BLOOD Glucose LEVELS AND THE MICROSCOPIC STRUCTURE OF KIDNEY WISTAR RAT DIABETES MELLITUS UNDER THE EFFECT OF Lawsonia inermis (Linn.) LEAVES ETHANOLIC EXTRACT

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ABSTRACT

Objective: Lawsonia inermis (Linn.) leaves are one of the alternative medicines to treat diabetes mellitus in Indonesia. We investigated the blood glucose level (BGL) of the L. inermis (Linn.) leaves ethanolic extract (LLEE) leaves and evaluated the histopathological alterations in diabetic rats. Methods: This study was an experimental study with posttest-only control group design. Alloxan (120 mg/kg, intraperitoneally)-induced diabetic rats. 35 of Wistar rats (Rattus norvegicus) were divided randomly into five groups, i.e. K: Normal control, P1: Diabetic control, and P2, P3, and P4 (200 mg/kg body weight [BW], 400 mg/kg BW, and 600 mg/kg BW of LLEE, orally) for 28 days. At the end of the treatment, the rats were sacrificed to obtain the kidney for histopathological evaluation using hematoxylin and eosin technique. BGLs were conducted using a glucose meter (GlucoDR). Results: One-way ANOVA test showed that dose 400 mg/kg BW of the LLEE was related to BGL of alloxan-induced diabetic rats (p=0.000). The histopathological of kidney showed glomerular inflammation (GI), epithelial membrane lining degeneration, vascular congestion, and interstitial tubule hemorrhage at diabetic control (P1). Meanwhile, treated with 600 mg/kg BW of LLEE (P4) showed increase cellular regeneration as normal architecture of the kidney.

Conclusion: The LLEE at dose 400 mg/kg BW effective decreased BGL and was able to restore the kidney destruction of alloxan-induced diabetic rats at dose 600 mg/kg BW.

Keywords: Lawsonia inermis (Linn.), Ethanolic extract, Blood glucose levels, Kidney, Diabetic rats.

INTRODUCTION

The treatment by utilizing natural resources such as plants has become a tradition of people around the world. Utilization of these natural resources was done by taking the extracts of plants and made as a medicine. The alternative treatment that uses herbs extract of plants nowadays was also become an option instead of modern medicine. The traditional medicines have more benefit, i.e. easy to find and cheap for the price. The traditional medicine from the plant that is less toxic and less severe side effect than chemical medicine. Since over the years, traditional medicine and traditional medical treatment have been existing in Indonesia [2]. One of the plants in Indonesia that is often used as a medicine is Henna/Lawsonia inermis (Linn.)/LL treated diabetes mellitus (DM) [3]. Hyperglycemia is one of the symptoms of DM. Based on the data that International Diabetes Federation, estimated, 415 million people can be at risk become DM in the world. The number of DM patient keep increasing along last three decades and estimated in 2040 will be 642 million [4]. The destruction of the body such as pancreas, heart, kidney, and eyes can happen on the DM patient. This complication can cause the death [5]. The increasing of reactive oxygen species (ROS) has the role in the pathogenesis of the hyperglycemia complication should be treated with antioxidant therapy [6]. The LL contained flavonoid which was included in the largest phenol group in the perfiee tissue, also controlled the activity and or the expression of rate-limiting enzymes which involved in the carbohydrate metabolism [3]. When flavonoid was given in the routine ways for the long period, it could reduce the level of plasma glucose to 60% [7]. Flavonoids might be reducing glucose level by a mechanism independent from insulin secretion, for example, by the inhibition of endogenous glucose production or by the inhibition of intestinal glucose absorption [8]. The aim of this study was to obtained the effect of L. inermis (Linn.) leaves ethanolic extract (LLEE) with various doses on blood glucose levels (BGLs) and the histopathological structure of rat kidney with alloxan induced.

METHODS

This study could be conducted after obtaining the approval from Faculty of Mathematics and Science ethic commission in University of Sumatera Utara (USU) with no of EC: 116/KEPH-FMIPA/2017. The study was conducted in Integrated Laboratory, Faculty of Medicine, Chemistry Laboratory of Faculty of Mathematics and Natural Sciences (FMIPA), and Biology Laboratory of FMIPA, USU.

Ethanol extraction of LL was done with maceration method that used ethanol solvent with ratio 1:10 (v/v). The dried LL was dissolved by ethanol solvent with ratio 1:10 (v/v). The dried LL was dissolved by ethanol solvent with ratio 1:10 (v/v). 

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if they died during the study. After adaptation for 1 week, Wistar rats were injected by alloxan at dose 120 mg/kg BW in the 0.1 M citric buffer intraperitoneally [11]. The treatments were given as follows [8,12]:
1. Group K without any treatment.
2. Group P1 alloxan positive control 120 mg/kg BW.
3. Group P2 alloxan 120 mg/kg BW and LLEE 200 mg/kg BW orally.
4. Group P3 alloxan 120 mg/kg BW and LLEE 400 mg/kg BW orally.
5. Group P4 alloxan 120 mg/kg BW and LLEE 600 mg/kg BW orally.

Treatment on the P2–P4 group was performed for 28 days starting from the BGLs >250 mg/dl [13]. BGLs were the measurement by the glucose meter (GlucoDr®) using the glucose test strips. The blood specimen to measure the BGLs was taken from rats’ tail [14]. The rats were sacrificed with light chloroform anesthesia on the 29 days. The result of sectioning tissue from kidney rats was the sampling of this study.

**Surgery of rats**
The surgical procedure performed through preparation, surgery, and sanitation by the routine procedure at Laboratory of Biologi, FMIPA, USU. The stages of rat surgery in this study were killed by neck dislocation. The rats put on the surgery table by using pin needle. The surgical lower abdomen was started by using scissors, the kidney was taken and cleansed from the fat of kidney.

**Preparation stages**
Prepared organ pots that have been labeled according to the number of rats treatment. Organ pots were filled with 10% buffer formalin to store the organs. Prepared insulin syringe 1 ml. The syringe has been labeled and rinsed with heparin sodium as an anticoagulant to hold the blood. Surgical equipment for this study as follows: surgical scissors, tweezer, pin needle, petri dish, watch glass, surgical table, pins, and glass beaker.

**The stage of surgical**
Rats were killed by neck dislocation after light chloroform anesthesia. Rats were positioned on the surgical board using pins or pin needles. The surgery started in rat stomach using surgical scissors. The kidney organ was removed, washed using 0.9% NaCl, cleaned, and put in the pot containing of 10% formalin buffer.

**Sanitation stage**
Sanitation stage was done by inserting the rest of rat organ that was unused into a plastic bag which will be destroyed. The rest of the surgical work area was cleaned and the surgical equipment was washed.

**Preparation histology slides of kidney rats**
The preparation histology slides of kidney rats were done in at the Pathology Anatomic Laboratory of Faculty of Medicine, USU, with routine histology method. The stages of preparation are as follows.

**Fixation**
The kidney tissue of Wistar rats washed to remove the blood. The cutting of tissue was no more than 1 cm x 1 cm. The tissue fixation was immersed in 10% formalin for 24 h.

**Dehydration**
The tissue was immersed until the entire tissue was submerged with alcohol. Dehydration performed using alcohols from low to high concentrations (70%, 80%, 96%, or absolute alcohol) for at least 12 h.

**Clearing**
The tissue was inserted into xylol 1 solution for 30 min, then inserted into the xylol 2 solution for 30 min. The process was to be the clearing the tissue. With this process, the tissue will be clear and transparent.

**Embedding**
The three containers containing paraffin were prepared in the oven. The tissue was immersed in an existing oven paraffin container 1 for 1 h. Then, insert the tissue into container 2 for 1 h. Next, insert the tissue to container 3 for 1 h.

**Blocking**
The pieces of L-shaped iron (Leuckhart) were prepared for blocking. Two pieces of iron were arranged on a sheet of metal to form a space like a cube. The tissue which too is seen was put in the bottom of the cube to be flat. Paraffin was poured into a cube-shaped piece of iron, waited for up to 12 h until the paraffin freezes.

**Sectioning**
Paraffin blocks containing tissue were cut by a microtome. The piece of slices was not folded by microtome stomped so that the slide well formed. Created multiples slide. The slide is inserted into the tube temperature of 45–50°C for deparaffinization. Then, the glass object was smeared with albumin. Gradually, paste the specimen into the glass object. Let stand for 12 h. Then, preparation was immersed with xylol 1 solution for 5 min to attract paraffin. Immerse with xylol II solution for 5 min. Next, rehydrate with alcohol from high to low concentration so the tissue contained water can be stained. Dip the glass of objects which containing the tissue into absolute alcohol 2 times for 2 min. Then, dip 90% alcohol as much as 2 times for 2 min in each. Continue on 80% alcohol for 2 times for 2 min in each. Last, alcohol concentration was 70% as much as 2 times for 2 min in each.

**Staining**
Object glass that contains tissue was inserted into hematoxylin solution for 5–10 min, rinsed with running water for 2–3 minutes, put into eosin solution, then rinsed over with running water. After that, dehydrated from low concentration alcohol to the high concentration alcohol. In every 2 minutes in the order of 70% alcohol, 70% alcohol, 90% alcohol, 80% alcohol, 80% alcohol, 90% alcohol, and lastly two times into absolute alcohol. Then inserted the object glass contains tissue into xylol solution, respectively for 2 minutes: xylol I and xylol II. Giving hematoxylin which was in the blue dye would stain the cytoplasm, nucleus, calcium due to the component that has alkaline. Meanwhile, eosin that was red dye would stain cytoplasm and connect the cells.

**Gluing (mounting)**
Dripped one drop of Canada Balsam on glass deck, close the glass objects which have the tissue, press repeatedly to make no bubble.

**Labeling**
Slides were given labeled.

**Data analysis**
The effect of LLEE to BGLs was obtained by processed and analyzed with as paired t-test using SPSS program version 16. The reading of preparation histopathology, after preparation, is ready to use observed with as paired t-test using SPSS program version 16. The effect of LLEE to BGLs was obtained by processed and analyzed with as paired t-test using SPSS program version 16. The reading of preparation histopathology, after preparation, is ready to use observed under a microscope with x400 magnification at the Pathology Anatomic Laboratory of Faculty of Medicine, USU. The histopathological of kidney cell was analyzed by qualitatively descriptive such as the degeneration of cells, congesting or widening of blood vessels, inflammation, necrosis cells, and thickening of membrane basal glomerulus. On the glomerulus (G), capsule Bowman (CB), proximal tubule (PT), and distal tubule (DT) analyzed of kidney appearance in each P1, P2, P3, and P4 groups compared.
to the normal control (K). The records of the histological appearance were obtained by photomicrography using digital photomicrography microscope.

**RESULTS**

The BGLs of Wistar rats before and after alloxan induced at P1-P4 groups had shown at Table 1.

At Table 1 can be seen, the BGLs before and after induced alloxan 120 mg/kg BW with a paired t-test on Groups P1-P4 showed the correlation (p<0.05).

Effect of LLEE on BGLs Wistar rat on the day 28th had shown at Table 2.

On Table 2, in the Groups P1, P3, and P4 of t-paired test analyzed showed decreasing BGLs between after alloxan induced compared to LLEE treated. The group receiving a dose of 400 mg/kg BW had lower decrease (significantly p=0.003).

The microscopic of the kidney tissue Wistar rats Group K can be seen in Fig. 1. The histopathological appearance of kidney in this group has not affected the hyperglycemia and also was not obtained the treatment LLEE.

The histopathology appearance of kidney on this Group (K) can be seen the normal structure of glomerulus (G) surrounded by the intact CB, DT, and PT without any inflammatory alterations.

In this study, the histopathological kidney hyperglycemia causes alloxan induced shows in Fig. 2.

In Fig. 2, microscopic groups of hyperglycemia rats after induced alloxan, 120 mg/kg BW shows the glomerular inflammation (GI).

Table 2: The BGLs of rats after induced alloxan and after LLEE treated on the day 28th

| Groups | BGLs after alloxan induced (mg/dl) | BGLs after LLEE treated (mg/dl) | p* |
|--------|-----------------------------------|-------------------------------|----|
| K      | 160±78.03                         | 112±25.05                     | 0.394 |
| P1     | 350±110.92                        | 309±71.14                     | 0.687 |
| P2     | 440±121.22                        | 450±93.66                     | 0.884 |
| P3     | 487±101.79                        | 236±66.57                     | 0.003** |
| P4     | 426±149.46                        | 256±41.25                     | 0.052** |

*Paired t-test <0.05, **significance. BGLs: Blood glucose levels, BW: Body weight, LLEE: Lawsonia inermis (Linn.) leaves ethanolic extract

In this study, the effect of natural antioxidants LLEE on the kidney tissue of Wistar rat alloxan-induced can be seen in Figs. 3-5.

Fig. 5 The histopathological appearance of kidney in Group P4 showed increased cellular regeneration as a normal cellular architecture. Giving LLE 200 mg/kg BW on the diabetic Wistar rats (P2) in Fig. 3 showed that there were GI, VC, and interstitial tubule hemorrhage. Microscopically, the damaging of cells that occurred in Wistar rats Group P2 almost same with the damaging of cell in the Group P1. In this study, LLEE with dose 200 mg/kg BW could not prevent or fix the damage of kidney cell on the hyperglycemia induced by alloxan.

Descriptive microscopic of the P3 group (LLEE 400 mg/kg BW) could be seen in Fig. 4. Histopathology of cell kidney showed that it was still seen glomerular (G) inflammation, but VC was less than Group P2. Compared with the image of histopathology Group P4 (treated with LLEE at 600 mg/kg BW) that was shown in Fig. 5. The microscopic of cell in this group showed that normal cell of kidney with increased cellular regeneration.

**DISCUSSION**

In this study, BGLs before and after induced alloxan 120 mg/kg BW with paired t-test on all group (P1–P4) showed the correlation (p<0.05). Alloxan compounds were one of the most toxic diabetogenic substances, especially to pancreatic beta cells, and when given to experimental animals such as rats can cause diabetic to the rats [11]. The alloxan toxicity mechanism begun with the entry of alloxan into the pancreatic β cells and when given to experimental animals such as rats can cause diabetic to the rats [11]. The alloxan toxicity mechanism begun with the entry of alloxan into the pancreatic β cells and the retrieval rate will determine the alloxan diabetogenic properties.

In this study, there was an association between the extracted ethanol of L. inermis (Linn.) leaves with the decreasing of BGLs in the group of an experimental rat with p<0.05 (Table 2). L. inermis Linn. is an one of the 40 types of plants used as medicines and listed in the list of drugs “Ebers Papyrus” written in 1550 BC. Based on phytochemical screening, the leaves of L. inermis (Linn.) contained phenolic compounds glycosylation proteins (coumarins, flavonoids, alkaloids, and tannins derivatives) [15]. A study by Fitrianda et al showed that the effect of flavonoids-containing herbs could decrease BGLs alloxan-induced diabetic Wistar rats [16]. Flavonoids can inhibit polyol pathways through inhibition of aldose reductase enzyme. The alkaloid is an...
Antioxidant that can reduce free radicals due to giving alloxan. Tannin has hypoglycemic activity by increasing glycogenesis process [15].

The image of kidney histopathology on this Group (K) can be seen normal structure, but after alloxan induced at 120 mg/kg BW shows the GI epithelial membrane lining degeneration, VC, and interstitial tubule hemorrhage. The kidney is the organ which has the role to keep the balancing of liquid and electrolyte also to control the blood pressure. The functional unit of the kidney is called nephron. Structural nephron contains glomerulus, capsule, and tubules. The condition of hyperglycemia can cause nephron damage. Diabetes control and complications trial and UK Prospective Diabetes Study show that hyperglycemia is the first cause of tissue damaging [17].

The study was similar to the previous study by Mohsen et al. showed that the destruction of the kidney. The study stated the proof of injection of alloxan 120 mg/kg BW subcutaneous after 1 week could cause the hyperglycemia condition in Wistar rats. After 8 weeks, decrypt histopathology from hyperglycemia rats' kidney showed the changing structure of vascular and interstitial from nephron tubules [18].

The width of complication on diabetes looks to be correlated with the concentration of blood glucose until over glucose that suspected as the main reason of the damaging of tissue. This phenomenon could be caused by the ability of hyperglycemia in vivo ways into the modification oxidative in vary subtract. Oxidative stress plays a crucial role in the development of hyperglycemia, which generates ROS. The result of ROS in the tissue can cause the overproduction of nitric oxide that will create interference in the various organs [19,20]. In the normal condition, the increasing of ROS or oxidation could be solved by antioxidant. However, the condition of hyperglycemia in the unbalance happened between the level of oxidant and antioxidant. The additional antioxidant was needed from the outside body to respond this thing.

Effect of natural antioxidants LLEE on the kidney tissue of Wistar rat alloxan induced showed in Figs. 3-5.

The use of herbs medicine could not be separated from the possibility of side effects due to the secondary metabolism of toxin substance that contains the herbs. This study was about the toxicity of LL that had been conducted before. In this study, there was no toxicity effect of the LL on tissue [12]. Dembinska-kiec et al. showed that flavonoids work as antioxidants by increasing the activity of SOD [21]. Flavonoids work as the radical scavenger, for oxygen singlet radicals and lipid peroxidation. Flavonoids suppressed the action peroxidase, thereby, inhibited the form of ROS by neutrophil. Flavonoids also inhibit the form of ROS by suppressing the action of enzymes production [22-24]. As a potential antioxidant, flavonoids tied the metals and stabilized the metals till the metals could not catalyze the reaction of free radical. Flavonoids dampen the reactivity of free radicals. Thus, it led to the molecule to be more stable. Flavonoids also led to donate ions hydrogen or electrons to the anions superoxidant until protecting lipoproteins, proteins, and DNA from oxidation [25]. The damaging of kidney cells organs due to hyperglycemia could be reversible. Repairing the function of the pancreas will create BGLs, the damaging of the kidney can overcome. The cells of kidney cells could repair and form the new cells.

CONCLUSION

Based on the results of this study that the LLEE dose 400 mg/kg BW effective decreased BGL and evaluated histopathological was able to restore the kidney destruction of alloxan-induced diabetic rats at LLEE 600 mg/kg BW.

CONFLICTS OF INTERESTS

The authors declare: There was no conflict of interest in this study.

AUTHORS' CONTRIBUTION

Mutia Indah Sari: literature research, designing the methodology of research, examining the blood specimen and reading the result of blood
Sari et al.

Asian J Pharm Clin Res, Vol 11, Issue 4, 2018, 257-261

specimen, analyzing the statistic data and writing the manuscript. Maya Anjelir Antika: collecting and preparing the sample, examining the blood specimen and reading the result of blood specimen, making the herbs, analyzing the statistic data. Dwi Rita Anggraini: literature research, analyzing the result of histopathology, analyzing the statistic data.

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