First detection of *Colletotrichum fructicola* (Ascomycota) on horsehair worms (Nematomorpha)

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Abstract

Fungal members of *Colletotrichum* (Ascomycota) were found to be associated with *Chordodes formosanus*, one of the three currently known horsehair worm (Nematomorpha) species in Taiwan. The fungi were identified as *Colletotrichum fructicola*, which is mostly known as a plant pathogen, through the use of the nuclear ribosomal internal transcribed spacer and partial large subunit (nrITS + nrLSU) and glyceraldehyde-3-phosphate dehydrogenase (GAPDH) DNA sequences. To our knowledge, this report represents both the first records for *Colletotrichum* associated with hairworms and for fungi on Nematomorpha. These findings expand the knowledge on the ecological relationships of both clades.

Keywords

Nematomorpha, Taiwan, *Colletotrichum*, horsehair worms, *Chordodes formosanus*, fungi
Introduction

The phylum Nematomorpha (which includes animals commonly known as “horsehair worms” or “hairworms”) is regarded as one of the most understudied animal groups, both from taxonomic and ecological perspectives (Schmidt-Rhaesa 2012). Most species have a complex life cycle that involves a larva encysting inside a freshwater intermediate host (i.e. usually an insect larva), a juvenile phase in which they parasitise terrestrial arthropods and a free-living adult freshwater stage (Schmidt-Rhaesa 2012, Bolek et al. 2015). However, some species have only freshwater hosts and a free-living freshwater adult stage (Schmidt-Rhaesa 2012). In addition to the freshwater ones, there are marine horsehair worms (all belonging to the genus Nectonema) that parasitise crustaceans as juveniles and live in surface seawater as adults (Schmidt-Rhaesa 2012, Kakui et al. 2021). Moreover, two recently-described Nematomorpha live in terrestrial wet environments in the adult phase (Anaya et al. 2019, Chiu et al. 2020).

Although we have some knowledge on Nematomorpha’s life history, there are very few studies on commensals, symbionts and parasites of hairworms. In addition, there are no reports of potential horsehair worm pathogens, both prokaryotic and eukaryotic, in literature (Schmidt-Rhaesa 2012, Bolek et al. 2015). The lack of data stems from two factors that make hairworms hard to observe: their generally reclusive behaviour (freshwater species tend to hide under rocks, fallen leaves and branches) and the absence of standardised protocols for sampling them. Moreover, few researchers study Nematomorpha due to their low medical and economical importance (Schmidt-Rhaesa 2012, Bolek et al. 2015).

Here we provide morphological and molecular evidence for the presence of fungi resembling Colletotrichum species (Ascomycota) living on and inside the body of Chordodes formosanus, one of the three described Taiwanese hairworm species (Chiu et al. 2020). The genus Colletotrichum mostly includes plant-associated (i.e. pathogen or endophytes) taxa with a broad host range. Some species also parasitise commercially-valuable crops (Cannon et al. 2012, Weir et al. 2012). However, species infecting sea turtles (Manire et al. 2002), cats (Winter et al. 2010), scale insects (Marcelino et al. 2008, Wynns et al. 2020) and humans (Cano et al. 2004, Lin et al. 2015) were described occasionally.

Materials and Methods

Four free-living adults of C. formosanus were collected in Wufengqi Waterfall area in Yilan County, Taiwan (24°49'59.6"N, 121°44'47.3"E) on 11 August 2020 (Suppl. material 1). After 10 days in a tank with a mixture of tap water and water collected from the collection site, fungi visibly started to develop on the hairworms. After one month and ten days, with the water replaced with tap water only, the fungi were widespread all over the worms’ cuticles (Fig. 1A and D). Despite this, three worms were alive at the time of fungal investigation.
The worms were investigated and two fungal structures (i.e. acervulus and perithecium) were dissected for further microscopic and molecular assessment (Fig. 1, Fig. 2 and Suppl. material 2). The investigation was performed with a dissecting microscope (Leica S9D) and a compound microscope (Nikon Eclipse N1). In addition, the regions of the worm with obvious fungal infection were sectioned with a tissue dissector (Leica CM3050 S Cryostat). Fungal perithecia were on and beneath the worm cuticle (Fig. 2). Asexual sporulating structures bearing conidia were found on the surface of the cuticle (Fig. 1D-F). Two of the hairworm specimens were deposited at the Biodiversity Research Museum, Academia Sinica, Taipei (collection IDs: ASIZ01000033 and ASIZ01000034).

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The aforementioned fungal structures were selected for DNA extraction and amplified with several universal primer sets for amplyfing four genes: ITS1F 5’ CTTGG TCATTTAGAGGAAGTAA 3’ and LR3 5’ CCGTGTTCAGACCGGG 3’ or ITS4 5’ TCCTCCGCTTATGATGTC 3’ (White et al. 1990, Vilgalys and Hester 1990), which targeted nuclear ribosomal internal transcribed spacer and partial large subunit (nrITS + nrLSU); GDF3 5’ GCCGTCACGACCCCTTCTTGA 3’ and GDR3 5’ TTCTCGTT GACACCCATCAGTAC 3’ (Chung et al. 2020) for targeting glyceraldehyde 3-phosphate dehydrogenase (GAPDH); CHS-79F 5’ TGGGGCAAGGATGCTTGGAAGAAG 3’ and CHS-345R 5’ TGGAAAGACCATCTGAGTTGTTG 3’ (Carbone and Kohn 1999) for chitin synthase (CHS-1); CL1C 5’ GAATTCAAGGAGGCTTCTTC 3’ and CL2C 5’ TTCTGATCATGAGGAG 3’ (Weir et al. 2012) for calmodulin (CAL). For DNA extraction, we placed tissues in Tris-EDTA (TE) buffer (50 µl) and stored them at -20°C.
Then, the frozen tubes were placed into an ultrasonic bath for 30 sec and in a thermal
cycler at 95°C for 10 min to break the cell wall.

PCR was undertaken by using Illustra™puReTaq Ready-To-Go PCR Beads (GE
Healthcare, United Kingdom) with 1 µl of forward and reverse primers (for a total of 2 µl), 2
µl of DNA and 21 µl of ddH₂O. The thermal cycler was set with an initial cycle at 94°C for 5
min, then 35 cycles with 94°C for 30 s, 52°C (ITS)/58 °C (other genes) for 1 min and 72°C
for 90 sec. Extension was done at 72°C for 10 min. The amplicons were sequenced by
both the forward and reverse primers.

The sequences derived from both directions were manually trimmed of the poor-quality
reads with MEGA X 10.1.8 (Kumar et al. 2018) and a consensus sequence was saved. The
sequences were submitted to NCBI GenBank with the following accession numbers: E5 =
MW714777, E6 = MW714778 for the ITS sequences; E5=MZ965243, E6=MZ965244 for
the GAPDH ones. CHS-1 and CAL were successfully amplified only for E6 and their
sequences have the following accession numbers: MZ965245 for CHS-1, MZ965246 for
CAL.

We then conducted a BLASTn search (Altschul et al. 1990) with default settings for finding
similar sequences in GenBank. After identifying a genus with high degree of sequence
similarity through BLASTn, we retrieved sequences of species inside that clade from
GenBank (Suppl. material 3) and we reconstructed a phylogenetic tree. Specifically,
sequences alignment was performed using the L-INS-i algorithm in MAFFT 7.471 (Katoh
and Standley 2013) and Maximum Likelihood phylogenies for concatenated genes were subsequently reconstructed using ModelTest and RAxML-NG implemented in raxmlGUI (Edler et al. 2021). Gene concatenation was undertaken by using SequenceMatrix (Vaidya et al. 2011). The trees were visualised with FigTree 1.4.4 (Rambaut 2018).

Results and Discussion

The fungus E6 (Fig. 3) had obpyriform perithecia, with colours ranging from black to brown, paler towards the ostiolar neck, without hairs (Fig. 1A-C). Asci were unitunicate with nonamyloid apex, with hyaline and unicellular ascospores, around 15 μm long (Fig. 1C). These features represent the sexual reproductive structures of the fungal genus *Glomerella* which is regarded as the sexual state of genus *Colletotrichum* (Cannon et al. 2012). The other fungus E5 appeared to be at its asexual stage and produced white acervuli bearing hyaline conidia (Fig. 1D-F). The phylogenetic results of the concatenated ITS and GAPDH tree showed that these fungi were *Colletotrichum fructicola* individuals (Fig. 3; Suppl. material 4), which is a taxon belonging to the *Colletotrichum gloeosporioides* species complex (Weir et al. 2012).

![Maximum Likelihood phylogenetic tree built by using concatenated ITS and GAPDH sequences. Specimens E5 and E6 were collected for this study and are emphasised in bold and red font. Bootstrap values ≥ 70 are shown. The names of species complexes are shown on the right. Strain number for sequences taken from GenBank are shown. Strains with the * mark are the ex-type strains. Accession numbers for the gene sequences used are available in Suppl. material 3. A picture of the species tree made with concatenated ITS, GAPDH, CHS-1 and CAL genes with specimen E6 is present in Suppl. material 4.](image)

In Taiwan, *Colletotrichum* species are mostly known for causing anthracnose in different kind of plantations (Sun et al. 2019, Damm et al. 2020, Wu et al. 2020, Chung et al. 2020). There is also a reported cutaneous infection on a human caused by *C. gloeosporioides* (Lin et al. 2015). From what concerns *C. fructicola*, it has been recognised as a pathogen of strawberry, mango and tea in the Island (Wu et al. 2020, Chung et al. 2020, Lin et al. 2021), but it has also been reported on other crops worldwide (Weir et al. 2012).
Fungi in the phylum Ascomycota are known to be resilient and they can pass from soil to aquatic environments (Jessup et al. 2004, Rypien et al. 2008, Sarmiento-Ramírez et al. 2010, Fisher et al. 2012); this trait is also present in *Colletotrichum* species, which have been reported both from seawater and freshwater organisms (Smith et al. 1989, Manire et al. 2002). In addition to this, all the horsehair worms are known not to feed in the adult stage (Schmidt-Rhaesa 2012). Non-feeding may make the hairworm hosts weaker as time goes by and allow the opportunistic fungi to grow on their cuticle. Further studies will be needed to determine the prevalence of *Colletotrichum* in the wild, apparently healthy Nematomorpha populations and if hairworms can contribute to the spread of the ascomycetous fungi to other organisms, as happens with other related fungal clades (Fisher et al. 2012). Given the possible arising of new diseases from opportunistic pathogens due to environmental change (Nnadi and Carter 2021) and to further understand the chronological order of infection and the pathogenicity of *Colletotrichum* on hairworms, inoculation experiments (for proving Koch’s postulate) combined with histological examination will be required. Besides these considerations, to our knowledge, this is the first report of fungi on horsehair worms. In addition, our report increases the already broad host range of the genus *Colletotrichum*.

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**Author contributions**

Mattia De Vivo (MDV) collected the animals and performed PCR on fungineal DNA. MDV and Ko-Hsuan Chen (KHC) analysed the data and wrote the manuscript. KHC, Wen-Hong Wang (WHW) and Jen-Pan Huang (JPH) designed the methodology. WHW extracted the fungi and their DNA and performed worms’ dissections. MDV, KHC and JPH conceived and coordinated the study. All authors contributed critically to the drafts and gave final approval for publication.
Conflicts of interest
The authors declare that they have no conflict of interest.

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Supplementary materials

Suppl. material 1: Free living specimens [doil]

Authors: Mattia De Vivo
Data type: Image
Brief description: Free living specimens of Chordodes formosanus, before the fungi started to be visible
Download file (3.51 MB)

Suppl. material 2: Further cross sections [doil]

Authors: Wen-Hong Wang and Ko-Hsuan Chen
Data type: Multimedia (PDF)
Brief description: Additional cross sections of the worms
Download file (126.78 kb)

Suppl. material 3: Accession list of the sequences used in this study [doil]

Authors: Mattia De Vivo
Data type: GenBank accession numbers (csv file)
Brief description: Accession numbers of the sequences used for this study
Download file (882.00 bytes)
Suppl. material 4: 4 genes tree

Authors: Mattia De Vivo
Data type: Tree (image)
Brief description: Picture of the concatenated phylogenetic tree based on all the sequenced genes (ITS, GAPDH, CHS-1 and CAL). Bootstrap values ≥ 70 are shown.
Download file (16.63 kb)