Cogongrass (Imperata cylindrica L.) Ethanol Extract on Sepsis Mice Model Body Weight and Sepsis Score

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Abstract

Sepsis causes damage for cells, behavioral phenotype regression, and will end in most patients' death. The ethanol extract of cogongrass (Imperata cylindrica L.) root (ECGR) acts as an antioxidant. This study aimed to observe the effect of giving ECGR to body weight (BW) and the sepsis score of the sepsis mice model by lipopolysaccharide (LPS) induction. This study was an in vivo study with a randomized post-test controlled group design at the animal laboratory of Universitas Padjadjaran, 2018. We used 4 (four) groups of male mice (Mus musculus) DDY strains. Group 1 as a control, group 2: LPS 10 μL/kgBW, group 3, and 4: LPS+ECGR (90 mg/kgBW, and a dose of 115 mg/kgBW, respectively). This treatment was performed for two weeks. Every three days, we measured their body weight. After two weeks, group 2, group 3, and 4 were injected with LPS for 8 hours to induce sepsis. Next, we measured body weight and sepsis score using murine sepsis score (MSS). Then statistical analysis was performed using ANOVA and the Kruskal-Wallis test. The results showed no differences in body weight were found in the treatment groups (3 and 4) compared with control, suggesting no effect of ECGR in decreasing mice body weight. The sepsis score was more than 21 in groups treated with LPS (2, 3, and 4), suggesting LPS can induce sepsis. There was a slight decrease in scores in-group 3 and 4 compared with group 2. This study concludes that the treatment of ECGR caused no harm to body weight and slightly decreased sepsis score in the sepsis mice model.

Key words: Body weight, cogongrass, murine sepsis score, reactive oxygen species
Introduction

Incidence of severe sepsis and septic shock has continued to increase over the past 30 years, and currently, more than 750,000 cases (about 3 out of 1,000 population). There is life-threatening organ dysfunction in sepsis due to the host’s response to the infection (inflammation). The process of organ damage will continue to occur during sepsis due to an imbalance in the redox cycle, potentially lethal organ dysfunction, and it can affect any age. Our previous study also found that sepsis induced by lipopolysaccharide (LPS) causes myocardial contractile dysfunction in mice.

Cogongrass (CG) or *Imperata cylindrica* L. is a plant that is often considered as weeds. Nevertheless, several recent studies have shown that this plant contains phenol compounds such as flavonoids and isoeugenin. These bioactive compounds can act as a potential anti-inflammatory and antioxidant.

Our previous study showed ethanol extract cogongrass (*Imperata cylindrica* L.) root (ECGR) depressed cholesterol level and triglycerides absorption in vivo studies. In this current study, we pursued to explore the effect of ECGR in mice model sepsis, where we used LPS to induce sepsis. In this current study, we measured the body weight and the sepsis score in the sepsis mice model after treatment with ECGR.

Methods

Experimental laboratory research was conducted using a randomized post-test controlled group design. The Health Research Ethics Committee of Universitas Islam Bandung, Bandung, West Java, Indonesia, has approved the ethical clearance with number 153/Komite Etik.FKIII/2018. This study was conducted at the animal laboratory of Universitas Padjadjaran, 2018.

Cogongrass root was obtained from Solo, Central Java, Indonesia, and was tested to its authenticity by Institute Teknologi Bandung Institute, Indonesia. The roots of CG were separated and washed clean with water, dried for two weeks, then macerated, and filtered. The filtration in the extract’s form was separated from the solvent by using a vacuum rotary evaporator. ECGR was diluted using carboxyl methylcellulose (CMC) 0.5% (Merck, U.S.A.) and divided in to 2 doses: 90 mg and 115 mg concentration/kgBW.

The mice (*Mus musculus*) DDY strains with the same breed, age of 8–10 weeks with body weight (30–35 grams), were selected as objects of this study. Mice were provided by PT Bio Farma, Bandung, West Java, Indonesia. These mice were divided into four experimental groups consisting of 8 mice/group. Therefore the total of mice used in this experiment was 32 mice. Group 1 as the negative control, mice that only treated CMC 0.5% (the solvent ethanol extracts of *Imperata cylindrica* L. root); group 2 (CMC 0.5%+LPS); group 3 and group 4, mice that treated ECGR in dose 90 mg/kg BW and dose 115 mg/kgBW, respectively + LPS. LPS is well known used to induce sepsis conditions in many previous types of researches.

The acclimatization of animal trials preceded the experiment for seven days in the laboratory. Thus they can adapt to their environment. Mice were placed within a cage at the animal laboratory of Universitas Padjadjaran with a controlled room temperature (setting of 12 hours of light and 12 hours of dark). Mice were given the standard food and drinking water ad libitum.

After acclimatization, groups 1 and 2 were treated 0.5% of CMC, groups 3 and 4 were treated ECGR+0.5% CMC each 90 mg/kgBW and 115 mg/kgBW for two weeks. The body weight of mice was measured every three days to find out whether there was an influence of the ECGR interfered with the bodyweight of mice or not. The ECGR was given once a day for two weeks, from 3 to 5 pm. After two weeks of treatment, groups 2, 3, and 4 were injected LPS 10 mg/kgBW (Sigma-Aldrich, St. Louis, U.S.A.). LPS was diluted within 50 μL PBS and injected intraperitoneal based on previous research. Next, we observed behavioral phenotype by using the murine sepsis score (MSS). There were seven variables assessed: appearance, level of consciousness, activity, response to the stimulus, posture, eyes, respiration rate, and respiration quality. The mice were in sepsis if the number of MSS in each mouse was more than 21 or had the respiratory quality or respiratory rate with a value of 3 or more.

Statistical analyses were performed using GraphPad statistical package. Variables were summarized using the mean±SD for normal distribution data and median+interquartile range (IQR) for skewed data. Normality distribution was assessed with the Shapiro-Wilk test. The p value was calculated using the analysis of variance (ANOVA) for normal distribution (parametric) and the Kruskal-Wallis test for skewed data.
(non-parametric). Two-tailed \( p \) values < 0.05 were considered statistically significant, and \( p < 0.01 \) = very significant.

**Results**

First, we measured the mice’s body weight before and after two weeks of treatment with ECGR. We showed no significant differences in mice’s body weight after treatment between each group (36.7; 38; 39.5; 39.4) gram with \( p = 0.35, p > 0.05 \).

Next, we measure the sepsis score using the MSS. Our result showed all mice in-group 2, 3, and 4, which were treated with LPS, were in sepsis condition (23, 21, 21), while in-group 1 (control) showed no sepsis (MSS=0). Using the Kruskal-Wallis test, we showed very significant differences in groups 2 and 4 compared with group 1 (\( p = 0.000; p = 0.007, p < 0.01 \), respectively) and significant differences in group 3 compared with group 1 (\( p = 0.018, p < 0.05 \)).

**Discussion**

In this study, we revealed that the treatment of ECGR did not interfere with the bodyweight of the mice; moreover, ECGR also slightly reduces sepsis score in the sepsis mice model, which is induced by LPS.

We showed that ECGR did not affect mice’s body weight, suggesting it does not interfere with the metabolism processes that can interfere with the bodyweight. Further study should be performed to measure body composition: fat mass, muscle mass, and body fat percentage.

MSS score is a method for assessing and comparing sepsis-associated outcomes, which consistently predicts sepsis mortality and progression in an animal model of sepsis.\(^{17–19}\) Using this score, in the current study, LPS increased MSS score of more than 21, suggesting LPS can induce sepsis. This result supported several previous studies.\(^{13–15,20,21}\)

Lipopolysaccharide (LPS) is an endotoxin located outside of the gram-negative bacteria membrane and one of the infection stimuli that is well known to be used in experiments in causing many profound immunological responses of the host; one mechanism the toll-like receptor (TLR) 4 pathway.\(^{1,15}\)

ECGR also slightly reduces sepsis score in the sepsis mice model, suggesting the possible role of ECGR maybe in some mechanisms in sepsis condition. The pathogenesis of sepsis involves the formation of reactive oxygen species (ROS). Endotoxins produced during sepsis, are capable of inducing ROS formation, such as superoxide, hydrogen peroxide, and hydroxyl.\(^{23}\) This ROS production will cause significant structural changes in the cell and ultimately cause multiple organ dysfunctions.\(^{24}\) Flavonoids in the ECGR play a role in the mechanism of ROS inhibition.\(^{8}\) Flavonoids work as an anti-inflammatory by inhibiting interleukin eight formations so that
the recruitment process of polymorphonuclear (PMN) cells to inflamed tissues can be inhibited.\textsuperscript{25} The flavonoids in the ECGR can also act as nitrite oxide (NO) scavenging, thereby reducing NO levels. Isoeugenin found in \textit{Imperata} roots can also inhibit inducible nitric oxide synthase (iNOS) so that the formation of NO free radicals can be suppressed.\textsuperscript{9} Our next project is to explore some possible mechanisms of the potential role of ECGR in sepsis condition. Also, we also observed the mice's behavioral phenotype by observing that mice in-group 1 had normal activities, such as eating, running, drinking, and other activities. On the contrary, the mice in group 2 showed suppressed activities, and most of the mice looked stationary. In-group 3 and 4 showed suppressing activities but more active than group 2, suggesting the potential effect of ECGR in improving the behavioral phenotype of mice induced by LPS.

\textbf{Conclusion}

The treatment of ECGR caused no effect on body weight and slightly decreased sepsis score in the sepsis mice model.

\textbf{Conflict of Interest}

The authors have read the manuscript and agreed to submit it in its current form for publication in the journal. There are no conflicts of interest to declare.

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