Antioxidant Activity of Haramounting Leaf Ethanol Extract (Rhodomyrtus tomentosa) In Preventing Heart Damage of Mice (Mus musculus L.) after exposure to Electronic Cigarette

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Abstract. Cigarette smoke is one factor that increases free radicals in the blood will affect liver health in the body. The purpose of this study is to find out the effect of giving leaves ethanol extract haramounting to histology of male mice liver exposed to electric cigarettes. The design of this study used Completely Randomized Design (CRD) consisting of + Group, - Group and three treatments of Haramounting Leaf Ethanol Extract (Rhodomyrtus tomentosa (Aiton) Hassk.) for 30 days at a dose of 100 mg/kg BW Body Weight, 200 mg/kg BW and 300 mg/kg BW for treatment use of five replications. Liver organ preparations were made using paraffin block method and Hematoxylin Erlich -Eosin (HE) staining. The results of the study show on histological observations The liver has a tendency that electric cigarette smoke can cause damage in liver cells. The results of this study indicate that ethanol extract leaves of Haramounting after exposure to electric cigarette smoke can effectively regenerate damaged liver cells.

1. Introduction
Epidemic of tobacco problems due to smoking is one of the biggest health threats the society currently faced by the world. Cigarettes an interesting phenomenon because besides its contribution as one of the health problems with a fairly high death rate, almost six million people per year with more composition out of five million deaths are the result of use direct cigarettes whereas more than six hundred thousand the remaining deaths are the result of non-smokers exposed to smokers [1]. Free radicals are a molecule which has unpaired electrons inside the outer orbitals are very reactive. Radical this tends to hold chain reactions that are if it occurs in the body it will get cause damage continuous and continuous. Human body have an endogenous defense system against free radical attacks mainly occur through events of normal cell metabolism and inflammation [2] Amount of free radicals can experience an increase due to factors stress, radiation, cigarette smoke and environmental pollution cause the body's defense system to exist inadequate, so the body needs it additional antioxidants from outside that can protect from free radical attacks.

Antioxidants are also used in food to control lipid oxidation. T-butyl hydroxy anisol (BHA) and di-butyl hydroxitoluen (BHT) compounds are used as food antioxidants, but the possibility of adverse
side effects is not used for therapeutic ingredients. Besides being able to protect the body from attacks by free radicals, natural antioxidants can also slow down the occurrence of chronic diseases caused by decreases in reactive oxygen species (ROS), especially hydroxyl radicals and superoxide radicals [3]. Chemicals used to prevent or slow down free radical damage is an antioxidant [4,5,6]. The endogenous antioxidant work to neutralize free radicals where in carrying out the process and also needed antioxidants from the outside (exogenous) in the form of vitamins and minerals has been obtained from food or supplements.

Haramonting (Rhodomyrtus tomentosa (Aiton) Hassk.) is a plant native to Southeast Asia and also spread in the territory of Indonesia. Haramonting can be used as an herbal remedy [7, 8], that the people of Kalimantan and South Thailand utilize this plant as an anti-diabetic medications, diarrhea, burns, and abdominal pain. Haramonting leaf extract contains methanol which can lower blood glucose in blood. Flavonoid content results in leaves and fruit haramonting may be free-radical scavengers (anti-oxidants) for human. Edible leaves contain steroids, terpenoids, alkaloids, phenols, flavonoids, and saponins [9]. According to the study [10] conducted in vivo and in vitro extract of leaf extract (Rhodomytrus tomentosa (Aiton) Hassk) has strong antioxidant activity. In previous studies into liver organs [11] against haramonting extracts on paracetamol [12] had an effect on the improvement in liver damage induced in paracetamol.

2. Methods
2.1 Research Design
This study uses a completely randomized design method that is divided in 5 treatments with 5 replications according to the Fereder formula, namely:

\[(t-1)(n-1) \geq 15\]

Description:
t = treatment group
n = repeat

Following this research treatment groupings can be seen in table 1

| Aquadest | Smoke Eletric Cigarette | Haramounting Exract 30 Days |
|----------|-------------------------|-----------------------------|
| K (-)    | √                       | 100 mg/kg BW                |
| K(+)     | √                       | 200 mg/kg BW                |
| P1       | √                       | 300 mg/kg BW                |
| P2       | √                       |                             |
| P3       | √                       |                             |

2.2. Preparation Sample for extracts
The extraction method uses the maceration process using ethanol solvent. Making extracts is done by soaking simplicia for 48 hours with the composition of 1000 grams of leaves is haramonting with solvent ethanol. Then it will be filtered using Whatmann paper and the evaporation process is carried out to separate the solvent with the extraction results.

2.3. Exposure to e-cigarette smoke and administration of extracts
The mice tested were given food and drink in an ad litbium, then given oral haramounting extract orally with doses of 100 mg / kgBW, 200 mg / kgBW and 300 mg/kgBW each distinguished durations of 30 days. First poured 8 drops of liquid on the 4 sides of each burning cotton morning. Then electric cigarette smoke was blown into the smoking chamber. Then prepared the equipment.
used in the presentation, namely smoking chamber and put the mouse in the cage per group containing five tail into the smoking chamber through the top of smoking chamber then closed again Repeated 4 times or according to the treatment then given extract of haramounting leaves after ± 6 hours of administration e-cigarette smoke and is done every day for 30 days.

2.4 Preparation for making histological preparations
Histological preparation of the liver using procedures [12] .First the liver that has been surgically removed is washed using 0.9% NaCl. After that, the organs are dissected and then fixed using a 10% formalin buffer. Then do dehydration with alcohol for 60 minutes. Then purified with xylol then infiltrated with paraffin in an oven with a temperature of 60-70 °C for 2 hour. Then the infiltrated organ is then planted in a small box containing paraffin. Next, cut organs transversely in the form of paraffin tape with using a microtome with a thickness of 4 μm and then affixed to the glass object which was smeared with glycerin which then deparafinated using xylol and then rehydrated with multilevel alcohol. Then cleaned with running water and stained with Hematoxylin-Eosin staining and finally covered with vaseline and glass cover.

2.5 Histopathological Analysis of the Liver
The liver was taken to make histopathological preparations using paraffin method. Histopathological studies were performed under a microscope with a magnification of 10x40. Liver histological preparations were observed under a light microscope in 5 different fields, with a magnification of 40x10 x. Each field counted 20 cells randomly so that in one preparation 100 liver cells were found. Then the mean score of histopatalogi liver change score was calculated in five fields of each - each mouse with the Scoring Histopathology Manja Roenigk model. The liver structure observed was normal hepatocytes and hepatocytes which suffered damage both necrosis [6], parenchymous degeneration and hydropic degeneration. Then the number of percentages that occur [13] are counted and calculated. The Assessment of Hepatocyte Damage Assessment Criteria for Spoiled Roenigk who have been modified can be seen in Table 2.

| Damage Rate            | Score |
|------------------------|-------|
| Normal                 | 1     |
| Parenkimatosa Degeneration | 2     |
| Hydropic Degeneration | 3     |
| Necrosis               | 4     |

2.6 Data Analysis
The data analysis used a complete randomized design (ANOVA) at 95% confidence level and, $\alpha = 0.5$ with bootstrap / duncan analysis. All data is analyzed using SPSS 23 program software.

3. Results and Discussions
The results will be discussed in two subsections, they are histopatology analysis and Hepatocyte Damage Rate.

3.1. Histopatology Analysis
Based on observations from histological liver that has been done on the liver of male mice after the exposure of electric cigarette smoke and given ethanol extract of Haramounting (Rhomodytrus tomentosa (Aiton) Hassk.) Can be seen in Figure 1.
Figure 1. Microscopic Hepatocytes Giving Haramounting Leaf Ethanol Extracts (*Rhodomytus tomentosa* (Aiton Hassk)) and Electric Cigarette Smoke Expansion with HE Coloring and 400x Enlargement (A) K· = Negative Controls (mice are given nothing but feed; (B) K + = Positive Control, Mice were exposed to electric cigarette smoke (C), (D), (E) P1 = Treatment of 1,2, and 3 mice were exposed to electric cigarette smoke and were given extracts of 100 mg /kg BW, 200 mg /kg BW and 300 mg /kg BW for 30 Day: (a) Central Veins, (b) Normal Hepatocytes, (c) Parenchymous Degeneration, (d) Hydrophic Degeneration and (e) Necrosis

In Figure 1 shows the results of the observation of the histology of the liver of male mice found cell changes. The percentage of normal hepatocytes in the K-group was more than the treatment. While the number of cells experiencing swelling, hydropic and necrosis is quite a lot. According to [9], the presence of hepatocyte swelling only occurs in the mitochondria and endoplasmic reticulum due to stimuli that produce oxidation. Hepatocyte damage in the form of swelling is reversible. Likewise with hydropic which is reversible. Hepatocytes that experience necrosis due to changes in the cell nucleus, this is natural because each cell will experience cell death. This is due to the presence of external factors, namely in the number of feeds and drinks in mice that are in accordance with the standard, the condition of the cage is less than ideal, the stress factor of mice, the influence of substances or other diseases, as well as other internal factors such as resistance and susceptibility of mice. In the treatment group, hepatocyte damage is likely due to the effect of treatment, which causes disruption of cell membrane permeability. The disruption of cell membrane permeability is due to the effect of giving ethanol extract of haramounting leaves and exposed to electric cigarette smoke where the content of haramounting leaf extract is high in flavonoids which interfere with free radicals. But in P1 treatment with haramounting leaf extract the amount of hepatocyte damage decreased compared to other treatments. This is because haramounting contains antioxidants that function as antioxidants to fight free radicals in the body including hepatocytes. This is in line with [10] Flavonoid compounds in
harmaounting extract can be free radical agents, reduce the amount of lipids, NO production and increase the activity of antioxidant enzymes in the body that will be exposed to organs.

3.2. Hepatocyte Damage Rate

The results of observations on the level of hepatocyte damage in each treatment on the transmission of electric cigarette smoke and haramounting leaf extraction (Rhodomyrtus tomentosa (Aiton). Hask) stastic analysis test. From the data the data experienced a decrease in cell death (necrosis) compared with exposed to electric cigarette smoke. Average Necrosis K - (5.4%), K + (43.6%), P1 (20.2%), P2 (31.2%) and P3 (32.4%). Can be seen in table 3.

| Treatment | Normal (%) | Parenkimosoa Degeneration (%) | Hidropyic Degeneration (%) | Necrosis (%) | Notation |
|-----------|------------|-------------------------------|---------------------------|--------------|----------|
| K-        | 94,4       | 0,2                           | 0,0                       | 5,4          | a        |
| K+        | 49,2       | 1,8                           | 5,4                       | 43,6         | b        |
| P1        | 51,6       | 17,4                          | 10,8                      | 20,2         | c        |
| P2        | 51,2       | 11,4                          | 6,2                       | 31,2         | c        |
| P3        | 46,0       | 15,2                          | 6,4                       | 32,4         | d        |

Information. K- = Negative Control (mice are given nothing but feed; K + = Positive Control, mice are exposed to electric cigarette smoke; P1 = Treatment 1, mice are exposed to electric cigarette smoke and extracts of 100mg/kg BW for 30 days; P2 = Treatment 2, mice were exposed to electric cigarette smoke and given 200mg/kg BW extract for 30 days; P3 = treatment 3, mice were exposed to electric cigarette smoke and given 300mg/kg BW extract for 30 days, different letters in different treatments showed significantly different.

Based on Table 3 it can be seen that there is a significant difference in the real level of hepatocyte damage between K- and K + and there is also a significant decrease in the number of presentate necaression in K+ with P1, P2 and P3 and Normal cell increase in K + with P1. This may be because the substances contained in electric cigarettes have not caused a harmful effect on the liver and also the content of ethanol extract of haramounting leaves which contain Falnovolid and Tanin which decreases the percentage of necrosis in the hepatocytes and also can protect the liver but can not restore the normal hepatocyte state. However, the percentage of P2 and P3 necrosis damage tends to increase with K- and K +. This is because the contents of the electronic cigarette are toxic to the liver and also the content of haramounting leaf extract which has flavonoid metabolite content.

This is in line with the study [14] states that the protection by haramounting leaf extract shows that haramounting leaf acetone extract has the ability to inhibit the formation of malondialdehyde (MDA) [15,16]; has a very strong reducing capacity; have capacity as a chealting agent. Hepatotoxicity due to chemical compounds is a potential complication that is almost always present in every chemical compound given because the liver is the center of metabolic disposition of all drugs and foreign substances that enter including electric cigarette smoke. Liver cell damage is rarely caused by a substance but often by metabolic toxic substances. The liver is an organ that is often damaged so that it can die through the path of apoptosis [17,18,19] or necrosis [20]. Metabolism of various compounds mainly occurs in the liver, resulting in very large organ damage. As a result the metabolic process does not run normally, it will cause various diseases. The cells contained in the liver will be deposited so they will change [21]

According to [20], hepatocytes are the type of cells that make up most of the liver. Hepatocytes are responsible for the central role of the liver in the metabolic process. These cells are located between
sinusoids filled with blood and bile ducts. According [22], if the liver cell is damaged due to various factors, a series of morphological changes will occur in the liver cells. These changes can be sublethal, degenerative or lethal in the form of necrotic.

One of the changes induced by free radicals is the change in the properties of cell membranes and cytoplasmic membranes in cell elements such as mitochondria and lysosomes caused by fat peroxide. After damaging the cell membrane, toxic effects that can also reach the nucleus and damage it, which causes the cell structure to become abnormal and eventually lead to necrosis [22]. Free radicals from the environment due to exposure to cigarette as occurs in passive smokers, causing antioxidants endogenous are no longer able to protect the body from oxidants, resulting in increased free radicals that trigger oxidative stress in cells [23, 24, 26].

The results of observations made on microscopic images of the liver given exposure [25] to electronic cigarette smoke and given hamamounting leaf extract (Rhodomyrtus tomentosa (Aiton) Hassk.) Found that there were normal hepatocytes and hepatocytes that experienced changes in parenchymous degeneration, phidrophic degeneration and necrosis.

4. Conclusions

The real level of protection effect on the liver after exposure to cigarette smoke and protection of hepatocyte damage occurred in the administration of ethanol hamamounting extract (Rhomodytrus tomentosa (Aiton) Hassk.) Significantly (p<0.05) at a dose of 100mg/kg BW even though it did not return the liver normal.

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