Molecular Confirmation of Forensic Important Flies Collected in Palembang, South Sumatra

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ABSTRACT

The identification of insect species is an important key in forensic entomology. Molecular examination of the nucleotide sequences of cytochrome c oxidase subunit II (COII) gene has been widely performed before the morphology identification. We analyzed the partial conserved COII nucleotide sequence of the forensic related flies. After the DNA extraction from collected larvae, thus PCR amplification of conserved sequencing of the partial COII sequence was implemented. Obtained sequences were analyzed for a phylogenetic tree and a distance matrix. Our data showed a very low intraspecific sequence distance among Calliphoridae and Sarcophagidae to the flies’ species of Chrysomyia megacephala. Further research should attempt to determine the larval growth of newly discovered flies, to apply the time of death in forensic crime setting.

Keywords: Calliphoridae, Sarcophagidae, cytochrome C oxidase subunit II, morphology

1. INTRODUCTION

Judicial and forensic medicine has several strategies on how to determine the time of death in summarizing the cause of finding bodies. Changes in the size of the cadaver skin bruising color can determine the hours after death [1], and the composition of the eyeball liquid potassium has been used in court for evidence [2]. The use of artificial intelligence by using metabolites and blood pH is promising as a biomarker measurement [3]. Molecular biology methods can determine the time of accurate and reliable than the age of the injury [4], and the ecology of microbes after the decomposition of corpses [5]. The order of arrival of the insect on the carcasses can also predict the estimated time of death [6]. Determining the right time can help the alibis of the suspects are precise and exact.

Some species of flies known to play a role in determining the time of death of a corpse. Flies of the family Calliphoridae came the first time of death whom larvae are cannibals against larvae of other species that infests the body [7]. Flies of the family Sarcophagidae came after Calliphoridae, larger and carcasses have been found in the larval stage, making them easier observed at the site of death [8]. Flies of the family Calliphoridae more often used in forensic examination, but the vast morphological variation makes it a bit challenging as the variant species of African countries [9]. The larval stage can determine the diagnosis of species, due to the specificity of the shape and numbers of spiracles posterior and anterior lobe [10]. Fly species both adult stage, pupa, and larvae can act as a species diagnosis [11,12].

Chrysomya megacephala is closely associated with forensic entomology [13, 14]. The suspect of one murder from a burned corpses infested with flies was managed to be investigated and arrested due to the use of flies' predictions 50 hours after death [15]. The same species of flies may have different morphologies on the strain state [16]. Morphometric geometry use pupae as a benchmark in identifying the species C. megacephala [17]. Limited expertise in taxonomy [18], can be overcome with the use of a reference database of C. megacephala DNA [19, 20]. Proper identification of the fly species carried out by various methods exist, will further determine the time of death of a corpse.

Some related research regarding flies in Palembang, Chrysomya megacephala species documented in the market [21] and landfills [22]. An experimental studies using animal carcasses showed the presence of the species using morphological examination [23]. The species of C. megacephala seemed to be the only one of forensic importance. This is the first molecular attempt to utilize the mitochondrial COII nucleotide sequence analysis to confirm the forensically important flies of Palembang city.
1.1. Materials and Methods

This experimental analytic with a cross-sectional study design was conducted in September 2018 at the lab of Medical faculty Sriwijaya University, Palembang. The samples were larva of flies that infested the dead rats (*Rattus norvegicus*) of 2-3 months old with weight around 150-200 grams. Before the morphological key identification by observing the posterior and anterior spiracles with a dissecting microscope, the DNA genomic was extracted using Qiagen Genomic DNA Kit 50 according to the manufacturer’s instructions. Ethical approval obtained from the Ethics Committee of the Faculty of Medicine, Sriwijaya University, Palembang.

1.1.1. Larva preparation and morphology staining

Larvae were prepared in the solution before the staining for the microscopic identification. Specimens were first dipped in the hot water for immobilization before dissection of the abdomen for internal organ removal. Later, the solution of KOH 10%, solution of acetic acid, and the solution series of alcohol 70-80-90-96% were performed 30 min respectively. The staining of specimens was using the clove oil solution for at least 30 min. Thus, the samples were ready to be observed using a microscope examination.

1.1.2. Polymerase chain reaction and automatic sequencing

Primer sequences for the COII gene were taken from the literature [19]. Amplifications of COII locus were performed using primers C1-J-2495 (5'–CAG CTA CTT TAT GAG CTT TAG G-3') and TK-N-3775 (5'–GAG ACC ATT ACT TGC TTT CAG TCA TCT-3') and PCRs were performed using a 2720 Thermal Cycler (Applied Biosystems, Foster City, CA, USA). The PCR reaction conditions consisted of an initial denaturation step at 95 °C for 5 min, annealing step at 93 °C for 1 min, extension step at 48 °C for 1 min, and final extension step at 72 °C for 2 min. Each reaction mixture was prepared using 50 ng of template DNA, 2.5 µL 10× AmpliTag Gold Buffer, 0.5U AmpliTag Gold DNA Polymerase (Applied Biosystems, Foster City, CA, USA), 10 pmol (each) upstream and downstream primers, 62.5 mM MgCl2, 5 mM (each) dNTPs, and sterile distilled water to a final volume of 25 µL. After purification of the PCR products, sequencing reactions were performed according to the manufacturer’s instructions using a BigDye v3.1 Cycle Sequencing Kit (Applied Biosystems, Foster City, CA, USA). The sequencing products were analyzed using an ABI 3730xl Genetic Analyzer (Applied Biosystems, Foster City, CA, USA). Assembled sequences were deposited into the NCBI GenBank database.

1.1.3. Phylogenetic analysis and sequence comparison

Phylogenetic trees were generated for 2 fly families by using the maximum likelihood method with 1,000 replicates of bootstrapping based on the Tamura-Nei model using MEGA6 software [24]. Initial trees for the heuristic search were obtained by applying the neighbor-joining method to a matrix of pairwise distances estimated using the maximum composite likelihood (MCL) approach. To make root for each tree, COII sequences for *Hemipyrellia ligurriens* (KF037968), *Sarcophaga ruficornis* (NC_041069), *Sarcophaga dux* (MH765524), and *Chrysomya rufificacies* (GQ912623) were introduced as outgroup taxa. Average intraspecific and interspecific sequence distances were calculated for sequence comparison. Sequences obtained in this study were later aligned with to previously published in GenBank data.

1.2. Our Contribution

This paper presents the first reports of DNA-based COII nucleotide sequences from flies’ species. *Hemipyrellia ligurriens*, *Sarcophaga ruficornis*, *S. dux*, and *Chrysomya rufificacies*, were four flies species that share the large similarity the Palembang flies. However, the morphology identification of the larva assumed the sample as *C. megacephala*. Microscopy showed of the spine between the first and second thoracic segments and the morphology of the posterior spiracle make it difficult to carry out differential identification sequences obtained from the mitochondrial COII gene fragment of Calliphoridae, and Sarcophagidae are now available and can be used as an additional tool for the identification of populations from the City of Palembang, South Sumatra, Ind, especially in forensic applications.

1.3. Paper Structure

The rest of the paper is organized as follows. The first section introduces the morphological identification, which used the anterior and posterior spiracles as microscopical keys. The second section described of the primers applies the COII conserved gene of flies. The third section presents the phylogenetic analysis from the sequence samples compared to the GenBank data. The last section concludes the findings.
2. RESULTS AND DISCUSSION

2.1. Morphological Results

Larval identification can be determined from the anterior and posterior spiracles. Larvae with papillae anterior spiracles showed 1 to 9 finger-shaped protrusions; posterior spiracles figured the peritreme incomplete and do not cover the button; there are three slits which are straight and wide (Figure 1). These 3 slits show larvae of *C. megacephala* at the third instar stage (Figure 2). On the contrary, the *C. rufifacies* has as many as 9-10 papillae anterior spiracles bulge fingers, which were arranged in a straight line. Posterior spiracles shaped slits width, also corresponding with larval instar stage, open peritreme, thin and slightly dark [24, 25]. It was concluded based on the anterior-posterior spiracles characteristics that samples found were identified as *C. megacephala*.

![Figure 1](image1.png)
**Figure 1** Microscopic examination of the larvae *Chrysomya megacephala*: the anterior spiracles with 9 protrusions (magnification 10x10x4.5). Caption: 1-9 end spine bands

![Figure 2](image2.png)
**Figure 2** Microscopic examination of the larvae *Chrysomya megacephala*: The posterior part with posterior spiracles of third instar larvae (magnification 10x10x4.5). Caption: (A) incomplete peritreme; (B) three slits, straight shape, and width; 1-3: slit numbers

2.2. PCR Results

Positive samples resulted in the appearance of the white band in 1324 basepair. Negative samples showed no white band, indicated that not all microscopically positive samples showed positive PCR tests (Figure 3). Positive samples showed faint and thick white bands, which were the number 16, 21, 24, 34, 43, 48, 49, 50, 51, 52 and 53. Out of 11 that have been observed whit thick white band, thus 5 samples were taken randomly for the sequencing process. Positive samples were furthered test for sequence analysis.

2.3. Phylogenetic Analysis Maximum Likelihood

The five samples randomly chosen were aligned with GenBank COII flies' genes. The five samples were 99-100% similarity to *Hemipyrellia ligurriens* (KF037968), *Sarcophaga ruficornis* (NC_041069), *Sarcophaga dux* (MH765524), and *Chrysomya rufifacies* (GQ912623). The GenBank data showed that four Palembang Indonesia species were highly similarity same as in Brazil, Portugal, Thailand, and Australia. *Chrysomya megacephala* is difficult to differentiate from the microscopic identification due to a high similarity, therefore, require experienced taxonomist. The results of sequencing are more useful for species identification as *C. megacephala* flies [26].

In summary, sequences obtained from the mitochondrial COII gene fragment of *Calliphoridae*, and *Sarcoptagidae* which now available online can be used as an additional tool for the confirmation of flies’ identification especially in forensic applications to the estimated time of the death investigation.
Figure 3 Agarose gel with ethidium bromide electrophoresis UV visualization of 50 samples. Positive PCR amplicons of COII were at 1324 bp. Caption: (M) marker; (C) negative control; (BP) basepair; (1K1.1-2K47) samples’ numbers

Figure 4 The phylogenetic tree using the conserved partial gene of COII showed confirmation of Calliphoridae and Sarchophagidae species. The five randomized samples of Palembang fly strain were the Calliphoridae family (Chrysomya rufifacies and 2 Hemipyrellia ligurriens); and the Sarchophagidae family (Sarchopaga dux and S. ruficornis)

3. CONCLUSION

We concluded that molecular reconfirmation is necessary for flies related to forensic identification, while morphology identification is quite not specific to public practice. Research should further investigate the larval growth of newly discovered flies related forensic flies in a laboratory setting, to understand the correlation with its finding in crime scene-setting.

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