The Immunomodulatory Effect of Cinnamon (Cinnamomum Burmanii) bark extract on the C-Reactive Protein (CRP) Level, Leukocyte Count and Leukocyte Type Count of Wistar Rats Exposed to Staphylococcus Aureus

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ABSTRACT
Introduction: Bacterial infection induces inflammation in human body. This process produces humoral and cellular immune responses. Cinnamomum burmanii grows very vast in Indonesia and contains cinnamaldehyde known to have an anti-inflammatory effect. Objective: To prove the effect of C. burmanii bark extract on CRP level, leukocyte count and differential blood count.
Methods: A posttest-only controlled group design with 25 Wistar Rats divided into 5 groups was employed. The CN-G group was given the standard feed, the CP-G group was given the standard feed and levamisole 2.5 mg/KgBW, while the CBE-100, CBE-200, and CBE-400 groups were respectively given the standard feed and cinnamon bark extract 100 mg/kgBW, 200 mg/kgBW and 400 mg/kgBW. The treatments were conducted for 7 consecutive days. On day 8, all rats were injected with the suspense of S. aureus intraperitoneally. The blood was then drawn on day 9, followed with CRP level measurement using the ELISA method. The total leukocyte count and differential blood count were manually measured.
Results: There is no significant difference in the value of CRP level (One Way ANOVA; p = 0.749) with the total counts of leukocytes (p=0.685), monocytes (p=0.769), and eosinophil (p=0.123) between groups. The neutrophils and lymphocytes of CBE-100 group are significantly different from the other groups.
Conclusion: C. burmanii extract has a potential benefit as immunomodulator.
Keywords: Cinnamomun burmanii, CRP, leukocytes, differential blood counts, Staphylococcus aureus

INTRODUCTION
Staphylococcus aureus (S. aureus) is commensal bacteria which may be found in 30% of human population. Simultaneously, S. aureus is human's main pathogen which causes infection in various organs, such as bacteremia, infective endocarditis, osteoarticular...
infection, skin and soft tissue, respiratory tract, and infection in users of invasive devices (Tong et al., 2015). Emerging strain which is resistant to antibiotic such as Methicillin-Resistant S. aureus (MRSA) makes it difficult to eradicate infectious diseases in Indonesia.

Bacterial infection in the body will be initiated with an inflammatory mechanism. Inflammation is a biological response to a tissue injury. Inflammatory process increases the permeability of blood vessel wall, which is followed with immune cell migration and increased activity of signaling proteins and enzymes (Liao et al., 2012). Tissue macrophages, monocytes, mastocytes, platelet and endothelial cells may produce proinflammatory cytokines as a response to inflammation. Interleukin-1 (IL-1), interleukin 6 (IL-6) and Tumor Necrosis Factor (TNF) alpha are cytokines released by macrophages and initiate an acute phase immune response. In this acute phase immune reaction, liver synthesizes a protein called C-Reactive Protein (CRP). CRP is an acute phase protein of which value may increase 1000 times at the site of inflammation or infection. CRP has long been used as a laboratory sign of infection (Kingsley and Jones, 2008; Sproston and Ashworth, 2018). CRP is capable of recognizing self-molecule and foreign molecule based on pattern recognition. The bond between CRP and phosphocholine on bacterial cell membrane will activate complement through the classical pathway (Sproston and Ashworth, 2018).

Cellular immune response is mediated by T lymphocyte (T cell). CD4+ plays an important role in cellular and humoral immune system. Naïve T cell will recognize antigen and be active in peripheral lymphoid organs which will be differentiated to effector T cells and memory cells. With this activation, it will form sufficient antibody response which will in turn activate neutrophil and macrophage, which are useful in defense against S. aureus infection (Yu et al., 2018). Blood cell type count illustrates the relative percentage of respective type of white blood cells.

Cinnamon (Cinnamomum sp.) is of the Lauraceae family covering 54 species. The most important Cinnamomum species are C. burmanii, C. zeylanicum and C. cassia. C. burmanii (Java cinnamon) is an indigenous species to Indonesia, vastly cultivated and an important commodity in trade. Its dry bark has long been used as food spice and coloring (P. Chen et al., 2014). Ranasinghe, et al. (2013) presents a comprehensive review of cinnamon's medical benefit, such as anti-bacteria and anti-parasite, reducing blood sugar and cholesterol, antioxidant, anti-nociception, anti-inflammation and so on (Ranasinghe et al., 2013).

METHODS

This experimental research employs a posttest-only control group design. The research subjects were 25 male Wistar rats which are randomly divided into 5 groups. The groups were CN-G group given with the standard feed, CP-G group given with standard feed and levamisole 2.5 mg/kgBW, CBE-100 group given with standard feed and cinnamon extract 100 mg/kgBW, CBE-200 group given with standard feed and cinnamon extract 200 mg/kgBW and CBE-400 group given with standard feed and cinnamon extract 400 mg/kgBW. The treatments were performed for 7 consecutive days and on day 8 the rats were injected with S. aureus suspension intraperitoneally. The blood was sampled 2 mL on day 9 through the rats’ lateral tail vein for CRP level examination. This research was conducted upon approval from the Health Research Ethics Commission (KEPK) of the Faculty of Medicine of Diponegoro University/RSUP Dr. Kariadi Semarang under number: 922/EC/FK-RSDK/IX/2016 issued on September 19, 2016.

C. burmannii bark extraction

C. burmanii bark was extracted using the sokletation extraction method with ethanol 70% as solvent. The final result of extract was obtained in the form of powder.

Staphylococcus aureus injection

S. aureus was cultured in blood agar media, then 1-2 colonies were taken for suspension into 2 mL NaCl 0.9% until turbidity pursuant to the standard 0.5 Mac Farland. The suspension was then injected into the rats intraperitoneally at a concentration of 108 for 0.2 mL.
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Blood sample was dissolved in Turk's solution and Hemocytometer counting chamber by two examiners. Leukocyte count and Leukocyte type count examined manually on blood sample using Neubauer method at a wavelength of 450 nm at the GAKI counted per 4 large fields of view. A kappa test was conducted to reduce subjectivity bias, resulting in p>0.8. Blood film was used to determine the type of leukocyte and colored using Hematoxylin-Eosin staining.

Data analysis
The data were presented in mean and standard deviation (SD). The data resulted were tested for data normality and homogeneity. The data of CRP level, leukocyte count, and lymphocyte and monocyte type count are normally distributed with homogenous variance, a one way ANOVA was then conducted with a post hoc test. The data of neutrophil and eosinophil type count examination was conducted by using the Mann withney test and the Kruskall Wallis test.

Leukocytes’s type

Figure 1. Leucocyte count in each group. Mann Withney Analysis: * p<0.05; ns not significant

Table 1. Mean of CRP Level, Leukocyte Count and Leukocyte Type Count

| Variable          | CN-G N=5 Mean±SD | CP-G N=5 Mean±SD | CBE-100 N=5 Mean±SD | CBE-200 N=5 Mean±SD | CBE-400 N=5 Mean±SD | P value |
|-------------------|-------------------|-------------------|---------------------|---------------------|---------------------|---------|
| CRP Level (mg/dL)| 0.81±0.45         | 0.84±0.36         | 0.70±0.15           | 1.00±0.53           | 0.77±0.38           | >0.05*  |
| Number of leukocyte (Σ) | 11203±2376.50 | 13140±3207.10 | 13460±8060.66       | 9070±6267.73        | 11798±5007.19       | >0.05*  |
| Neutrophil        | 18.40±7.33        | 22.20±2.86        | 33.60±9.32          | 19.40±2.60          | 18.20±3.11          | 0.11**  |
| Lymphocyte        | 71.60±3.36        | 70±4.42           | 59±6.89             | 74.20±4.44          | 74.20±3.49          | >0.05** |
| Monocyte          | 7.40±2.30         | 6±1.58            | 5.40±3.65           | 6.60±2.30           | 7±2.92              | >0.05*  |
| Eosinophil        | 2.60±2.07         | 1.80±1.10         | 2±0.71              | 1.40±1.52           | 0.40±0.55           | 0.08*** |

Note: *Saphiro Wilk; ** Levene; ***Anova; “ Kruskall Wallis

CRP level examination
CRP level was examined using the ELISA method at a wavelength of 450 nm at the GAKI Laboratory, Faculty of Medicine, Diponegoro University, Semarang.

Leukocyte count and Leukocyte type count
Leukocyte count and leukocyte type count were examined manually on blood sample using Neubauer Hemocytometer counting chamber by two examiners. Blood sample was dissolved in Turk’s solution and counted per 4 large fields of view. A kappa test was conducted to reduce subjectivity bias, resulting in p>0.8. Blood film was used to determine the type of leukocyte and colored using Hematoxylin-Eosin staining.

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counts are not normally distributed, a Kruskal-Wallis non-parametric test was then conducted, followed with a Mann-Whitney test. The result was deemed significant if p value< 0.05.

RESULTS

After C. burmanii extract treatment and injection with S. aureus, the mean of CRP level, leukocyte count and leukocyte type count were displayed in Table 1.  

| CN-G: Control Negative-Group; CP-G: Control Positive-Group; CBE-100: group administered with 100 mg/kgBB Cinnamomum burmanii; CBE-200: group administered with 200 mg/kgBB Cinnamomum burmanii CBE-400: group administered with 400 mg/kgBB Cinnamomum burmanii |

Table 1 shows that statistically, the CRP level, leukocyte count, monocyte count and eosinophil type count are not significantly different between the treated groups (p>0.05). Meanwhile, the neutrophil and lymphocyte counts show significant difference between the groups (p<0.05). To identify which groups are significantly different, an LSD post hoc test was conducted for lymphocyte type count data and a Mann Whitney test was conducted for neutrophil type count data.

Neutrophil type count

The neutrophil type count of CBE-100 groups is of the highest value. Statistically, using the Mann Whitney test, the lymphocyte type count of CBE-100 group is significantly different from that of CN-G group (p=0.01) and of CBE-200 group (p=0.02) (figure 1).

Lymphocyte type count

The mean lymphocyte type count of CBE-100 group in Table 1 seems to be of the lowest value among the groups. Statistically, according to the data of LSD post hoc test, the lymphocyte type count of CBE-100 group is significantly different from that of CN-G, CP-G, CBE-200 and CBE-400 groups (p<0.00) (figure 1).

DISCUSSION

Immune system is a very complex defense mechanism requiring cooperation between immune system, nervous system and endocrine. Its main function is to protect the body from external biological invasion and to regulate body’s internal environment. The immunomodulatory concept shows the non-specific work of a substance on the immune system (cellular or effector molecule production by activated cells) (Balekar et al., 2014).

Imumomodulator works in various elements of the immune system through various mechanisms. In this research, we find that CBE administration does not significantly influence humoral immune system (CRP), but indirectly influences the composition of cellular immune system (neutrophil and lymphocyte type counts) in pathogen infection (S. aureus injection).

C-Reactive Protein (CRP) is an acute phase protein which significantly increases in an inflammatory process. In the statistical analysis, we find that there is no significant difference in the CRP level between the research groups, but the CRP level of CBE-100 and CBE-400 groups is generally lower than that of CN-G group, proving that CBE influences the CRP level decrease. The meta analysis study conducted by Vallianou et al. shows that administration of C. burmanii extract and its derivatives may decrease CRP level (Vallianou et al., 2019).

CRP is excreted in 4-6 hours after stimulus, and will be multiplied every 8 hours thereafter and reach its peak level in 36-50 hours. As an inflammatory sign, CRP plays a role in non-specific immunity which may with assistance of Ca2+ bind various molecules, such as phosphorylcoline expressed on damaged cell surface and polysaccharides and peptosaccharides on bacteria, parasite and fungi. Cynnamaldehyde in C. burmanii extract may inhibit IL-1β and TNFα secretion in lipopolysaccharide (LPS), which influences CRP expression. This secretion inhibition is mediated by modulation via JNK pathway, p38, activation of ERK1/2 and Iκ-Bα(Chao et al., 2008; Mendes et al., 2016).

The administration of C. burmanii bark extract in this research does not influence total leukocyte count and monocyte and eosinophil type counts. There is significant different in neutrophil and lymphocyte type counts between CBE-100 group and other groups. Neutrophil is one type of leukocyte serving in innate immune system, particularly in phagocyte function. Lymphocyte is a subtype of leukocyte serving in adaptive immune system. Immune system modulation resulting from C. burmanii bark extract administration influences immune response at the time of exposure to S. aureus infection (Nassan et al., 2015).

T cell is an important immune cell in adaptive immune system in improvement if phagocytosis mechanism and will through memory process provide long-term immunity. When Naïve T cells meet an antigen, they will become an active lymphoblast which will be proliferated and differentiated to be effector cells, which directly kill pathogen or secrete cytokines, such as IFN-γ, IL-2 and IL-4. IL-2 improves the lymphocyte resistance and growth factor, while IL-
4 induces antibody production. Cytokine modulation in administration of C. burmanii and its derivative products may influence leukocyte count and leukocyte type count (Lee et al., 2011). The research conducted by Roth-Walter et al. in 2014 states that cinnamaldehyde at low concentration increases NF-kB and at high concentration shows otherwise. The anti-inflammatory effect of cinnamaldehyde which is mediated by NF-kB blockage inhibits cell viability and influences proliferation and apoptosis induction in immune cells (Roth-Walter et al., 2014).

This change in leukocyte population is associated with administration of C. burmanii bark extract which is induced by TNF-α through adhesion molecule expression (Liao et al., 2012). The research conducted by Mendes et al. in 2016 states that cinnamaldehyde, an important constituent in C. burmanii, activates Transient Receptor Potential Ankyrin 1 (TRPA1) to modulate leukocyte cells (Mendes et al., 2016). Besides influencing migration, C. burmanii extract may influence secretion of some cytokines serving in proliferation and leukocyte differentiation, such as IL-2 and IL-4 (Lee et al., 2011). Chen et al. in 2014 states that procyanidin oligomer content in C. burmanii has the effect of suppressing splenocyte proliferation and also reducing interferon γ (IFN γ) and IL-2 contents (L. Chen et al., 2014).

The interesting fact regarding C. burmanii extract is that immunomodulatory function is determined by its dose and species (Vetal et al., 2013). The difference in the content of different types of constituent causes C. burmanii bark extract to be immunostimulant and immunosuppressant (Balekar et al., 2014). This is apparent from the result of variable of the three treatment groups administered with C. burmanii bark extract. In this result, we use C. burmanii extract and ethanol 70% as solvent. Ethanol 70% or 96% renders maximum effect on cinnamaldehyde and trans-cinnamaldehyde contents. Different extraction method and solvent type choice influence the final content of active agent obtained during extraction process (Wardatun et al., 2017).

CONCLUSION

Cinnamon (C. burmanii) bark extract is a potential immunomodulator. Therefore, more extensive further research is needed to examine the main effect of Cinnamon (C. burmanii) bark extract on immune system, particularly in the process of expression of genes and cytokines which play a role in cellular immune system proliferation process.

CONFLICT OF INTEREST

Authors declare no conflict of interest within this manuscript.

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