Record of *Dicranosepsis javanica* (de Meijere, 1904) (Diptera, Sepsidae) from Meghalaya (India), with morphological and genetic data

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Abstract. The present article reports *Dicranosepsis javanica* (de Meijere 1904) (Diptera: Sepsidae) from Meghalaya and makes a note of its Oriental distribution range. The specimens were sampled in January 2014 by sweep net in the village of Schnongrim, Jaintia Hills, Meghalaya, situated in the north-eastern part of India. A short morphological description is provided for the male specimen, while the female taxonomic characters are presented for the first time. The taxonomic identification of species was confirmed by DNA Barcoding.

Key words: *Dicranosepsis javanica*; Diptera; Sepsidae; Meghalaya; India.

INTRODUCTION

The species belonging to Sepsidae family are found worldwide, having a wide distribution range (Ozerov, 2005). The adults can be found on dung, flowers, decomposed organic matter, manure, excrements, rotting fruits, and other decaying substrates (Ozerov, 2005). Given this feeding preference, these species are also known as the black scavenger flies. Concerning their ecological importance, sepsids have a significant role in the decomposition of animal dung, especially cow dung.

From the Oriental region, 100 Sepsidae species are known to date (Ozerov & Krivosheina, 2011), while in India this family comprises 45 species from 11 genera (Chakraborty et al., 2014). According to Ozerov (2003) and Iwasa & Thinh (2008, 2012), the genus *Dicranosepsis* Duda 1926 comprise nearly 37 species, being very well represented in the Oriental region, by 29 species. Furthermore, according to Ozerov (2003), the females of this genus are difficult to be differentiated. However, it is known that the males and the females of the same species are often found in the same breeding environment (Pont & Meier, 2002). *Dicranosepsis javanica* was first described by de Meijere in 1904, from Tosari (Indonesia). The species is also mentioned as being one of the widely distributed oriental species (Iwasa & Thinh, 2012). To date, the diagnosis of the species was performed only for males, the female of the species remaining undescribed do to the so far apparent lack of morphological differentiation. Numerous authors (Duda, 1926; Steyskal, 1966; Munari, 1982; Iwasa, 1984, 1986; Iwasa & Tewari, 1990; Iwasa & Jayasekera, 1994; Ozerov, 1996; Iwasa, 1999; Ozerov, 2003, 2005; Iwasa & Thinh, 2008, 2012; Ozerov & Krivosheina, 2011) have dealt with the study of this oriental genus, providing taxonomic key to species, revised key, new records, taxonomic and faunistic notes. Regarding the species distribution in the Oriental region, records were cited from China (Guangdong, Taiwan), India, Indonesia...
Lavinia IANCU, Khlur MUKHIM, Liviu MOSCALIUC

(Java I.), Myanmar, Nepal, Pakistan, Philippines (Luzon), Sri Lanka, Thailand, and Vietnam (Iwasa, 1984; Xiao & D. U. Yz, 2008; Letana, 2014).

MATERIAL AND METHODS

The specimens were sampled during an expedition in Shnongrim, Jaintia Hills, Meghalaya, India, in January 27, 2014. At the sampling time, an average temperature of 18°C and a relative humidity of 58% were recorded from the Meghalaya–Agricultural Meteorology Division (www.imdagrimet.gov.in/dwf/Meghalaya).

The sampling was performed by net sweeping several times in tall dry grass, in a human modified environment of clearings in secondary forest, few cattle and swine being present, at an altitude of 1159 m, with the GPS coordinates lat. 25.350° long. 92.516° (Fig. 1).

Taxonomic identification and DNA barcoding

The specimens were sorted and preserved in 75% ethanol, according to Gullan & Cranston (2010) recommendation. The male and female genitalia were photographed and drawn after being kept in lactic acid for several minutes. The taxonomic identification was performed under a stereomicroscope (Leica S6D), using Iwasa & Thinh (2008) and Letana (2014) identification keys.

The taxonomic identification was supported alongside by DNA barcode. The genomic DNA was extracted by using the DNeasy Blood & Tissue Kit and protocol (Qiagen) and its concentration was measured with a NanoDrop1000 (Thermo Scientific).

The PCR amplification of COI (Cytochrome c oxidase I) gene sequences was performed in a total volume of 50 µl, using a Mastercycler ProS System (Eppendorf). The pair of primers (10 pmols each) used in the reaction were: forward LCO1490: 5’-GGTCAACAAATCATAAAGATATTGG-3’ and reverse HC02198: 5’-TAAACTTCAGGGTGACCAAAAAATCA-3’ (Folmer et al., 1994). The PCR reaction was composed from 200 ng insect genomic DNA, 1 unit of Taq DNA polymerase, 1x Taq buffer, 2.5 mM MgCl2, and 0.1 mM dNTP (Thermo Scientific). The amplification initial incubation step was of 1 min at 94°C, followed by 5 cycles

Fig. 1 – Habitat of the sampled specimens.
of 30 sec at 94°C (denaturation), 1.5 min at 45°C (annealing) and 1 min at 72°C (extension), followed by 35 cycles of 30 sec at 94°C, 1.5 min at 51°C, 1 min at 72°C and a final extension step of 5 min at 72°C.

The DNA COI gene fragments (710 bp) were analysed by electrophoresis on 1% agarose gels and sequenced (Macrogen). The specimens are preserved in the “Grigore Antipa” National Museum of Natural History scientific collection with the voucher id 155045.

Sequence analyses

For the species genetic identification, the nucleotide sequences were analysed and quality trimmed with BioEdit Software (Hall, 1999) and compared for similarity within nucleotide databases by using BLAST-NCBI (http://www.ncbi.nlm.nih.gov).

The resulted COI gene sequence of 655-bp (female) and 612-bp (male) of D. javanica (Diptera: Sepsidae), were deposited in the GenBank database and the accession numbers [KX360168] and [KY564172] were assigned.

Abbreviations

The terminology is the one used by McAlpine (1981), as follows: head (oc, occellar setae; vti, inner vertical setae; vte, outer vertical setae); thorax (npl, notopleural setae; dc, dorsocentral setae; sa, supra alar setae; pa, postalar setae; m, mesopleural setae; ap sc, apical scutellar setae; pprn, postpronotal setae); abdomen (katep, katepisternum; aneps, anepisternum; anepm, anepimeron; mer, meron); legs (v, ventral seta, av, anteroventral seta); ovipositor (e, epiproct; c, cercus; s, sternite; t, tergite).

RESULTS AND DISCUSSION

According to the material available, the description and illustration of the morphological elements with taxonomic significance are presented below.

**Dicranosepsis javanica** (de Meijere, 1904)

**Male description and genetic confirmation** (Figs 2–4, 6 B, D)

The male of this species was described for the first time by Meijere, in 1904. Body wholly black and shining, measuring approximately 4 mm. (Fig. 2 A) Head. Round, with 10 peristomal setae disposed in the line, 1 oc, 1 vti, 1 vte, pvt vestigial, and the genal lower part without long wavy hairs. Wings. Hyaline, entirely irisated, with dark brown venation, and without dark spot near apex. R4+5 almost parallel with M1+2, no prominent posterior lobe on alula, upper and lower calypter whitish, margin of upper calypter with hairs, while the halters are whitish with the base darkened. (Fig. 2 B) Thorax. Anepimeron with a shining area and the katepisternum shining anteroventrally. Scutellum and postscutellum slightly dusted, 1 pprn, 2 npl, 2 dc, 1 sa, 1 pa, 1 ap sc present. Abdomen. Sternite 3, 4, and 5 oblong shape. The epandrial process is bifurcated at the tip. Legs. Middle and hind femur without distinct setae. Fore femur with 3–4 spines at the middle part, without av setae in the basal part (Fig. 3 A–C). Hind tibia (Fig. 3 F) with a blackish trace of osmeterium anterodorsally. Mid tibia without brownish ring at the distal end; fore tibia (Fig.3 D–E). The tarsomeres of the fore legs are cylindrical and the fore coxa is straight.

Regarding the genetic confirmation, BLAST analysis of D. javanica male sequence revealed 96% coverage, 99% identity and a maximum and total score of 1074 compared with the reference sequence [EU435787].

**Female description and genetic confirmation** (Figs 5, 6 A, C, 7 A–B).
Fig. 2 – Stereomicroscopic images of *Dicranosepsis javanica* male. (A) habitus, lateral view; (B) wing venation. Scale bars: 1 mm.

Fig. 3 – *Dicranosepsis javanica* male legs. (A, C) fore femur anterior view; (B) fore femur posterior view, details; (D) fore tibia anterior view; (E) fore tibia posterior view; (F) hind tibia with osmeterium–like trace. Scales (mm): A-B – 0.1; C-F – 0.5.
From the same sample, two female specimens were identified. The body length was of 2.9 mm (Fig. 5 A), and the wing length measured between 2.0 – 2.2 mm (Fig. 5 B).

Fig. 4 – *Dicranosepsis javanica* male. Hypopygium. (A) posterior view; (B) lateral view. Scale (mm): 0.4.

Fig. 5 – Stereomicroscopic images of *Dicranosepsis javanica* female. (A) habitus lateral view; (B) wing venation; (C) head, lateral view; (D) head, oblique view. Scales (mm): A-B – 1; C-D – 0.5.
Head. Round, with black, shining frons; face and gena brown-black, slightly dusted; facial orbits wider than in male; clypeus brownish; occiput black and dusted; antenna brown-black; arista black and bare; vibrissa with 3 setae; 12 peristomal bristles in a line disposed on the margin of gena, (Fig. 5 C, D).

Thorax. Black and partially dusted, with mesonotum slightly dusted, without bristles; meron slightly dusted; anepisternum shining; anepimeron dusted and slightly shining centrally, while in male is shining in the anterior half part; katepisternum dusted posterodorsally and shining anteroventrally; anatergite, katatergite, metanotum, metanepisternum, and metepimeron dusted; scutellum and postscutellum slightly dusted 1 pprn, 2 npl, 1 dc, 1 sa, 1 pa, 1 ap sc. Dusting pattern resembling male (Fig. 6 A, B).

Abdomen. With no distinct bristles, it is wholly black and shining; the 2nd tergite slightly narrowed anteriorly; sternite 3, 4, and 5 oblong shape as in male (Fig. 6 C, D): s3–4 pairs of bristles, s4–3 pairs of bristles, s5–2 pairs of bristles along the outer edge. Cercus small and cylindrical (Fig. 7 A). Epiproct present (Fig. 7 B). Hipoproct present, not sclerotized.

Legs. Yellow, with tarsomeres 3, 4, and 5 brownish; fore femur without distinct setae, but with small bristles; basitarsus of fore leg with one v seta in the proximal
first part; hind leg basitarsus with 2 av setae; fore, middle and hind coxa yellow and brownish in the proximal part.

Both male and female specimens were confirmed by DNA barcoding. For the female specimen, the BLAST analysis of the nucleotide sequences presented 94% cover, 99% identity and a maximum and total score of 1103 with the COI gene fragment of *D. javanica* [EU435787] registered in the online database.

The present article aimed to provide supplementary data on *D. javanica*, by adding morphologic and genetic information from this south-eastern part of Asia, straightening the current knowledge regarding this oriental species. In the present case, the specimens were identified taxonomically and confirmed genetically by DNA barcode.

The Oriental distribution of *D. javanica* seems to be somehow fragmented, missing reports from Cambodia, Bangladesh, and Laos (Fig. 8). Given the species importance as decomposers, as well as their ability of expanding the distribution range, due to human activities and livestock relocation, signaling their presence in different geographical regions constitutes an important aspect.

![Fig. 7 – Female ovipositor drawing at camera lucida. (A) dorsal view (e–epiproct, h–hypoproct, c– cercus, s– sternite); (B) lateral view. Scale (mm): 0.1.](image-url)
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Fig. 8 - Dicranosepsis javanica, Oriental region distribution map.
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