamide 4%, arbutin 3%, bisabolol 1%, and retinaldehyde 0.05% offers a safe and effective option for treatment of epidermal melasma.

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Epidermal melasma is a common hyperpigmentation disorder that can be challenging to treat. The pathogenesis of melasma is not fully understood but has been associated with increased melanin and melanocyte activity.1,2 Melasma is characterized by jagged, light- to dark-brown patches on areas of the skin most often exposed to the sun—primarily the cheeks, forehead, upper lip, nose, and chin.3 Although it can affect both sexes and...
all races, melasma is more common in Fitzpatrick skin types II to IV and frequently is seen in Asian or Hispanic women residing in geographic locations with high levels of sun exposure (eg, tropical areas). Melasma presents more frequently in adult women of childbearing age, especially during pregnancy, but also can begin postmenopause. Onset may occur as early as menarche but typically is observed between the ages of 30 and 55 years. Only 10% of melasma cases are known to occur in males and are influenced by such factors as ethnicity, hormones, and level of sun exposure.

Topical therapies for melasma attempt to inhibit melanocytic activation at each level of melanin formation until the deposited pigment is removed; however, results may vary greatly, as melasma often recurs due to the migration of new melanocytes from hair follicles to the skin’s surface, leading to new development of hyperpigmentation. The current standard of treatment for melasma involves the use of hydroquinone and other bleaching agents, but long-term use of these treatments has been associated with concerns regarding unstable preparations (which may lose their therapeutic properties) and adverse effects (eg, ochronosis, depigmentation). Cosmetic agents that recently have been evaluated for melasma treatment include nicotinamide (a form of vitamin B₃), which inhibits the transfer of melanosomes from melanocytes to keratinocytes; arbutin, which inhibits melanin synthesis by inhibiting tyrosinase activity; bisabolol, which prevents anti-inflammatory activity; and retinaldehyde (RAL), a precursor of retinoic acid (RA) that has powerful bleaching action and low levels of cutaneous irritability.

This prospective, single-arm, open-label study, evaluated the efficacy and safety of a novel cream formulation containing nicotinamide 4%, arbutin 3%, bisabolol 1%, and retinaldehyde 0.05% in the treatment of epidermal melasma.

**Study Product Ingredients and Background**

**Nicotinamide**—Nicotinamide is a water-soluble amide of nicotinic acid (niacin) and one of the 2 principal forms of vitamin B₃. It is a component of the coenzymes nicotinamide adenine dinucleotide and nicotinamide adenine dinucleotide phosphate. Nicotinamide essentially acts as an antioxidant, with most of its effects exerted through poly(adenosine diphosphate–ribose) polymerase inhibition. Interest has increased in the role of nicotinamide in the prevention and treatment of several skin diseases, such as acne and UV radiation–induced deleterious molecular and immunological events. Nicotinamide also has gained consideration as a potential agent in sunscreen preparations due to its possible skin-lightening effects, stimulation of DNA repair, suppression of UV photocarcinogenesis, and other antiaging effects.

**Arbutin**—Arbutin is a molecule that has proven effective in treating melasma. Its pigment-lightening ingredients include botanicals that are structurally similar to hydroquinone. Arbutin is obtained from the leaves of the bearberry plant but also is found in lesser quantities in cranberry and blueberry leaves. A naturally occurring glucosidase, arbutin reduces tyrosinase activity without affecting messenger RNA expression. Arbutin also inhibits melanosomal maturation, is nontoxic to melanocytes, and is used in Japan in a variety of pigment-lightening preparations at 3% concentrations.

**Bisabolol**—Bisabolol is a natural monocular sesquiterpene alcohol found in the oils of chamomile and other plants. Bisabolol often is included in cosmetics due to its favorable anti-inflammatory and depigmentation properties. Its downregulation of inducible nitric oxide synthase and cyclooxygenase-2 suggests that it may have anti-inflammatory effects.

**Retinaldehyde**—Retinaldehyde is an RA precursor that forms as an intermediate metabolite in the transformation of retinol to RA in human keratinocytes. Topical RAL is well tolerated by human skin, and several of its biologic effects are identical to those of RA. Using the tails of C57BL/6 mouse models, RAL 0.05% has been found to have significantly more potent depigmenting effects than RA 0.05% (P<.001 vs P<.01, respectively) when compared to vehicle.

Although combination therapy with RAL and arbutin could potentially cause skin irritation, the addition of bisabolol to the combination cream used in this study is believed to have conferred anti-inflammatory properties because it inhibits the release of histamine and relieves irritation.

**Methods**

This single-center, single-arm, prospective, open-label study evaluated the efficacy and safety of a novel cream formulation containing nicotinamide 4%, arbutin 3%, bisabolol 1%, and RAL 0.05% in treating epidermal melasma. Clinical evaluation included assessment of Melasma Area and Severity Index (MASI) score, photographic analysis, and in vivo reflectance confocal microscopy (RCM) analysis.

The study population included women aged 18 to 50 years with Fitzpatrick skin types I through V who had clinically diagnosed epidermal melasma on the face. Eligibility requirements included confirmation of epidermal pigmentation on Wood lamp examination and RCM analysis and a MASI score.
of less than 10.5. A total of 35 participants were enrolled in the study (intention to treat [ITT] population). Thirty-three participants were included in the analysis of treatment effectiveness (ITTe population), as 2 were excluded due to lack of follow-up postbaseline. Four participants were prematurely withdrawn from the study—3 due to loss to follow-up and 1 due to treatment discontinuation following an adverse event (AE). The last observation carried forward method was used to input missing data from these 4 participants excluding repeated measure analysis that used the generalized estimated equation method.

At baseline, a 25-g tube of the study cream containing nicotinamide 4%, arbutin 3%, bisabolol 1%, and RAL 0.05% was distributed to all participants for once-daily application to the entire face for 30 days. Participants were instructed to apply the product in the evening after using a gentle cleanser, which also was to be used in the morning to remove the product residue. Additionally, participants were given a sunscreen with a sun protection factor of 30 to apply daily on the entire face in the morning, after lunch, and midafternoon. During the 30-day treatment period, treatment interruption of up to 5 consecutive days or 10 nonconsecutive days in total was permitted. At day 30, participants received another 30-day supply of the study product and sunscreen to be applied according to the same regimen for an additional 30-day treatment period.

Clinical Evaluation—At baseline, demographic data and medical history was recorded for all participants and dermatologic and physical examination was performed documenting weight, height, blood pressure, heart rate, and baseline MASI score. Following Wood lamp examination, participants’ faces were photographed and catalogued using medical imaging software that allowed for measurement of the total melasma surface area (Figure 1A). The photographs also were cross-polarized for further analysis of the pigmentation (Figure 1B).

A questionnaire evaluating treatment satisfaction was administered to participants (ITTe population \(n=33\)) at baseline and days 30 and 60. Questionnaire items pertained to skin blemishes, signs of facial aging, overall appearance, texture, oiliness, brightness, and hydration. Participants were instructed to rate their satisfaction for each item on a scale of 1 to 10 (1 = bad, 10 = excellent). For investigator analysis, scores of 1 to 4 were classified as “dissatisfied,” scores of 5 to 6 were classified as “satisfied,” and scores of 7 to 10 were classified as “completely satisfied.” A questionnaire evaluating product appreciation was administered at day 60 to participants who completed the study \(n=29\). Questionnaire items asked participants to rate the
study cream’s ease of application, consistency, smell, absorption, and overall satisfaction using ratings of “bad,” “regular,” “good,” “very good,” or “excellent.”

Treatment efficacy in all participants was assessed by the investigators at days 30 and 60. Investigators evaluated reductions in pigmentation and total melasma surface area using ratings of “none,” “regular,” “good,” “very good,” or “excellent.” Local tolerance also was evaluated at both time points, and AEs were recorded and analyzed with respect to their duration, intensity, frequency, and severity.

Targeted hyperpigmented skin was selected for in vivo RCM analysis. At each time point, a sequence of block images was acquired at 4 levels of skin: (1) superficial dermis, (2) suprabasal layer/dermoepidermal junction, (3) spinous layer, and (4) superficial granular layer. Blind evaluation of these images to assess the reduction in melanin quantity was conducted by a dermatopathologist at baseline and days 30 and 60. Melanin quantity present in each layer was graded according to 4 categories (0%–25%, 25.1%–50%, 50.1%–75%, 75.1%–100%). The mean value was used for statistical evaluation.

Results
Efficacy evaluation—The primary efficacy variable was the mean reduction in MASI score from baseline to the end of treatment (day 60), which was $2.25 \pm 1.87$ ($P < .0001$). The reduction in mean MASI score was significant from baseline to day 30 ($P < .0001$) and from day 30 to day 60 ($P < .0001$). The least root-mean-square error estimates of MASI score variation at days 30 and 60 were 1.40 and 2.25, respectively.

The mean total melasma surface area (as measured in analysis of clinical photographs using medical imaging software) was significantly reduced from 1398.5 mm² at baseline to 1116.9 mm² at day 30 ($P < .0001$) and 923.4 at day 60 ($P < .0001$). From baseline to end of treatment, the overall reduction in mean total melasma surface area was 475.1 mm² ($P < .0001$) (Figure 2). Clinical and cross-polarized photographs taken at day 60 demonstrated a visible reduction in melasma surface area (Figure 3), which was confirmed using medical imaging software.

In vivo RCM analyses at each time point showed reduction in pigmentation in the 4 levels of the skin that were evaluated, but the results were not statistically significant.

Participant satisfaction—There was strong statistical evidence of patient satisfaction with the treatment results at the end of the study period ($P < .0001$). At baseline, 75.8% (25/33) of participants were dissatisfied with the appearance of their skin as compared with 15.2% (5/33) at day 60. Additionally, 18.1% (6/33) and 6.1% (2/33) of the participants were satisfied and completely satisfied at baseline compared with 33.3% (11/33) and 51.5% (17/33) at day 60, respectively. Participant satisfaction with signs of facial aging also increased over the study period ($P = .0104$). At baseline, 60.6% (20/33) were dissatisfied, 12.1% (4/33) were satisfied, and 27.3% (9/33) were completely satisfied; at the end of treatment, 30.3% (10/33) were dissatisfied.

![Figure 2. Mean surface area of melasma measured at baseline (1398.5 mm²), day 30 (1116.9 mm²), and day 60 (923.4 mm²), showing a mean total reduction of 475.1 mm² from baseline to day 60.](image-url)
36.4% (12/33) were satisfied, and 33.3% (11/33) were completely satisfied with the improvement in signs of facial aging.

Increased patient satisfaction with facial skin texture at baseline compared to day 60 also was statistically significant ($P=.0157$). At baseline, 39.4% (13/33) of the participants were dissatisfied, 30.3% (10/33) were satisfied, and 30.3% (10/33) were completely satisfied with facial texture; at day 60, 15.1% (5/33) were dissatisfied, 30.3% (10/33) were satisfied, and 54.6% (18/33) were completely satisfied. Significant improvement from baseline to day 60 also was observed in participant assessment of skin oiliness ($P=.0210$), brightness ($P=.0003$), overall appearance ($P<.0001$), and hydration ($P<.0001$).

**Product appreciation**—At day 60, 89.7% (26/29) of the participants who completed the study rated the product’s ease of application as being at least “good,” with more than half of participants (55.2% [16/29]) rating it as “very good” or “excellent.” Overall satisfaction with the product was...
rated as “very good” or “excellent” by 48.3% (14/29) of the participants. Similar results were observed in participant assessments of consistency, smell, and absorption (Figure 4).

Safety evaluation—A total of 52 AEs were observed in 23 (69.7%) participants, which were recorded by participants in diary entries throughout treatment and evaluated by investigators at each time point. Among these AEs, 48 (92.3%) were considered possibly, probably, or conditionally related to treatment by the investigators based on clinical observation. The most common presumed treatment-related AE was a burning sensation on the skin, reported by 30.3% (10/33) of the participants at day 30 and 13.8% (4/29) at day 60. Of the reported AEs related to treatment, 91.7% (44/48) were of mild intensity and 93.8% (45/48) required no treatment or other action. There were no reported serious AEs related to the investigational product. Blood pressure, heart rate, and weight remained stable among all participants throughout the study.

The intensity of the AEs was described as “light” in 91.7% (44/48) of cases and “moderate” in 8.3% (4/48) of cases. The frequency of AEs was classified as “unique,” “intermittent,” or “continuous” in 45.8% (22/48), 39.6% (19/48), and 14.6% (7/48) of cases, respectively. Of the 48 AEs, 3 (6.3%) occurred in 1 participant, necessitating interruption of treatment, application of the topical corticosteroid cream containing mometasone, and removal from the study.

The amount of melanin present in each layer of the skin. These limitations suggest that scales used in future in vivo RCM analyses of melasma should be narrower.

Epidermal melasma is one of the most difficult dermatologic diseases to treat and control. Maintenance of clear, undamaged skin remains a treatment target for all dermatologists. This novel cream formulation containing nicotinamide 4%, arbutin 3%, bisabolol 1%, and RAL 0.05% has proven to be an effective, safe, and tolerable treatment option for patients with epidermal melasma.

REFERENCES