Immunomodulator effect of ketamine on rat models of sepsis (fecal induced peritonitis): Eosinophil and monocyte modulation

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Abstract

Background & Objective: Sepsis is a result of dysregulated inflammation and it affects the inflammatory mediator cell production. Ketamine has an immunomodulatory effect and it can be used in the sepsis immunotherapy. We conducted this study to determine the effect of ketamine on the number of eosinophils and monocytes in the rat model of sepsis.

Methodology: This research was a true animal experiment using 30 white rats (Rattus norvegicus), which were divided into six groups. Control negative group (NC) were not treated and the control positive group (PC) were the sepsis groups. Sepsis was produced by fecal induced peritonitis (FIP). Animals in the four treatment groups (A, B, C, and D) received intraperitoneal (IP) ketamine at 5 mg/kg body weight at different single hours (0, 3, 5 hours) and intermittently (every 2 hours), respectively, after receiving FIP. The eosinophil and monocyte counts were obtained 6 hours after FIP. The data were analyzed using One-Way ANOVA and Tukey HSD statistical test using SPSS software version 16.0 (p < 0.05).

Results: Sepsis significantly decreased the number of eosinophils but increased monocyte production. Ketamine treatment right after FIP (Group A) prevents the reduction of eosinophils in the sepsis group (p < 0.05) from 8 cells to 66 cells. Ketamine treatment in sepsis groups (Group A, B, and D) significantly reduced the monocyte production to close to normal (equivalent to NC group).

Conclusion: Sepsis significantly reduces the number of eosinophils and increases monocyte production in the rat models of sepsis. Ketamine, given soon enough after onset of sepsis helps to prevent the sepsis-induced changes in eosinophil and monocyte counts.

Key words: Eosinophil; Fecal induced peritonitis; Immunomodulation; Ketamine; Monocyte; Sepsis

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1. Introduction

Sepsis is a systemic dysregulated inflammatory response to infection. It is characterized by an overproduction of pro-inflammatory cytokines and has been associated with a high mortality rate. The number of patients with sepsis admitted to Intensive Care Units (ICU) increases every year. In Indonesia, a study shows that out of 4,774 patients admitted to the ICU, there were 504 patients diagnosed with sepsis. The mortality rate of sepsis is up to 70.2%. Medical record data collected between 2012-2013 at Dr. Saiful Anwar General Hospital, Malang found 1,026 patients diagnosed with sepsis and 788 (76.8%) of them died.

Various diagnostic markers of sepsis have been developed. C-reactive protein (CRP) and...
procalcitonin (PCT) are commonly used biomarkers. Recent research has shown that eosinophil is a specific sepsis marker and can be used as a diagnostic tool.\textsuperscript{6,7} The research showed an association between eosinophils and bacterial infections in ICU patients. An abnormal fall in the number of eosinophils (eosinopenia) can be used as a predictor of mortality in septic patients.\textsuperscript{8} Eosinopenia occurs because of the rapid sequestration of eosinophils in the peripheral circulation, suppression of eosinophil production and suppression of mature eosinophils migration. The decrease of eosinophil also happens due to the release of glucocorticoids from the adrenals stimulated by endotoxin and lipopolysaccharide. Glucocorticoids will inhibit the eosinophil release from the bone marrow, as well as their adhesion, migration, and chemotaxis. The eosinophils chemotactic inhibition through interleukin (IL)-3, IL-5, and GM-CSF inhibition mechanism. IL-5 controls eosinophil growth, differentiation, and survival rate. Sepsis also activates the inflammatory mediator cells such as monocytes and macrophages. Overexpression of inflammatory response leads to the abnormality of the immune system and result in organ-system dysfunction. In bacterial sepsis, endotoxin and lipopolysaccharide from the bacteria will induce macrophages, dendritic cells, and neutrophils secrete pro-inflammatory cytokines such as IL-1, IL-6 and tumor necrosis factor-\(\alpha\) (TNF-\(\alpha\)).\textsuperscript{8,10}

Ketamine is an anesthetic agent commonly used in sepsis patients.\textsuperscript{9} A study by Gurfinkel \textit{et al.} \textsuperscript{10} showed that ketamine improves the survival rate of severe burn injury followed by sepsis. Ketamine also has a good effect on maintaining cardiovascular and hemodynamic stability; and it has immunomodulating activities.\textsuperscript{10} Research carried out by Shanked \textit{et al.}\textsuperscript{11} showed that ketamine significantly increased the survival rates in rats with \textit{Escherichia coli}-induced sepsis.\textsuperscript{12} It inhibits pro-inflammatory cytokine production through downregulation of pro-inflammatory gene expression pathway.\textsuperscript{13} Eosinophil and monocyte counts can be used as therapeutic targets in sepsis. This study was conducted to determine the effect of ketamine on the eosinophil and monocyte counts in rats models of sepsis.

\section{2. Methodology}

This animal experimental study was conducted at the clinical pathology laboratory of the Faculty of Medicine, Brawijaya University, Malang, Indonesia in December 2016-January 2017. The research was approved by the Health Research Ethics Committee of Faculty of Medicine, Brawijaya University (No.452/EC/KEPK/12/2016). This study used 30 white male rats (\textit{Rattus norvegicus}), aged 5 months, weight 200-250 grams, showed active movements, and their hair did not fall out. The samples were divided into six treatment groups. The mice in negative control (NC group) were not treated. Positive control (PC group) was the sepsis group induced with fecal induced peritonitis (FIP). FIP was produced by dissolved feces from rat with normal saline until the concentration became 200 mg/ml. Feces 1 mg/g were administrated intraperitoneally (IP) to induce peritonitis. The mice received ketamine 5 mg/kg body weight intraperitoneally (IP) right after FIP (0 hours) in Group A, 3 h after FIP in Group B, 5 hours after FIP in Group C and intermittently every 2 hours (0, 2 and 4 hours after FIP) in Group D. Peripheral blood mononuclear cells were isolated from the heart at 6 h after FIP. The number of eosinophils and monocyte was calculated using a hematology analyzer.

\subsection{2.1. Statistical analysis}

The data (the number of eosinophils and monocytes) were analyzed statistically using SPSS software (version 16.0, IBM Statistic, United States). The data were analyzed using the homogeneity test and normality test with \(p > 0.05\). The data were then analyzed using the One-Way ANOVA test followed by the Tukey HSD test with \(\alpha = 0.05\). The significant difference happens when \(p < 0.05\).

\section{3. Results}

This research showed varied results. The number of eosinophils in the normal group (NC) was 56.4 cells. The administration of FIP in the sepsis group (PC group) significantly decreases the number of eosinophils to 8.8 cells \((p = 0.016)\). The administration of ketamine significantly increased the number of eosinophils in group A (66.2 cells) \((p = 0.008)\) compare to sepsis group. The number of
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Figure 1: Number of eosinophils in each group

Group control (-): normal; Group control (+): sepsis; Group A: administration of ketamine 0 h after FIP; Group B: administration of ketamine 3 h after FIP; Group C: administration of ketamine 5 h after FIP; Group D: administration of ketamine intermittently every 2 h. The letters above the error bar indicate when there is no statistically significant difference between the groups. Hence, the groups in this study that have a statistically significant difference are those labelled with an “ab” compared to those labelled with a “c.”

Figure 2: The number of monocyte in each group

Group control (-): normal; Group control (+): sepsis; Group A: administration of ketamine 0 h after FIP; Group B: administration of ketamine 3 h after FIP; Group C: administration of ketamine 5 h after FIP; Group D: administration of ketamine intermittently every 2 h. The letters above the error bar indicate when there is no statistically significant difference between the groups. Hence, groups are statistically significantly different when their labels share no letters.
eosinophils in group B was 30.3 cells, while in group D it was 24.2 cells. Both groups were not statistically different from the sepsis group, but did show an increase in the number of eosinophils. Group C had fewer eosinophils than the PC group (sepsis group) (Figure 1).

The results showed a significant increase in the number of monocytes in the PC group (sepsis model) compare to normal group. The number of monocytes in the NC (normal) group was 3.4 cells. FIP induction significantly increased the number of monocytes in the PC group to 13.6 cells ($p = 0.000$). The administration of ketamine in the sepsis group significantly caused a decrease in the number of monocytes in groups A, B, and D compare to sepsis group ($p < 0.05$). The number of monocytes in group A and B was 5.6 and 5.8 cells, while in the group D was 3.8 cells ($p > 0.05$). The $p > 0.05$ indicated that at the number of monocyte in groups A, B, and D were statistically similar to the normal group. The number of monocyte in Group C was 11 cells and it did not differ from the sepsis group significantly (Figure 2).

### 4. Discussion

Giving FIP in the PC group significantly decreased the number of eosinophils ($p = 0.016$). The decrease in the eosinophil count is called eosinopenia. The result supports the study by Abidi et al. which states that sepsis significantly decreases the number of eosinophils. The eosinophil number in the rats, that were immediately given ketamine after FIP, were higher than the sepsis group. Ketamine prevents the decrease of eosinophil count in the rat model of sepsis and keeps it close to the normal condition. Ketamine administration in the rat that were given ketamine 3 h after FIP, 5 h after FIP and intermittently every 2 h did not have a significant difference from the sepsis group ($p = 0.841$). Giving ketamine in the rat that were given ketamine 5 h after FIP did not prevent the decrease of eosinophil count because the drop in the number of eosinophils at 5 hours after FIP had occurred by the time the ketamine had been given.

The results showed that there was a significant increase in the number of monocytes in the PC group (sepsis group) from 3.4 cells to 13.6 cells ($p = 0.000$). The results of this study are similar to the research conducted by Lee and Kim. An increase in the number of monocytes occurs as an innate immune response. The administration of ketamine in the rats that were immediately given ketamine after FIP (5.6 cells), 3 h after FIP (5.8 cells), and intermittently every 2 h (3.8 cells) significantly caused a decrease in the number of monocytes close to the normal group ($p > 0.05$). Ketamine can inhibit monocyte differentiation through N-methyl-D-aspartate-dependent mechanisms.

The study showed that ketamine 5 mg/kg body weight (IP) can potentially serve as sepsis immunotherapy. Ketamine 5 mg/kg body weight can modulate both eosinophil and monocyte production. Ketamine was able to modulate the number of cells to stay in the homeostasis condition. The administration time shows a significant effect. The administration of ketamine right after FIP gives the best result in normalizing the number of eosinophils and monocytes to that of the normal condition. There was, however, no significant difference between one time administration and intermittent repeated administration of ketamine.

### 5. Conclusion

Sepsis significantly reduces the number of eosinophils and increases monocyte production in the rat models of sepsis; and ketamine, given soon enough, can help prevent the sepsis-induced changes in eosinophil and monocyte counts.

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### 7. Authors’ contribution

AAN, WWJ: Conceived and designed the experiments; conducted the study; analyzed and interpreted the data

AN, ODI: Conducted the study; analyze and interpreted the data; manuscript preparation and editing
8. **Conflict of interest**

The authors declare no conflict of interest.

9. **References**


