



Low-level laser therapy for the treatment of androgenetic alopecia in Thai men and women: a 24-week, randomized, double-blind, sham device-controlled trial

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Abstract

Low-level laser/light therapy (LLLT) has been increasingly used for promoting hair growth in androgenetic alopecia (AGA). Our institute developed a new home-use LLLT device, RAMACAP, with optimal penetrating energy, aiming to improve therapeutic efficacy and compliance. To evaluate the efficacy and safety of the new helmet-type LLLT device in the treatment of AGA, a 24-week, prospective, randomized, double-blind, sham device-controlled clinical trial was conducted. Forty subjects with AGA (20 men and 20 women) were randomized to treat with a laser helmet (RAMACAP) or a sham helmet in the home-based setting for 24 weeks. Hair density, hair diameter, and adverse events were evaluated at baseline and at weeks 8, 16, and 24. Global photographic assessment for hair regrowth after 24 weeks of treatment was performed by investigators and subjects. Thirty-six subjects (19 in the laser group and 17 in the sham group) completed the study. At week 24, the laser helmet was significantly superior to the sham device for increasing hair density and hair diameter ($p = 0.002$ and $p = 0.009$, respectively) and showed a significantly greater improvement in global photographic assessment by investigators and subjects. Reported side effects included temporary hair shedding and scalp pruritus. In conclusion, the novel helmet-type LLLT device appears to be an effective treatment option for AGA in both male and female patients with minimal adverse effects. However, the limitations of this study are small sample size, no long-term follow-up data, and use of inappropriate sham devices, which do not reflect the true negative control. Trial registration: <http://clinicaltrials.in.th/index.php?tp=regtrials&menu=trialsearch&smenu=fulltext&task=search&task2=view1&id=2061>, identifier TCTR20160910003.

Keywords Low-level laser therapy · Low-level light therapy · Photobiomodulation · Androgenetic alopecia · Female pattern hair loss

Introduction

Androgenetic alopecia (AGA) is a common hair loss disorder, which affects approximately 80% and 50% of Caucasian men and women by age 70, respectively [1, 2]. It is characterized by the gradual transformation of terminal to vellus-like hairs resulting in hair thinning with a characteristic pattern. Clinically, men are presented with anterior hairline recession and hair thinning on the vertex. In women, this disorder, also called female pattern hair loss (FPHL), manifests with diffuse

hair thinning, mostly on the mid-scalp, and preservation of the frontal hairline. Patients with AGA commonly report a negative psychological impact on the quality of life [3]. Oral finasteride and topical minoxidil are current medications approved by the Food and Drug Administration (FDA) for treating AGA. These two drugs generally provide a favorable result. However, patients experiencing a poor response or unacceptable side effects are in need of additional treatment options. Low-level laser/light therapy (LLLT) has been increasingly used as an alternative treatment for AGA.

LLLT generally utilizes non-thermal effects of low-intensity light at red or near-infrared wavelengths to alter biological activity in cells, termed photobiomodulation or photobiostimulation. Its beneficial therapeutic effects have been shown in a wide range of medical conditions, including wound healing, nerve regeneration, pain reduction, and hair loss [4, 5]. Although the precise mechanisms of action are not yet well established, strong evidence suggests that LLLT acts

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on mitochondria leading to an increase of reactive oxygen species (ROS) levels, adenosine triphosphate (ATP) production, and induction of transcription factors, which activate genes and produce proteins useful to the cell [4–7].

Hair growth-stimulating effect of LLLT was accidentally discovered by Endre Mester in 1967. Since then, several studies from animal models to clinical trials have cumulatively supported its benefits with minimal adverse events, establishing a novel treatment option for hair loss [4, 5]. LLLT appears to stimulate anagen reentry of telogen hair follicles and prolong the duration of anagen phase [8–10]. The modulation of hair cycle reveals an increase of hair density and diameter as well as a decrease of hair shedding, resulting in clinical improvement of alopecias. Even though LLLT has been introduced to treat hair loss for over a decade, many questions remain about the better light source between laser and light-emitting diode (LED), the most effective wavelength, optimal therapeutic regimens, and maintenance regimens, and how to choose the right candidate.

The first LLLT device was cleared for use as a safe modality in the treatment of AGA by the US FDA in 2007 for men and in 2011 for women. After that the market has been flooded with numerous LLLT devices claimed to promote hair growth. However, only a small number of these devices have proved their efficacy and safety in a well-designed randomized controlled study. With regard to LLLT devices used for hair loss, there are three basic categories: hood, brush-based (combs, hairbrushes, or headbands), and hat-based (helmets or caps) systems. These devices use coherent diode lasers, incoherent LEDs, or a combination of both as the source of light in the wavelength range of 650–900 nm. Home-use and hands-free devices are principally designed for improving treatment adherence.

In Thailand, a few portable LLLT devices with high-quality clinical evidence are commercially available. The laser comb and laser headband have parting teeth allowing the laser light to reach the scalp surface, but manual movement is needed to treat an extensive scalp area. The helmet combining laser and LED works hands free. However, the query on scalp absorption is mentioned as its main power comes from LED light, which is not collimated and more divergent than laser light. Therefore, a new model of helmet-type LLLT device was invented and named “RAMACAP” under the collaboration between our institute, the Division of Dermatology, Ramathibodi Hospital, and the National Innovation Agency (NIA), a public organization established by the Royal Thai Government, aiming to provide an easy-to-use device with sufficient energy to reach intended targets, in particular, follicular mitochondria. RAMACAP is a portable laser helmet with full scalp coverage and working entirely hands free. The purpose of this study was to assess its efficacy and safety for the treatment of AGA in both men and women.

Methods

Study design

A 24-week, prospective, randomized, double-blind, sham device-controlled clinical trial was conducted at a university-based hospital (Ramathibodi Hospital, Mahidol University, Bangkok, Thailand). The study protocol was approved by the Mahidol University Institutional Review Board on January 20, 2016 (protocol number 01–59-22; clinicaltrials.in.th identifier: TCTR20160910003). Date of first enrollment was January 26, 2016.

Study participants

Men aged over 18 years with AGA classified as type III vertex, IV, and V by the Norwood-Hamilton classification, and women aged over 18 years with FPHL classified as type I, II, and III by the Ludwig classification were eligible for the study. The exclusion criteria included use of finasteride or dutasteride within 18 months, use of other topical or systemic drugs with hair growth-promoting properties within 6 months, a history of hair transplantation, and patients with a scalp or systemic disease that may affect hair growth. Prior to participation, each eligible subject must provide a written informed consent.

Intervention

RAMACAP is a combat helmet-shaped device containing single-mode laser diodes, which emit at the wavelength of 660 ± 10 nm (Fig. 1). It consists of three layers: (1) the outer protective layer is made from acrylonitrile butadiene styrene plastic; (2) the middle layer is flexible printed circuit boards with 224 pieces of red laser diodes, arranging into four equal sectors; and (3) the inner layer is made from clear thermoplastic elastomers, in order to protect direct contact between laser diodes and human skin/hair. The biocompatibility of inner layer is verified by tests for in vitro cytotoxicity according to ISO 10993-5 and tests for irritation and skin sensitization according to ISO 10993-10.

The maximum optical power for each laser diode is 5 mW with pulse wave at 4.2 Hz. As a longer distance from the source leads to a lesser light intensity to transmit and absorb, laser bulbs of this model nearly touch the scalp by wearing. The distribution of laser bulbs is also designed for an even coverage across the scalp with a power density of 3.5 mW/cm^2 on the scalp surface. An illumination time of 19.04 min is required to deliver a fluence of 4 J/cm^2 , which appears to be an appropriate dosage for the purpose of stimulating hair growth. Regarding the light source, laser is selected over LED since laser beams are collimated and less divergent, supposing to have better light transmission to intended targets under the

Fig. 1 RAMACAP, a helmet-type portable LLLT device containing 224 red diode laser bulbs (660 nm)



scalp surface. This LLLT system meets FDA requirements for class 3R (IIIa) laser products.

Subjects were divided by using stratified randomization into experimental and control groups. Subjects were firstly stratified by sex and randomly assigned in a 1:1 ratio to use either a laser or a sham device. Laser and sham devices were identical in appearance with the exception of the light source. The sham device contained 224 red LED pods emitting 0.5 mW of power at the wavelength of 650 ± 20 nm. Subjects were instructed to use an assigned device at home for 20 min per session, three times per week, over a 24-week period, as well as to maintain the same hair color, length, and style throughout the study. Investigators and subjects remained blinded to the type of assigned device until study completion.

Assessments

The primary efficacy endpoint was the change from baseline in hair density and diameter on the target area of the scalp at weeks 8, 16, and 24, photographing with a Folliscope® and measuring with Folliscope 2.8 software (LeadM Corporation, Seoul, Korea). The measurement point was a 1-cm-diameter round area either on the vertex of male subjects or on the mid-scalp of female subjects, mapped by the intersection point of three distances from fixed anatomical landmarks (tip of the nose, right, and left tragus).

Secondary efficacy endpoints included global photographic assessment (GPA) for hair regrowth by investigators and subjects. Global photographs were taken at baseline and week 24, using a Nikon D5100 DSLR camera (Nikon Corporation, Tokyo, Japan) mounted on the stereotactic head-positioning device for exposure fixation. The photographs were also

controlled for camera angles and lighting. Paired photographs were evaluated by a panel of three dermatologists, blinded to study assignment, and the subjects themselves, based on 7-point rating scale (-3 = marked deterioration, -2 = moderate deterioration, -1 = mild deterioration, 0 = no change, 1 = mild improvement, 2 = moderate improvement, 3 = marked improvement). All possible adverse events were recorded at every follow-up visit.

Statistical analyses

The sample size calculation was based on data from the previous LLLT clinical study in the Asian population [13]. The mean change from baseline to study endpoint in hair density of subjects in the LLLT group was an increase of 17.2 hairs/cm² with a standard deviation of 12.1 hairs/cm², and those of subjects in the sham control group was a decrease of 2.1 hairs/cm². To achieve a power of 80% and a two-sided significance level of 5%, the minimum sample size required was seven in each group (the experimental group and the control group).

Statistical analysis was performed using SPSS version 18.0 (SPSS Inc., Chicago, IL, USA). The data were not analyzed separately for male and female subjects as the preliminary analysis showed no statistically significant difference in outcomes between genders. Differences in demographic data and global assessment scores were measured using the independent *t* test or chi-squared test depending on data characteristics. Fleiss's kappa was used to evaluate agreement among three investigators. Changes from baseline in hair density and hair diameter between groups were compared by repeated measures analysis of variance and subsequently analyzed by Fisher's protected least significant difference test.

Results

A total of 40 subjects, consisting of 20 men and 20 women, were enrolled and underwent stratified randomization into the experimental group and the control group (10 men and 10 women in each group). Nineteen subjects (10 men and 9 women) in the experimental group and 17 subjects (9 men and 8 women) in the control group completed the study. Four subjects withdrew from the study due to personal reasons that were not treatment-related. Analyses for demographic data and efficacy assessments were performed with the intention-to-treat population. A per-protocol analysis was applied for safety evaluation. There were no statistically significant differences between two groups regarding demographics, baseline characteristics, and the severity of hair loss (Table 1).

At week 24, the laser device was significantly superior to the sham device for increasing hair density and hair diameter ($p = 0.002$ and $p = 0.009$, respectively). Laser device-treated subjects showed a significantly greater increase in hair density at weeks 16 and 24, as well as in hair diameter at week 24, compared with sham device-treated subjects (all $p < 0.05$; Fig. 2). The mean change from baseline to week 24 in hair density was 10.21 ± 3.25 hairs/cm² in the laser group versus 3.95 ± 1.32 hairs/cm² in the sham group. The mean change from baseline to week 24 in hair diameter was 6.11 ± 2.15 μ m in the laser group versus 3.76 ± 1.24 μ m in the sham group.

Table 1 Demographics and baseline characteristics of the participants

	RAMACAP (N = 20)	Sham device (N = 20)
Age (years)		
Mean (SD)	38.9 (11.3)	35 (14.1)
Age of onset of hair loss (years)		
Mean (SD)	31.1 (11.1)	29 (15.3)
Duration of hair loss (years)		
Mean (SD)	7.8 (3.7)	6 (2.2)
Classification		
Norwood-Hamilton (10 men), n (%)		
III vertex	3 (30)	2 (20)
IV	5 (50)	6 (60)
V	2 (20)	2 (20)
Ludwig (10 women), n (%)		
I	4 (40)	5 (50)
II	5 (50)	4 (40)
III	1 (10)	1 (10)
Hair density (hair/cm ²)		
Mean (SD)	112.4 (18.9)	115.4 (16.7)
Hair diameter (μ m)		
Mean (SD)	53.8 (11.6)	54.2 (15.1)

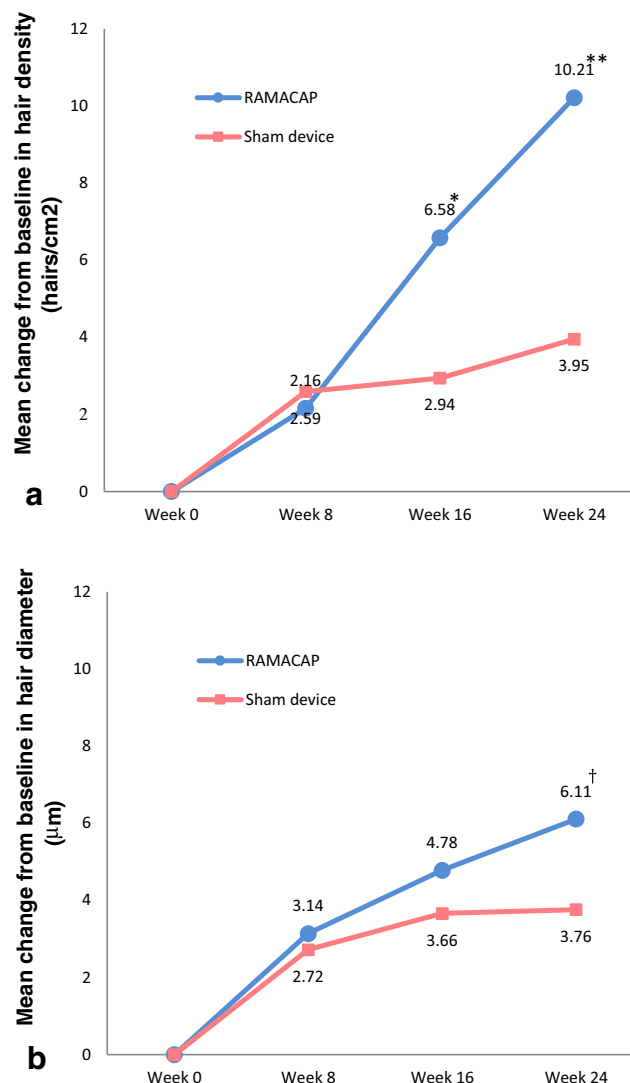
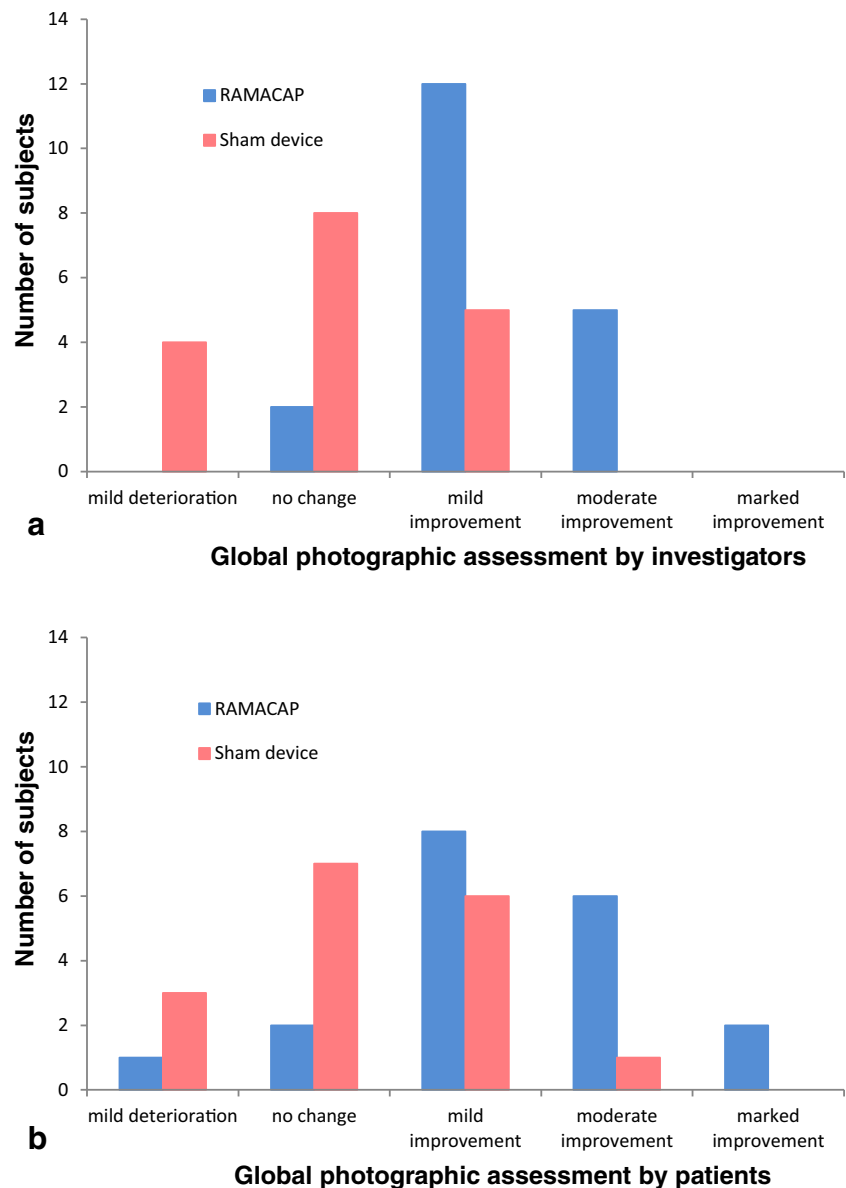


Fig. 2 The mean change from baseline in hair density (**a**) and hair diameter (**b**) at weeks 8, 16, and 24 of laser helmet-treated patients and sham device-treated patients. * $p = 0.001$, ** $p = 0.002$; † $p = 0.009$

Global photographic assessments for hair regrowth after 24 weeks of treatment by investigators and subjects are demonstrated in Fig. 3. Subjects treated with the laser device showed a significantly greater improvement in hair regrowth compared with subjects treated with the sham device for both investigators' evaluation and subjects' self-assessment ($p = 0.0002$ and $p = 0.0026$, respectively). There was substantial agreement on individual scores rated by a panel of three investigators (Fleiss' kappa = 0.65). Examples of global photographs of laser device-treated subjects, who were rated as having moderate improvement, are shown in Fig. 4.

Incidence of adverse reactions was not significantly different between two groups. There was no serious adverse event reported in any subject. One female subject in the laser group complained of increased hair shedding, which occurred at 2 weeks after starting treatment and spontaneously resolved within 6 weeks. Mild scalp itching was described in two laser-

Fig. 3 Global photographic assessment based on paired global photographs from baseline to week 24 by investigators (a) and patients (b). Subjects treated with the laser device showed a significantly greater improvement for both evaluations ($p = 0.0002$ and $p = 0.0026$, respectively)



treated subjects and one sham-treated subject without the need for treatment. No subject experienced an adverse event that resulted in the discontinuation or interruption of the study.

Discussion

LLLT continues to gain popularity in the treatment of AGA due to the ease of use and absence of significant side effects. Apart from anecdotal evidence and non-controlled studies, LLLT showed superior hair growth-promoting effects in men and women with AGA compared to the sham-operating control group in eight randomized controlled clinical trials with the end point varying from 16 to 26 weeks. LLLT devices used in these studies include (1) 655-nm laser combs [11, 12]; (2) a 635-nm and 655-nm laser comb [12]; (3) a helmet

using 630-nm LED, 660-nm LED, and 650-nm laser [13]; (4) a helmet using 655-nm LED and 655-nm laser [14, 15]; (5) a helmet using 660-nm LED and 650-nm laser [16]; (6) a 650-nm laser dome fixed in a sport-style hat [17]; and (7) a 655-nm and 808-nm laser scanner [18]. A recent systematic review and network meta-analysis, evaluating the efficacy of non-surgical treatments for AGA, indicated that LLLT is a highly effective treatment option for AGA; however, the quality of evidence is very low as determined by the risk of bias assessment. Most trials reported funding and support from the device manufacture [19]. High-quality randomized controlled studies are required to confirm this benefit.

The cellular and molecular action mechanisms of LLLT are now much better understood than in the past. Cytochrome C oxidase (CCO), the terminal enzyme in the mitochondrial respiratory chain, has been most often identified as the primary



Fig. 4 Baseline and week 24 global photographs of laser helmet-treated patients who were rated as having moderate improvement

photoacceptor or chromophore in LLLT. However, CCO has a weak absorption at wavelengths longer than 900 nm, which suggests the existence of other chromophores. Intracellular water in light-sensitive ion channels, such as the transient receptor potential vanilloid, has been proposed as another candidate chromophore, especially at mid- and far-infrared wavelengths, based on its high absorption capacity in the infrared spectrum [20–22]. The most widely accepted mechanism of LLLT is that nitric oxide, a competitive inhibitor of oxygen binding, is photodissociated from its binding sites in CCO, allowing an immediate influx of oxygen, which drives the electron transport chain to generate ATP and ROS. Afterward, changes in cellular redox state will activate redox-sensitive transcription factor nuclear factor kappa B and activator protein-1. These transcription factors enhance gene transcription and protein synthesis, contributing to cell proliferation and migration [5, 6, 22].

Regarding LLLT and hair growth-promoting effects, the speculated mechanism mentioned earlier is the direct stimulation of hair follicle stem cells in the bulge region, thereby inducing differentiation and proliferation via increasing the level of heat shock proteins (HSPs), such as HSP27 [4, 9, 23]. Increasing evidence suggests that the transition to anagen phase might be a result of combination effects between (1) direct stimulation on cell proliferation of outer root sheath keratinocytes (ORSKs) and dermal papilla cells (DPCs) and

(2) the release of paracrine growth factors from DPCs, which in turn stimulate proliferation of ORSKs. The ORSK proliferation is upregulated by activating the extracellular signal-regulated kinase (ERK) pathway, while the DPC proliferation is stimulated via the Wnt/ β -catenin and ERK pathways. Several potential paracrine mediators from the dermal papilla induced by LLLT are suggested, hepatocyte growth factor, leptin, vascular endothelial growth factor, fibroblast growth factor 7, lymphoid enhancer-binding factor 1, noggin, Wnt10a, and Wnt10b [5, 10, 24–26].

Biphasic dose response has been demonstrated in a variety of in vitro studies and animal experiments on LLLT [6, 27]. This response follows the Arndt-Schulz rule, which states that for every substance, small doses stimulate, moderate doses inhibit, and large doses kill. Even though convincing instances have not been reported in patients, systematic reviews of randomized controlled clinical trials on LLLT mentioned that some ineffective trials might be explained by overdosing on energy [27]. In addition, effects of LLLT appear to depend on irradiance (or power density; W/cm^2) and illumination time rather than fluence (or energy density; J/cm^2) [6, 27]. Different irradiances but the same fluence show different results. Insufficient irradiance or too short an illumination time will have no response. Too high irradiance or too long an illumination time will have inhibitory effects. Only an optimal balance between power density and time will produce stimulating

effects. It is also believed that the absolute figures of this curve are different at different wavelengths and tissue types [27]. Although there is scanty data on the Arndt-Schulz curve in terms of stimulating hair growth, light within the red spectrum at doses of 2 to 4 J/cm² is suspected to be appropriate, regardless of the light source [5, 27]. Shukla et al. showed that the exposure of mice to 632.8-nm laser, emitting 5 mW/cm², at a dose of 1 J/cm² led to a significant increase in the number of anagen hair follicles, whereas those at a dose of 5 J/cm² led to a significant decrease, compared to the control group [8].

This study demonstrated superiority of RAMACAP, the laser helmet, over the sham device by significantly increasing in hair density after 16 weeks and hair diameter after 24 weeks. Analysis of global photographic assessment data also showed a significantly greater improvement in laser helmet-treated patients by both investigator and subject scores. The safety profile was similar to those of devices in the previous clinical trials. There were no serious adverse events reported throughout the study period. One patient in the laser helmet group experienced transient hair shedding during the first 2 months of treatment. Temporary increased hair shedding was also reported in another clinical trial of a 655-nm laser comb. By contrast, it arose in about one third of studied subjects [28]. Increased hair shedding induced by LLLT is most likely related to the synchronization of hair cycle by stimulating telogen hair follicles to reenter the anagen phase, leading to an increase of the exogen phase. An additional adverse effect was mild scalp pruritus, which occurred in both groups. Wearing a helmet might create a warmer environment and higher humidity on the scalp, possibly leading to itchiness.

From the preliminary analysis, when comparing male versus female responses to the laser helmet, the results revealed no statistically significant differences between genders with respect to changes in hair density and diameter, as well as rating scores of GPA for hair regrowth. Likewise, there has been no published article demonstrating a difference in treatment outcomes between male and female patients receiving LLLT for hair loss. This might be explained by the fact that the main mechanism of hair regrowth induced by LLLT is hypothesized to be the stimulation of hair follicle stem cells, ORSKs, and DPCs. The cellular functions of these cells appear to be identical among men and women.

The outcomes of LLLT for hair loss might depend on interindividual differences in hair density, hair length, hair color, and skin color, which are factors affecting absorption and transmission of the light [5]. Long hair, thick layering of hair, dark hair, and dark skin will absorb and reflect more light, resulting in a lower transmission. A bioengineering study investigating the effects of hair color on LLLT demonstrated that light transmission was reduced by 32% for blond hair

and 37% for black hair [5]. We also observed that subjects who had apparent hair thinning experienced a greater improvement compared to those with milder severity.

In the sham control group undergoing treatment with the 650-nm LED helmet, some patients also showed mild improvement in hair regrowth. This unexpected result could be explained by inadequate light source used in the control group. Hair growth-stimulating effect of LED lights has been previously reported in the literature [4, 10]. However, since the delivered energy of our sham device was extremely low, therapeutic responses were almost invisible.

The main limitations of this study are the small number of subjects and no long-term follow-up data. A small sample size may reduce the power of the study and increase the margin of error as well as the study subjects may not represent the entire population. The 6-month study duration could not evaluate long-term efficacy and safety. In addition, using a more adequate sham device which yields a true negative effect in the control group could reduce the risk of potential bias in the study. A well-planned large randomized controlled trial with a longer study period is needed to support the benefits and determine whether tachyphylaxis or inhibitory effects could occur. Moreover, studies on optimal therapeutic and maintenance regimens are also essential.

In conclusion, the novel helmet-type LLLT device named RAMACAP might be an effective and well-tolerated treatment option for AGA, at least during the first 6-month treatment period, without age and gender restrictions. The device can be considered as an adjunct to a standard medication or monotherapy for people who refuse to receive pharmacological treatments.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflicts of interest.

Ethical approval This study was conducted in accordance with the principles of the Declaration of Helsinki and in compliance with the International Conference on Harmonization-Good Clinical Practice and local regulatory requirements. The study was reviewed and approved by the appropriate Independent Ethics Committees, and written informed consent was obtained from all subjects prior to study initiation.

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