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1 First report of marine horsehair worms (Nematomorpha: Nectonema) parasitic in 2 isopod crustaceans 3 Keiichi Kakui<sup>1</sup> · Jun Fukuchi<sup>2</sup> · Daisuke Shimada<sup>1</sup> 4 5 6 <sup>1</sup>Faculty of Science, Hokkaido University, Sapporo 060-0810, Japan 7 <sup>2</sup>Seto Marine Biological Laboratory, Kyoto University, Nishimuro 649-2211, Japan 8 Corresponding author: K. Kakui. e-mail: kakui@eis.hokudai.ac.jp 9 ORCID (K. Kakui): 0000-0003-4630-9065 10 ORCID (J. Fukuchi): 0000-0001-6857-9723 11 ORCID (D. Shimada): 0000-0002-6913-3659 12 13 **Abstract** 14 *Nectonema*, the only horsehair worm (Nematomorpha) genus found in marine environments, 15 was previously known to be parasitic only in decapod crustaceans. We report *Nectonema* sp. 16 as the first record of a marine nematomorph parasitic in isopod crustaceans. This is also the 17 third record of marine nematomorphs from the North Pacific. Six infected isopods 18 (Natatolana japonensis) collected from 1425 m depth in the Sea of Japan each contained one 19 to seven (mean 2.33) nematomorphs in the body cavity in the pereon. There was no 20 correlation between the host body length and number of parasites. For *Nectonema* sp., we 21 describe and illustrate morphological features of the parasitic juvenile stage and present 22 nucleotide sequences for the cytochrome c oxidase subunit I gene (COI or cox1; 451 nt), 18S 23 rRNA gene (1777 nt), and region spanning the internal transcribed spacer 1 (ITS1) and the 24 28S rRNA gene including the 5.8S rRNA gene and ITS2 (1218 nt in total). In an 18S 25 maximum-likelihood tree that included 24 nematomorph species, Nectonema sp. grouped 26 with N. agile from the northwestern Atlantic; the 18S gene from these two taxa was divergent 27 by 11.8% K2P distance, suggesting they are different species. *Nectonema* species may have a 28 broader range of host groups than previously suspected, but may have been previously 29 misidentified as nematode parasites. 30 31 **Keywords** Cymothoida · Deep sea · Endoparasite · Hairworm · Nectonematida · Peracarida 32

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- 35 Nectonema is the only nematomorph genus found in marine environments. There are five
- 36 species described in this genus to date: *Nectonema agile* Verrill, 1879 (type species) from
- 37 Atlantic Ocean and Mediterranean and Black Seas; *Nectonema melanocephalum* Nierstrasz,
- 38 1907 from Indonesia; *Nectonema munidae* Brinkmann, 1930 from Bergen and Norway;
- 39 Nectonema svensksundi Bock, 1913 from Spitzbergen, Norway; and Nectonema zealandica
- 40 Poinar and Brockerhoff, 2001 from New Zealand (Schmidt-Rhaesa et al. 2013). The
- occurrence records of this group in the North Pacific have been limited, with only two reports
- from Japan: Oku et al. (1983) and Yoshida (2016) reported unidentified *Nectonema*
- 43 individuals from the brachyuran *Erimacrus isenbeckii* (Brandt, 1848) and the anomuran
- 44 Pagurus brachiomastus (Thallwitz, 1891), respectively.
- The life cycle of nematomorphs contains a larval stage, a parasitic juvenile stage,
- and a free-living adult stage (Hanelt and Janovy 2004). In *Nectonema*, juveniles have so far
- been reported only from crustaceans in the order Decapoda (shrimps, hermit crabs, crabs,
- etc.), among which more than 27 species have been reported as hosts (Schmidt-Rhaesa et al.
- 49 2013). In a single host species, *Nectonema* individuals at various developmental stages can be
- found (Huus 1932; Schmidt-Rhaesa 1996).
- Here we report the first record of *Nectonema* (as *Nectonema* sp.) parasitic in a
- 52 species in the order Isopoda, a morphologically diverse crustacean group with more than
- 53 10,000 described species. This is also the third record of marine nematomorphs from the
- North Pacific. We describe the morphology of this nematomorph and present nucleotide
- sequences for its cytochrome c oxidase subunit I (COI) gene, 18S rRNA (18S) gene, and ITS
- cluster, including the 3' region of internal transcribed spacer 1 (ITS1), the 5.8S rRNA gene,
- 57 ITS2, and the 5' region of the 28S rRNA gene.

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## Materials and methods

- 60 Isopods were collected from plastic jars that contained cut sardine bait and had been placed in
- 61 baited traps (Saito et al. 2014: fig. 1J) during a cruise of R/V Soyo-maru (National Research
- 62 Institute of Fisheries Science, Japan); the traps were recovered on 16 July 2014 at station
- 63 Kago-4 (40°00.59′ N 135°57.63′ E), 1425 m depth, Sea of Japan. The isopods were picked
- 64 from the jars by Ken Fujimoto and kept alive at 4°C until 30 July. Infected individuals were
- anesthetized with 35% MgCl<sub>2</sub> and dissected to extract nematomorphs. One infected isopod
- and one extracted nematomorph were photographed live. Isopods were fixed and preserved in

ethanol; nematomorphs removed from them were fixed in DESS solution (Yoder et al. 2006) or 99% ethanol.

Body length (BL) was measured from the anterior edge of the cephalothorax to the tip of the pleotelson for the isopods, and from the anterior to posterior tip of the body for the nematomorphs; the cephalothorax width (CW) of the isopods was measured at the widest portion of the cephalothorax.

Isopods were dissected with needles under a Nikon SMZ1500 stereomicroscope; detached appendages were mounted on glass slides in glycerin and observed with a Nikon E600 microscope. Two nematomorphs in DESS solution were transferred into a 1:3:6 mixture of glycerin, absolute ethanol, and deionized water and placed in a thermostatic chamber at 40°C for two days, after which they were mounted on glass slides in glycerin and observed with an Olympus BX51 microscope. Illustrations of nematomorphs were prepared with Inkscape 1.0 from digital micrograph images. Morphological terminology for *Nectonema* here follows Poinar and Brockerhoff (2001).

Total DNA was extracted from part of the body from each of four nematomorphs by using a NucleoSpin Tissue XS Kit (TaKaRa Bio, Japan). Primers used for PCR and sequencing are listed in Table 1. PCR amplification conditions for the ITS cluster and COI with TaKaRa Ex Taq DNA polymerase (TaKaRa Bio) were 94°C for 1 min; 35 cycles of 98°C for 10 s, 50°C for 30 s, and 72°C for 1 min; and 72°C for 2 min. Conditions for 18S amplification with KOD FX Neo (Toyobo, Japan) were 94°C for 1 min; 45 cycles of 98°C for 10 s, 65°C for 30 s, and 68°C for 75 s; and 68°C for 3 min. PCR products for 18S were separated on a 2% agarose gel, excised with a micro spatula, and purified with a MagExtractor PCR & Gel Clean Up Kit (Toyobo). All nucleotide sequences were determined by direct sequencing with a BigDye Terminator Kit ver. 3.1 and a 3730 DNA Analyzer (Life Technologies, USA). Fragments were concatenated by using MEGA7 (Kumar et al. 2016).

The 18S dataset for a phylogenetic analysis included our 18S sequence from *Nectonema* sp. and 25 sequences from 23 nematomorph species and two outgroup taxa (a nematode and a tardigrade) taken from public databases (DDBJ 2021; Online Resource 1). Methods for alignment of all 18S sequences (1806 positions in the aligned dataset; Online Resource 2) and selection of the optimal substitution model (GTR + I + G) were as described by Homma et al. (2020). The Kimura (1980) 2-parameter (K2P) distance between the aligned *Nectonema* sequences was calculated with MEGA7. A maximum likelihood (ML) analysis was conducted in RAxML-NG (Kozlov et al. 2019), with nodal support values obtained by

analysis of 1000 bootstrap pseudoreplicates. The ML tree was drawn by using FigTree v1.4.4 (Rambaut 2021).

Our nematomorph specimens were deposited in the Invertebrate Collection of the Hokkaido University Museum (ICHUM), Sapporo (ICHUM-6178–6191); host isopods were deposited in the Seto Marine Biological Laboratory (SMBL-V0598–0603). The sequences we determined were deposited in the International Nucleotide Sequence Database through the DNA Data Bank of Japan, under the accession numbers LC605980–605988.

The six infected isopods we observed (three males, BL 14.7–15.8 mm; three females, BL

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## **Results and discussion**

13.5–16.9 mm) were identified as *Natatolana japonensis* (Richardson, 1904) in the suborder Cymothoida (Fig. 1). All infected individuals harbored nematomorphs internally in the cavity of the pereon. The nematomorph infections ranged from one to seven individuals per host (mean intensity = 2.33) (Table 2). The prevalence of infection is unknown because we lack data on the total number of isopods in the baited traps. No clear correlation was detected between host size (BL) and the number of parasites (N) (N =  $1.51 \times BL - 20.6$ ; R<sup>2</sup> = 0.543). The nematomorphs ranged in BL from 23.7 mm to 94.0 mm (mean, 58.3 mm; N = 12 intact individuals; Online Resource 3). An abbreviated description of individual ICHUM-6181 (Fig. 2) is as follows. Anterior and posterior ends rounded. Cuticle colorless. Epidermis with single cell layer. Cuticular natatory bristles not observed. Cephalic papillae absent. Septum distinct. Anterior chamber translucent, with ca. 10 giant cells. Anterior region not pigmented. Mouth opening at anterior apex. Oral cavity with eight (?) bifurcated hooks protruding from cuticular wall. Sclerotized proboscis with two anterior scissor-shaped spines. Pharynx sclerotized, connecting with intestine posterior to septum. Body filled with mesenchyme between septum and subposterior region. Gonads not observed. Subposterior region with posterior opening (gonopore?) surrounded by elongate epidermal cells. COI and ITS-cluster sequences were determined from four nematomorphs (ICHUM-6182-6184, 6188). The four COI sequences (accession numbers LC605980-605983; 451 nt long, translating to 150 amino acids) were identical, as were the four ITS-cluster sequences (accession numbers LC605984-605987; 1218 nt long). No Nectonema COI and ITS-cluster sequences have previously been deposited in public databases (DDBJ 2021). The 18S sequences (accession numbers LC605988, 605989) determined from two nematomorphs (ICHUM-6184, 6188; 1777 nt long) were identical, and were 11.8% divergent in K2P distance from the only *Nectonema* 18S sequence available in public databases (N. agile; Bleidorn et al. 2002). In the ML tree for 18S (Fig. 3), Nectonema sp. and N. agile formed a clade with 100% bootstrap support, which was the sister group to a Gordiida clade having 87% bootstrap support; the relationships within Gordiida differed from those in previous studies (e.g., Bleidorn et al. 2002; Tobias et al. 2017), which may have resulted from the difference in the dataset used. Morphology-based species identification is difficult in *Nectonema*, and Schmidt-Rhaesa (2005) noted that the five known species may have been described based on specimens at different developmental stages, making the descriptions not directly comparable. It should be noted that our *Nectonema* samples differ from juveniles of three species (*N. agile*, N. munidae, and N. zealandica; no juveniles have been reported in the other two species) in having ca. 10 giant cells and lacking cephalic papillae. Molecular identification is currently unavailable for this group because no molecular markers have been determined for all named Nectonema species. Identification of our material to species was not possible, as no adult specimens were obtained. On the basis of the 11.8% K2P divergence, *Nectonema* sp. is likely not conspecific with N. agile. Furthermore, its occurrence in an isopod indicates that Nectonema may use other, undetected groups besides Decopoda for hosts. In the parasitic stage, nematomorphs resemble nematodes, and infections of unusual hosts in the past might have been misidentified as nematode infections. To understand true diversity of marine nematomorphs, host surveys targeting non-decapods as well as decapods and integrative taxonomic approaches will be necessary. Acknowledgments We thank Ken Fujimoto, Shiro Sawadaishi, and the crew of R/V Soyo-maru for collecting specimens and providing facilities; Matthew H. Dick for reviewing the manuscript and editing our English; and two anonymous reviewers for improving this manuscript with their

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166	Competing interests
167	We declare no competing interests.
168	
169	Author contributions
170	KK conceived and designed the study, collected samples, and conducted the molecular
171	analysis. JF identified the host isopods. KK and DS made morphological observations of the
172	nematomorphs. DS made drawings. KK wrote the first draft of the manuscript, and all
173	authors commented on the first draft and read and approved the final draft.
174	
175	Supplementary information
176	This work includes the following online supplementary information.
177	Online Resource 1. Species included in the phylogenetic analysis based on 18S
178	rRNA sequences.
179	Online Resource 2. Aligned sequences used for phylogeny reconstruction.
180	Online Resource 3. Details of the animals studied.
181	
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261 Fig. 1 Nectonema sp. parasitic on Natatolana japonensis. a, b N. japonensis containing 262 Nectonema sp., dorsal and ventral views, fresh specimen (SMBL-V0598). c Nectonema sp., 263 fresh specimen (ICHUM-6178) 264 265 Fig. 2 Nectonema sp. (ICHUM-6181). Anterior (a, b, e, f) and posterior (c, d) portions and 266 sclerotized proboscis with two anterior scissor-shaped spines (b') of Nectonema sp., in 267 glycerin (ICHUM-6181), as microphotographs (a, c, e, f) and line drawings (b, b', d). Arrowheads in e indicate giant cells. Abbreviations: ac, anterior chamber; bh, bifurcated 268 269 hook; cu, cuticle; du, duct (gonoduct?); ep, epidermis; gc, giant cell; in, intestine; me, 270 mesenchyme; mo, mouth opening; mu, muscle; ph, pharynx; po, posterior opening 271 (gonopore?); pr, proboscis; se, septum; sp, scissor-shaped spine 272 273 Fig. 3 ML tree for 18S sequences (1806 positions) from nematomorphs, including *Nectonema* 274 sp. from isopods. Values near nodes are bootstrap values ≥ 80%; black circles indicate 100% 275 bootstrap support. Outgroup taxa (the tardigrade Milnesium tardigradum and the nematode 276 Deontostoma magnificum) are not shown 277

**Table 1**. List of PCR and cycle sequencing (CS) primers used in this study.

Gene	Primer	Sequence	Reaction	Source
COI	F1	CCTACTATGATTGGTGGTTTTTGGTAATTG	PCR	Kanzaki and Futai (2002)
	R2	GTAGCAGCAGTAAAATAAGCACG	PCR, CS	Kanzaki and Futai (2002)
18S	SR1	TACCTGGTTGATCCTGCCAG	PCR	Nakayama et al. (1996)
	SR12	CCTTCCGCAGGTTCACCTAC	PCR	Nakayama et al. (1996)
	18S-b3F	CCTGAGAAACGGCTACCACAT	CS	Kakui and Shimada (2017)
	18S-b4F	TGCGGTTAAAAAGCTCGTAGTTG	CS	Kakui et al. (2011)
	18S-b4R	TCCAACTACGAGCTTTTTAACC	CS	Kakui et al. (2011)
	18S-b5F	GATCGAAGGCGATYAGATACC	CS	This study
	18S-b6F	CCTGCGGCTTAATTTGACTC	CS	Kakui et al. (2011)
	18S-a6R	AACGGCCATGCACCAC	CS	Kakui et al. (2011)
	18S-b8F	GGTCTGTGATGCCCTTAGATG	CS	Kakui et al. (2011)
ITS cluster	NC5	GTAGGTGAACCTGCGGAAGGATCATT	PCR, CS	Zhu et al. (1998)
	NC2	TTAGTTTCTTTTCCTCCGCT	PCR, CS	Zhu et al. (1998)
	NC13	ATCGATGAAGAACGCAGC	CS	Zhu et al. (1998)

**Table 2.** Size of isopod host (*Natatolana japonensis*) and nematomorph (*Nectonema* sp.) individuals. All measurements are in millimeters. BL, body length; CW, cephalothorax width. SMBL, Seto Marine Biological Laboratory.

Isopods				N	Iematomorphs
SMBL-	Sex	BL	CW	No. of parasites	BL
V0598	Female	16.9	3.8	7	26.9–94.0 (mean 58.1; n = 6*)
V0599	Male	14.7	3.4	2	40.2–84.9 (mean 62.6; n = 2)
V0600	Male	15.8	3.8	1	87.8
V0601	Female	14.9	3.3	2	23.7–68.6 (mean 45.2; n = 2)
V0602	Female	13.5	3.2	1	46.0
V0603	Male	15.1	3.6	1	42.1**

\*One damaged specimen (length 50.1 mm; part of body missing) excluded. \*\*Data from damaged specimen lacking part of body.





