**Chrysomya chani** Kurahashi (Diptera: Calliphoridae), a blow fly species of forensic importance: morphological characters of the third larval instar and a case report from Thailand

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**ABSTRACT**

Blow flies are worldwide important insects from a forensic point of view. In Thailand, beside from the two most common species, *Chrysomya megacephala* (F.) and *Chrysomya rufifacies* (Macquart), *Chrysomya chani* Kurahashi was also found to be of forensic importance. We present a case of a human female cadaver in its bloated stage of decomposition, discovered at Pachangnoi Subdistrict, northern Thailand. Entomological sampling during the autopsy displayed an assemblage of numerous dipteran larvae. Macroscopic observations showed the coexistence of third instar larvae of the three blow flies *C. megacephala*, *Chrysomya villeneuvi* Patton, an unknown blow fly species and one muscid, *Hydrotaea* sp. The minimum post-mortem interval was estimated to be six days, based on the developmental rate of *C. megacephala*. The ID of the unknown larva, which is the focus of this report, was revealed later as *C. chani* by DNA sequencing, using a 1205 bp of cytochrome c oxidase subunit I (COI). The occurrence of *C. chani* on a human body revealed the need to analyse and describe the morphology of its immature stage, to enable forensic entomologists to identify this fly species in future cases. The morphological examination of the third instar was performed, revealing peculiar characteristics: protuberant tubercles encircling abdominal segments; 9–11 lobes on the anterior spiracle; six prominent pairs of tubercles along the peripheral rim of the eighth abdominal segment; a heavily sclerotized complete peritreme of the posterior spiracles. A key to differentiate the third instar of blow flies of forensic importance in Thailand is provided.

**Keywords**

Forensic science; forensic entomology; *Chrysomya chani*; larval morphology, identification; Thailand

**Introduction**

Blow flies (Diptera: Calliphoridae) are insects of forensic importance, since they are the first colonizers of human cadavers, often arriving promptly after death, thereby showing a great potential in forensic investigations worldwide [1]. In Thailand, 10 blow fly species (*Chrysomya megacephala*, *Chrysomya rufifacies*, *Chrysomya villeneuvi*, *Chrysomya chani*, *Chrysomya bezziana* Villeneuve, *Chrysomya pinguis* (Walker), *Chrysomya nigripes* Aubertin, *Lucilia cuprina* (Wiedemann), *Lucilia porphyrina* (Walker) and *Hemipyrella ligurriens* (Wiedemann)) have been revealed to be forensic relevance in the last decade [2–5]. *Chrysomya chani* was only reported once so far, infesting human remains in a forest [2]. Some larvae from that case were reared to the adult stage and identified by their adult morphology. Hereby, we present a second case and describe the larval morphology of this so far unknown species in forensic entomology. By providing a key for the third larval stage of *C. chani* and describing the molecular identification (of any developmental stage or even just fragment) of that species, we establish an identification tool for this species.

**Case report**

**Case history**

In August 2006, the remains of a 38-year-old female were discovered in Pachangnoi Subdistrict (N 19°19’24.24”; E 100°27’17.28”), Pong District, Phayao Province, northern Thailand (Figure 1), and transferred to the Department of Forensic Medicine, Faculty of Medicine, Chiang Mai University. Forensic autopsy revealed the impact of blunt and sharp forces on the head and the abdomen. The bloated body was infested by fly maggots. Two blow flies, *C. megacephala* and *C. villeneuvi*, and one muscid species belonging to the genera *Hydrotaea* were identified. The minimum post-mortem interval (PMImin) was estimated to be six days, based on the developmental rate of *C. megacephala*. However, identification of one fly species could not be achieved due to the limited information on fly...
larvae morphology of species of forensic importance in Thailand at that time.

**Morphological identification**

Initial macroscopic examination of the fly larvae sampled showed two groups of maggots: third instars of non-hairy maggots and third instars hairy maggots. The non-hairy maggots were identified as the blow fly, *C. megacephala* and the muscid *Hydrotaea* sp. The hairy maggots were mainly identified as third instars of the blow fly, *C. villeneuvi* [2]. Interestingly, we sampled about 20 third instars specimens that were unfamiliar to us. All larvae sampled were preserved in 70% alcohol. The largest larva was measured using Vernier calipers. In 2017, we decided to reanalyse the unidentified larvae of this case. Some specimens were cut using a sharp blade at two sites, across the third thoracic segment and across the middle of the eighth abdominal segment, to examine the cephaloskeleton and posterior spiracle, respectively. The anterior and posterior ends were transferred into a small eppendorf tube consisting 10% KOH. These tubes were then put into a beaker (half-filled with boiling water) which was placed on a hotplate (Barnstead/Thermolyne, Model: SP46920-26, USA) for 10 min. Specimens were washed twice with distilled water. To remove the alcohol, specimens were placed in Cellosolve (ethylene glycol monoethyl ether) and left for 5 min. The specimens were then transferred onto a glass slide and one drop of Euparal® was added. Specimens were arranged in their appropriate positions and covered with a cover slip. Prepared anterior and posterior ends were examined and photographed under a light microscope (Olympus CX41, Tokyo, Japan, with Olympus DP22 digital camera). For photographing, the focus stacking was shot by taking a series of images with the same composition and gradually changing the area of sharp focus. The number of pictures taken depended on the thickness of the specimen. For preserved larvae, stacking pictures were taken using a Nikon D7100 digital camera with a Nikkor lens Af-s macro 60 mm f2.8G. Each picture was merged in the program Helicon Focus 6.6.0 using method C (pyramid stacking). Terminology for general larval morphology followed Courtney et al. [6], for peripheral tubercles of larvae followed Liu and Greenberg [7] and for modifications of larval cephaloskeleton followed Szpila et al. [8].

The largest specimen was 12.8 mm long. The prominent features are the protuberant tubercles encircling body segments, of which the prominent tubercles originate from the first abdominal segment and are present until the eighth abdominal segment (Figures 2(A–C) and 3(A)). Spine bands between segments are obvious in the thoracic segments, with the most prominent between the first and second thoracic segments (Figure 2(A)). These spine bands between thoracic segments are relatively sclerotized (Figures 2(A,B)). However, spine bands between the abdominal segments are unnoticeable (Figure 2(C)).

A very distinct ultrastructure of the surface integument is seen on the body surface, which is covered with dense variable size of denticles (Figures 2(A–C) and 3(B)). The anterior spiracle consisted of 9–11 lobes (n = 13) arranged in a single row (Figures 3(B,C)). The
Cephaloskeleton (Figure 3(C)) has large and heavily sclerotized mouthhooks, curved downwards. The posterior base of the mouthhooks is large and broad. An accessory sclerite is moderately sclerotized, adjacent to the base of the mouthhooks. The dental sclerite is apparent, curved backward apically and connected to the base of the mouthhooks. An intermediate sclerite is present. The parastomal bar is slender and slightly curved upward apically. The dorsal bridge is slender apically and bent downward, with the same length as the anterior margin of parastomal bar. The dorsal cornua, vertical plate and ventral cornua are heavily sclerotized. The dorsal cornua are much longer than the ventral cornua. The ventral cornua have an opening or window (Figure 3(C), arrow).

The posterior end of the third instar shows six prominent pairs of tubercles along the peripheral rim of the eighth abdominal segment (Figures 2(A,C) and 3(E)), of which all six pairs (inner dorsal, median dorsal, outer dorsal, outer ventral, median ventral and inner...
ventral tubercles) are almost equally in their protuberance. Viewed posteriorly, there is a remarkably sculpture encircling the posterior spiracles and adjacent to these tubercles (Figure 3(E)). Higher magnification of the posterior spiracles revealed thick, heavily sclerotized complete posterior spiracular peritremes enclosing three spiracular slits (Figure 3(F)). The button (or ecdysial scar) is indistinct (Figure 3(F), arrow).

Figure 3. Light micrographs of third instar larvae of *Chrysomya chani*. (A) Whole body, lateral view; anterior end at left, posterior end at right. Arrows indicate the initial protuberant tubercles encircling body segments at the first abdominal segment (arrow). (B) Anterior end showing cephaloskeleton (c), anterior spiracle (a) and spine bands (s) between the first and second thoracic segments. Anterior spiracle consisted of 9–11 papillae arranging in a single row. Arrow displays surface integument, covering with dense variable size of denticles. (C) Higher magnification of the cephaloskeleton and anterior spiracle (a). Arrows indicate window of ventral cornua. Abbreviation of cephaloskeleton: as, accessory sclerite; db, dorsal bridge; dc, dorsal cornua; den, dental sclerite; is, intermediate sclerite; mh, mouthhooks; pb, parastomal bar; vc, ventral cornua; vp, vertical plate. (D) Posterior view of the eighth abdominal segment showing prominent six pairs of tubercles along the peripheral rim. Posterior spiracle (ps) is apparent. Abbreviation of tubercles: IDT, inner dorsal tubercles; MDT, median dorsal tubercles; ODT, outer dorsal tubercles; OVT, outer ventral tubercles; MVT, median ventral tubercles; IVT, inner ventral tubercles. (E) Posterior view of the eighth abdominal segment showing remarkably sculpture encircling the posterior spiracles (ps) and adjacent to these tubercles (arrow). (F) Higher magnification of posterior spiracles displaying thick, heavily sclerotize complete posterior spiracular peritreme (p) enclosing three spiracular slits. Arrow indicates indistinct button (or ecdysial scar).
A key for identification of the third instar of blow flies of forensic importance in Thailand is provided as follows:

1 Abdominal segments with large, elongate tubercles (Figures 4(A) and 5(A)).......................... 2
2 Abdominal segments lacking large, elongate tubercles (Figure 4(B))........................................ 3
2 Tubercles bear numerous small spines at tip (Figure 5(B)); anterior spiracle with 9–12 lobes; posterior spiracle large, with heavily sclerotized incomplete peritreme (Figure 5(C)) .......................................................... C. rufipes (Macquart) Tubercles bear numerous small spines throughout (Figure 5(D)); anterior spiracle with 13–15 lobes; posterior spiracle large, with heavily sclerotized incomplete peritreme (Figure 5(E)) ............................................. C. villeneuvi Patton
3 Abdominal segments bear protuberant tubercles (Figure 2(A–C)); anterior spiracle with 9–11 lobes; six prominent pairs of tubercles along the peripheral rim of the eighth abdominal segment (Figures 3(D,E)); posterior spiracle large, with moderately sclerotized complete peritreme (Figure 3(F)) ............................................. C. chani Kurahashi
4 Peritreme incomplete.................................................................... 5
5 Peritreme complete...................................................................... 7
5 End of upper peritreme gradually enlarged (Figure 6(A)); spines between the first and second thoracic segment large, multipointed (Figure 6(B)); some specimens with brown patch on dorsal integument (Figure 6(C)), but some not (Figure 6(D)).................................................................................. C. nigripes Aubertin
6 Anterior spiracle with 4–6 lobes; posterior spiracle large, with moderately sclerotized incomplete peritreme (Figure 7(A)); spines between the first and second thoracic segment large, single point (Figure 7(B))............................................................................... C. bezziana Villeneuve
7 With prominent outer ventral tubercle at the rim of the eighth abdominal segment; anterior spiracle with 9–12 lobes; posterior spiracle lightly sclerotized with incomplete peritreme (Figure 8(A)); spines between the first and second thoracic segment large, arrange singly or rows (Figure 8(B)); accessory sclerite heavily sclerotized (Figure 8(C)). ........................................................................... Lucilia sinensis Aubertin
8 Posterior spiracle small, lightly sclerotized (Figure 8(D)); anterior spiracle with 3–6 lobes; spines between the first and second thoracic segment small, arranged mostly in group or row (Figure 8(E)); accessory sclerite un sclerotized (Figure 8(F))........................................................................... Lucilia cuprina (Wiedemann) Posterior spiracle large, moderately sclerotized (Figure 8(G)); anterior spiracle with 5–9 lobes; spines between the first and second thoracic segment small, arranged mostly in rows (Figure 8(H)); accessory sclerite un sclerotized (Figure 8(I)) ............................................. Lucilia porphyrina (Walker)

Specimens of larvae used in the key were from laboratory colony and/or forensic death scenes, by which adults were confirmed by morphology.

**Molecular identification**

Some samples of these unknown species have been preserved in 70% ethanol since 2006 with the label of date and time of collection. To confirm the state of the species, molecular identification was performed in 2017 using these samples kept for more than 10 years.

For DNA extraction, polymerase chain reaction (PCR) amplification and DNA sequencing, the genomic DNA was extracted from one larva according to the dilution protocol of the Phire Animal Tissue Direct PCR Kit (Thermo Scientific). After measuring the DNA concentration, 100 ng/mL of the extracted DNA was subsequently used for PCR reaction.

DNA amplification was performed according to the PCR protocol of the kit. Partial COI sequences were amplified using the primers TY-J-1460 (5’-TACAATT-TATGCCCTAAAACITCCAGGC-3’) and CI-N-2800 (5’-CAATTCAAGCTGTGAAGCATC-3’) [9]. Cycling condition was initially denaturation at 98 °C for 5 min, followed by 40 cycles of denaturation at 98 °C for 5 s, annealing at 61.7 °C for 5 s, extension at 72 °C for 30 s, followed by extension at 72 °C for 1 min. PCR products were electrophoretically separated in an 1% agarose gel, stained with RedSafeTM (Intron Biotechnology). For sequencing, the unpurified PCR products were sent to the First BASE Laboratories Sdn Bhd (Selangor, Malaysia).

To perform the sequence alignment and phylogenetic analysis, the obtained DNA sequences from both directions were edited and assembled using BioEdit software version 7.0.9.0. [10]. For the highest similarity search, the sequence was compared with the available sequence database via a Basic Local Alignment Search
Figure 4. Light micrographs of third instar larva of Chrysomya villeneuvi and Lucilia cuprina. (A) Third instar larva of C. villeneuvi showing large, elongate tubercles on the abdominal segments. (B) Third instar larva of L. cuprina showing smooth abdominal segments.

Figure 5. Light micrographs of larva of Chrysomya rufifacies and Chrysomya villeneuvi. (A) Third instar larva of C. rufifacies showing large, elongate tubercles on the abdominal segments. (B) Tubercle of C. rufifacies larva bears numerous small spines at tip. (C) Posterior spiracle of C. rufifacies showing large, heavily sclerotized incomplete peritreme. (D) Tubercle of C. villeneuvi larva bears numerous small spines throughout. (E) Posterior spiracle of C. villeneuvi larva showing large, heavily sclerotized incomplete peritreme.
Figure 6. Third instar larva of *Chrysomya nigripes*. (A) Posterior spiracle showing gradually enlarged of end of upper peritreme (arrow). (B) Spines between the first and second thoracic segment. (C) A specimen with brown patch on dorsal integument (arrow). (D) A specimen without brown patch of dorsal integument.

Figure 7. Light micrographs of larva of *Chrysomya bezziana* and *Chrysomya megacephala*. (A) Posterior spiracle of larva of *C. bezziana*. (B) Spines between the first and second thoracic segment of *C. bezziana*. (C) Posterior spiracle of larvae of *C. megacephala*. (D) Spines between the first and second thoracic segment of larvae of *C. megacephala*. 
Tool (BLAST) search at the National Center for Biotechnology Information [http://blast.ncbi.nlm.nih.gov/Blast.cgi]. Using MEGA6 software [11], neighbour-joining tree [12] was constructed using Kimura 2-parameter (K2P) model [13] with 1000 bootstrap replications. Additionally, reference sequences retrieved from GenBank covering a fragment length equal or longer than ours were aligned, trimmed and added to the analyses.

Based on the BLAST search, our larval sample collected from the human corpse was 100% identical to C. chani (GenBank accession no: KR921606), confirming to be C. chani, based on 1205 bp of COI (Figure 9). Additionally, analysis of phylogenetic analysis revealed that C. chani was grouped within C. megacephala, C. pinguis, C. thanomthini, C. bezziana and C. nigripes [14].

**Discussion**

To our knowledge, biological information of C. chani is rather limited [15,16]. This species was first described as a new species from Singapore in 1979, of which male genitalia and female ovipositor have been illustrated [15]. This is the first report on morphological characteristics of the third larval instar of C. chani. Altogether, morphological and molecular tools now strongly affirmed C. chani as a forensically important species in Thailand. It is our belief that this fly is much more common on human bodies in Asia than expected so far, and that the problem of identification is the main reason for this. The case 11 years ago illustrates this dilemma.

Several morphological characteristics of third larval instar C. chani were similar to those reported in third instar of the hairy maggots, C. rufipes [17], C. ville-neuvi [18] or Chrysomya albiceps (Wiedemann) [19], namely the protuberant tubercles which encircle the abdominal segments, the six pairs of prominent tubercles along the peripheral rim of the eighth abdominal segment, the heavily sclerotized spine bands between the thoracic segments, the sculpture of the surface integument, the number of papillae on the anterior spiracles and the heavily sclerotized peritreme of the posterior spiracles.

Remarkably, the morphology of the male genitalia of C. chani showed to be very similar to those of C. megacephala, C. pinguis and C. thanomthini Kura-hashi and Tumrasvin (Sontigun et al., unpublished data). In addition, our phylogenetic analysis based on nearly full length of COI (Figure 9) and cytochrome c oxidase subunit II (COII) genes (Sontigun et al.,
unpublished data) revealed that C. chani was placed within the non-hairy group (C. megacephala, C. pinguis or C. thanomthini) even though several morphological characteristics of third instar C. chani were similar to third instar of the hairy maggots (C. rufifacies, C. villeneuvi or C. albiceps). This finding was similar to the results of Singh et al. [14], who performed the phylogenetic analysis based on 2 386 bp of combined COI (1 536 bp) and nuclear carbamoylphosphate synthetase genes, and Zajac et al. [20] based on the COI barcode region (about 700 bp). In contrast, when using 28S nuclear rRNA gene (about 1 000 bp), C. chani was grouped with the hairy maggot blow flies, namely C. rufifacies, C. villeneuvi, and C. albiceps [20]. These findings indicate that the choice of the gene to be analysed may cause variation in the outcome and the subsequent taxa arrangements. Based on the phylogenetic incongruence between the mitochondrial and nuclear genes for C. chani, the combination of multiple genes should be analysed to resolve the phylogenetic relationships. In the previous studies, the phylogenetic trees of the blow flies were frequently differed when constructed using different loci of genes [21–23]. Furthermore, the phylogenetic placement of taxa was found to depend on the analysed taxa, gene length and tree-building methods [14,24]. Although the single gene was sufficient to identify blow flies it is unlikely to separate some closely related species [25–29] and it cannot be used to resolve the phylogenetic relationships. Therefore, the use of multiple genes in different loci is required to enhance confidence in species identification, particularly of closely related species, and resolve the phylogenetic relationships between species of blow flies of forensic importance.

The main characters used for identification of third instar C. chani observed in this study (e.g. presence of accessory sclerite on cephaloskeleton, number of papillae on anterior spiracle, heavily sclerotized posterior spiracle, complete peritreme) was related to those previously described in a key for the identification of flies of forensic importance in Malaysia [16]. Interestingly, some characters observed in this study are morphologically similar to Hemilucilia segmentaria (F.), a blow fly species of the Central and South America. Such characters are the particular prominent tubercles along the peripheral rims of the eighth abdominal segment and the sculpture encircling the posterior spiracles (Figure 6 in [30]). Despite sharing morphological similarity in these characters, our molecular analysis of the COI gene (Figure 9) revealed that these two species are rather distance in genetic consideration, based on a genetic divergence of 8.6%.

Chrysomya chani is an endemic species in Bangladesh (Chittagong), India, China (Guangdong, Hainan I.), Indonesia (Kalimantan), Malaysia (Borneo; Sabah, Kuala Lumpur, Pahang, Perak,

Figure 9. Neighbour-joining tree based on COI gene (1 205 bp) of Chrysomya chani collected from the human remain from Thailand (pink dot) and other blow fly species. Voucher codes or accession number are presented in parentheses. Bootstrap values (>50%) were shown nearby the node of the trees.
Selangor), Nepal, Philippines (Luzon I., Mindanao I., Palawan I., Samar I.), Singapore, Sri Lanka, Thailand and Vietnam [31,32]. In northern Thailand, the habitat of this species is mainly natural forested areas at 335–1142 m above sea level [33], while in India, adults have also been collected from secondary forests [32]. Adults are found on decomposing animal matter in tropical rain forests [34]. Research in Selangor, Malaysia using monkey (Macaca fascicularis Raffles) carcasses indicates that C. chani occurs both in outdoor and indoor scenarios. Under outdoor conditions, adults were sampled from Days 6 to 13, indicating a preference for the decomposition stages of decay and advanced decay, while at indoor sites the species was sampled from Days 4 to 30, indicating a preference for bloated to advanced decay [35]. Assessment in Malaysia, using carcasses of the New Zealand White rabbit (Oryctolagus cuniculus (L.)), indicated that C. chani adults are mainly active during the bloated stage of decomposition (Days 2–3). Second and third larval instars were sampled during active decay (Days 3–5); while the third larval instar also was collected during the advanced stages of decay (Days 6–8) [36]. PMI_{min} estimation of six days in this case correlated with previous information. Interestingly, Omar et al. [37] showed that C. chani females prefer to oviposit not in the natural orifices of the animal carcasses but in the fur all over the bodies. In Shenzhen, China, C. chani was one of the main species, along with C. megacephala and C. rufifacies, when colonizing pig carcasses in summer [38].

Despite the limitations regarding biological information of C. chani, our findings may have an important implication for the use of this species in forensic investigations. Morphological details of the third instar and molecular analysis in the current study will improve identification success in future and finally elucidate its real forensic relevance. Developmental rates for the immature stages are strongly needed to be helpful in forensic investigations, particularly to estimate the PMI_{min}.

**Note**

1. Rearing of the third instar to adult for identification is recommended to differentiate from the closely related species, C. pinguis. Alternatively, a DNA analysis can be performed.

**Compliance with ethical standards**

This article does not contain any studies with human participants or animals performed by any of the authors.

**Disclosure statement**

No potential conflict of interest was reported by the authors.

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