Antihyperpigmentation Effect of The Combination of Turmeric (*Curcuma domestica* Val.) and Bitter Melon Leaves (*Momordica charantia* L.) Ethanol Extracts on Guinea Pig Skin

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**Abstract**

Turmeric (T) and Bitter Melon leaves (BM) extract has been proven in decreasing melanin contents in in vitro study, but their single extracts effects were lower than the positive control. A study confirmed the combination of plants extracts had melanogenic effect better than the positive control and their single extracts. This study aimed to investigate the anti-hyperpigmentation effect of the combination of T and BM extract on guinea pig skin and compared with the positive control group. This study used a post-test control design. Twenty-five guinea pigs were divided into 5 groups. The negative control group was given by dimethyl sulfoxide; the positive control group was given by a pharma cream that consists of hydroquinone, tretinoin, and fluocinolone acetonide. The combination of extracts was given to experimental groups with doses 500 µg/mL of T and 200 µg/mL of BM; 750 µg/mL of T and 400 µg/mL of BM; 1,000 µg/mL of T and 600 µg/mL of BM, respectively. All groups exposed to UV-B light in 2 minutes/day for 2 weeks. Each experimental group was given 1 ml combination extract once a day for 2 weeks and in the last step, skin biopsies were done. The histopathological examination was conducted by staining with Fontana-Masson and Nuclear Fast Red. The average percentage of melanin area were compared in all group and analyzed with the Kruskal Wallis test followed by Mann-Whitney test with 95% of confidence level. The result showed group-2 and 3 had the better effect than pharma cream.

**Keywords:** Antihyperpigmentation; Turmeric; Bitter melon; Guinea pig; Melanin

**Abstrak**

Ekstrak rimpang kunyit dan ekstrak daun pare terbukti mampu menurunkan kandungan melanin secara in vitro namun efek keduanya lebih rendah dibandingkan kelompok kontrol positif. Penelitian sebelumnya membuktikan kombinasi ekstrak tanaman mempunyai efek antimelanogenik yang lebih besar dibandingkan kelompok kontrol positif dan masing-masing ekstrak tunggal. Penelitian ini bertujuan membuktikan adanya efek antihiperpigmentasi kombinasi ekstrak etanol rimpang kunyit dan daun pare pada kulit marmut jantan serta membandingkan dengan krim farma. Desain penelitian ini menggunakan post-test only control group design. Ekstraksi menggunakan metode maserasi dengan etanol 70%. Dua puluh lima ekor marmut dipaparkan sinar UVB 2 menit/hari selama 2 minggu kemudian dibagi menjadi 5 kelompok. Kelompok kontrol negatif diberi dimetil sulfoksida sedangkan kelompok kontrol positif diberi krim farma yang mengandung hidrokuinon 4%; tretinoin 0,05%; dan fluocinolone acetonide 0,01%. Kelompok perlakuan diberi 1 ml ekstrak per hari selama 2 minggu dengan kombinasi ekstrak pada kelompok 1 sebesar 500 µg/mL kunyit dan 200 µg/mL daun pare, kelompok 2 sebesar 750 µg/mL kunyit dan 400 µg/mL daun pare, dan kelompok 3 sebesar 1000 µg/mL kunyit dan 600 µg/mL daun pare. Setelah perlakuan dilakukan biopsi jaringan. Pemeriksaan histopatologi dilakukan dengan pewarnaan Fontana-Masson dan Nuclear Fast Red. Perbedaan rata-rata persentase luas melanin pada selulur kelompok penelitian diuji menggunakan Kruskal-Wallis test yang dilanjutkan Mann-Whitney test dengan taraf kepercayaan 95%. Data menunjukkan bahwa kelompok perlakuan kombinasi 2 dan 3 mempunyai efek antihiperpigmentasi yang lebih baik dibandingkan krim farma.

**Kata kunci:** Antihiperpigmentasi; Kunyit; Pare; Marmut; Melanin
INTRODUCTION

Indonesia is a tropical country that many obtain exposure to ultraviolet light. This exposure will trigger the skin to produce Reactive Oxygen Species (ROS). If ROS continue to be produced, it will activate DNA transcription nuclear factor which triggering tyrosinase enzyme to convert tyrosine in several steps into melanin. The excessive melanin production can lead into hyperpigmentation in the epidermal layer skin and cause the skin become darker.

The melanin formation can be prevented with synthetic substances that are usually contained in cosmetics, for example are kojic acid and hydroquinone. Kojic acid is more stable than hydroquinone. However, these compounds cause some side effects. According to Miyazawa, kojic acid is carcinogenic and its use in high concentrations can damage the skin. Meanwhile, long-term use of hydroquinone can cause some side effects, such as contact dermatitis, irritation, hyperpigmentation post-inflammation, and okronosis.

These side effects are the reason for finding inhibitors of melanin production that is safe for skin, for example from natural compounds which have lower side effects. The natural antioxidant compounds have been shown could inhibit melanogenesis or melanin formation, for example are curcumin and flavonoids. One of the plants containing curcumin is Turmeric (T) and containing flavonoids is Bitter Melon leaves (BM).

The anti-hyperpigmentation research with curcumin and bitter melon leaves showed anti-hyperpigmentation effect in in vitro study. Last research by Sugiharto, et al showed that turmeric extract at a concentration of 25 mg/mL was able to degrade melanin by 45.67-53.87% in B16-F1, which is lower than kojic acid’s effect (59.03-66.76%). Tsai, et al showed the bitter melon leaves extracts at a concentration of 200 ug/mL was able to reduce melanin by 11.29% in the B16-F10 cell culture, which is lower than kojic acid’s effect (41.20%). The other research by Rizza et al showed that several Mediterranean plant extracts possessed an inhibitory effect on tyrosinase enzyme. Each extract showed a similar inhibiting activity but less intensive than kojic acid and hydroquinone. Otherwise, when the extracts were combined, it could produce a significant higher activity than kojic acid and hydroquinone.

The in vitro studies based on the provision of treatment outside of a living organism without describing the effects on living organisms. So, in this study we conducted research on guinea pig skin with expectations to describe the activity of the substances in humans by the biological similarity. This study aimed to investigate antihyperpigmentation combination of T and BM extract on guinea pig skin and compared to pharma cream as the positive control group. The anti-hyperpigmentation effect is based on the observation of the amount of melanin in the epidermis layer.

METHOD

The method adopted from Hastiningsih and Indiradewi’s research. This research was an experimental research with a post-test control group design. This study had approved by the medical research ethics committee of Medical Faculty of Diponegoro University Semarang, to be carried out on animal testing with ethical clearance number 828/EC/FK-RSDK/2016. Twenty-five male guinea pigs adapted for one week. The lower back hair of guinea pigs shaved with the size 3x3 cm. Then, guinea pigs were exposed to ultraviolet B light in 2 minutes/day for 2 weeks and divided into five groups. The negative control group was given by dimethyl sulfoxide (DMSO), the positive control group was given by a pharma cream that consists of hydroquinone 4%, tretinoin 0.05%, and fluocinolone acetonide 0.01%. The pharma cream is a triple fixed combination agent for hyperpigmentation that discovered by...
The 1-3 of treatment groups were treated with combination of both extracts (500 µg/mL T and 200 µg/mL BM), (750 µg/mL T and 400 µg/mL BM), and (1,000 µg/mL of T and 600 µg/mL BM). The dose selection based on the conversion of in vitro concentration for every single extract to in vivo of guinea pig skin. Each treatment was given by 1 ml extract once a day for 2 weeks. On day 15, the guinea pigs were euthanized with ether and tissue biopsy was done with 2 mm depth (up to subcutaneous) in the pigmented lesions with a length of 2 cm. The skin tissue was embedded in the 40% formalin solution and prepared for calculating the amounts of melanin.

Materials and Tools
Bitter melon leaves and turmeric were purchased from the bitter melon and turmeric plant cultivation at Boja, Central Java. Their fresh and dry parts used as the requirement for this study. This research used ethanol (Brataco, 70%), dimethyl sulfoxide (Sigma, p.a), Formalin (Hepilab), formalin buffer (Hepilab), xylene (Merck, 99%), sodium thiosulfate solution (Merck, 0,1 mol/L), nuclear fast red (Sigma, nuclear fast red in 5% of aluminum sulfate).

The amount of melanin’s analysis used Opti-lab Pro camera and Olympus CX41 microscope with 400x magnification. The image results were edited using Adobe Photoshop CS3 software version 10:01.

The extraction process
Plant materials were cleaned with water. Simplicia dried powder were made by using a blender and then sieved with a sieve size of 25 mesh. Before maceration process, the water content in the dry simplicia should be less than 10%. The extraction process was done by maceration method with ethanol 70%. The simplicia powder was weighed approximately 500 grams put in a jar and added 3,75 L of 70% ethanol, closed and left for 5 days, protected from light while stirring occasionally at least three times a day for 5 days. The filtrate was mixed and concentrated by using a rotary evaporator at temperatures <50ºC to obtain a thick extract.

Histological Preparation
The guinea pig skin tissues inserted into 10% formalin phosphate buffer solution for 1 day. Then, the tissue sections were trimmed and soaked consecutively in alcohol concentration of 30%, 40%, 50%, 70%, 80%, 90%, 96%, respectively 3 times for 25 minutes. The third step is clearing phase, the skin tissue was soaked in clearing agent (alcohol: xylene 1:1) for 30 minutes and dipped in pure xylene until transparent. After infiltrating 4 times with pure paraffin, the skin tissue was embedded in liquid paraffin until it shaped into a block (± 1 day) so could be easily slice with a microtome. Tissue cutting process was done using a Leica microtome 820 with 5 μ thickness, taken sliced serially into 5 μ, 10 μ, 15 μ, and then the tissue slice was stucked on sticker-smeared object glass.

Staining with Fontana-masion and nuclear fast red
The slides were soaked twice in xylene, each for 5 minutes, then soaked in 100%, 95%, and 70% ethanol, and in dH₂O respectively for 2 minutes. After that, the slides were soaked again in Silver Nitrate Fontana solution for 2 hours and incubated at 56°C in an oven. The slides were washed three times with dH₂O, dropped by 1% Gold Chloride solution and left in for 5 minutes. The slides were reashed three times with dH₂O and dropped 5% Sodium thiosulfate solution, left in for a minute. Then slides were washed with dH₂O and stained with Nuclear Fast Red for 5 minutes. After that, the slides were washed twice with dH₂O and dehydrated with 70%, 95% and 100% ethanol respectively for 20 seconds. Thus, the slides were cleared twice using xylene respectively for 2 minutes and mounted on a xylene-based
medium. Finally, the slides were stained with melanin and showed black melanin granules with pink cell nucleus and pale pink cytoplasm.

**Procedures of result observation**

The amount of melanin was calculated by digital analysis method. Every preparation was photographed three times using an Optilab Pro camera and Olympus CX41 microscope with 400x magnification and saved in JPEG format. The image was edited using Adobe Photoshop CS3 software version 10:01, to pick epidermal tissue at the left, middle and right side of the preparation with the Polygonal Lasso tool. The field of view was taken, an area which indicated most melanin was marked with a black area.

**Calculation procedure of the amount of melanin**

The calculation of the amount of melanin in pixel units was done with the Image version 1.47t software using the red channel with threshold adjustment. The amount of normal melanin was calculated per field of view as follows:

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\text{pixel of melanin} / \text{pixel of epidermis} \times 100\%
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**Data analysis**

The number of melanin differences in the entire test group was analyzed with the Kruskal Wallis test followed by Mann-Whitney test with 95% of confidence level.

**RESULTS AND DISCUSSION**

Guinea pigs were used in this study because they were easy to obtain, inexpensive, easy to handle, had many biological similarities with humans, and had some kinds of melanin, i.e. eumelanin, phaeomelanin, and albinos. Male Guinea pigs were used to avoid the influence of Melanocyte Stimulating Hormone (MSH), estrogen, and progesterone that are widely available in females than males. Therefore, production of melanin is not much influenced by the internal triggers include hormonal stimulation but due to UVB rays.

The guinea pigs were exposed to UVB rays during the day for conditioning such as exposure to sunlight. UVB radiation is intended to stimulate the production of melanin by melanocytes in the basal stratum (germ). The treatment conducted for 2 weeks at night to avoid the sunlight because of hyperpigmentation treatment should avoid sunlight in order to achieve maximum results. Anti-hyperpigmentation effects in this study reflected by the average percentage of melanin area on each group at the end of the treatment. The histopathological results of the combination of turmeric and bitter melon extract can be seen in figure 1.

This study reinforces the previous study conducted by Sugiharto et al who proved the ability of curcumin to reduce melanin by 45.67% at 25 μg/ml in B16-F1 cell culture, which curcumin is the largest content in turmeric. This study also reinforces the findings of Tsai et al which proves that bitter melon extracts are significantly suppressed activity tyrosinase and melanin levels in B16-F10 melanocytes.

The average percentage of melanin area (figure 2) of combination treatment group 1 was 3.98%, combination group 2 was 2.52%, and combination group 3 was 0.89%. The higher concentration of the extract combination has better ability of melanin formation inhibitor which characterized by smaller percentage of melanin area. All combination treatment groups of turmeric and bitter melon extract showed smaller percentage of melanin area than DMSO group (p <0.05), it proved that the combination group revealed anti hyperpigmentation effect. Meanwhile, the combination group 1 had anti hyperpigmentation effect that comparable to the positive control (p>0.05), the combination group 2 and 3 showed average percentage of melanin area smaller than the positive
The anti hyperpigmentation effect result of ethanol extract of turmeric on guinea pig skin contained in the research report. The combination group 1 showed anti hyperpigmentation effect higher than 1000 µg/ml of turmeric extract (2.01%) (p <0.05). Meanwhile, the combination group-3 had the average percentage of melanin area which was significantly smaller than 500 µg/ml of turmeric extract.
(p <0.05), which meant that the combination group 3 has a better anti hyperpigmentation effect than 500 μg/ml of turmeric extract.

The anti-hyperpigmentation effect of ethanol extract of bitter melon leaves on guinea pig skin has been published in Cendekia Eksata Journal. The combination group 1 showed higher anti hyperpigmentation effect than 200, 400, and 600 μg/ml of bitter melon leaves extract group which reduced melanin by 2.01; 1.06; and 0.62 % respectively (p <0.05). The combination group 2 showed higher anti hyperpigmentation effect than 400 μg/ml of bitter melon leaves extract (0.62%) (p <0.05).

These findings were consistent with the results of Rizza et al (2012) that evaluated skin whitening effects of Mediterranean herbal extracts by in vitro and in vivo models. The single extract of caper buds, blood orange, rice grains, and olive leaf showed less intensive inhibiting activity than kojic acid and hydroquinone. Otherwise, when all the extracts were combined, it has the higher activity significantly than kojic acid and hydroquinone.

The possibility of antihyperpigmentation effect of turmeric and bitter melon leaves extract combination is due to compounds contained in the extract combination. The compounds are flavonoids, saponins, alkaloids, terpenoids, phenols, and tannins. The ethanol extract of bitter melon leaves contains phenolic compounds, polyphenols, tannins, saponins, alkaloids, vitamin C, gallic acid and catechin. The active compound suspected to play a role in decreasing the amount of melanin is curcumin through the antioxidant mechanism. Whereas the active compound suspected to play a role in decreasing the amount of melanin is gallic acid, salicylic acid, cinnamic acid, myricetin, quercetin, and lutein through the antioxidant activity, cell protection, and anti melanogenic activities.

CONCLUSIONS

The combination of turmeric and bitter melon leaves extract (group-2 and 3) showed better effect than the pharmaceutical cream.

RECOMMENDATIONS

This study proved that the combination of natural ingredients had a better effect than the pharmaceutical cream, in the hope of having fewer side effects, so it needs to be tested further on the toxicity test on animal’s skin, as well as clinical trial phases 1 and 2.

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