Topical application of purple cabbage (Brassica oleracea L. var. capitata f. rubra) ethanol cream extract of dermic collagen on male Wistar rats (Rattus norvegicus) exposed to ultraviolet B

Komang Jegek Triangga Apsari,¹ I Gusti Ayu Dewi Ratnayanti,² I Gusti Komang Nyoman Arijana,² I Wayan Sugiritama²

ABSTRACT

Background: Purple cabbage or also known as red cabbage (Brassica oleracea L. var. capitata f. rubra) contains polyphenols, especially anthocyanin (cyanidin-3-diglucoside-5-glucoside). Anthocyanin has a protective effect against radical oxidative superoxide (ROS). Thus it can prevent skin damage caused by ultraviolet (UV)-B radiation.

Aim: The research aims to prove the effectiveness of distribution of purple cabbage ethanol cream extract prevented the decrease of dermal collagens on Wistar rats (Rattus norvegicus) exposed to UV-B rays.

Method: This research was arranged with the randomize post-test only control group design. 30 rats were divided into 6 groups containing 5 rats each, there are control group (P1) without distribution of any cream, Treatment Group (P2) with a distribution of base cream (placebo), (P3) sunblock 33 SPF, (P4) 5%, (P5) 10 %, and (P6) 20 % smeared by purple cabbage ethanol cream extract. All groups exposed to UV-B with a total dose of 840 mJ/cm² for 4 weeks. Then a sampling rats skin was done for examining the level of collagen skin with Sirius red staining. The level of collagen was calculated by the percentage of the area of the pixel of collagen and was compared with the pixels of entire dermal tissues.

Result: The result demonstrated that the mean of collagens in The First Group/P1 (57.74%) and P2 (60.84%) decreased more significantly than P3 (82.17%), P4 (68.23%), P5 (76.93%), and P6 (84.54%) following UV-B exposure. The Post Hoc result showed that there was no significant difference in the level of collagen in the control group and placebo group and the sunblock group with the purple cabbage's ethanol cream extract group of 20% (p>0.05).

Conclusion: The distribution of purple cabbage ethanal cream extract prevented the decrease of dermal collagens on male Wistar rats skin exposed to UV-B.

Keywords: Anthocyanin, antioxidant, purple cabbage ethanol cream extract, dermal collagen, UV-B ray.

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INTRODUCTION

Indonesia is a tropical country which has a high temperature.¹ Living in a tropical country means more exposure to sunlight. Sunlight plays an essential role in human health, especially for bone development.²³ However, the exposure also has a negative impact on skin health. Chronic exposure to ultraviolet rays can cause damage to the structure and function of the skin thus accelerating the aging process of the skin; it is called Photaging.⁴⁵⁶ Ultra Violet (UV) radiation is divided into three types: UV-A, UV-B, UV-C.⁷ UV-A and UV-B play a role in photaging while UV-C is absorbed directly by the ozone layer in the atmosphere.⁷⁸ In the skin that undergoes photaging, it can show clinical features of the rough surface, nodule, smooth and rough wrinkles, yellowish, dry and telangiectasia.⁹¹⁰ The pathobiological effects of ultraviolet light (UV-A and UV-B) produce free radicals and cause DNA damage (Deoxyribonucleic Acid) as the source of formation of ROS (Reactive Oxygen Species).⁶¹¹ Increased of ROS as a result of free radicals because UV-B rays can cause increased lipid peroxidation.¹² This ROS compound also plays a role in collagen metabolism, because it can destroy the collagen Matrix Metalloproteinase (MMPs), resulting in skin collagen decreased.¹³¹⁴ Its decrement is caused by UV rays and is chained by two most responsible mechanisms AP-1 induction and lower regulation of type II TGF-β (Tumor Growth Factor Beta).¹³¹⁵¹⁶¹⁷ Antioxidants are factors that can reduce ROS, both topical and systemic.¹⁸ Based on the source of the acquisition there are two kinds of antioxidants, namely natural antioxidants and synthetic antioxidants (synthetic).¹⁹ The human body does not have antioxidant reserves in excess level, so if there is exposure to excessive radicals, then the body needs exogenous antioxidants.²⁰ The presence of concerns about the possible unknown side effects of synthetic antioxidants causes natural antioxidants to be a much-needed alternative.²⁰
Methods

Subject

Based on the calculation using Federer’s formula, the subject of the study used 30 rats of Wistar strain (Rattus norvegicus), male, 3-4 months old, weighing 150-200 grams, divided into six groups, each with five rats. The subject did not appear to be active or sick or die including exclusion criteria.

Design of Experiment

This research design used the randomized post test only control group design.

Subject Treatment

Subjects were divided into 6 (six) groups with each leveling to 5 rats. The control group exposed only UV-B rays (P1), Placebo Group (P2), Group Sunblock cream SPF 33 (P3), Group cream extract ethanol of purple cabbage 5% (P4), Group extract ethanol of purple cabbage 10% (P5), and Group cream extract ethanol of purple cabbage 20% (P6). UV-B light from Polish Ultraviolet broadband, Philips type UVB-311nm (pl-s 9w/01) lamp. Rats from all treatment groups were given UVB exposure three times a week for 4 weeks beginning with a dose of 50 mJ/cm² for 50 seconds in the first week, followed by 70 mJ/cm² for 70 s in the second week and 2 weeks later with 80 mJ/cm² for 80 seconds, so the total UVB received is 840 mJ/cm² for 4 weeks. The rats were left first for 24 hours after irradiation tended to exclude the effects of acute effects.

Application of Substances

All rats were shaved on their backs before being applied with the cream by their respective treatment groups. Each cream according to treatment was administered as much as 0.05mg/cm², applied two times daily about 20 minutes before irradiation (to give a topical absorption time into the skin) and 4 hours after irradiation (ROS formation started 4 hours after exposure). Topical application of the material remains on a day without irradiation.

Histopathological Observation

The rats were euthanasia first with anesthesia by inserting into a jar containing ether-cotton. Subsequent samples were taken from the back tissue of Wistar rats with 2 mm deep subcutaneous, 2 cm long and 2 cm wide using scalpel no.10 and anatomical scissors. Samples were then prepared for histopathologic examination with preparations using Sirius red staining. Collagen texture was shown in bright red on the picture of the preparations taken using the Optilab Pro camera (Miconos, Indonesia) and the Olympus Cx41 microscope with an objective magnification 400 times. The result of preparing photos is stored in JPEG format, then calculated by using Adobe Photoshop CS3 and Image J software.

Statistic Analysis

Data were processed using IBM SPSS 21.0. The analysis begins with a descriptive test to look at the characteristics of the data, test the normality of data using Shapiro-Wilk, homogeneity test using Levene's test. Comparative analysis between groups using One Way ANOVA. Post-Hoc analysis was performed after it was found that there were significant differences between the treatment group and normal, homogeneously distributed data then Post-Hoc test with LSD (Least Significant Difference-test) test. There was a significant difference in the group if (p <0.05).

Results

Treatment effect analysis was tested based on a percentage of collagen between groups after treatment. The result of significance analysis with One Way ANOVA test was shown in Table 1. The test showed that the value of F = 43.28 and p-value = 0.000. This means that there was a difference in the collagen in the six groups (p<0.05). A further test with the post-hoc test was Least Significant Difference-test (LSD). The test results were presented in Table 2.

The above test results showed that there was no significant difference in the level of collagen in the control group and placebo group and the sunblock
group with the cream extract ethanol extract group of 20%.

**Histological Appearance**
The histological preparation of Wistar rats’ skin with Sirius red staining appears as shown on the Figures 1.

**DISCUSSION**

**Characteristics of the Sample**
This study used Wistar rats (*Rattus norvegicus*) as experimental animals because they have the equation of organ structure with humans, other than that the short and not thick fur owned by Wistar rats facilitate research using skin texture as research samples. A group with the cream extract ethanol extract group of 20%.

**Collagen On Male Wistar Rats**
In this study showed that in the control and placebo groups after the post-hoc test, p > 0.05 was obtained, which means there is no difference in the level of collagen (Table 2). So the effect of prevention of the reduction of the level of pure collagen is due to the giving of purple cabbage ethanol extract and decreasing the level of collagen in all treatment groups resulting from UV-B. In the Control and
Placebo Groups showed damage to the composition and structure of collagen tissue, so the collagen fibers were not intact (fragmented) and did not absorb the red staining of Sirius red well (Figures 1).

The decrease in the level of collagen dermis caused by ultraviolet rays which can form free radicals that can activate mitogen-activated protein kinase pathways (MAPK) as collagenase producer (MMP-1) that can destroy collagen. The oxygen molecule (O2) present at the bottom of the epidermis is the main target of UV-B rays that enter the skin. UV rays that penetrate the skin can as a donor of an electron in an oxygen molecule that causes oxygen to become unstable, then become aggressive free radicals. In the dermal layer, ultraviolet B radiation causes collagen damage at a higher rate than naturally caused aging damage.6,9,10,11

Sunlight damages collagen fibers and causes abnormal accumulation or elastin accumulation. When this elastin accumulates, an enzyme called metalloproteinase will be produced in large quantities.38,11 Usually, this enzyme works to repair skin damaged by light by producing and forming collagen, but this process does not always work well, and some of these proteins actually destroy collagen, the result is the emergence of a collection of irregular collagen fibers known as solar scars.5,11 When the skin keep this imperfect repair process repeatedly, it would forms wrinkles.5,11 In photoaging skin, collagen fibers are disorganized.5,8 The results of this study are in accordance with the results of Gilchrest and Krutmann's research, which found that the skin that experienced photoaging decreased the number of precursor collagen type I and III.5,6 Similarly, the Moertolo study showed that UV-B exposure with a total dose of 840 ml/cm2 for 4 weeks may result in a decrease in the level of collagen in the skin of Wistar rats (Rattus norvegicus).37 It is known that UV-B causes more DNA cell damage. Damage that can be caused in the form of DNA lesions in cyclobutane pyrimidine dimer. Clinically the disorder is erythema or redness. The end result of glycation or advanced glycation end product (AGE) that accumulates in long-lived proteins such as extracellular matrix also turns out the function as a sensitizer for ultraviolet, which can damage dermal fibroblasts cells. Ultraviolet light is also shown to increase collagen degradation through activation of MMPs Ultraviolet rays can stimulate the synthesis of MMP-1 and MMP-3 through the release of TNF-α (Tumor Necrosis Factor Alpha) by keratinocytes and fibroblasts. UV-B directly affects DNA damage mainly in two large lesions of cyclobutane dimer and pyrimidine-pyrimidone photoproduct, which directly affects nucleic acid synthesis. Although nuclear DNA can repair itself, DNA damage is rarely improved entirely and has the potential to become malignant cells.38

In this study also showed that in the Treatment Group given 5%, 10% and 20% of purple cabbage ethanol extract can prevent the decrease in the level of dermal collagen caused by UV-B exposure (Figures 1). In this study also showed that in the Treatment Group given 5%, 10% and 20% of purple cabbage ethanol extract can prevent the decrease in the level of dermal collagen caused by UV-B exposure.39 A study by Moertolo showed that the presence of phenol compounds, one of the class of flavonoid (anthocyanin class of cyanidin-3-glucoside and peonidin-3-glucoside) in black rice which has antioxidant activity in preventing the decrease of collagen group treatment more quickly than control group.37 Similar research has also been conducted by Dianasari about giving cream of purple corn extract and Widiyowati research about green tea extract cream giving that anthocyanin content as an antioxidant can preventing collagen reduction in the faster treatment group than the control group.27,40

Several previous studies have shown that polyphenolic compounds contained in purple cabbage...
have essential effects on the body, such as protection against cardiovascular disease, diabetes mellitus, anti-inflammation, anticancer, and antioxidants. In accordance with some studies both in animals and humans that prove that polyphenol compounds can prevent the formation of free radicals and lipid peroxidation due to exposure to ultraviolet light. The mechanism of action of polyphenols in inhibiting damage caused by UV rays encompasses three effects, namely sunscreen effects, anti-inflammatory effects, and antioxidants. Most of the natural polyphenols are pigments, generally yellow, red or purple and absorb UV radiation. When administered topically, polyphenols will prevent the entry of radiation into the skin layer. Radiation that can be absorbed by polyphenols includes the entire UV-B spectrum and some UV-A and UV-C. Through the ability of this absorption natural polyphenols can act as a sunscreen. The strength of polyphenols to work as sunscreens can reduce inflammation, oxidative stress and DNA damage caused by UV radiation in the skin. In topical administration, the photoprotective ability of polyphenols is obtained through the sunscreen effect.

In this research, the level of collagen between the sunblock group and the group of ethanol extract of purple cabbage 20% after the post-hoc test was obtained p>0.05 (Tabel 2), which means no significant difference. This suggests that the effect is more vital to prevent a decrease in the level of collagen compared to the group given 5%, 10% cream, placebo and control groups which means the addition of cream doses can add cream effectiveness as strong as sunscreen as a prototype antioxidant that can provide protection from UV-B exposure. According to Irsyadah’s research, the use of sunscreen on some SPF (Sun Protection Factor) values can provide the protection from UV-B exposure. Application of cream extract of purple cabbage ethanol can be used to protect the skin from the effects of UV radiation that can cause both erythemas in acute effects and chronic effects that can cause premature skin aging.

The purple cabbage (Brassica oleracea L. var, capitata f. Rubra) also contains vitamin C, which is an antioxidant that serves to keep the cells, including skin cells. In addition, vitamin C also plays an active role in the production of collagen. Vitamin C works as a collagen protector of the skin by donating two electrons derived from the double bond between the second and third carbon so that ROS is not formed. In this process, vitamin C will oxidize to produce dehydroascorbic acid, which can then be reduced back to ascorbic acid with the help of the enzyme 4-hydroxyphenylpyruvate dioxygenase. In collagen biosynthesis, vitamin C has a role as a cofactor of prolyl hydroxylase enzyme and lysyl hydroxylase. In collagen found many prolines and lysine, hydroxylation both will stimulate the formation of new collagen. Similar research results by Soejanto, vitamin C content found in red pomegranate extract cream can prevent the decrease in the level of collagen.

**CONCLUSION**

Based on the result of this research, it can be concluded that 5%, 10% and 20% of the purple cabbage (Brassica oleracea L. var, capitata f. Rubra) ethanol cream extract can prevent the decrease of dermal collagen of male Wistar (Rattus norvegicus) dermis exposed to UV-B.

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