Growth Rate Differences of *Chrysomya* sp. Larvae on *Rattus novergicus* Wistar Strain Corpse Exposed and Unexposed to Ephedrine Toxic Dose

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ABSTRACT

Post Mortem Interval (PMI) is used as a parameter to define the onset, cause, manner, and time of death to help maintenance of law and identify crime victims by the age of the larvae found. Larval growth is influenced by the temperature, humidity, and contaminant (drug or toxic). This experiment used two media of dead rats (200 grams), one given ephedrine of LD$_{50}$ = 266 mg/kg and the other without ephedrine. Both were put in 2 cages each containing 50 *Chrysomya* sp. Ten larvae were randomly taken every day for 14 days to be measured their length, weight, and duration of growth. The 3rd larvae stage in the media with ephedrine had weight gain on 5th – 6th day morning, while larvae in media without ephedrine had static weight gain on 5th day morning – afternoon and weight decrement on 6 – 7th day morning. Larva’s peak ratio of length/weight in the media with ephedrine was higher than that of larvae in the media without ephedrine. Larvae on media with ephedrine grew faster in 1st, 2nd, 3rd larvae stage, and pupal stage compared to larvae without ephedrine. Therefore, this study indicates that ephedrine can accelerate the growth rate of *Chrysomya* sp. larvae.

Keywords: *Chrysomya* sp., Ephedrine, Larval Growth

INTRODUCTION

The FDA has reported a lot of ephedrine intoxication cases, leading FDA to withdraw ephedrine supplements in 2000. However, ephedrine is still sold freely and readily available at pharmacies due to its indications that are still needed [1]. Ephedrine can help relax the bronchial muscles so that it can be used in the treatment of bronchial asthma. Ephedrine is also able to boost appetite and also methamphetamine precursor materials due its similar structure [2, 3].

Death according to Indonesian government regulation no. 18/1981 Article 1 paragraph 9 is the cessation of heart function, breathing, or function of the brain that is declared by medical experts authorized. Meanwhile, according to declaration of Indonesian Doctor Association no: 231 / PB / A.4 / 07/1990 point 4, death is the cessation of spontaneous breathing and heart function which are irreversible with any evidence of brain stem death. Death can occur through a natural and unnatural process. Some examples of unnatural cause of death are homicide, accident, suicide, poisoning, and drug addiction. The incidence of deaths which are caused by the unnatural things is still a lot, still growing and increasingly worrying [4]. One branch of medicine that helps uncover cases of unnatural death is forensic entomology.

Forensic entomology is an applied science that is a combination of two fields of science, forensic and entomology. Entomology is the study of insects while forensics is the application of science to address crime and law enforcement. Thus, forensic entomology can be defined as the study of insects for the purposes of handling the problems of crime, particularly involving corpses. The most important application of forensic entomology is the approximation of Post Mortem Interval (PMI) or...
the time of death by looking at the minimum period of insect activity on the corpses [5].

Flies play an important role in medico legal forensic entomology through their breeding. In forensic entomology, the proper identification of specimens that relate to various types of insects, especially Chrysomya sp. (egg, larva, pupa, and adult form) that were found on the bodies, is very important on post mortem interval identification. Chrysomya sp. is often discovered because this species usually comes first on a dead animal or human [5].

This study is aimed to determine the speed of larval growth in media of Rattus norvegicus Wistar strain that were exposed to toxic doses of ephedrine. R. norvegicus Wistar strain is used because it has similar metabolic physiology to human.

MATERIALS AND METHODS

Research design

This study was a laboratory experimental research with true experimental design that aimed to compare length, weight, and growth rate differences of Chrysomya sp. larva on growth media containing rat corpse with ephedrine and without ephedrine. This research was conducted at the Laboratory of Parasitology, Faculty of Medicine, Brawijaya University in November 2014.

Growth Media

In this study, rats that were used as a growth media weighted 200 grams. One cage was filled with dead rat which was previously administrated with 120 mg (LD$_{50}$ = 600 mg/kg) of ephedrine orally then was euthanized by cervical dislocation after the onset of toxicity such as hyperkinesia acquired. Another cage was filled with dead rat which was euthanized by cervical dislocation without any treatment. Incision was made in the ventral midline from neck until anus so that visceral organ of rat could be clearly exposed. Then, these two cages were placed in a room with 25 – 28°C temperature.

Research samples

Chrysomya sp. was cultured in dead rat media with and without ephedrine. Fifty Chrysomya sp flies supplied by Laboratory of Parasitology, Brawijaya University, were put in each cage. Ten largest Chrysomya sp. larvae were selected from media with and without ephedrine which then were measured for their length, weight, and growth rate as sample.

Results and Discussion

The difference in length and weight of Chrysomya sp. larvae is presented in Figure 1. The length of larvae in media without ephedrine exposure had average peak length of 17.634 mm when on 3rd larva stage (day 5b) and later became pupae with an average length of 8.648 mm. In comparison, larvae in media with ephedrine was able to reach average peak length of 17.122 mm when on 3rd larva stage (day 6a) and pupae reached an average length of 9.164 mm.

On day 3b, lengths of larvae that grew in media without ephedrine were almost equal to the length of larvae in media with ephedrine, that were 6.266 mm and 6.846 mm (p = 0.050). As the days before, there was no significant length difference in both media on day 4a to 5a (p > 0.05). On day 5b, larvae in media without ephedrine gained an average length of 17.634 mm higher than that of larvae that grew in media with ephedrine, which was 15.650 mm (p = 0.003). Observation on day 6a also showed a significant difference (p = 0.001). However, the average length of larvae that grew in media containing ephedrine was 17.122 mm higher than that of larvae in media without Ephedrine, which was 14.840mm.

On day 6b, results showed no statistically significant difference (p = 0.073). On day 7a, larvae that grew in media containing ephedrine had been transformed into pupae. Meanwhile, the larvae that grew in media without ephedrine were still in 3rd larva stage. On day 7b to 13a average length was constant where the larvae had turned into pupae. The average maximum length of pupae that grew in media with ephedrine was 9.164 mm. Meanwhile, pupae that grew in media without ephedrine was 8.648 mm (p = 0.355).

Figure 2 showed that the growth of larvae in media containing ephedrine was able to achieve an average peak weight during 3rd larva stage (day 5b) with 77.12
mg of weight, while the larvae in media without ephedrine could reach the peak during 3rd larva stage (day 6a) with 75.46 mg of weight. On day 4a, the average weight of larvae in the media without ephedrine was 21.300 mg, which was higher than larvae in the media containing ephedrine with 14.220 mg of weight (p = 0.033).

On day 4b, larvae in media without ephedrine had an average weight of 37.620 mg which was higher than that of larvae in media containing ephedrine with 16.980 mg of weight (p = 0.003). This was because on day 4b larvae in the media with ephedrine had been transformed into 3rd larva stage. Day 5a and 5b showed the average weight differences of larvae in both media (p <0.05). However, on day 6a, there was no average weight differences of larvae in both media (p = 0.079). While on the day 6b, average weight of larvae in media containing ephedrine was statistically higher than those without (p = 0.046). On day 7a there was also a significant difference in weight (p = 0.002), but the larvae in
Ephedrine has sympathomimetic action as α and β receptor agonist and also increases the release of norepinephrine from sympathetic neurons [9]. Because of increasing activity of sympathetic nervous system, this may affect larvae staging and disrupt fly’s metamorphosis metabolism hormone. Ephedrine which increases the release of adrenergic nature, such as dopamine, can affect 20-hydroxyecdysone (20E) hormone contained in insect body. This hormone is an ecdysone active metabolite hormone that plays an important role in the coordination of developmental transitions, such as larval molting and metamorphosis [10, 11]. In human, hypothalamus signal which represents emotion, stress, and trauma, all can affect hypothalamus control toward growth hormone secretion. Furthermore, experiment has revealed that catecholamine, dopamine, and serotonin, all of which are released by various nervous systems within hypothalamus, can increase growth hormone secretion rate [12].

In this study, ephedrine in media influences larval growth and development process at each stage. Figure 1 shows comparison of overall average length of *Chrysomya* sp. larvae. On day 2 afternoon, media without ephedrine had 1st larva stage, whereas media containing ephedrine did not have any larvae, so that T test could not be performed. This was possibly due to starting time delay for flies to place their eggs in media containing ephedrine. On day 3 morning, average length of larvae in media without ephedrine was higher than media containing ephedrine (p = 0.000). On day 5b larvae in media without ephedrine had higher average length than larvae grown in media containing ephedrine (p = 0.003) did. However, on day 6 morning, the average length of larvae in media containing ephedrine was higher than that of larvae in media without ephedrine (p = 0.001). This is because ephedrine affects *Chrysomya*’s ecdysone hormone. On 7th day morning, larval stage showed a statistically significant difference (p = 0.001) because larvae in media with ephedrine had turned into pupa.

Figure 2 shows a comparison of overall average weight of *Chrysomya* sp. larvae. On day 3, 1st larva stage there was no a statistically significant difference (p > 0.05) in each media. On day 4, average weight of larvae in media without ephedrine was higher than that of larvae in the media with ephedrine (p = 0.033). On day 5, average weight of larvae in media without ephedrine was higher than that of larvae in media with ephedrine (p < 0.05). Further, on day 6 afternoons, average weights of larvae in media containing ephedrine was significantly higher than in media without ephedrine.

Observations of larvae were conducted starting from the length of larvae. It was found that on the first day, there were some larval length differences. This was because the newly hatched 1st larva stage did not digest dead rat. The 1st larva stage is still too small and their mouth are not strong enough to pierce tissue [13]. The maximum length of larva was obtained when the larva...
had entered the 3rd larva stage or the post feeding stage, which after this stage, larvae would metamorphose into pupae. The body length of larva would decline when it entered the pupa stage. This is the underlying reason why the length of larvae can be used as a growth rate parameter until the pupa stage [13].

The first stage of feeding is commonly called as 1st larva stage. A molting process indicates 2nd larva stage transformation, the second phase of feeding. At this stage, the larvae eat more than 1st larva stage. The outer skin of larvae then inflate until molting process occurs for second time to turn the larvae into 3rd larva stage [3]. In 3rd larva stage, there are two stages, namely feeding stage and post-feeding stage. Feeding stage larvae is the last stage of feeding. At this last stage, larvae eat greedily and have body size increment. When going into the pupa stage, 3rd larva stage larvae will stop eating and get away from the food source to change into pupae namely Post Feeding stage. In this study, larvae in media containing ephedrine stayed away from the food source (dead rat). Later, 3rd larva stage will turn into pupae. This process is influenced by hormones that play a role in metamorphosis. This process changes soft and yellowish white larval skin becomes blackish brown hard skin. Morphological and physiological changes occur, such as a decrease in movement, longitudinal muscle contraction, outer skin or cuticle shrinking, and cuticle color darkening.

This study showed a difference between Chrysomya sp. larvae which were exposed and unexposed by toxic doses of ephedrine. 1st larva stage which were normally have 36 hours life cycle was found to be 24 hours. The 2nd larva stage was 24-hour became 12 hours, and 3rd larva stage were 12 hours earlier than larvae which were unexposed by ephedrine. In both media, pupal stage took 156 hours until it turns into fly, but in media containing ephedrine it took 12 hours earlier. It is because the life cycle of larvae can be altered by the presence of contaminants that affect growth hormone. In addition to larval examination, forensic toxicology tests are required to determine any contaminants that exist in bodies that may affect the growth of larvae so there is no error in determining Post Mortem Interval (PMI).

This study shows that ephedrine affects the weight of larvae, in the sense that the media containing ephedrine has larvae which are heavier than larvae breed in the media without ephedrine. On day 6 morning average weight of larvae in media with ephedrine was heavier than those in the media without ephedrine. Peak of larval length per weight ratio in the media with ephedrine was achieved on day 6 morning with length/weight = 17.12/75.46 = 0.226875. This ratio was higher than the peak in media without ephedrine, that was 17.364/77.12 = 0.225155. The possible cause of this event is the aggressiveness of larvae to feed which increases, causing stratum muscularium of larvae increase in media with ephedrine exposure. A change in nutrition is one factor which influences the growth of larvae.

Life expectation of larvae can be measured by determining the larval stage. The size and length of larvae cannot be used as reference age of larvae due to the overlapping of different sizes on stage, especially 2nd and 3rd larva stage. In addition, post-feeding stage larvae have decreased body weight and length and then turn into pupae. Staging is determined through the observation of slit on larval posterior spiracles. Larvae undergo molting twice at 1st to 2nd larva stage and from 2nd to 3rd larva stage. Then, the larvae will undergo pupariation into pupae until later become adult flies [13]. To see the effect of ephedrine in detail at every stage, the examination of the posterior spiracles is performed. On day 5 afternoon, larvae in media without ephedrine still have 2 pieces of slit on posterior spiracles while larvae in media containing ephedrine already have 3 pieces of slit on posterior spiracles.

Some difficulties were encountered in this study. The number of samples used was 10 larvae per day. Multiplying the number of samples might cause a shortage of samples for subsequent observations because of difficulty to ensure the number of flies that laid eggs in the media. This study did not specifically look at the difference of larval growth in organs with high drug distribution.

**CONCLUSION**

It can be concluded that there are significant differences in the length of the Chrysomya sp. larvae. exposed to ephedrine toxic doses compared with the unexposed ones; there are significant differences in larval weight between larvae exposed to toxic doses ephedrine compared to unexposed ones; and ephedrine accelerates growth rate of Chrysomya sp. at 2nd to 3rd larva stage for 12 hours, 3rd larva stage to pupae for 12 hours, and transformation from pupae to adult flies for 12 hours.

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