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Title	Diexanthema hakuhomaruae sp. nov. (Copepoda: Siphonostomatoida: Nicothoidae) from the Hadal Zone in the Northwestern Pacific, with an 18S Molecular Phylogeny
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2	zone in the northwestern Pacific, with an 18S molecular phylogeny
3	
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### 31 Abstract

32	Purpose Diexanthema copepods are ectoparasites on deep-sea isopods. This genus currently contains six
33	species, all reported from the North Atlantic. Our study describes a new species of Diexanthema found on
34	isopods from 7184–7186 m depth in the Kuril-Kamchatka Trench, northwestern Pacific.
35	Methods We observed the copepod's morphology, made camera-lucida drawings, and compared our
36	species with congeners. We determined partial sequences for its 16S rRNA and 18S rRNA genes and
37	constructed an 18S-based maximum-likelihood copepod tree to place it phylogenetically. We identified the
38	host isopod species through morphology and cytochrome $c$ oxidase subunit I (COI, cox1) and 18S
39	sequences.
40	Results/conclusion We described the copepod as <i>Diexanthema hakuhomaruae</i> sp. nov. and identified its
41	host as Eugerdella cf. kurabyssalis Golovan, 2015 (Desmosomatidae). This is the first Diexanthema
42	copepod from the Pacific and also from hadal depths. Diexanthema hakuhomaruae most closely resembles
43	D. bathydiaita Richie, 1975, parasitic on Nannoniscus sp. (Nannoniscidae) in the Atlantic, but differs from
44	the latter in having a smooth body surface and leg 5 in the ventrolateral region of the urosome. In the 18S
45	tree, D. hakuhomaruae was the sister group to the Rhizorhina clade, which is consistent with the
46	morphology-based hypothesis that they are closely related.
47	

48 Keywords Crustacea · ectoparasite · integrative taxonomy · Japan · parasites

### 49 Introduction

50	Nicothoidae is a family of parasitic copepods, with 22 genera and about 140 species [1]. Nicothoids utilize
51	various crustaceans as hosts, including Ostracoda, Leptostraca, Decapoda, Amphipoda, Cumacea, Isopoda,
52	Mysida, and Tanaidacea [2]. Nicothoid morphology is highly diverse. Rhizorhina copepods, for example,
53	have an unsegmented spherical body, whereas Paranicothoe copepods have a multi-segmented body
54	consisting of a prosome and urosome. Recent molecular phylogenetic analyses have called into question
55	the monophyly of this family [3, 4].
56	The nicothoid genus Diexanthema is characterized by the following female features: a spherical
57	prosome lacking a head process; an unsegmented urosome; antennules with two or fewer articles;
58	maxillipeds absent; and caudal rami shorter than the urosome, or absent [1, 5–7]. It currently contains six
59	named species, Diexanthema apoda Boxshall and Harrison, 1988, Diexanthema bathydiaita Richie, 1975,
60	Diexanthema corrugatum Boxshall and Harrison, 1988, Diexanthema desistoma Richie, 1975,
61	Diexanthema nudum Boxshall and Harrison, 1988, and Diexanthema ritchiei Boxshall and Harrison, 1988,
62	all of which are parasitic on deep-sea isopods in the North Atlantic Ocean [8, 9] (Table 1). Boxshall and
63	Lincoln [10] placed this genus in the "Rhizorhina group," along with the genera Rhizorhina and
64	Choniorhiza. The same authors proposed two other groups in Nicothoidae, the "Nicothoe group" and the
65	"Sphaeronella group," containing five and eight genera, respectively. The validity of this grouping has not
66	yet been tested with molecular data.

67

In 2022, we collected an undescribed Diexanthema species parasitic on a desmosomatid isopod

68	from the hadal zone in the Kuril-Kamchatka Trench, northwestern Pacific Ocean. This is the first
69	Diexanthema copepod from outside the North Atlantic, and the first from hadal depths. Here we describe
70	this species, provide partial sequences for multiple genes to aid future DNA barcoding, and infer its
71	phylogenetic position in Siphonostomatoida based on an 18S-rRNA (18S) tree to test the validity of the
72	"Rhizorhina group."
73	
74	Materials and methods
75	Five copepods, each from a different host individual, were collected on 6 October 2022 during cruise KH-
76	22-8 of R/V Hakuho-maru (Japan Agency for Marine-Earth Science and Technology; JAMSTEC), with
77	small plankton nets attached inside an Agassiz trawl (cf. fig. 2B in Akiyama et al. [11]), at depths of 7184-
78	7186 m. Copepods were attached to the pereonite-3 sternite or the basis of pereopods 3, 4, or 6 of the host
79	isopods. They were photographed and then fixed and preserved in 80% ethanol.
80	Three copepods were detached from their host with chemically sharpened needles, and two were
81	retained intact on the host for future non-destructive observation or molecular analysis. Two of the three
82	detached copepods were used for morphological observations and one for DNA extraction (see below).
83	The former were transferred through an ethanol series (70, 60, 50, 40, and 30% ethanol, each step for ca. 5
84	min) at room temperature, mounted on cavity slides (T8-R004; Toshin Riko, Japan) in 30% ethanol, and
85	observed with an Olympus BX53 microscope. Illustrations were prepared with Adobe Illustrator CS6 from
86	draft line drawings made with a camera lucida. In copepods, body length (BL) was measured from the

87	anterior to posterior ends of the body (prosome + urosome), and prosome width (PW) and urosome width
88	(UW) at the widest portion of the prosome and urosome, respectively. In host isopods, BL (from the
89	anterior edge of the cephalothorax to the tip of the pleotelson) and the pereonite-2 width (P2W) were
90	measured. All measurements are presented in the text in micrometers unless noted otherwise. The
91	specimens studied were deposited in the Invertebrate Collection of the Hokkaido University Museum
92	(ICHUM), Sapporo, Japan, under catalog numbers ICHUM8451–ICHUM8455.
93	DNA was extracted from the whole body of one copepod and percopod 1 of one host by using the
94	NucleoSpin Tissue XS Kit (Macherey-Nagel, Germany). For the cytochrome c oxidase subunit I (COI)
95	gene, PCR primers used for the amplification and cycle sequencing were LCO1490 and HCO2198 [12].
96	For the 18S rRNA gene, amplification primers were SR1 and SR12 [13], and six primers (18S-b3F, 18S-
97	b4R, 18S-b5F, 18S-b6F, 18S-a6R, and 18S-b8F [14, 15]) were used in cycle sequencing. For the copepod
98	16S rRNA gene, the newly designed primers Copepod16S_F (CGCCTGTTTATCAAARACWY) and
99	Copepod16S_R (TCGATTTGAACTCAAATCAWG) were used for amplification and cycle sequencing.
100	PCR amplification conditions for COI with TaKaRa Ex Taq DNA polymerase (TaKaRa Bio, Japan) were
101	as described by Munakata et al. [16]; those for 18S with KOD FX Neo (Toyobo, Japan) were as described
102	by Okamoto and Kakui [17]; and those for 16S with KOD ONE PCR Master Mix (Toyobo) were 45 cycles
103	of 98°C for 10 s, 50°C for 5 s, and 68°C for 1 s. PCR products for 16S were separated on a 2% agarose gel,
104	excised with a micro spatula, and purified with the MagExtractor PCR & Gel Clean Up Kit (Toyobo). All
105	nucleotide sequences were determined with a BigDye Terminator Kit ver. 3.1 and a 3730 DNA Analyzer

106	(Life Technologies, USA). Fragments were concatenated by using MEGA7 [18]. The sequences we
107	determined were deposited in the International Nucleotide Sequence Database (INSD) through the DNA
108	Data Bank of Japan.
109	The copepod 18S dataset for a phylogenetic analysis comprised the copepod dataset from Kakui
110	and Munakata [4] and the one Diexanthema sequence we determined, representing 50 siphonostomatoid
111	species and one outgroup taxon (Misophthriopsis okinawensis, a misophrioid). The sequences were aligned
112	(1528 positions in the aligned dataset; see Online Resources 1 and 2) as described by Munakata et al. [19];
113	methods for selecting the optimal substitution model (GTR+F+R3), the maximum likelihood (ML)
114	analysis, and drawing the tree were as described by Kakui and Shimada [20].
115	
116	Results and Discussion
117	Host identification
118	All host individuals were females with developing oostegites. An abbreviated description of their
119	morphology is as follows. Pereonite 1 slightly longer than pereonite 2. Pleotelson with small posterolateral
120	spines. Coxae I-II each with acute anterior projection (longer on coxa I longer than on coxa II). Pereopod
121	1 stout: ischium with five robust distodorsal, unequally bifid setae; carpus enlarged, ventral margin convex
122	in proximal two thirds but straight in distal third, with minute distal, unequally bifid, distally setulate seta
123	
	(UBDS) at base of penultimate seta, and row of four robust UBDS of irregular size. Operculum truncate,

125	According to these character states, our individuals were Eugerdella kurabyssalis Golovan, 2015,
126	described from the Northwest Pacific Basin east of the Kuril-Kamchatka Trench [21] and later reported
127	also from the Kuril-Kamchatka Trench [22]. Jennings et al. [22] molecularly detected Eugerdella cf.
128	kurabyssalis Golovan, 2015 among Eugerdella specimens collected from the Kuril-Kamchatka Trench. It
129	closely resembles <i>E. kurabyssalis</i> but shows minor differences from the latter, such as a smaller, less
130	pronounced coxa [22], and has been collected from deeper depths than E. kurabyssalis (E. cf. kurabyssalis
131	from 7081–7123 m; <i>E. kurabyssalis</i> from 4830–6051 m [21, 22]). In a BLAST search [23] of the public
132	database, the partial COI sequence (LC741552; 655 bp long) we determined for one host individual
133	(ICHUM8452) was most similar to the COI sequence from E. cf. kurabyssalis (MN179516; query cover
134	91%, identity score 99.83% [22]). Our isopods were collected from 7184–7186 m depth, similar to the
135	depth range reported for <i>E</i> . cf. <i>kurabyssalis</i> . Although we could not judge whether the coxa in our
136	specimens is smaller and less pronounced than in <i>E. kurabyssalis</i> , given the similarity in COI sequence and
137	sampling depth, we concluded that the host isopods were E. cf. kurabyssalis.
138	
139	Taxonomy
140	<i>Diexanthema hakuhomaruae</i> sp. nov.
141	(Figs. 1, 2)
142	

143 Diagnosis (females). Body smooth, lacking furrows or minute hairs; anterior hood absent; caudal ramus

145	absent; semicircular irregular branching structure present; rootlet absent; mandible present; legs 1-4
146	absent; leg 5 present, small lobe shape, with three spiniform setae in ventrolateral region of urosome.
147	Type host. Eugerdella cf. kurabyssalis Golovan, 2015 (Isopoda: Asellota: Desmosomatidae).
148	Type locality. Station A5, Kuril-Kamchatka Trench axis, northwestern Pacific (41°14.024' N
149	145°01.931' E to 41°14.126' N 145°01.108' E), 7184–7186 m depth.
150	Attachment sites. Pereopod basis and pereon sternite.
151	Material examined. Holotype: female (ICHUM8451), BL 356, PW 346, UW 86, one vial
152	containing extracted copepod and host (BL 1797, P2W 483). Paratypes: four females (ICHUM8452, BL
153	397, PW 359, UW 96, used in DNA extraction, one slide and one vial containing host [BL 1925, P2W
154	489]; ICHUM8453, BL 342, PW 326, UW 87, one vial containing extracted copepod and host [BL 1801,
155	P2W 475]; ICHUM8454, BL 365, PW 385, UW 89, one vial containing copepod attached to host [BL
156	1984, P2W 499]; ICHUM8455, BL 400, PW 366, UW 95, one vial containing copepod attached to host
157	[BL 1820, P2W 490]). All specimens were collected at the type locality on 6 October 2022 by R/V
158	Hakuho-maru.
159	Representative DNA sequences. One 16S (INSD accession number LC741550; 424 bp long) and
160	one 18S (LC741551; 1762 bp long) sequences were determined from paratype female ICHUM8452. One
161	COI (LC741552; 655 bp long, encoding 218 amino acids) and one 18S (LC741553; 2174 bp long)
162	sequences were determined from host female ICHUM8452.

Etymology. The specific name (a noun in the genitive case) is from R/V Hakuho-maru, the vessel

- 164 from which the type specimens were collected.
- 165 Description (female, based on the holotype). Body (Figs. 1, 2) transparent (white in ethanol),
- 166 smooth, lacking furrows or minute hairs; prosome globular but slightly flattened dorsoventrally, containing
- 167 bulging ovaries (Fig. 1b); anterior hood absent; urosome wide, oblong. BL/PW 1.03, BL/UW 4.14,
- 168 PW/UW 4.02. Caudal ramus present, small lobe shaped, with two spiniform setae (outer seta longer than
- 169 inner). Rod/horn-like antennule and antenna absent; semicircular irregular branching structure present
- 170 posterior to oral sucker. Rootlet absent. Mandible present, styliform. Pore (maxillary gland pore?; arrow in
- 171 Fig. 2e) present on ventrolateral swelling. Legs 1–4 absent. Leg 5 present, small lobe shaped, located in
- 172 ventrolateral region of urosome, with three spiniform setae. Genital area without ornamentation; openings
- 173 of seminal receptacles and genital apertures indistinct.
- 174 *Male and copepodid*. Unknown.
- 175 *Variation.* One female paratype (ICHUM8453) observed in addition to the holotype showed all
- 176 diagnostic characters seen in the holotype. Ranges (mean with standard deviation in parentheses) of BL,
- 177 PW, UW, BL/PW, BL/UW, PW/UW for five individuals in the type series were 342-400 (372±23), 326-
- 178 385 (356±20), 86–96 (91±4), 0.95–1.11 (1.05±0.05), 3.93–4.20 (4.10±0.09), and 3.74–4.32 (3.94±0.22),

179 respectively.

180

181 Genetic divergence and phylogenetic analysis

182	We attempted to determine COI, 16S, and 18S sequences for <i>D. hakuhomaruae</i> sp. nov. but were unable to
183	PCR-amplify the COI region. Among copepod 16S sequences in the INSD database, one from Tripaphylus
184	elongatus (C. B. Wilson, 1932) (as Paeon elongatus, FJ447423 [24]) was most similar to our 16S sequence
185	in a BLAST search, but the query cover and identity score were low (46% and 76.73%, respectively). To
186	date, no other nicothoid 16S sequences have been deposited in public databases [25].
187	In the ML tree (Fig. 3), D. hakuhomaruae formed a fully supported clade with the Rhizorhina
188	clade. The other relationships were identical to those provided in Kakui and Munakata [4], except for
189	minor differences in ultrafast bootstrap values.
190	
191	Remarks
192	Diexanthema hakuhomaruae sp. nov. is the seventh species described in Diexanthema. Females lack
193	segmented antennules, have mandibles, and lack legs 1-4, features shared with females of <i>D. bathydiaita</i> .
194	The former differs from the latter in having a smooth body surface (body covered with minute hairs in <i>D</i> .
195	bathydiaita) and leg 5 located ventrolaterally on the urosome (laterally in D. bathydiaita). Their host
196	groups are different at the family level: Desmosomatidae for <i>D. hakuhomaruae</i> , Nannoniscidae for <i>D.</i>
197	bathydiaita.
198	Diexanthema hakuhomaruae sp. nov. lacks rootlets observed in D. apoda. This species appears to
199	use its oral sucker to attach to its host isopod.
200	Our 18S tree showed a close relationship between Diexanthema and Rhizorhina. This suggests

201 that, although we lacked *Choniorhiza* sequences, the "*Rhizorhina* group" proposed by Boxshall and

202 Lincoln [10] may reflect close phylogenetic relationships among its members.

203 Individuals of *Diexanthema hakuhomaruae* sp. nov. were, as with *Rhizorhina* individuals (KK

- 204 unpublished data), easily deformed by changes in solution; for example, the body of one individual shrank
- when transferred into a 1:3:6 mixture of glycerin, absolute ethanol, and deionized water, making
- 206 morphological observation difficult (transferred into 30% ethanol, it recovered its spherical shape). This
- suggests that differences in body form should be treated with caution in *Diexanthema* (and *Rhizorhina*)
- 208 taxonomy. In addition, most appendages are strongly reduced or completely lacking in these nicothoid
- 209 genera, and often few morphological differences are observable among congeners. As adopted in other

210 parasitic groups [26], a "turbo taxonomy" [27] approach, i.e., providing concise morphological

- 211 descriptions along with DNA-sequence and host data in establishing new species, may be advisable in the
- 212 taxonomy of several character-poor nicothoid genera.

213

#### 214 Conclusions

215 We described *Diexanthema hakuhomaruae* **sp. nov.** parasitic on the desmosomatid *Eugerdella* cf.

216 *kurabyssalis* Golovan, 2015 collected from a hadal depth in the Kuril-Kamchatka Trench, northwestern

- 217 Pacific. This species represents the first *Diexanthema* species from the Pacific, and the first from hadal
- 218 depths. Our 18S-based tree confirmed a close relationship between *Diexanthema* and *Rhizorhina*,
- 219 previously suggested by morphology. Continued molecular studies, with wider taxon sampling and

220 additional molecular markers, should further elucidate the phylogenetic relationships among nicothoid

221 copepods, with consequent advances in taxonomy.

222

223	Supp	lementary	' Inforr	nation
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224 Online Resource 1 Aligned 18S sequences used in the maximum-likelihood analysis, with alignment-

ambiguous sites retained.

226 Online Resource 2 Aligned 18S sequences used in the maximum-likelihood analysis, reduced to 1528

227 positions by removing alignment-ambiguous sites.

228

#### 229 **Declarations**

- 230 Author contributions: KK conceived and designed the study, made morphological observations on the
- 231 copepods, and conducted the molecular analysis; MO made morphological observations on the isopods;
- 232 KK, JF, and MO collected samples, wrote the manuscript, and read and approved the final draft.
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- 237 **Competing interests:** The authors have no competing interests to declare that are relevant to the content of
- this article.

240	Со	nsent: Not applicable.
241	Dat	ta availability: The raw data (sampling locality; sampling date; museum deposition numbers, INSD
242	acc	ession numbers, and the depository for specimens) are included in the manuscript.
243		
244	Re	ferences
245	1.	Walter TC, Boxshall G (2023) Nicothoidae Dana, 1852-1853.
246		https://www.marinespecies.org/aphia.php?p=taxdetails&id=135530. Accessed 13 March 2023
247	2.	Boxshall GA, Halsey SH (2004) An introduction to copepod diversity. The Ray Society, London
248	3.	Kakui K (2016) Descriptions of two new species of Rhizorhina Hansen, 1892 (Copepoda:
249		Siphonostomatoida: Nicothoidae) parasitic on tanaidacean crustaceans, with a note on their
250		phylogenetic position. Syst Parasitol 93:57-68. https://doi.org/10.1007/s11230-015-9604-x
251	4.	Kakui K, Munakata M (2022) A new Sphaeronella species (Copepoda: Siphonostomatoida:
252		Nicothoidae) parasitic on Euphilomedes sp. (Ostracoda: Myodocopa: Philomedidae) from Hokkaido,
253		Japan, with an 18S molecular phylogeny. Syst Parasitol. https://doi.org/10.1007/s11230-022-10075-z
254	5.	Ohtsuka S, Boxshall GA, Harada S (2005) A new genus and species of nicothoid copepod (Crustacea:
255		Copepoda: Siphonostomatoida) parasitic on the mysid Siriella okadai Ii from off Japan. Syst Parasitol
256		62:65-81. https://doi.org/10.1007/s11230-005-5483-x
257	6.	Bamber RN, Boxshall GA (2006) A new genus and species of the Langitanainae (Crustacea:

Ethics approval: Not applicable.

- 258 Peracarida: Tanaidacea: Tanaidae) bearing a new genus and species of nicothoid parasite (Crustacea:
- 259 Copepoda: Siphonostomatoida: Nicothoidae) from the New Caledonia slope. Spec Divers 11:137–
- 260 148. https://doi.org/10.12782/specdiv.11.137
- 261 7. Boyko CB (2009) Nomenclatural issues with Paranicothoe Carton, 1970 and Pseudonicothoe Avdeev
- 262 & Avdeev, 1978 (Crustacea: Copepoda: Nicothoidae), with comments on the female isopod type
- 263 specimen of *Paranicothoe cladocera* Carton, 1970. Proc Biol Soc Wash 122:206–211.
- 264 https://doi.org/10.2988/08-49.1
- 8. Ritchie L (1975) A new genus and two new species of Choniostomatidae (Copepoda) parasitic on two
- 266 deep sea isopods. Zool J Linn Soc 57:155–178. https://doi.org/10.1111/j.1096-3642.1975.tb01415.x
- 267 9. Boxshall GA, Harrison K (1988) New nicothoid copepods (Copepoda: Siphonostomatoida) from an

amphipod and from deep-sea isopods. Bull Br Mus Nat Hist (Zool) 54:285–299

- 269 10. Boxshall GA, Lincoln RJ (1983) Some new parasitic copepods (Siphonostomatoida: Nicothoidae)
- 270 from deep-sea asellote isopods. J Nat Hist 17:891–900. https://doi.org/10.1080/00222938300770701
- 271 11. Akiyama T, Shimomura M, Nakamura K (2008) Collection of deep-sea small arthropods: gears for
- collection and processing of samples on deck. TAXA 24:27–32. https://doi.org/10.19004/taxa.24.0 27
- [in Japanese with English abstract]
- 12. Folmer O, Black M, Hoeh W, Lutz R, Vrijenhoek R (1994) DNA primers for amplification of
- 275 mitochondrial cytochrome c oxidase subunit I from diverse metazoan invertebrates. Mol Mar Biol

276 Biotechnol 3:294–299

- 13. Nakayama T, Watanabe S, Mitsui K, Uchida H, Inouye I (1996) The phylogenetic relationship
- 278 between the Chlamydomonadales and Chlorococcales inferred from 18SrDNA sequence data. Phycol
- 279 Res 44:47–55. https://doi.org/10.1111/j.1440-1835.1996.tb00037.x
- 280 14. Kakui K, Katoh T, Hiruta SF, Kobayashi N, Kajihara H (2011) Molecular systematics of Tanaidacea
- 281 (Crustacea: Peracarida) based on 18S sequence data, with an amendment of suborder/superfamily-
- level classification. Zool Sci 28:749–757. https://doi.org/10.2108/zsj.28.749
- 283 15. Kakui K, Fukuchi J, Shimada D (2021) First report of marine horsehair worms (Nematomorpha:
- 284 *Nectonema*) parasitic in isopod crustaceans. Parasitol Res 120:2357–2362.
- 285 https://doi.org/10.1007/s00436-021-07213-9
- 286 16. Munakata M, Tanaka H, Kakui K (2021) Heterocypris spadix sp. nov. (Crustacea: Ostracoda:
- 287 Cypridoidea) from Japan, with information on its reproductive mode. Zool Sci 38:287–296.
- 288 https://doi.org/10.2108/zs200127
- 289 17. Okamoto N, Kakui K (2022) Integrative taxonomy of Zeuxo (Crustacea: Peracarida: Tanaidacea) from
- Japan, with the description of a new species. Biologia 77:2497–2506. https://doi.org/10.1007/s11756-
- 291 022-01121-8
- 292 18. Kumar S, Stecher G, Tamura K (2016) MEGA7: Molecular evolutionary genetics analysis version 7.0
- for bigger datasets. Mol Biol Evol 33:1870–1874. https://doi.org/10.1093/molbev/msw054
- 294 19. Munakata M, Tanaka H, Kakui K (2022) Taxonomy and natural history of Cavernocypris hokkaiensis
- sp. nov., the first ostracod reported from alpine streams in Japan. Zoosyst Evol 98:117–127.

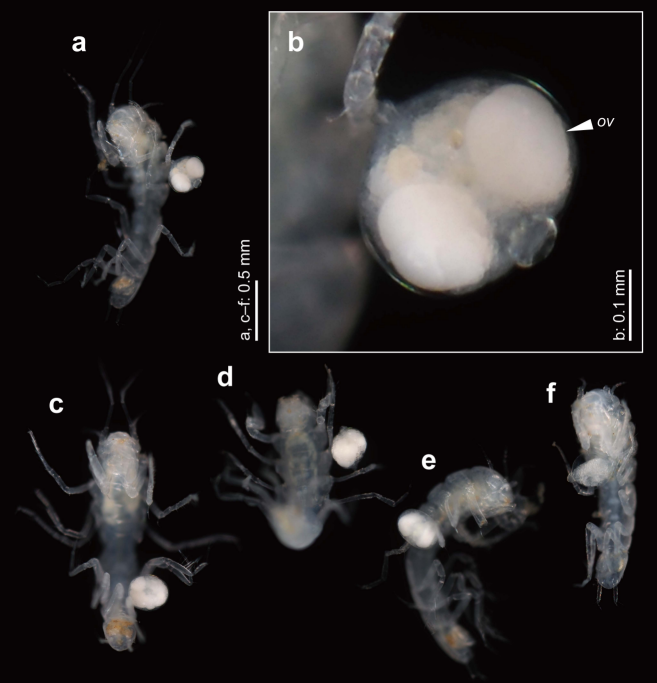
- https://doi.org/10.3897/zse.98.80442
- 297 20. Kakui K, Shimada D (2022) Dive into the sea: first molecular phylogenetic evidence of host
- 298 expansion from terrestrial/freshwater to marine organisms in Mermithidae (Nematoda: Mermithida). J
- 299 Helminthol 96:e33. https://doi.org/10.1017/S0022149X22000256
- 300 21. Golovan OA (2015) Description of two ubiquitous species of Desmosomatidae (Isopoda: Asellota)
- 301 from the Northwest Pacific Basin east of the Kuril-Kamchatka Trench. Zootaxa 4039:201–224.
- 302 https://doi.org/10.11646/zootaxa.4039.2.1
- 303 22. Jennings RM, Golovan O, Brix S (2020) Integrative species delimitation of desmosomatid and
- 304 nannoniscid isopods from the Kuril-Kamchatka trench, with description of a hadal species. Prog
- 305 Oceanogr 182:102236. https://doi.org/10.1016/j.pocean.2019.102236
- 306 23. Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ (1990) Basic local alignment search tool. J
- 307 Mol Biol 215:403–410. https://doi.org/10.1016/S0022-2836(05)80360-2
- 308 24. Dippenaar SM (2009) Estimated molecular phylogenetic relationships of six siphonostomatoid
- 309 families (Copepoda) symbiotic on elasmobranchs. Crustaceana 82:1547–1567.
- 310 https://doi.org/10.1163/001121609X12511103974538
- 311 25. DDBJ (2022) DNA Data Bank Japan. https://www.ddbj.nig.ac.jp/ Accessed 29 November 2022
- 312 26. Summers MM, Al-Hakim II, Rouse GW (2014) Turbo-taxonomy: 21 new species of Myzostomida
- 313 (Annelida). Zootaxa 3873:301–344. https://doi.org/10.11646/zootaxa.3873.4.1
- 27. Butcher BA, Smith MA, Sharkey MJ, Quicke DLJ (2012) A turbo-taxonomic study of Thai Aleiodes

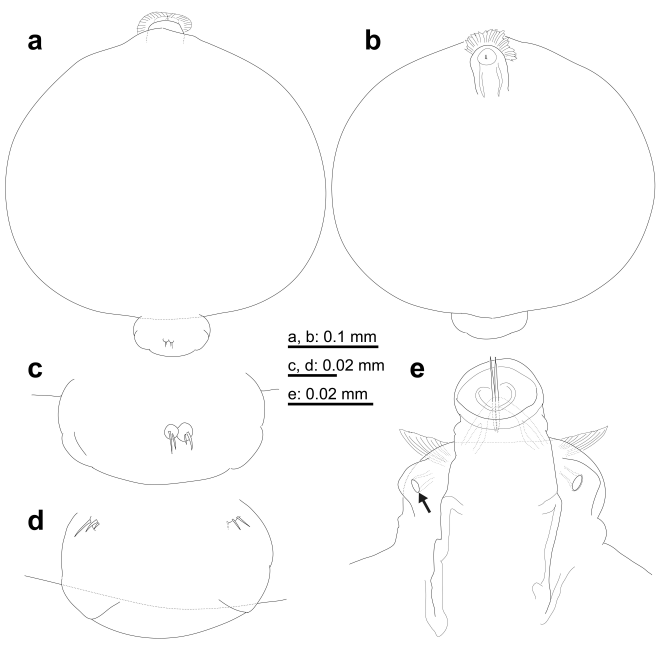
- 315 (Aleiodes) and Aleiodes (Arcaleiodes) (Hymenoptera: Braconidae: Rogadinae) based largely on COI
- barcoded specimens, with rapid descriptions of 179 new species. Zootaxa 3457:1–232.
- 317 https://doi.org/10.11646/zootaxa.3457.1.1

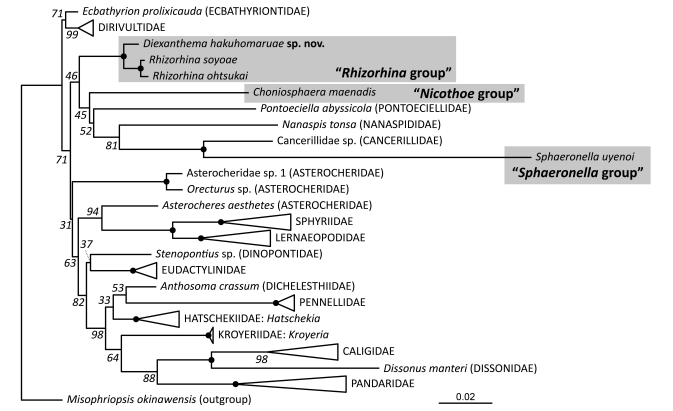
## 318 Figure and table legends

319	Fig. 1 Diexanthema hakuhomaruae sp. nov. (females) attached to the host, Eugerdella cf. kurabyssalis
320	Golovan, 2015 (females with developing oostegites), fresh specimens. a, holotype on host (ICHUM8451);
321	b, same, enlarged view; c-f, paratypes on hosts (ICHUM8452-ICHUM8455). Abbreviation: ov, ovary
322	
323	Fig. 2 Diexanthema hakuhomaruae sp. nov., holotype, female (ICHUM8451). a, b, body, dorsal and
324	ventral views, respectively; c, d, urosome, dorsal and ventral views, respectively; e, head region, ventral
325	view (arrow, right maxillary gland pore?)
326	
327	Fig. 3 Maximum-likelihood (ML) tree based on 18S sequences (1528 positions). Numbers near nodes are
328	ultrafast bootstrap values. Black circles indicate 100% ultrafast bootstrap support. Clades containing more
329	than two confamilial terminal taxa were collapsed (terminal triangles), except for those in Nicothoidae
330	(shaded) and Asterocheridae. Group names follow Boxshall and Lincoln [10]. The scale bar indicates
331	branch length in substitutions per site
332	

333 Table 1 Information on six *Diexanthema* species from North Atlantic Ocean







Species	Isopod host	Locality	Depth (m)
D. apoda	Munnopsidae: Bathybadistes	Rockall Trough, off	c. 2910
	spinosissima (Hansen, 1916)	Scotland	
D. bathydiaita	Nannoniscidae: Nannoniscus sp.	Off North America	1624–1796
D. corrugatum	Munnopsidae: Acanthocope sp.	Rockall Trough	c. 2900
D. desistoma	Desmosomatidae: Chelator insignis	Off Africa	1330–1470
	(Hansen, 1916)		
D. nudum	Desmosomatidae: Mirabilicoxa sp.	Rockall Trough	c. 2540
	(probably M. acuminata Hessler, 1970)		
D. ritchiei	Ischnomesidae: Gracilimesus	Porcupine Seabight,	3490-3550
	tenuispinis (Hansen, 1916)	off Ireland	

Table 1 Information on six Diexanthema species from North Atlantic Ocean