The Hepatoprotector Effect of Uncaria gambir Roxb Extract in Wistar Rats Induced by Paracetamol

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Abstract

Background: Drug induced liver injury (DILI) is known as the damage of liver cells due to chronic administrations of drug. The chronic administration of paracetamol could be trigger the damage of liver cells. The hepatoprotector agents are still limited worldwide. Gambier (Uncaria gambir Roxb) is an Indonesia traditional medicine which have many benefits as antioxidant, anti septic, anti diarrhoea, etc. that commonly used in society.

Method: The present study was conducted to investigate the hepatoprotector effect of gambier in wistar rats induced by paracetamol. The wistar rats were divided into seven groups and received the treatment orally for 12 days. Group I (aquadest), II (curcuma, 400 mg/kgBW), III (gambier, 26 mg/200gr), IV (gambier, 53 mg/200gr), V (gambier, 106 mg/200gr), VI (gambier, 212 mg/200gr) and VII (gambier, 424 mg/200gr). Termination, blood and liver organ collection were done after all group induced by paracetamol for two days. Histopathology changes of liver were examined using Hematoxycilline (HE) staining. AST and ALT levels were analyzed.

Results: There were significant differentiation of AST levels among the groups, especially between group I and group IV and between group II and group IV. The ALT levels were statistically significant between group II and group V using Mann-Whitney test (p<0,05). In histopathology examination, there were significant differentiation between group I with another group, not only group II but also group III-VII (p<0,05). In the treatment group, group III and IV had been showed the improvement of liver cells damage than group I by using One-way Anova, post hoc Bonferroni (p<0,05).

Conclusion: Uncaria gambir Roxb has hepatoprotector activity start at dose 53 mg/200grBW in rats. The hepatoprotector activity was not superior than curcuma.

Keywords: hepatoprotector, Uncaria gambir Roxb, AST, ALT, histopathology

1. INTRODUCTION

The incidences of liver injury worldwide and in Indonesia are steadily increasing over the years. There were two billion people suffering of Hepatitis B worldwide, 170 people get hepatitis C and 1.5 million were die due to hepatitis every year 1. In Indonesia, there were 38,2% cases of hepatitis in the hospital located in Tangerang2. There are several factors which induce liver injury such as viral infection, alcohol and chronic administration of drugs. 3 Drug induced liver injury (DILI) are commonly caused by rifampisin, isoniazid, ceftriaxone and analgetic drug, paracetamol. Previous study showed 35% of 100 patients underwent DILI. 4 Paracetamol has been already known as the popular analgetic drug due to wide therapeutic index and the potency to treat mild-moderate pain in several diseases; nevertheless the hepatotoxic effect are increasing in chronic and large doses administration of paracetamol. 5 The availability of hepatoprotector drugs to prevent DILI was still limited to Curcuma6. 

In Indonesia, especially North Sumatera, gambir (Uncaria gambir Roxb) is the traditional medicine widely used for diarrhoea, dysentery, gargle for throat sore etc. Prior studies analyzed the effect of antihyperlipidemia and anti atherosclerosis of this traditional medicine 7. The
antioxidant activity usually induced by catechin in gambir. The usage of gambir is common in North Sumatera but there were limited data showed the efficacy of gambir as hepatoprotector⁶⁹.

2. MATERIALS AND METHODS

This research is experimental study with post-test only design. The study was conducted during March to October 2019 and has been approved by Health Research Ethics Committee Polytechnic Health Ministry Of Health Medan No.337/KEPK Poltekkes Kemenkes Medan/2019

Materials

Our study used gambier which is an herbal medicine contain catechins as antioxidant. We used gambier (Uncaria gambir Roxb) in powder form, product by Toyo Brother®, Batch No: 0133517, branded name “Sari Uncaria). The composition of “Sari Uncaria” were consisted of water (3,22%), tannin (21,48%), catechine (81,37%). The composition analyze process were done in Nature Organic Chemistry Lab, Chemistry Dept, Institut Teknologi Bandung ¹⁰. We also used paracetamol tablet (Dexa Medica), curcuma tablet (Dexa Medica), aquadest and rats pellets.

Animals

Male healthy Wistar rats (Rattus norvegicus) with fulfilled inclusion criteria in this study,i.e,150-200 g BW, age 2-3 months, were obtained from animal house of Universitas Sumatera Utara. The aggressive and sick/died rats were excluded from this study.

Experimental design

The animals were divided randomly into seven groups of eight rats each and treated as follows:

1. Group I (NC): negative control rats (standard pellets and water ad libitum) for 14 days
2. Group II (PC): positive control, rats were administered with 400 mg/kg of curcuma, standard pellets, and water ad libitum for 14 days.
3. Group III (P1): Rats were administered with 26mg/200grBB of gambier, standard pellets, and water ad libitum for 14 days.
4. Group IV (P2): Rats were administered with 53 mg/200grBB of gambier, standard pellets, and water ad libitum for 14 days.
5. Group V (P3): Rats were administered with 106mg/200grBB of gambier, standard pellets, and water ad libitum for 14 days.
6. Group VI (P4): Rats were administered with 212 mg/200grBB of gambier, standard pellets, and water ad libitum for 14 days.
7. Group VII (P5): Rats were administered with 424 mg/200grBB of gambier, standard pellets, and water ad libitum for 14 days.

All the groups were administered paracetamol with dose 1 gr/kgBW in day 13-14. Animal termination and extraction of blood and liver samples were done in Pharmacology Lab, Dept.of Pharmacology, Universitas Sumatera Utara. The histopathology of liver were examined by using Hematoxylin-Eosin (HE) Staining and ALT/AST were analyzed by using the photometric method. The histopathology of liver was analyzed using scoring HE index. Scoring were determined based on microscopis examination. The liver cells were divided into three category: steatosis cells, ballooning cells and inflammation cells.

Data were analysed with Kruskall Wallis, Mann Whitney, One-Way Anovna and post hoc Bonferroni test using IBM SPSS Statistic 21.

3. RESULTS

3.1 Differentiation of AST values

In our study, there was differentiation of AST values among the groups (Kruskal-wallis test). Data was showed in table 1:

| No. | Groups | Median (Min-Max) | p    |
|-----|--------|-----------------|------|
| 1.  | NC     | 137,2 (65-218)  | 0,39⁴ |
| 2.  | PC     | 91,0 (70-291)   |      |
| 3.  | P1     | 96,2 (47-146)   | 0,016¹,⁴ |
| 4.  | P2     | 57,4 (50-66)    | 0,009⁴,²,⁴|
| 5.  | P3     | 122,6 (110-138) |      |
| 6.  | P4     | 118,2 (89-138)  |      |
| 7.  | P5     | 109,2 (98-130)  |      |

⁴Kruskal-wallis test  ⁵Mann-whitney

Based on table 1, there was a significant differentiation of AST values between NC group and P1 group (Mann-whitney test, p<0.05). The AST values were lowest in P2 group. This data showed, the administration of gambir (53 mg/200 grBW) had an effect to protect the liver from cell's destruction due to paracetamol induction. Our study also showed a significant differentiation of AST values between NC group and P2 group (p<0.05). A lower AST of P2 group compared to PC group indicated the effect of gambir (53 mg/200grBBW) was stronger to protect the liver cells than curcuma (p<0.05).

3.2 Differentiation of ALT values

There was a significant differentiation of ALT values among the groups (p<0.05). Data was showed in table 2 below:

| No. | Groups | Median (Min-Max) | p    |
|-----|--------|-----------------|------|
| 1.  | NC     | 84(60-109)      | 0,026⁴ |
| 2.  | PC     | 81(69-132)      |      |
| 3.  | P1     | 102 (93-283)    | 0,047²,⁵  |
| 4.  | P2     | 107 (67-138)    |      |
| 5.  | P3     | 60 (54-85)      |      |
| 6.  | P4     | 71 (59-78)      |      |
| 7.  | P5     | 67(47-109)      |      |

⁴Kruskal-wallis test  ⁵Mann-whitney
Based on Table 2, there was no significant differentiation of ALT values in NC group compared to PC and gambir group in all doses, but we found a significant differentiation of ALT values between PC group and P3 group (106 mg/200grBW) \((p<0.05)\). Our data showed there was no significant increasing level of ALT for each groups after induced by paracetamol.

### 3.3 Histopathology of Liver Cells

We analyzed the histopathology of liver cells by Haematoxicilin-Eosin Scoring Index system. Scoring was determined by microscopic examination in pada 10 lapangan pandang dengan menggunakan lensa perbesaran 10 x 40. Scoring was counted depend on mean total of destroying liver cells. The classification of destroying liver cell consisted of three category, which were steatosis cell (score 0-3), ballooning cell (score 0-3) and inflammation cell (score 0-3). The final scoring of destroying liver cells were counted based on the mean of steatosis, ballooning and inflammation cells score. The lower the score, indicated less destroying liver cells and higher the score showed the contrary result. The figure 1 showed the scoring of liver cells destruction among the group:

![Scoring of liver cells destruction](image)

**Figure 1: Scoring of liver cells destruction. Values are Mean±SD, \(*\)\(^{**}\)\(^{***}\)\(^{p<0.05}\) vs negative control**

Based on figure 1, the highest scoring was NC group while PC, P1-P5 groups had lower score. This result indicated PC, P1-P5 group had effect to protect the liver (hepatoprotector) which induced by paracetamol. There was a significant differentiation of scoring among the groups \((p<0.05,\) One-Way Anova\). We did the post-hoc test Bonferroni to analyze which groups had a significant differentiation scoring. Statistical analysis showed there was significant differentiation of scoring between NC groups with PC, P1-P5 groups \((p<0.05)\) but there was no differentiation of scoring between PC and gambir (P1-P5) groups \((p>0.05)\). In gambir groups, P1 (26 mg/kgBW) had showed the significant effect as hepatoprotector than NC, followed by P2-P5 \((p<0.05)\). Nevertheless, data showed gambir (P1-P5) were not superior than NC group (curcuma). In histopathology liver cells examination by Haematotxicillin staining, we found all of steatosis, ballooning and inflammation cells in NC group (figure 2).

![Histopathology liver cells in NC group](image)

**Figure 2: Histopathology liver cells in NC group. (Microscopic 400x), H=hepatosit Si=sinusoid B=Ballooning S=Steatosis Arrow= limfosit (inflammation cells)**
In PC group, we also found steatosis, ballooning and inflammation cells (figure 3) while in P1 and P2 group, we only found steatosis cells, there were no inflammation and ballooning cells (figure 4).

Figure 3: Histopatology liver cells in PC group. (microscopic 400x), H= hepatosit  Si=sinusoid   B=Balloning   S=Steatosis  Arrow= limfosit (inflammation cells)

Figure 4: Histopatology liver cells in P1, P2 group. (Microscopic, 400x)  H=hepatosit,  S=steatosis,  Si=sinusoid

4. DISCUSSION
In testing the efficacy of Uncaria gambir as a hepatoprotector, liver function was destroyed by induction of paracetamol. The mechanism of liver damage by paracetamol is the formation of a toxic metabolite called N-acetyl-p-benzoquinoneimine (NAPQI). This NAPQI compound has hepatotoxic effect on the liver. In normal conditions, NAPQI can be bound by glutathione in the body into mercapturate compounds so that it is non-toxic, but in high doses, NAPQI can deplete glutathione reserves in hepatocytes and bind to liver’s mitochondria so that oxidative stress can occur in the liver. Continuous oxidative stress can cause injury, apoptosis or necrosis in hepatocytes.11-12

In this study, we analyzed the effect of Uncaria gambir on the activity of AST and ALT with different doses. The result showed there was a decrease in the value of AST in gambir administration of 53mg / 200grBW dose compared to the negative control group and positive control group that received curcuma while we increased the dose of Uncaria gambir, there was no significant decrease in AST value. This
study indicated that at higher doses of Uncaria gambir, the hepatoprotector activity was reduced.

There was no significant difference for the ALT value among the groups although there was a significant difference of ALT values between P2 and P3 group. The AST measurement is the most sensitive assay for assessing liver cell injury, whereas an increase in ALT is sensitive for heart failure. AST enzymes are more abundant in the mitochondria compared to cytoplasm, whereas ALT enzymes are more abundant in the cytoplasm, in further or chronic damage, the mitochondrial membrane emits more AST.

It is commonly recognized that DILI is mediated by two chief mechanisms: intrinsic and idiosyncratic hepatotoxicity. Intrinsic hepatotoxicity is caused by some drugs include paracetamol which cause liver damage directly by its metabolite. Paracetamol can induced acute liver failure (ALF), in morphological features, we can see necrosis with little or no inflammation, ballooning degeneration of hepatocytes which is a form of liver parenchymal cell death. Necrosis can be accompanied by steatosis. In our study, we found all three morphological features, which are ballooning, inflammation and steatosis cells in negative control group, it proved the paracetamol induction caused acute liver failure. In positive control group and treatment group, we also found all of these cells in less amount. Surprisingly, we didn’t find the inflammation and ballooning cells in P1 (26 mg/200grBW) and P2 (53 mg/200grBW). This present study showed gambier has an effect to protect the liver cells by induction of paracetamol.

The decrease in liver damage in rats given Uncaria gambir occurs because it has catechins as antioxidants. Catechins are able to neutralize free radicals, and also able to increase other antioxidants in the body such as Glutation S-Transferase. This antioxidant is capable to protect liver cells from free radicals produced during the metabolic process of paracetamol into NAPQI. NAPQI is a toxic metabolite which can be detoxified by gluthatione S-transferase to a mercapturate compound but in a toxic dose NAPQI will use up the gluthionate S-transferase reserve in hepatocytes. Catechin is also known to have antioxidant effects by scavenging superoxide and pro-oxidant effects through cytochrome P450 2E1 (CYP 2E1)9. CYP2E1 is an enzyme that plays a role in the bioactivation of paracetamol. CYP2E1 participates in the formation of ROS such as superoxide and hydrogen peroxide. ROS can mediate the toxic effects of xenobiotics. Paracetamol knockout mice were found to be less toxic than normal mice. Even though catechin also has a pro-oxidant effect, but catechins in gambier are a polyphenolic component that is associated with various health benefits, both in vitro, in vivo and kinis. The characteristics of this therapy are related to the effect of catechin as a high antioxidant and the effect of free radical scavenging.

Our study is relevant to previous studies which showed that the administration of gambier leaves extract was able to reduce CCL4 induced liver fibrosis in rats. This is also in line with other studies of gambier extract using Carbon tetrachloride (CCL4) as an inducer of liver cell damage, showed that gambier has a hepatoprotector effect by significantly reduce AST and ALT activity, as well as being able to reduce the body weight ratio of mice. In our study, we found the effective dose of gambier is 53 mg/200 grBW as hepatoprotector since the administration of 53 mg/200 gr BW could lower AST and protect the liver cells by only steatosis cells appeared in histopathology examination. This present study showed no superior than curcuma as hepatoprotector. Meanwhile, there was a significant differentiation of AST values, but there was no significant differentiation in histopathology examination. Previous studies showed curcuma already has an hepatoprotector effect.

5. CONCLUSION

Uncaria gambir Roxb have hepatoprotector activity at dose 53 mg/200grBW. The hepatoprotector activity was not superior than curcuma.

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