Important declarations

Please remove this info from manuscript text if it is also present there.

Associated Data

Data supplied by the author:

Images and videos of the specimens S2 https://filesender.renater.fr/?s=download&token=7ed9a906-859e-49fc-b4e5-a33922604019

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The authors declare that they have no competing interests.

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On *Dracograllus miguelitus* sp. nov. (Nematoda: Draconematidae) from an inactive structure: insights into its taxonomy, biodiversity and ecology at hydrothermal vents

William Johnson da Silva Corresp., 1, Daniela Zeppilli 1, Valentin Foulon 2, Pierre-Antoine Dessandier 1, Marjolaine Matabos 1, Jozée Sarrazin 1

¹ Ifremer, BEEP, Univ Brest, Plouzané, France., Plouzané, Plouzané - France, France

² ENIB - École Nationale d'Ingénieurs de Brest, Plouzané, France., Plouzané, Plouzané - France, France

Corresponding Author: William Johnson da Silva Email address: William.Johnson.Da.Silva@ifremer.fr

Background. Hydrothermal vent fields are habitats to a diverse array of benthic organisms, including several nematode species, which represent a significant portion of the biodiversity in these environments. Despite their ecological importance, most research on hydrothermal vents has focused on macro-invertebrates. As a result, vent nematode biodiversity remains largely unexplored, especially in peripheral and inactive structures, underscoring the need for further investigation. A sampling program conducted in 2017 and 2018 along a gradient of venting activity led to the collection of a number of Draconematidae species in various habitats. In this paper, we introduce *Dracograllus miguelitus* sp. nov., the first species of the genus described at a hydrothermal vent field, sampled from a visually inactive sulphide structure.

Methods. The samples were collected at the Lucky Strike vent field, on the Mid-Atlantic Ridge, using the suction sampler of the Remotely Operated Vehicle Victor6000. Specimens were retrieved from an edifice covered by a black layer of manganese oxy-hydroxides, with no local visible hydrothermal activity, at a depth of 1.639 meters. Samples were sieved on a 32 µm mesh onboard, sorted and, for nematodes, identified to species level back in the lab. Fluorescent images were obtained using the ApoTome Fluorescence Microscope Module, and 3D observations were possible through the depth change method. Results. We established *D. miguelitus* sp. nov. as a new species based on the combination of the following characters: four cephalic adhesive tubes (CATs), an elongated loop-shaped amphid with varying branch sizes between males and females, and a circular amphid in juveniles. Additionally, females display a minute setae emerging from the vulvar aperture. In males, the posterior adhesive tubes (PATs) are arranged in four longitudinal rows: two sublateral rows, each containing 10-12 PATs, and two subventral rows, consisting of 10 PATs in each. In females, sublateral and subventral rows with 13 PATs each. So far, *D.*



miguelitus sp. nov. is the first species of the genus to be described from a hydrothermal environment and the deepest one. Beyond the formal description of this new species, we provide ecological and taxonomic backgrounds on Draconematidae at hydrothermal vents, with insights into the genus distribution, biogeography, and nomenclatural issues. Conclusion. This discovery contributes to the knowledge of Draconematidae biodiversity, and highlights the importance to investigate nematode communities at species-level, data that is often missing at vent studies. Additionally, it underscores the significance of preserving inactive hydrothermal habitats, which are threatened by deep-sea mining activities.

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- 34 35
- William Johnson¹, Daniela Zeppilli¹, Valentin Foulon², Pierre-Antoine Dessandier¹, Marjolaine
 Matabos¹, Jozée Sarrazin¹
- 38
- 39 ¹ Ifremer, BEEP, Univ Brest, Plouzané, France.
- 40 ² ENIB École Nationale d'Ingénieurs de Brest, Plouzané, France.
- 41
- 42 Corresponding Author:
- 43 William Johnson¹
- 44 1625 Rte. de Sainte-Anne, Plouzané, 29280, France
- 45 Email address: <u>William.Johnson.Da.Silva@ifremer.fr</u>
- 46

47 Abstract

48 Background. Hydrothermal vent fields are habitats to a diverse array of benthic organisms,

- 49 including several nematode species, which represent a significant portion of the biodiversity in
- 50 these environments. Despite their ecological importance, most research on hydrothermal vents
- 51 has focused on macro-invertebrates. As a result, vent nematode biodiversity remains largely
- 52 unexplored, especially in peripheral and inactive structures, underscoring the need for further
- 53 investigation. A sampling program conducted in 2017 and 2018 along a gradient of venting
- 54 activity led to the collection of a number of Draconematidae species in various habitats. In this
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- 63 observations were possible through the depth change method.
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- 65 following characters: four cephalic adhesive tubes (CATs), an elongated loop-shaped amphid
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- 69 containing 10-12 PATs, and two subventral rows, consisting of 10 PATs in each. In females,
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- 72 Beyond the formal description of this new species, we provide ecological and taxonomic
- 73 backgrounds on Draconematidae at hydrothermal vents, with insights into the genus distribution,
- 74 biogeography, and nomenclatural issues.
- 75 Conclusion. This discovery contributes to the knowledge of Draconematidae biodiversity, and
- 76 highlights the importance to investigate nematode communities at species-level, data that is often
- 77 missing at vent studies. Additionally, it underscores the significance of preserving inactive
- 78 hydrothermal habitats, which are threatened by deep-sea mining activities.
- 79

80 Introduction

A significant portion of the benthic diversity associated with hydrothermal vents is represented 81 by nematodes, which play crucial roles at the ecosystem level such as bioturbation and organic 82 matter degradation (Vanreusel, Van den Bossche & Thiermann, 1997; Vanreusel et al., 2010a; 83 Vanreusel et al., 2010b). These environments are characterized by a hard substratum with high 84 85 contents of metal compounds such as copper, zinc and iron, resulting from the precipitation of 86 polymetallic sulphides contained in the vent fluids (Hoaglang et al., 2010). Unlike other deepsea ecosystems, hydrothermal vents exhibit a unique combination of low diversity and high 87 biomass, largely driven by chemosynthetic energy sources (Tunnicliffe, 1991). Nematodes thrive 88 in these conditions, highlighting their remarkable ability to adapt to habitat heterogeneity and 89 extreme environments (Vanreusel et al., 2010b). 90

- 91
- Surviving to the peculiar environmental conditions of the deep-sea high pressure, low
 temperatures, food scarcity- poses a significant challenge for faunal communities. Hydrothermal
- 94 vents introduce further selective pressures linked to the type, origin, and intensity of
- 95 hydrothermal activity and resulting environmental conditions (Koschincky et al., 2008).
- 96 Consequently, nematode communities at vents differ from those in the surrounding deep-sea.
- 97 They harbor species that possess adaptations and strategies that are essential to survive in these
- 98 harsh environments (*Vanreusel et al., 2010a*). Species diversity vary significantly across sites
- 99 with different levels of hydrothermal activity (Gollner, Miljutina, Bright, 2013) and differences
- 100 in species composition underscore their ability to occupy various niches, making them important
- 101 contributors to the functioning of hydrothermal ecosystems (*Vanreusel et al., 2010b*).
- 102

Some examples of these adaptations can be observed in the Draconematidae family *Filipjev, 1918.* These nematodes are easily recognizable by their S-shaped body morphology,
which is common to most species. This unique shape has earned them colloquial names of
"walking nematodes" or "dragon nematodes." In addition to their distinct morphology, many

107 Draconematidae exhibit specialized structures that are closely tied to their locomotion and

108 habitat use. Their cephalic (CATs) and posterior (PATs) adhesive tubes are linked to glands that secrete adhesive substances. These secretions allow them to "stick" parts of their bodies to the 109 substratum, enabling alternative movements with intervals of "attachment and release" of both 110 anterior and posterior body regions (Stauffer, 1924; Cobb, 1929; Clasing, 1980; Tchesunov et 111 112 al., 2014). The Draconematidae family comprises 16 genera and 89 valid species (Nemvs, 2024). most of which are commonly found in coastal regions, typically associated with biological 113 structures such as worm tubes, algae and coral reefs (Decraemer, Gourbault & Backeljau, 1997). 114 The unexpected discovery of Draconematidae species in high abundances at hydrothermal vents 115 was first reported in the Guaymas Basin on the East Pacific Rise (2 000 m water depth) by 116 117 (Dinet, Grassle & Tunnicliffe, 1988). Since then, additional records of the family in deep-sea habitats, including hydrothermal vents, have been reported. Several genera typical of deep-sea 118 environments were collected, such as *Cephalochaetosoma* (syn. *Bathychaetosoma*) and *Dinetia* 119 from the subfamily Draconematinae, as well as *Prochaetosoma* from the subfamily 120 Prochaetosomatinae (Kito, 1983; Decraemer, Gourbault & Backeljau, 1997; Rho, Kin & Kin, 121 122 2007; Rho & Kim, 2011, and references therein). On the East Pacific Rise (EPR), Dinetia sp. were associated with *Bathymodiolus* mussel beds (*Flint et al., 2006*). Similarly, at the Lucky 123 Strike vent field, along the northern Mid-Atlantic Ridge (MAR), both Dinetia and 124 125 Cephalochaetosoma were associated with Bathymodiolus mussels (Husson et al., 2017). More recently, an experimental colonization study showed that *Cephalochaetosoma* represented 126 between 76% and 90% of the nematode community on inorganic substrata deployed in intense 127 vent emission areas (Zeppilli et al., 2015). 128 129 130 The genus Dracograllus Allen & Noffsinger, 1978 represents the largest genus within the

family, with 25 valid species (Min et al., 2016; Nemys, 2024), most of them reported in shallow 131 waters, and, as for several Draconematidae species, associated with biogenic structures 132 (Verschelde & Vincx, 1993). Even without apomorphic characters, the genus can be 133 134 distinguished from other genera by several features, including a non-enlarged cuticle in the head region, the absence of bilateral cephalic acanthiform setae on the head capsule (except for D. 135 stekhoveni), the absence of precloacal copulatory thorns, and the presence of paravulvar setae in 136 some species (Allen and Noffsinger, 1978; Decraemer, 1988; Decraemer, Gourbault & 137 138 Backeljau, 1997). Up to now, no species of Dracograllus had been formally described from deep-sea or hydrothermal habitats, as their distribution is generally limited to depths shallower 139 than 100 meters. However, several recent studies have reported Dracograllus specimens at 140

141 greater depths (Vanhove et al., 1999; Gad, 2009; Zeppilli et al., 2013, 2014; Spedicato et al.,

- 142 2020), although none have been formally described so far.
- 143

Most part of the vent ecological studies have focused on the microbial and macrofaunal compartments on active hydrothermal structures, neglecting the smaller meiofauna and also, the fauna from regions adjacent to the vents and inactive structures. However, although they received less attention, there is an increased interest in studying inactive vents, because they are

- 148 the main target for mineral extraction (*Menini et al., 2023*). Recent studies have shown
- 149 differences in faunal diversity between hydrothermally active and inactive habitats with a much
- 150 higher diversity in the latter (*Cowart et al., 2020*). Few studies in the vent periphery have shown
- that nematode diversity extends outside the active zones (*Vanreusel et al., 2010b*). In this
- 152 context, it becomes critical to better understand their diversity patterns in vent ecosystems
- including in their sphere of influence (*Levin et al., 2016*). Such knowledge is essential for
- 154 developing environmental management plans to mitigate the impacts of deep-sea mining.
- 155

156 To assess meiofaunal benthic biodiversity associated with different vent environmental conditions, a sampling was carried out at 1700 m depth at the Lucky Strike vent field in three 157 contrasting habitats: an active vent site, a 'visually' inactive structure, and an area peripheral to 158 venting activity (Cowart et al., 2020). In this study, we describe for the first time a new species 159 of *Dracograllus* sampled from a deep-sea inactive sulfide structure, and supply updates on the 160 161 taxonomy, ecology, and distribution of the genus. Additionally, we provide a dichotomous key to aid in the identification of *Dracograllus* species. Finally, we examine the implications of our 162 results for the conservation of hydrothermal ecosystems, focusing on species composition, 163 interactions and ecosystem functions in the context of the challenges posed by the mining 164 165 industry.

166

167 Materials & Methods

168 Study area and sampling collection

The Lucky Strike (LS) vent field is located in the northern part of the Mid-Atlantic Ridge 169 (MAR), south of the Azores (Fig.1A), with a mean depth of 1.700m (De Busserolles et al., 170 171 2009). LS consists of three volcanic cones that harbor over 25 active hydrothermal edifices surrounding a central lava lake (Humphris et al., 2002; Ondréas et al., 2009). Each active site -or 172 edifice- is made of several smokers as well as patches of diffuse venting areas that extend in the 173 174 periphery. To characterize the meiofaunal communities at the vent field scale, three habitat types 175 were sampled: an active area, the periphery away from hydrothermal activity and a visually inactive edifice (Fig.1B). 176

177

178 The active habitat was located on the Montségur edifice (37°17.28'N, 32°16.53'W), in the southern region of LS, and consisted of cracks on a flat hydrothermal slab at the base of the 179 180 edifice. The peripheral habitat was approximately 30 meters from Montségur (37°17.28'N, 32°16.52'W), and covered by a few centimeters of sediments. Finally, the visually inactive 181 182 structure, peripheral to the active Sintra edifice, lied about 400 meters north of Montségur (37°17.48'N, 32°16.50'W), and consisted in an indurated sulfide structure covered at its base by 183 a thin black layer of manganese oxy-hydroxides (Fig.1C-D). Sampling was conducted during the 184 2018 Momarsat cruise (*Cannat, 2018*) using the suction sampler of the Remotely Operated 185 Vehicle Victor6000. Neither the peripheral nor the inactive habitats exhibited visible 186 hydrothermal activity or typical vent fauna. Once onboard, samples were sieved on 300 and 20 187

188 μ m mesh sizes, and the fraction between 20-300 μ m was preserved in 4% borax buffered

- 189 formalin.
- 190

191 Sample preparation and image acquisition

- 192 Nematodes were extracted from the sediment by the use of colloidal silica (Ludox), with specific
- 193 gravity of 1.39 (*Pfannkuche & Thiel, 1988*). Specimens were fixed in formalin, and after *the De*
- 194 *Grisse, 1989* protocol, they were transferred to glycerol and mounted onto permanent slides
- 195 (Somerfield & Warwick, 1996). Drawings and measurements were made using a light
- microscope Leica DM 2500 LED with the aid of a drawing tube and a Leica DMC 4500 camera.
- 198 For the fluorescent observations, a Zeiss Axio Imager.Z2 microscope equipped with an
- 199 Colibri.7 light, an ORCAFlash4.OLT (Hamamatsu, Hamamatsu-city, Japan) camera and a
- 200 Apotome.2 slider module (for optical sections) was used. Autofluorescence and Phloxine B stain
- 201 (Exitation 561 nm, Emission 571 nm) were used to observe internal and external structures in
- 3D. Thus, four fluorescent channels were used: Blue filter Zeiss 49 DAPI Ex. G365 nm, Em.
- 203 445/50 nm, Green filter Zeiss 38 HE GFP Ex 470/40 nm, Em LP 515 nm, Orange filter Zeiss
- 43HE dsREd Ex. 550/25 nm Em. 605/70 nm, Red filter Zeiss 50Cy5 Ex. 640/30 Em. 690/50
- 205 nm. Combinations of one to five channels (with brightfield) were used for optical section,
- 206 increased depth of field and 3D depending on the observations. Images were processed using
- 207 Zeiss Zen Pro and Arivis 4D Pro software.
- 208

In one of the earliest reviews of Draconematidae, *Allen & Noffsinger, 1978* provided key
recommendations regarding specimen measurements, morphological analysis, and species
delimitation. Building on their guidance, this study incorporates the following recommendations
and observations:

- 213 214
- Measurements on the CATs on the right side of the nematode;
- For the measurements of the swollen esophageal and cephalic region:
- 216 o Length: measured from the anterior tip of the lip region to just posterior to the swollen esophageal region (in most Draconematidae, the body is constricted in this region);
- Width: body diameter measured at the widest part of the swollen esophageal region;
 - Rostral width measured at the base of the rostrum, just anterior to the first body annule.
- 222 223

221

- For comprehensive details about measurements and possible variations along development stages, see *Allen & Noffsinger*, *1978* and *Clasing*, *1980*.
- 226

227 Nomenclatural acts

- 228 The electronic version of this article in Portable Document Format (PDF) will represent a
- 229 published work according to the International Commission on Zoological Nomenclature (ICZN),
- and hence the new names contained in the electronic version are effectively published under that
- 231 Code from the electronic edition alone. This published work and the nomenclatural acts it
- contains have been registered in ZooBank, the online registration system for the ICZN. The
- 233 ZooBank LSIDs (Life Science Identifiers) can be resolved and the associated information viewed
- through any standard web browser by appending the LSID to the prefix http://zoobank.org/. The
- LSID for this publication is: [AA6564D7-6BA7-405E-94D3-B659E62B8BDB]. The online
- version of this work is archived and available from the following digital repositories: PeerJ,
- 237 PubMed Central SCIE and CLOCKSS.
- 238

239 Results and discussion

- 240 This is the first study to describe a new *Dracograllus* species from an inactive vent structure. It
- also corresponds to the greatest depth recorded among all known congener valid species. This
- 242 finding provides insights into the diversity of potential habitats for Draconematidae, with genera
- 243 and species distributed across a wide range of environments from shallow to deep regions.
- 244
- 245 SYSTEMATICS
- 246 Class CHROMADOREA Inglis, 1983
- 247 Subclass CHROMADORIA Pearse, 1942
- 248 Order Desmodorida *De Coninck*, 1965
- 249 Suborder Desmodorina De Coninck, 1965
- 250 Superfamily Desmodoroidea Filipjev, 1922
- 251 Family Draconematidae Filipjev, 1918
- **Diagnosis.** (Emended from *Leduc & Zhao, 2016*): Body short, S-shaped, usually with more or
- 253 less enlarged pharyngeal and mid-body region. Cuticle annulated except for the head capsule
- 254 (helmet) and tail terminus. Annules sometimes with spines, minute vacuoles, or a longitudinal
- 255 lateral field in mid-body region or tail region. Cephalic sensilla in three circles (6+6+4): six inner
- 256 labial papillae, six outer labial setae, and four cephalic setae. Rostrum present (except in
- 257 *Dinetia*). Amphideal fovea spiral to loop-shaped, rarely reduced or an internal longitudinal bar.
- 258 CATs present, located dorsally on the cephalic capsule. Somatic setae arranged in eight
- 259 longitudinal rows. Buccal cavity small to well developed, usually with a dorsal tooth, with or
- without subventral teeth. Pharynx cylindrical, dumbbell shaped, or with posterior bulb.
- 261 Secretory–excretory system absent. At least the anterior-most PATs are arranged in four
- longitudinal rows, two subventrally and two ventrosublaterally, located on the posterior third of
- the body. PATs with or without differentiated tips, usually straight, rarely long and flexible.
- 264 Copulatory apparatus with two spicules and trough-shaped gubernaculum. Three caudal glands
- extending beyond anus/cloaca.
- 266

267 Subfamily Draconematinae *Filipjev*, 1918

- **Diagnosis.** (Emended from *Decraemer, Gourbault & Backeljau, 1997*): Pharynx dumbbell-
- shaped. Swollen anterior body region is usually conspicuous and short. CATs with clearly open
- tips, mainly blister-shaped, and enlarged bases of insertion, located on the rostrum. Cephalic
- acanthiform setae may be present on the helmet. Buccal cavity narrow, unarmed. PATs shorts
- and straights (except for *Draconema trispinosum*, characterized by some longer tubes), with bell-
- shaped tips. All PATs are anterior to cloaca (except in *Dracograllus eira* and *Dracograllus*
- *minutus*). Copulatory thorns are rare. Paravulvar setae present or not. Tail cylindro-conoid with
- 275 numerous nodules. Non- annulated tail region, with different length between species.
- **276 Type genus.** *Draconema Cobb, 1913.*
- 277

278 Genus Dracograllus Allen & Noffsinger, 1978

Diagnosis. (Emended from *Leduc & Zhao, 2016*): Draconematidae, Draconematinae. Swollen
anterior body region 18–26% of total body length. Usually eight CATs on the helmet, rarely 10
to 15. Non-enlarged cuticle on the rostrum. Amphids lateral, usually loop-shaped, with branches

usually of unequal length, rarely spiral. Precloacal copulatory thorns are usually absent. PATs all

anterior to cloacal opening (except in *Dracograllus eira Inglis, 1968* and *Dracograllus minutus*

- 284 *Decraemer*, 1988), and usually extending posterior to anus in females (four exceptions).
- 285 Paravulvar setae may be present.
- 286
- **287** Type species. Dracograllus cobbi Allen & Noffsinger, 1978.
- 288
- 289 Remarks on *Dracograllus* taxonomy. The genus *Dracograllus* is placed within the subfamily
- 290 Draconematinae, along with *Draconema*, *Paradraconema*, and also the genus *Tenuidraconema*.
- 291 On the other hand, the subfamily Prochaetosomatinae includes the deep-sea typical genera:
- 292 Bathychaetosoma, Cephalochaetosoma, and Dinetia, as well as Prochaetosoma. This
- 293 classification is based on characteristics such as the cylindrical shape of the pharyngeal sphincter
- with a terminal bulb, suggesting that the typical dumbbell-shaped pharynx was possibly
- secondarily lost in *Tenuidraconema* (Decraemer, Gourbault & Backeljau, 1997).
- 296

297 A molecular phylogenetic analysis of the family Draconematidae by *Rho & Min (2011)* 298 revealed that the genus *Dracograllus* was the first lineage to diverge in the family, followed by 299 five branching orders: Dracograllus - Megadraconema - Draconema - Paradraconema and 300 *Prochaetosoma*. Consequently, the genus was excluded from the subfamily Draconematinae, as previously suggested. More recently, Leduc & Zhao (2016) examined the phylogenetic position 301 302 of species within Desmodoroidea and confirmed the basal placement of *Dracograllus* within the Draconematidae, consistent with Rho & Min (2011) findings. Additionally, Leduc & Zhao 303 304 (2016) found that sequences from *Dracograllus*, along with two other specimens of the genus, 305 clustered with high posterior probability and bootstrap support, further reinforcing its basal 306 position in the Draconematidae and providing new insights into the group's evolutionary relationships. 307

308

309 Taxonomic issues. Taxonomic issues within *Dracograllus* include synonymy, redescriptions,

- and taxon transfers, often arising from descriptions based on immature specimens or
- 311 discrepancies in the number of longitudinal PAT rows. For example, *Dracograllus eira* was
- 312 originally described as *Draconema eira* by *Inglis (1968)*, later synonymized with *Dracograllus*
- 313 *eira*. Similarly, *Chaetosoma falcatum Irwin-Smith*, *1918* underwent multiple reclassifications
- before being recognized as *Dracograllus falcatus Allen & Noffsinger, 1978*. Another synonym is
- 315 Tristicochaeta falcata Johnston, 1938. Since Allen & Noffsinger (1978), and now, the valid name
- 316 is *Dracograllus falcatus*. See valid species and nomen nudum section.
- 317

318 Allen and Noffsinger (1978) first described *Dracograllus filipjevi* from holdfasts of kelps 319 from Japan (Oarai, Ibaraki-ken, Honshu Island). It was characterized by: (1) larger body size

- 320 (500-700 µm long), (2) the absence of cephalic acantiform setae, (3) the presence of
- 321 longitudinally areolated body cuticle with dot-like punctations, (4) the presence of some
- 322 cuticular collar in swollen pharyngeal region, and 9 sublateral and 8-9 subventral PAT in males,
- and 12-13 and 9-10 in females. *Rho, Kin & Kin, (2006)* also found *D. filipjev* associated with
- 324 calcareous algae in Daebo-ri, Guryongpo, Korea, at 3-5 m depth. However, the Korean
- specimens did not align well with the original description in the number of PATs in male with 8to 9 tubes, compared to 9 to 11 in the original description. Given that these characteristics are
- 327 crucial for the taxonomy of Draconematidae, this discrepancy supported the redescription made
- 328 by *Rho*, *Kin & Kin*, (2006).
- 329

Analyzing the Draconematidae from Guryongpo (Daebo-ri, Korea), *Rho & Kim (2011)* reported several species of the genus *Dracograllus*. However, these species are considered invalid, meaning they do not comply with certain taxonomic criteria required for formal recognition (see Article 16.1 of the International Code of Zoological Nomenclature). According to this article, every new name published after 1999 must clearly indicate its new status using specific terms such as 'fam. nov.,' 'gen. nov.,' 'sp. nov.,' or an equivalent expression (e.g., 'species nova,' 'new species').

337

338 List of valid species.

- 339 Dracograllus antillensis Decraemer & Gourbault, 1986
- 340 Dracograllus chitwoodi Allen & Noffsinger, 1978
- 341 Dracograllus cobbi Allen & Noffsinger, 1978
- 342 Dracograllus cornutus Decraemer, 1988
- 343 Dracograllus demani Allen & Noffsinger, 1978 [Decraemer, 1988; Verschelde & Vincx, 1993]
- 344 Dracograllus eira (Inglis, 1968) Allen & Noffsinger, 1978; [Decraemer, 1988; Verschelde &
- 345 Vincx, 1993]
- 346 Syn. Draconema eira Inglis, 1968
- 347 Dracograllus falcatum (Irwin-Smith, 1918)
- 348 Syn. Chaetosoma falcatum Irwin-Smith, 1918

- 349 Syn. Notochaetosoma falcatum (Irwin-Smith, 1918) Cobb, 1929
- 350 Syn. Drepanonema falcatum (Irwin-Smith, 1918) Cobb, 1933
- 351 Syn. Claparediella falcatum (Irwin-Smith, 1918) Filipjev, 1934
- 352 Syn. Draconema falcatum (Irwin-Smith, 1918) Kreis, 1938
- 353 Syn. Tristicochaeta falcata (Irwin-Smith, 1918) Johnston, 1938
- 354 Syn. Dracograllus filipjevi Allen & Noffsinger, 1978
- 355 Syn. Dracograllus gerlachi Allen & Noffsinger, 1978
- 356 Syn. Dracograllus gilbertae Verschelde & Vincx, 1993
- 357 Dracograllus grootaerti Decraemer, 1988
- 358 Dracograllus kreisi Allen & Noffsinger, 1978
- 359 Dracograllus laingensis Decraemer, 1988
- 360 Dracograllus mawsoni Allen & Noffsinger, 1978
- 361 Dracograllus minutus Decraemer, 1988 [Gourbault & Decraemer, 1992]
- 362 Dracograllus miguelitus sp. nov. Johnson et al., 2024
- 363 Dracograllus ngakei Leduc & Zhao, 2016
- 364 Dracograllus papuensis Decraemer, 1988
- 365 Dracograllus pusillus Decraemer, 1988
- 366 Dracograllus solidus (Gerlach, 1952) Allen & Noffsinger, 1978
- 367 Syn. Draconema solidum Gerlach, 1952
- 368 Dracograllus spinosus Decraemer, 1988
- 369 Dracograllus stekhoveni Allen & Noffsinger, 1978
- 370 Dracograllus timmi Allen & Noffsinger, 1978 [Gourbault & Decraemer, 1992]
- 371 Dracograllus trispinosum (Allen & Noffsinger, 1978) Decraemer, 1988
- 372 Syn. Dracotoranema trispinosum Allen & Noffsinger, 1978
- 373 Dracograllus trukensis Min et al., 2016
- 374 Dracograllus wieseri Allen & Noffsinger, 1978
- 375
- 376 Nomen nudum.
- 377 Dracograllus brevitubulus Rho & Kim, 2011 (unaccepted > nomen nudum)
- 378 *Dracograllus geomunensis Rho & Kim, 2011* (unaccepted > nomen nudum)
- 379 *Dracograllus gosanensis Rho & Kim, 2011* (unaccepted > nomen nudum)
- 380 Dracograllus jaewani Rho & Kim, 2011 (unaccepted > nomen nudum)
- 381 *Dracograllus jongmooni Rho & Kim, 2011* (unaccepted > nomen nudum)
- 382 *Dracograllus sungjooni Rho & Kim, 2011* (unaccepted > nomen nudum)
- 383 *Dracograllus chiloensis Clasing, 1980* (uncertain > taxon inquirendum)

384

- 385 **Description of** *Dracograllus miguelitus* sp. nov.
- 386 (Table 1; Figs. 2-5; S1-4)
- 387

388 Type material. All specimens are deposited in the Muséum National d'Histoire Naturelle de

389Paris, France. Male holotype, two juvenile paratypes and the female paratype in the inventory

- number MNNH BN511 I1 L1 B. Two male paratypes and female paratypes in MNHN BN511 I2 L1 A.
- 392

Other material. Other specimens are held in the collection of the Laboratoire Environnement
 Profond of the Biologie et Ecologie des Ecosystèmes marins Profonds research unit- Ifremer,
 Plouzané, France.

396

Etymology. The specific epithet is in honor of 'Pedro Miguel', nephew of the first author.

398

Type locality and habitat. Lucky Strike vent field - MAR. Samples were collected from a hard

substratum covered by a thin layer of volcaniclastic sediment, on a visually inactive vent

structure at 1.639 m depth. Environmental conditions exhibited background or slightly higher

seawater temperature (i.e. 4.8-5.7°C) and higher pH (i.e. 7.8-7.9) than the surrounding deep-sea
but the activity was very low compared to active habitats where recorded temperature varied

404 between 5.2°C to 9.5°C, reaching a maximum of 22.1°C, and pH varying from 7.2 to 7.6.

405

406 Measurements. Table 1

407

Holotype male. Habitus typical for the genus. 612 µm long, swollen anterior body region 408 representing 19% of total length (Figs. 2A, C). Amphid elongate loop-shaped with non-equal 409 410 branch sizes, and with one more ventrally than another, amphideal fovea 7.1 µm (Figs. 2C and 3C). Helmet strongly cuticularized (Fig. 2D), with punctations and granular appearance in the 411 lateral part (Fig. 3A). Annulation without ornamentation along the body, except for the tail tip 412 and helmet, with minute punctations (Figs. 2C, 2H, 3D). Four CATs on the rostrum (22.5-26.0 413 414 µm long), arranged dorsally in two transverse rows, all with enlarged bases (Figs 2C, 3A-B). Some specimens exhibited depressions resembling CAT insertions; however, none of these 415 depressions contained tubes. The setae in the cephalic region and along the body possess a 416 cuticular collar at their insertion and alternation of short and long setae, this collar is projected 417 418 outside of the cuticle, as a pedicel setae (PS), with 1.2-2.1 µm long (Figs. 2C, 3A-B), also in the paratype male (Fig. 6E). Cephalic and cervical region with 8 longitudinal rows of setae on each 419 side, between 26 and 32 µm long, and also some irregular minute setae (6-12 µm long) (Fig. 2C). 420 Slender cervical region without lateral differentiation (Fig. 2C and 3A). Buccal cavity narrow, 421 unarmed (Fig. 2B). Cardia short. Pharynx dumbbell-shaped with a weakly developed isthmus 422 423 from the muscular posterior large endbulb (Figs. 2D, 3A-B). Intestine narrow, mostly cylindrical, with a granular appearance, gradually widening posteriorly and lying dorsally to the reproductive 424 system (Fig. 3B-D). Reproductive system with a single and outstretched anterior testis 425 426 (monorchic) with a well-developed germinative region (Fig. 2G and 3B). Spicules 50.2 µm long, 427 moderately arcuate (in some specimens more arcuate than in others), proximal region with an

offset knob-like capitulum (Figs. 2E-F and 3F). Gubernaculum 13.6 µm long, lying parallel with
the distal end of the spicules, with a minute distal and lateral wing-like expansions (Fig. 3F).

PATs weakly slender, with tongue-like tips (Figs. 2E, 3B, D and E), difficult to observe 431 432 due to their thickness. All PATs located anterior to the cloacal opening (Figs. 2E, 3D). PATs are arranged in four longitudinal rows: two sublateral rows each with 10-12 PATs, and two 433 subventral rows consisting of 10 PATs each one (Figs. 2E, 3D), intermingled setae are present 434 between the PATs. Dorsal posterior part with small and irregular distributed setae between the 435 setae following the rows of the body, more visible than in the ventral side, where only few and 436 437 minute intermingled setae are present (Figs. 3D). Two pairs of setae are present in the annulated tail region. One pair of long setae in the last annules of the dorsal part of the tail, each setae with 438 a collar at the base, close to each other (48-50 um long), ventrally, one pair of shorter setae, with 439 collar at the basis (11-12 µm long; Figs 2H, 4B-D). Additional five setae are present in the tail. 440 441 in each somatic row of setae. One to two minute setae in the middle of the non-annulated tail tip 442 $(1.5 \,\mu m \log)$, in some specimens, some of the setae were absent, but the minute insertion of them is visible. 443

444

445 **Paratype female.** Similar to males in most aspects, but differ in the following characteristics: greater length, with 765.5 µm long (748-788 µm long in paratypes female and 630-735 µm long 446 in paratypes males: Fig 4A). Amphid also elongated loop-shaped, however slightly wider than in 447 males. In addition, amphid in females is positioned more anteriorly than in males (Figs. 4C, 5B). 448 Pedicel setae at both lateral and ventral side of the head, distributed for all body length and more 449 450 developed than in males (2.5-3.2 µm long; Figs. 4C, 5B). Swollen pharyngeal region 18% of total body length. Some of the setae appear to be lost, with only the cuticular collars remaining, 451 distributed irregularly and in smaller various sizes than in the head (Fig. 6A-B). Just some 452 collars, without setae, are also present in the swollen head region and also on the helmet. Both 453 454 the anterior and posterior regions of the pharynx vary between sexes. The anterior pharyngeal bulb in females has a diameter of 24-26 µm, and the posterior bulb has a diameter of 34-38 µm 455 (compared to 20-22 µm long and 29-33 µm long respectively in males). 456

457

458 Reproductive system didelphic-amphidelphic with reflexed ovaries, both located ventrally relative to the intestine. Uterus filled with a mass of ovoid reproductive cells (Fig. 4F, 459 circles). The region surrounding the vulvar aperture protrudes outward, with the cuticle giving a 460 labial appearance (Fig. 4E). Two pairs of paravulvar setae present, one anterior and one posterior 461 to the vulvar aperture, with length between 6.09-6.69 µm long. Also a setae emerging from the 462 463 vulvar aperture (5.03 µm long; Figs. 5A and E). Well-developed contractor muscles in the vagina (Fig. 4F). PATs all anterior to the anus, more slender, but shorter than in males (56.8-58.8 versus 464 63.4-68.4 in males), with weakly developed bell shaped tips with a tongue-like valve. PATs 465 arranged in four longitudinal rows: two sublateral rows each consisting of 13 adhesion tubes with 466 467 intermingling and irregular somatic setae and two subventral rows of 13 adhesion adhesion tubes

- also with intermingling somatic setae (Fig. 4G). First SIAT on the females with 58.9 µm long
 and 62.5 on males. All PATs weakly broadened at insertion base. Tail gradually tapering
 posteriorly to a cylindro-conoidal non-annulated tail tip (Figs. 4H, 5A and D). One pair of long
 setae in the last annules of the dorsal part of the tail (Fig. 4H, circle and 5D). Each setae with a
 collar at the base and close to each other (50.44 µm long). Ventrally, there is one pair of short
 setae with also a collar at the base (11-13 µm long). Two pairs of setae are present on the last
- 474 annules of the tail, each setae featuring a collar at its base and positioned close to one another,
- 475 measuring 52-56 μm in length (Fig. 5D). Additionally, there is another pair of shorter setae, also
- 476 with collars at their bases, measuring 12-14 μ m in length (Figs. 4G, 4H, 5D). Five more setae are
- 477 distributed along the tail, in similar size as in males, in each somatic row of setae. One or two
- 478 minute setae (1.5 μ m long) are located at the non-annulated tail tip. In some specimens, one of
- these setae (and also for those on the annulated tail) is absent, though its minute insertion point
- 480 remains visible (Fig. 4H). Non-annulated tail tips are long, constituting 56-59% of total tail
- 481 length. Caudal glands not evident.
- 482

483 Juveniles paratypes

- Juvenile third stage. Body shape similar to adults. Body length 426.3 μm long, head diameter
 19.3 μm long and a pharynx 75.3 μm long, with a minimally developed isthmus (Fig. 6A).
- 486 Swollen region representing 24% of the total length. Amphideal fovea is smaller in both length
- 487 (9.2 um) and width (4 um), circular and closed-shape (Fig. 6C). Several pedicel setae with 2.18
- μ m long, in some specimens as long as in some in adults (Fig. 6C). Two CATs in the dorsal
- region of the helmet, all at the level of the amphid ($15.89-22.24 \mu m \log g$) (Fig. 6C). 5 PATs in
- 490 both subventral and sublateral row (Fig. 6A, 6D). Tail slender, with the non-annulated tail tip
- 491 corresponding to 50.7 % of the tail length, similar to that in adults (Fig. 6D). The cuticle of the
- 492 non-annulated tail tip end ornamented with minute punctations. All observed juveniles exhibit a
- 493 globular appearance on the lateral sides of the body between the body wall and the cuticle,
- sometimes with brownish or yellowish coloration in the pharyngeal and anterior region.
- 495

Juvenile fourth stage. Body shape similar to adults. Body length 514.3 μm long, with head

- 497 diameter 24.3 μ m long. Swollen region representing 25.2% of the total length (Fig. 6B).
- 498 Amphideal fovea 80.6 μm, circular and closed, similar to the third stage. Presence of pedicel
- 499 setae with 2.82 μ m long. Three CATs located on the helmet with the longer one and more
- 501 μ m long and first SvAT with 44.3 μ m long (Fig. 6B).
- 502

503 Diagnosis.

- 504 D. miguelitus sp. nov. is characterized by the presence of four CATs located in the dorsal side of
- the helmet at the level of the amphid. All CATs situated anterior to the cuticular annulations.
- 506 Amphid elongate loop-shaped with different branch sizes in males and females and circular in
- 507 juveniles. A collar is present at the base of some setae, as a pedicel-like structures. Paravulvar

- setae are present in two pairs, one anterior and one posterior to the vulvar aperture. Additionally,
- a minute setae is visible parallel to the vulvar aperture. In the males, PATs are arranged in four
- 510 longitudinal rows: two sublateral rows each with 10-12 CATs, and two subventral rows
- 511 consisting of 10 PATs each one. In the females, sublateral and subventral rows with 13 CATs
- 512 each one. Two pairs of setae in the annulated part of the tail, one pair with long setae (40-50 μ m)
- and one pair with shorter setae (11-13 μ m). The non-annulated tail tip corresponds to 40-44% in
- 514 males, 56-59% in females, and 50-55% in juveniles.
- 515

516 Differential diagnosis and relationship

- 517 *Dracograllus miguelitus* **sp. nov.** is immediately distinguished from its congeners by possessing
- 518 only four cephalic adhesive tubes (CATs) located at the level of the amphid, in contrast to
- 519 species with six CATs (*D. minutus*), eight CATs (*D. antillensis, D. chitwoodi, D. cobbi, D.*
- 520 *demani*, *D. eira*, *D. filipjevi*, *D. gilbertae*, *D. grootaerti*, *D. kreisi*, *D. laingensis*, *D. mawsoni*, *D.*
- 521 ngakei, D. papuensis, D. pusillus, D. solidus, D. spinosus, D. timmi, D. trispinosum, D. trukensis,
- and *D. wieseri*), or more, such as *D. cornutus* (11 CATs), *D. falcatus* (12 CATs), *D. gerlachi* (13
- 523 CATs), and *D. stekhoveni* (14 CATs).

524 525 The absence of cuticular ornamentation further differentiates D. miguelitus sp. nov. from species with spines (D. antillensis, D. chitwoodi, D. grootaerti, D. minutus, and D. trukensis) or 526 527 dot-like punctations (D. filipjevi, D. gerlachi, D. kreisi, D. pusillus, and D. trispinosum). The 528 cuticle of D. miguelitus sp. nov., with the collars at the bases of its setae, resembles those 529 observed in D. cobbi, D. mawsoni, D. filipievi, and D. timmi, though the setae in these species 530 are significantly smaller compared to those in *D. miguelitus*. The presence of paravulvar setae 531 distinguishes D. miguelitus sp. nov. from several species, including D. chitwoodi, D. cobbi, D. cornutus, D. filipjevi, D. grootaerti, D. minutus, D. ngakei, D. pusillus, D. solidus, D. spinosus, 532 D. stekhoveni, D. timmi, D. trispinosum, and D. trukensis, all of them lacking setae at the vulva. 533

534

Dracograllus miguelitus **sp. nov.** is geographically closest to *D. demani* and *D.* 535 *trispinosum* but can be distinguished from these species by several morphological features. See 536 537 Tables 2 and 3 for the morphometrical and descriptive comparison between all the valid species 538 In addition to the number of cephalic adhesive tubes (CATs) – four in *D. miguelitus* sp. nov. 539 compared to eight in both D. demani and D. trispinosum – the new species differs in the number 540 of sublateral adhesive tubes (10-12 in *D. miguelitus* **sp. nov.** versus 6 in *D. demani* and 10 in *D.* trispinosum), subventral adhesive tubes (10 in both D. miguelitus sp. nov. and D. demani, but 541 seven in D. trispinosum), and spicule length (50 µm in D. miguelitus sp. nov., compared to 37 542 543 µm in D. demani and 61 µm in D. trispinosum). Additionally, the non-annulated tail tip of D. 544 *miguelitus* sp. nov. is longer (44% of body length in the male holotype and 56% in the female paratype) compared to the shorter tail tips in *D. demani* and *D. trispinosum* (32% and 26%. 545 respectively). Morphometrical and descriptive comparison between all the valid species (Tables 546 547 2 and 3).

548

549 **Comments on the imaging approach**

550 Several challenges related to the study of marine nematodes have been discussed here, and we

- would like to emphasize one of the most important ones: the difficulty in observing and
- 552 measuring their morphological structures. Therefore, it is crucial to investigate advanced

- 553 imaging methods to facilitate identification and capture additional morphological features
- 554 (Foulon et al., 2025 in press). Fluorescence based 3D microscopy was already used for
- nematode taxonomy, such as on the occasion of the redescriptions of *Craspodema reflectans*
- 556 (Cyatholaimidae) and *Longicyatholaimus maldivarum* (Cyatholaimidae) by *Semprucci* &
- 557 Burrattini (2015) and Semprucci et al., (2017), respectively. Additionally, an introduction to the
- application of confocal techniques for observing marine nematodes is provided in *Semprucci et*
- 559 *al.*, (2016). In our study, 3D fluorescence imaging has proven effective in several key aspects of
- identification, particularly for the Draconematidae family. We observed that various structures,
- that are difficult to study with traditional microscopic techniques, were analyzed with relative
- ease in our study. These included the insertion of the CATs, the cuticular ornamentation or
- annulations, and the number of rows of setae. Videos of the 3D fluorescence captures and
- additional pictures are available in the supplementary material (S1-4).
- 565

566 Biodiversity, distribution and ecology

In terms of species diversity, *Dracograllus* is the largest genus in the Draconematidae family, 567 with 26 valid species (Nemvs eds., 2024). These species are distributed across a variety of marine 568 environments, and despite their large distribution, the genus is predominantly associated with 569 shallow, tropical and subtropical regions (*Min et al., 2016*). The distribution and general 570 ecological characteristics of all *Dracograllus* valid species are presented in Table 4. The 571 Dracograllus genus includes species distributed across the Atlantic (13 species). Pacific (9 572 species), and Indian (4 species) oceans. Some species, such as D. eira, occur in multiple oceanic 573 regions, underscoring their adaptability to diverse oceanic regimes. Distribution of all valid 574 575 species, and species occurrences including non-identified Dracograllus specimens are provided in Figure 7. The Pacific Ocean is the region where the highest number of Dracograllus species 576 have been both recorded and described, likely reflecting a bias due to a more extensive sampling. 577 Examples include D. cornutus, D. falcatus, D. filipjevi, D. gerlachi, D. grootaert, D. laingensis, 578 579 D. mawsoni, D. minutus, D. papuensis, D. pusillus, D. spinosus, D. timmi, D. trukensis, and D.

- 580 *wieseri* from a variety of habitats and environmental conditions, particularly in coastal regions.
- 581 More recently, *Leduc & Zhao (2016)* described *D. ngakei*, a species from intertidal coarse sand
- and gravel sediments in New Zealand, including molecular and morphological data.
- 583
- In the Atlantic Ocean, species such as *D. antillensis*, *D. chitwoodi*, and *D. kreisi* are
- typically found in shallow marine environments, often associated with sandy beaches on
 intertidal or subtidal zones. *Decraemer & Gourbault (1986)* found approximately 500
- intertidal or subtidal zones. *Decraemer & Gourbault (1986)* found approximately 500
 individuals of *D. antillensis* in samples from Guadeloupe, a notably high number for a single
- individuals of *D. antillensis* in samples from Guadeloupe, a notably high number for a single
 species, especially when compared to the abundances typically observed in *Dracograllus* and
- species, especially when compared to the abundances typically observed in *Dracogratus* at
 even within the broader Draconematidae family. *Allen & Noffsinger* (1978) described D.
- 509 even within the broader Diaconematidae family. Allen & Nojjsinger (1978) described h
- *trispinosus* at 20 meters depth, revealing the species occurrence in subtidal zones.
- 591

592 The Indian Ocean hosts fewer described species, yet their habitats share similarities with those in other regions. For instance, D. demani has been reported in tidal coral sands along the 593 Malindi coast and also, in similar sandy habitats in the Pacific Ocean. Likewise, D. eira is known 594 from both the Atlantic Ocean, where it inhabits subtidal sandy zones, and the Indian ocean, 595 596 where it has been recorded in mangrove-associated sediments, demonstrating its ability to thrive in a range of coastal habitats. D. solidus, another widespread species, has been documented in 597 the Atlantic, Pacific, and Indian oceans, consistently associated with coarse sand in subtidal 598 599 regions.

600

601 Only a few studies have investigated the spatial distribution of *Dracograllus* in deep-sea ecosystems. Four morphotypes were found at the summit of the GSM in the Atlantic, which is a 602 flat plateau covering more than 1 400 km², with 293 to 511 meters depth (Pfannkuche, Sommer 603 & Kähler, 2000). They exhibit significant abundance compared to other Draconematidae genus, 604 605 with non-overlapping occurrences between each morphotype. The specific sedimentary processes on the GSM (Levin & Nittrouer, 1987), combined with the erosion of old coral reefs, 606 create a coarse sedimentary environment, with small and morphologically complex biogenic 607 structures covering the substratum (Pasenau, 1971; Nellen, 1998). This wide variety of 608 ecological niches would explain their high abundance and the co-occurrence of several species. 609 Similarly, Zeppilli et al., (2014) reported several Dracograllus specimens at the Condor 610 Seamount (CS), at 206 m depth, in the Azores archipelago (Northern Atlantic). The summit of 611 this structure exhibited a highest species richness and dominance for several genus, clearly 612 differing from the surrounding deep-sea habitats or along other CS habitats. The summit was the 613 614 only area of the study where *Dracograllus* sp1 was present, a flat region, covered by biogenic structures such as sponge sediments or corals, as observed for most Dracograllus species. 615 Similar results regarding Draconematidae species were obtained on other biogenic and 616 sedimentary habitats (Willems et al., 1982; Ndaro & Olafsson, 1999; Raes & Vanreusel, 2006; 617 618 Raes, et al., 2007 and Raes, Decraemer & Vanreusel, 2008).

619

620 Draconematidae species were recently observed in samples collected from two deep hydrothermal vent fields, TAG and Snake Pit (SP) (Spedicato et al., 2020) located on the Mid-621 622 Atlantic Ridge. Dracograllus sp. was present in 50% of the SP samples, occurring in reddish sediments covered by polychaete tubes. In contrast, these features were absent or less evident at 623 TAG, where only Cephalochaetosoma was recorded. Environmental conditions differed 624 significantly between the vent fields. The total sulfur content in the sediment profiles (0-5 cm) 625 was higher at SP than at TAG and the oxygen penetration about ten times lower at SP. High 626 concentrations of sulfur can lead to death due to the inhibitory action of H₂S on cytochrome c 627 oxidase, an essential enzyme for aerobic respiration. This mechanism blocks the electron 628 transport chain, disrupting ATP production and resulting in metabolic collapse (Bagarinao, 629 1992). However, some nematode species have developed strategies to cope with sulfide toxicity, 630 such as the oxidation of H₂S into elemental sulfur and its deposition in the epidermis, a process 631

632 observed in Oncholaimus campylocercoides (Thiermann, Vismann & Giere, 2000). This ability

- to accumulate and later remove elemental sulfur may enable nematodes to colonize sulfide-rich
- environments, exploiting niches where most organisms cannot survive. Moreover, body
- 635 elongation and a higher surface-to-volume ratio may help them cope with low O_2 levels in
- habitats with limited oxygen availability (*Vanreusel et al., 2010b*).
- 637

The non-overlapping distribution of Dracograllus species at deeper sites suggests that 638 each species may have specific habitat requirements, influenced by both the nature and 639 composition of the substratum, as well as the level of hydrothermal activity. The type-habitat of 640 D. miguelitus sp. nov., is characterized by low hydrothermal influence compared to active sites 641 at the Lucky Strike (LS) vent field (*Chavagnac*, 2018). However, residual venting activity is still 642 present, evidenced by the presence of manganese oxide-hydroxide and high CH4 concentrations 643 measured above the substratum in one of the samples. Environmental conditions, including 644 645 sediments rich in sulfide minerals, can stimulate microbial communities, which are essential as primary producers in these deposits (Van Gaever et al., 2009). This creates a higher food 646 resource availability and provides structural conditions suitable for the occurrence of 647 Draconematidae species, including D. miguelitus sp. nov., classified as microbial feeders. In 648 649 summary, the residual hydrothermal activity likely promotes microbial growth, a significant food source for bacterivores and microbial feeders like D. miguelitus sp. nov. and most 650 Draconematidae species. Moreover, the presence of biogenic structures like microbial mats, 651 appear to play a role in the distribution of *Dracograllus* species. These species are capable of 652 using adhesive tubes to anchor to these structures and may also feed on them (Raes et al., 2007). 653 Prior to the description of D. miguelitus sp. nov., the Snake Pit species were the deepest-known 654

- representatives of the genus, found at depths between 3.480 3.570 m.
- 656

657 Biogeography and evolutionary perspectives

- 658 Several species of *Dracograllus*, and Draconematidae in general, have been found in only a few
- 659 locations beyond their type habitats. However, nematologists agree that cosmopolitanism is
- 660 common among various species and groups of marine nematodes (Decraemer, Gourbault, N., &
- 661 *Helléouet, 2001*). As reflected by Gad (2009), based on Draconematidae species, one important
- 662 starting point to determine the origins of these species is to identify their closest relatives and
- 663 where they occur. In fact, some of the closely related species of *Dracograllus miguelitus* **sp. nov.**
- 664 (*D. demani* and *D. minutus*) inhabit coastal and sublittoral environments of the Mediterranean
- 665 Sea. These Mediterranean regions could be the source of this species, as surface currents
- transport waters from Gibraltar toward the Azores and upper regions of the North Atlantic
- 667 (Dietrich et al., 1975; Brenke, 2002). Drifting-buoy experiments have confirmed that
- 668 Mediterranean water eddies travel westward from Gibraltar across the Atlantic (Richardson,
- 669 *1996*). Such westward flows also occur at approximately 900 m depth, facilitating the transport
- of fauna, including meiofauna, which may drift as eggs, juveniles, or adults attached to marine
- 671 snow (Pingree, García-Soto & Sinha, 1999; Gad & Schminke, 2004). This may also be the case

672 for some *Prochaetosoma* species, as congeners have also been recorded in the Mediterranean.

673 Other potential source regions, such as the coasts of Mauritania or Morocco, remain unconfirmed

due to the absence of Draconematidae records from these areas. If such transport occurred, it

- would also depend on Mediterranean water flows (*Gad*, 2009).
- 676

Studies by Gad (2009) on the Great Meteor Seamount, Zeppilli et al., (2014) on the 677 Condor Seamount, and Spedicato et al., (2020) on the TAG and Snake Pit vent fields identified 678 closely related species in various deep-sea habitats along the northern MAR. Fifteen 679 Draconematidae species, spanning several genera—including Draconema, Paradraconema, 680 *Eudraconema*, *Prochaetosoma*, *Cephalochaetosoma*, and *Tenuidraconema*—were recorded on 681 the GMS plateau, located 500 km from the Lucky Strike vent field. Remarkably, fourteen of 682 these species were new to science, including four *Dracograllus* morphotypes (Table 5). In the 683 CS (~300 km from LS), a rich and exclusive nematode community was documented, with 35 684 685 species distributed across genera such as Akanthepsilonema, Apenodraconema,

686 *Bathychaetosoma, Dracograllus*, and others. Similarly, *Spedicato et al., (2020)* observed several 687 Draconematidae specimens from three genera: *Cephalochaetosoma, Dinetia*, and *Dracograllus*.

688

689 Both hydrothermal vents and seamounts can be considered true oases of life compared to the surrounding deep-sea environment (*McClain et al.*, 2010), emphasizing their importance for 690 biogeographic studies, particularly for taxa that exhibit some degree of habitat exclusivity, as 691 observed in Draconematidae in the North Atlantic. Another intriguing aspect of Draconematidae 692 in these regions is their morphological variability, which may reflect underlying biogeographic 693 694 processes (Costello & Chaudhary, 2017). For example, in Dracograllus species from the GMS, individuals from the southern part of the plateau possess a fully divided cephalic capsule 695 (helmet), whereas those from the northern part have a partially divided one (Gad, 2009). 696 Additionally, there are variations in the number of SIATs and SvATs. Several other distinctive 697 698 traits were reported, including the presence of eight strong spines around the vulva in Draconema sp. 1, a long and conical cephalic capsule in Cephalochaetosoma sp. 10, and extra-699 wide annules in the pharyngeal region of *Prochaetosoma* sp. 12. None of these distinctive traits 700 were observed in Draconematidae species from LS (Tchesunov, 2015; W Johnson, 2025, 701

702 *unpublished data*).

703

704 The intrageneric variation in the helmet among *Dracograllus*, along with the nonoverlapping distributions of several Draconematidae genera and species across the CS, GMS, 705 Snake Pit, and LS, and this may be related to an ongoing speciation process (George, 2004; Gad, 706 707 2004; Gad, 2009), similar to what was observed by George & Schminke (2002) and Gad & Schminke (2004) in copepods and macrofaunal species, respectively. In fact, when closely 708 related species exhibit significant morphological variations within small geographic regions, it 709 710 suggests that species may be arising through micro-allopatric speciation, where populations 711 diverge due to localized environmental differences, leading to subtle-but sometimes crucial-

morphological distinctions (*Rundle & Nosil, 2005*). As these populations adapt to specific

recological niches, genetic divergence and reproductive isolation may drive the emergence of new

species, highlighting the importance of understanding local biodiversity and the environmental

715 factors influencing species differentiation.

716

Given the known limitations of morphology-based taxonomy—such as cryptic diversity
and convergent evolution—future studies integrating molecular markers, such as COI or 18S,
will be crucial for validating the observed patterns and refining our understanding of species

connectivity and dispersal (*Palmer, 1988a; 1988b; De Ley et al., 2005; Bhadury et al., 2006;*

- 721 Derycke et al., 2010; Curini et al., 2012; Ahmed et al., 2015; Martínez García et al. 2023).
- 722 Despite these challenges, our findings, together with the limited existing data on species
- 723 distributions, suggest that both oceanic currents and local conditions and adaptations may play a
- role in shaping Draconematidae distributions. This highlights the need for further
- interdisciplinary approaches to fully elucidate the evolutionary and ecological processes
- 726 governing meiofaunal diversity in deep-sea environments.
- 727

728 Inactive vent structure remarks and conservation implications

Hydrothermal vents have been the focus of numerous ecological studies since their discovery in 729 730 1977. These investigations have significantly enhanced our understanding of the structure and dynamics of benthic communities and the role of environmental conditions at various spatial and 731 732 temporal scales (Godet, Zelnio, Van Dover, 2011). These habitats are known for their unique biogeochemical characteristics, which include commercially valuable mineral resources such as 733 734 iron, copper, and zinc (Van Dover, 2019). However, the prospect of mining these sites poses serious environmental threats, including permanent alterations in the local topography and 735 removal of habitats (Boschen et al., 2013). Furthermore, mining could release toxic metals, 736 disrupt ecological functions, and hinder the recruitment and recovery of sessile invertebrates, 737

738 particularly in regions where hard substrata are limited (*Van Dover, 2019*).

739

740 While the fauna of active sites has been the focus of most vent studies, that of inactive sites is virtually unknown. Few studies report the presence of filter-feeders on the relief created 741 742 by these mineral-rich mounds (Boschen et al., 2013; Van Dover, 2019). Moreover, it is suspected that these mineral deposits may host totally different communities than those found at active 743 vents. This is supported by an eDNA study by Cowart et al., (2020) on the Lucky Strike vent 744 field, which observed significantly higher diversity (OTUs - Operational Taxonomic Units) in 745 both inactive and peripheral regions compared to active ones, as well as notably distinct 746 747 communities among the active, inactive, and peripheral areas. While inactive vent systems differ from active ones, both of them face significant threats from deep-sea mining, with potentially 748 severe consequences for biological communities and ecosystem functioning. This challenge is 749 750 exacerbated by the limited knowledge on these habitats and their associated communities, 751 particularly meiofaunal organisms, which are often overlooked in ecological studies. These



- knowledge gaps may hinder the development of effective management and conservationstrategies (*Menini et al., 2023*).
- 754

By documenting the species present at inactive vents, researchers can better assess their ecological roles and connections with neighboring active systems. The discovery of *Dracograllus miguelitus* **sp. nov.** at an inactive vent structure exemplifies the biodiversity hidden in these understudied environments and highlights the urgent need for species-level research. Such findings are crucial for balancing conservation priorities with industrial ambitions, ensuring that management strategies are grounded in a comprehensive understanding of ecosystem

- 761 dynamics and connectivity.
- 762

763 Conclusions

The discovery of a new nematode species not only provides valuable taxonomic and ecological

data on a poorly studied genus but also underscores the ecological significance of inactive
 hydrothermal structures. These habitats increasingly warrant attention in the face of deep-sea

767 mining threats. Future research should aim to further investigate the biodiversity and ecological

768 roles of nematodes and other meiofauna in inactive vent ecosystems, integrating these findings

- 769 into conservation and management strategies.
- 770

771 While these findings advance our understanding of vent nematode biodiversity, the study is

772 limited to a single structure. Broader exploration across diverse hydrothermal regions and

inactive structures and areas is essential to fully understand the genus distribution, biogeography,

- and ecological roles. Notably, the presence of *D. miguelitus* **sp. nov.** on an inactive structure may
- result from dispersal events from nearby areas. This highlights specific adaptations to both
- substratum type and heterogeneity, as well as hydrothermal influences, which require further in-
- 777 depth study. In conclusion, this study emphasizes the importance of incorporating species-level
- data into hydrothermal vent research and highlights the urgent need for proactive conservation
- measures to safeguard the biodiversity of all types of hydrothermal habitats in the face of
- 780 increasing anthropogenic pressures.
- 781

782 Dichotomous key to Dracograllus valid species

783 The dichotomous key was constructed based on previous studies (Allen & Noffsinger, 1978;

- 784 Decraemer, Gourbault & Backeljau, 1997; Min et al., 2006). The complete list for the
- 785 description of valid *Dracograllus* species is listed in the reference section.
- 786
- **1.** Four CATs on rostrum ... *D. miguelitus* sp. nov.
- -More than four CATs on rostrum ... 2
- 789
- 790 2. Without sublateral cephalic acanthiform setae on rostrum ... 3
- -With one pair of bilateral cephalic acanthiform setae on mid-rostrum ... **D.** stekholveni

792		
793	3.	Males with 3-4 preanal corniform setae; 10-11 CATs. Females with 10 CATs: SIATn
794		18, SvATn 13–16 <i>D. cornutus</i>
795		-Males without preanal corniform setae; more CATs. Females with larger numbers of
796		PATs 4
797		
798	4.	Males with 7–8 short stiff setae in subventral rows just anterior to SvATl; spicules 39 μ m
799		long. Females with 24 SIAT, including 2 tubes posterior to the anus. Both sexes with a
800		swollen pharyngeal region representing 22% of the total body length D. gerlarchi
801		-Males with 3–4 short stiff setae in subventral rows just anterior to SvATI; spicules 71
802		μ m long. Females with 21 SIAT, including 3 tubes posterior to the anus. Swollen
803		pharyngeal region 13–14% of total length <i>D. falcatus</i>
804	-	
805	5.	Six CATs on rostrum; males with 5 SIAT, 2–3 SvAT; total length 290 µm; spicules 18
806		μ m <i>D. minutus</i>
807 808		-Eight CATs on the rostrum. Number of PAT higher in males; spicule typically long 6
808 809	6.	All CATs adjacent to or posterior to the amphideal fovea 7
810	0.	-All CATs anterior to the amphideal fovea <i>D. eira</i>
811		-All CATS anterior to the ampindear lovea D. eu a
812	7.	Males with preanal corniform setae; slender, conspicuously long and short SIAT
813		alternating in both sexes <i>D. trispinosum</i>
814		-Males without preanal corniform setae; SIAT without alternating long and short tubes in
815		both sexes 8
816		
817	8.	Several somatic setae in the posterior body region with spiny cuticular insertion and non-
818		annulated tail tip representing 59% of tail length D. spinosus
819		-Somatic setae with spiny insertion collar 9
820		
821	9.	Some somatic setae pedicellate; pedicels $1-8 \ \mu m \ long \ \dots \ 10$
822		-Somatic setae without pedicels 13
823		
824	10.	Males with 5–9 SIAT; females with 6–12 SIAT, all anterior to the anus; 9–14 SvAT11
825		-Males with 12–24 SIAT; females with 15 SIAT (1 posterior to anus) and 16 SvAT D .
826		mawsoni
827		
828	11.	Males with 5–7 SIAT; females with 9–13 SvAT 12
829		-Males with 9 SIAT; females with 14 SIAT D. cobbi
830		

831 832	12.	Males and females with pedicellate setae in ventrosublateral row just anterior to SIAT; spicules $45-53 \mu m$; females with $6-8 \text{ SIAT} \dots D$. <i>demani</i>
833 834		-Without pedicellate setae in ventrosublateral rows; spicules 36 μ m; females with 12 SIAT D. kreisi
835 836	13.	Annulated body cuticle without ornamentation14
837 838		-Annulated body cuticle ornamented with spines, dots, and vacuoles15
839 840	14.	Amphids long, inverted U-shaped in both sexes; males with 10 SIAT, 11 SvAT; spicules 29 μ m; females with 11–13 SIAT and 9–11 SvAT <i>D. papuensis</i>
841 842		-Amphids sexually dimorphic: loop-shaped in males, elongated unispiral in females; males with 7 SIAT, 11 SvAT; spicules 46 µm; females with 8–11 SIAT and same for
843 844		SvAT D. solidus
845	15.	Body cuticle with vacuolar and granular ornamentation D. wieseri
846 847		-Body cuticle ornamented with dots and spines16
848	16.	Body annules ornamented with two rows of dots 17
849 850		-Spiny ornamentation of the body cuticle 18
851 852	17.	Amphids long, oblique loop-shaped in females; tail slender (tail/abd = 5.6) <i>D</i> . <i>chitwoodi</i>
853 854		-Amphids inverted U-shaped in females; tail/abd = 3.9 <i>D. timmi</i>
855	18.	Amphids long, inverted U-shaped, as long as the rostrum 19
856		-Amphids short and wide, inverted U-shaped D. antillensis
857 858	19.	Short body (L = 310 μ m); faint rostrum ornamentation; body annules with minute spines; spicule 26 μ m; males with 6 SvAT <i>D. pusillus</i>
859		-Body > 400 μ m; spiny rostrum ornamentation; longer spicules; more than 6 SvAT in
860		males20
861	• •	
862 863	20.	Long swollen pharyngeal region; amphids inverted U-shaped in males and elongated unispiral in females <i>D. laingensis</i>
864		-Shorter, wider swollen pharyngeal region; amphids U-shaped in both sexes; spicules 68
865		μm D. grootaerti
866		
867	21.	Body annules with dot-like punctations; no anal flap; females with two pairs of
868		paravulvar setae (anterior and posterior to vulva, 5–6 μm) D. filipjevi
869		-Anal flap present; females with different number/position of paravulvar setae 22
870		

		I
871	22.	Large spaces between body annules; females without paravulvar setae D. gilbertae
872		-Body annules closely spaced; females without paravulvar setae 23
873		
874	23.	Body annules without ornamentation; paravulvar setae absent D. ngakei
875		-Paravulvar setae present; body annules with ornamentation 24
876		
877	24.	Body annules with numerous ridges and spiny protrusions, denser in lateral fields. Males
878		with large loop-shaped amphids with ventral branch longer than dorsal, extending to the
879		first body annule D. trukensis
880		
881	Abb	reviations
882	The te	erminology used for the description and measurements was according to Min et al., 2016,
883	Leduc	e & Zhao, 2016 and the classical approach by De Man, 1880. The abbreviations used are as
884	follow	VS:
885		
886	L	body length;
887	Α	ratio body length / body maximum width;
888	b	ratio body length / pharynx length;
889	c	ratio body length / tail length; c': ratio tail length / anal body diameter;
890	V%	position of the vulva as a percentage of the total body length from anterior;
891	Mdb	maximum body diameter;
892	(mdb	
893	mdb]	
894	ph	length of pharynx;
895	abd	anal body diameter;
896	t	tail length;
897	tmr	length of non-annulated tail terminus;
898	spic	length of the spicule measured along median line;
899	gub	length of gubernaculum;
900	CATS	
901		1
902	1SIAT	
903	SIAT:	
904 005	1SvA Svat	
905 006	SvAT PATs	
906	ГАІS	posterior adhesion tubes;
907		

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- 913

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Figure 1

Figure 1. Study site and sampling approach

A, Location of the Lucky Strike (LS) vent field along the Mid-Atlantic Ridge (MAR). B, location of the tree contrasting edifices. C-D, quadrats, faunal sampling and substratum view at the inactive habitat at LS. Source: Victor6000, Momarsat 2018, Ifremer. LS map modified from Zeppilli et al., 2019.

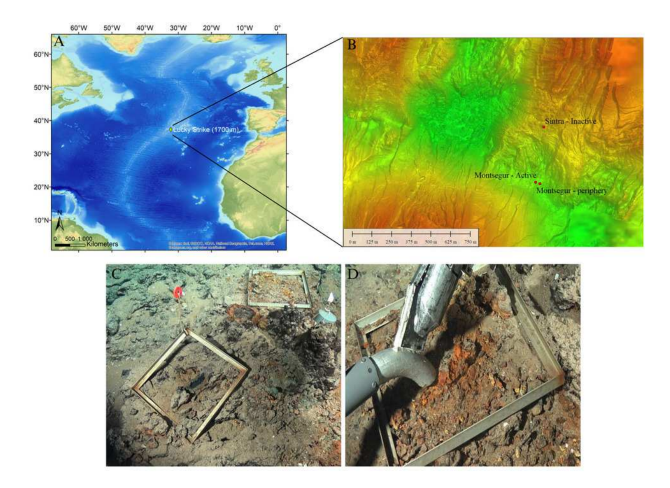


Figure 2

Figure 2. Dracograllus miguelitus sp. nov., 3D fluorescence microscopy

Male holotype: general view (blue, green and red fluorescent channels) (A). Anterior view of the buccal cavity (blue and red fluorescent channels - maximum intensity projection) (B). Swollen anterior region (blue, green and red fluorescent channels - maximum intensity projection), showing the CATs, the amphideal fovea and four of the longitudinal rows of setae and the pedicel setae (C). Internal view of the head region (blue fluorescent channel - optical section), with the well-cuticularized helmet, some of the CATs and the pharynx (D). Posterior male region (blue, green and orange fluorescent channels - maximum intensity projection), with both sublateral and subventral rows of PATs, their insertion (circles), spicule and gubernaculum (E). Ventral view of the posterior regions (blue, green and orange fluorescent channels - maximum intensity projection), with the arcuate spicules (F). Mid-mody (blue and red fluorescent channels - optical section) showing intestine and testis, circle indicates reproductive cells (G). Posterior tail region (blue fluorescent channel - maximum intensity projection), with the non-annulated tail region, and the setae associated (H). Arrows/Abrev: Buc. Cav, buccal cavity; CATS, cephalic adhesive tubes; PS, pedicel setae; Hel, helmet/cephalic capsule; Ph, pharynx; SIAT, sublateral adhesive tubes; SvAT, subventral adhesive tubes; Spic, spicule; Gub, gubernaculum; Test, testis; T. set, tail setae.

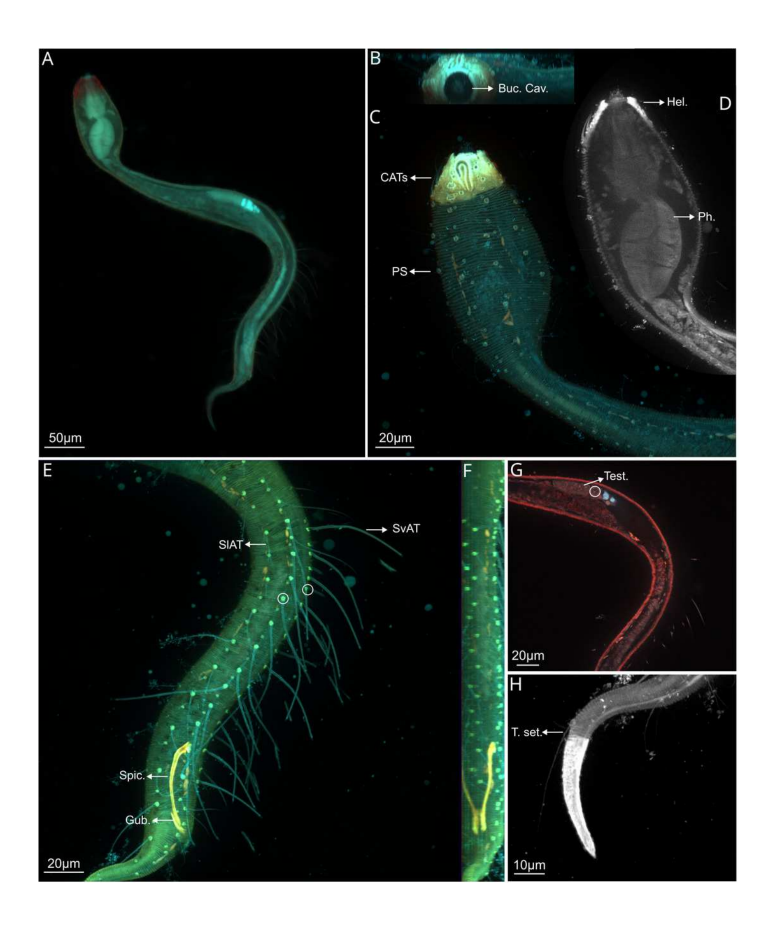


Figure 3

Figure 3. *Dracograllus miguelitus* sp. nov.

Male holotype: head (A), general view (B), fovea (C), posterior region, rows of PATs, not all tubes include (D), cervical cuticle regions and the 3 first SvATs and first SIAT, spicule and gubernaculum (F).

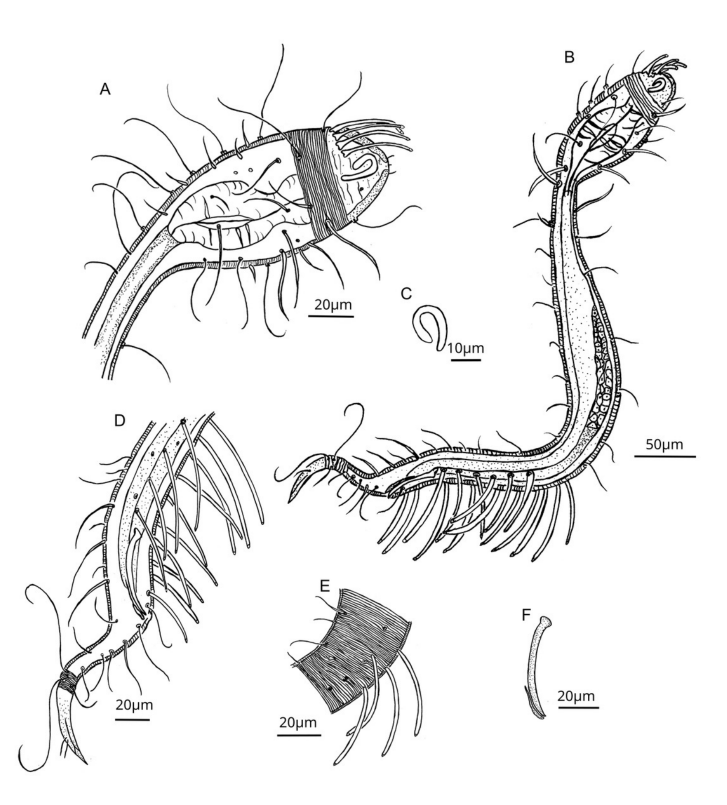
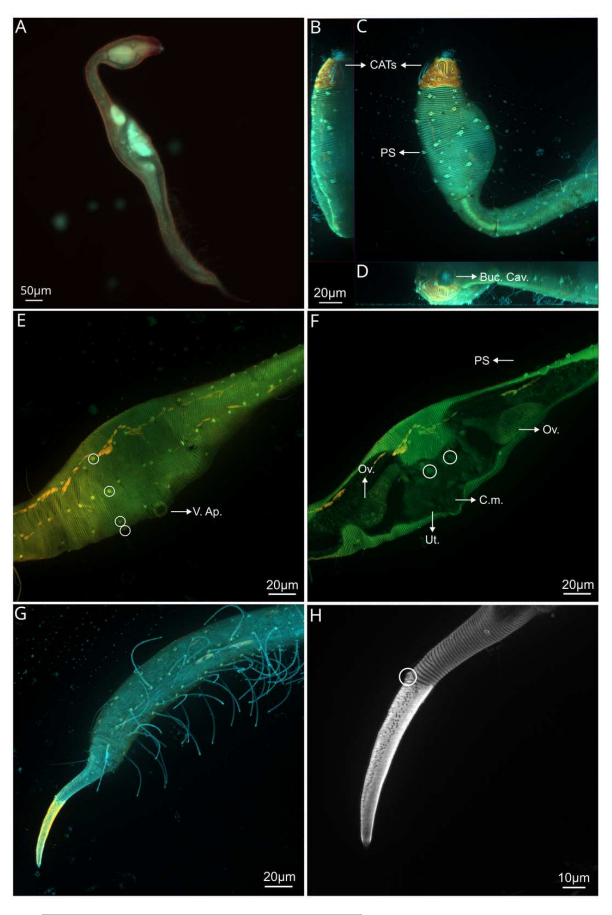


Figure 4

Figure 4. Dracograllus miguelitus sp. nov., 3D fluorescence microscopy

Female paratype: general view (blue, green and red fluorescent channels) (A). Dorsal view of the head (blue, green and orange fluorescent channels - maximum intensity projection), with the four CATs (B). Head and cervical region (blue, green and orange fluorescent channels - maximum intensity projection), with evident fovea and pedicel setae, note the clear helmet ornamentation (C). Face view of the buccal cavity (blue, green and orange fluorescent channels - maximum intensity projection), also CATs and fovea (D). Mid body region (green and orange fluorescent channels - maximum intensity projection), with the vulvar aperture, and four of the longitudinal rows of setae (circles) (E). Internal view of female reproductive system (green and orange fluorescent channels - maximum intensity projection), reproductive cells within circles (F). Posterior body region (blue, green and orange fluorescent channels - maximum intensity projection), with some of the both sublateral and subventral rows (G). Posterior tail region (blue fluorescent channel - maximum intensity projection), with the non-annulated tail region and a setae insertion (circle) (H). Arrows/Abbrev: CATs, cephalic adhesive tubes; PS, pedicel setae; Buc.Cav., buccal cavity; V. Ap., vulvar aperture; Ov., ovaries; Ut., uterus; C.m., constrictor muscles.

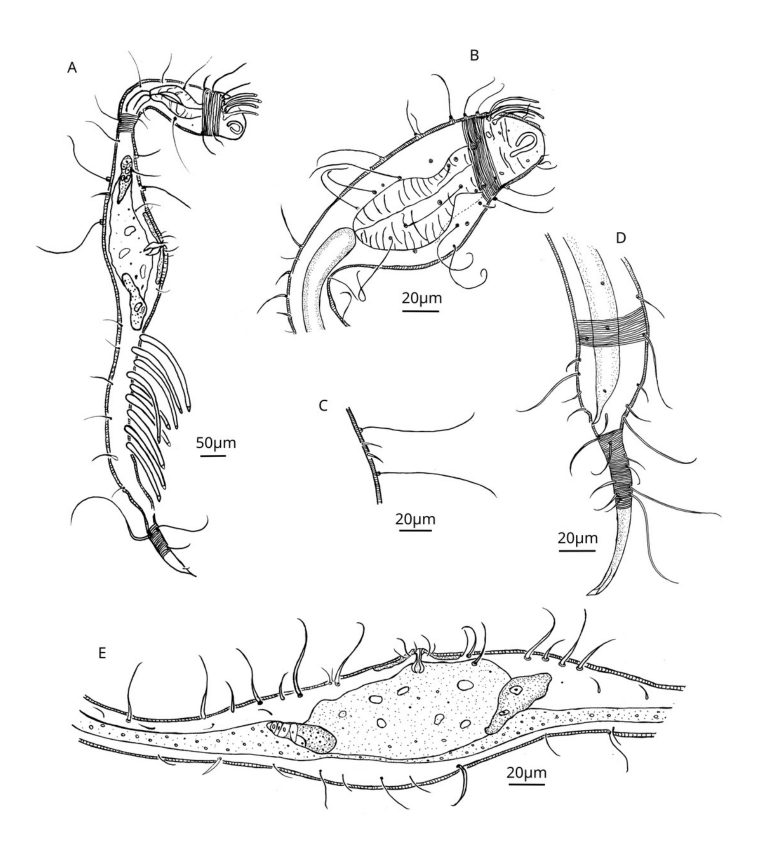


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Figure 5

Figure 5. Dracograllus miguelitus sp. nov .

Female paratype: general view (A); Head (B). Cuticle at cervical region (C). Posterior and tail region, female reproductive system (E).

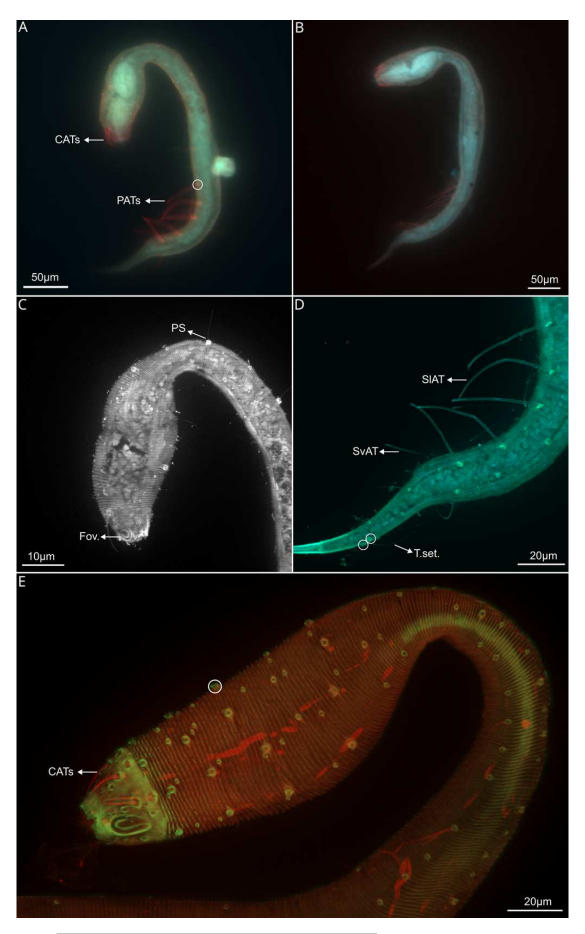


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Figure 6

Figure 6. *Dracograllus miguelitus* sp. nov., 3D fluorescence microscopy of the juveniles and a male paratype

General view of juvenile third stage (blue, green and red fluorescent channels) (A), showing both cephalic and posterior adhesive tubes. General view of the juvenile fourth stage (blue, green and red fluorescent channels) (B). Head and cervical region of juvenile third stage (blue fluorescent channel - maximum intensity projection) (C), arrows indicate the closed shape of fovea and the pedicel setae. Posterior and tail region of the third-stage juvenile (blue and green fluorescent channels - maximum intensity projection, with arrows indicating the rows of adhesive tubes and circles highlighting the insertion points of the tail setae (D). Head and cervical region of the paratype male (green and red fluorescent channels maximum intensity projection) (E), CATs on arrows and pedicel setae within the circle. Arrows/Abbrev: CATs, cephalic adhesive tubes; PATs, posterior adhesive tubes; PS, pedicel setae; Fov., fovea; SIAT, sublateral adhesive tubes; SvAT, subventral adhesive tubes; T.set., tail setae



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Figure 7

Figure 7. Global occurrence locations of the genus Dracograllus

Black dots indicate type localities of valid species (holotype names in bold, non-bold for paratypes). Colored circles represent occurrence locations of unidentified individuals or those classified as morphotypes (green points indicate these occurrences in shallow waters, 0-200 m, and blue points in deep-sea habitats, >200 m). Horizontal lines group morphotypes with overlapping occurrences or geographically close localities (e.g., all species described for the Papua New Guinea region). For the precise locations, habitats, sampling details, and remarks on each valid and undetermined species globally, refer to Tables 4 and 5, respectively. *Nomen nudum* and invalid species in general not shown, but available in the genus review section.

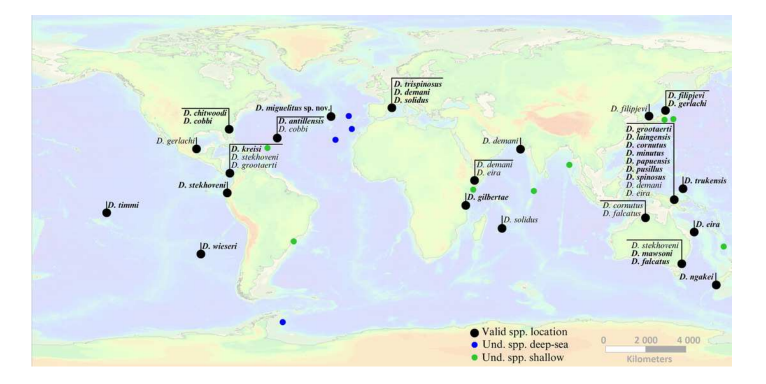


Table 1(on next page)

Table 1. Morphometric measurements (µm) of *Dracograllus miguelitus* sp. nov.

J3 and J4, juveniles third and fourth stages respectively; L, body length; a , ratio of body length to maximum body width; b , ratio of body length to pharynx length; c , ratio of body length to tail length; c' , ratio of tail length to anal body diameter; Amph./cbd (%), percentage of amphideal length relative to the corresponding body diameter; V%, position of the vulva as a percentage of the total body length from the anterior; mdb, maximum body diameter; (mdb), minimum body diameter at mid-body level; mdb ph, maximum body diameter in the pharyngeal region; ph, pharynx length; abd: body diameter; t: tail length; tmr: length of nonannulated tail tip; spic: length of the spicule measured along the median line; gub, length of the gubernaculum; CAT, cephalic adhesion tubes; 1SIAT1, length of the first subventral adhesion tube; SvATn, number of sublateral adhesion tubes; PAT, posterior adhesion tubes; P. setae, paravulvar setae length; V. b. diam., vulvar body diameter. *: At the level of the amphid; **: in each row.

	Μ	lales	F	'emales	Juve	Juveniles	
Parameter		Paratypes		Paratypes	Paratype	Paratype	
	Holotype	(n=2)	Paratype	(n=2)	J3	J4	
L	612	630-735	765.5	748-788	426.3	514.3	
а	10.6	13.7-14.9	11.6	12.0-12.8	14.2	10.1	
b	6.9	7.4-7.5	7.3	7.7-8.1	5.7	6.3	
С	6.2	6.2-7.0	8.1	8.4	6.4	6.9	
с′	4.4	4.7-5	4.5	3.8-4.1	3.9	4.4	
Head diam. *	32.4	27.2-33.92	31.1	29.6-34.37	19.3	24.3	
Amphid Length	15.5	14.2-15.2	14.4	12.7-13.5	9.2	10	
Amphid width	7.12	6.8-6.9	8.2	7.5-7.9	4	4.1	
Amph./cbd(%)	22.0	20.4-25.0	26.6	25.4-25.5	16.7	20.1	
Amphid from ant.	5.04	4.1-5	3.5	3.3-3.7	1.5	1.9	
Pharynx Leng.	88.5	84.0-99.5	93.3	92-103	75.3	80.6	
Phar. bulb diam. (ant.) Phar. bulb diam.	21.6	20.3-22.8	24.2	26.1-26.7	20.8	22.5	
(post.) Max. body diam.	30	29.3-33.5	36.7	34.1-38.5	24.2	28	
Phar	57.4	45.7-56.3	58.8	58.4-61.6	40.6	54.3	
Max. body diam. Mb	44.9	44.8-49.3	49.9	61.8-70	30.1	41.4	
Min. body diam.	11.7	16.5-18.59	20.8	20.1-25	20.2	20.6	
Spic. Leng.	50.2	47.1-54.9	-	-	-	-	
Gub. Apoph. Lengt.	13.6	12.7-14.5	-	-	-	-	
abd	22	19.9-22.2	18.3	19.7-22.9	17.1	17.3	
T. Leng.	97.2	100.8-105.2	84.2	88.6-94.3	66.6	75.9	
Non.Ann.T. Leng.	43.0	41.1-44	47.4	46.9-53	36.7	38.5	
Non.Ann.T. Leng. %	44.2	40.8-41.8	56.3	56-59	50.7	55.2	
T. Leng. / abd	4.4	4.7-5.0	4.5	3.8	3.8	4.3	
Longest tail setae	48.2	49	47.0	45.1-45.8	42.8	44.6	
CATn	4	4	4	4	3	4	
CATI	22.5 - 26.0	23.130.7	23.7	23.88-24.73	15.5	21.2	
1SIATI	62.5	63.4-68.4	58.9	56.8-58.8	47.4	50.4	
SlATn**	10	10-12	13	13	5	7	
1SvATl	50.0	53.3-54.3	51.5	50.9-50.2	42	44.3	
SvATn**	10	10	13	13	5	7	
V. to ant.	-	-	317.1	298.5-302.4	-	-	
V. (%)	-	-	41.4	39.9-41.3	-	-	
P. setae	-	-	5.0 - 5.9	5.5-6.1		-	
V. b. diam.	-	-	65.3	64.7-68.8	-	-	

1 Table 1. Morphometric measurements (µm) of *Dracograllus miguelitus* sp. nov.

2 J3 and J4, juveniles third and fourth stages respectively; L, body length; a, ratio of body length to

3 maximum body width; *b*, ratio of body length to pharynx length; *c*, ratio of body length to tail

4 length; *c'*, ratio of tail length to anal body diameter; Amph./cbd (%), percentage of amphideal

- 5 length relative to the corresponding body diameter; V%, position of the vulva as a percentage of
- 6 the total body length from the anterior; mdb, maximum body diameter; (mdb), minimum body
- 7 diameter at mid-body level; mdb ph, maximum body diameter in the pharyngeal region; ph,
- 8 pharynx length; abd: body diameter; t: tail length; tmr: length of non-annulated tail tip; spic:
- 9 length of the spicule measured along the median line; gub, length of the gubernaculum; CAT,
- 10 cephalic adhesion tubes; 1SIAT1, length of the first sublateral adhesion tube; SIATn, number of
- 11 sublateral adhesion tubes; 1SvAT1, length of the first subventral adhesion tube; SvATn, number
- 12 of subventral adhesion tubes; PAT, posterior adhesion tubes; P. setae, paravulvar setae length; V.
- 13 b. diam., vulvar body diameter. *: At the level of the amphid; **: in each row.

Table 2(on next page)

Table 2. Morphometrical comparison for all valid species of *Dracograllus* genus including *Dracograllus miguelitus* sp. nov.

L, body length; CATn, number of cephalic adhesive tubes; SIATn, number of sublateral adhesive tubes; SvATn, number of subventral adhesive tubes; Ms, type series males; M, male holotype; Fs, type series females; F, female holotype. L and spicule measurements expressed in µm.

1 Table 2. Morphometrical comparison for all valid species of the *Dracograllus* genus including

2 Dracograllus miguelitus sp. nov. L, body length; CATn, number of cephalic adhesive tubes;

3 SIATn, number of sublateral adhesive tubes; SvATn, number of subventral adhesive tubes; Ms,

4 smallest and bigger value among type series males; M, based on the male holotype; Fs, smallest

5 and bigger value among type series females; F, based on the female holotype. L and spicule

 $6 \qquad measurements expressed in \ \mu m. \ Non-annulated \ tail \ tip \ \%, a \ percentage \ of \ the \ non-annulated \ region$

7 in the total tail length.

Specie	L	CAT n	SlATn	SvATn	Spicul e	Non- annulated tail tip %
<i>D. antillensis</i> Decraemer &	Ms: 410-510	8	Ms: 6-10	Ms: 9-14	36-40	Ms:14-20
Gourbault, 1986	Fs: 410-510	8	Fs: 7-9	Fs: 8-12	30-40	Fs: 29-32
<i>D. chitwoodi</i> Allen & Noffsinger, 1978	Fs:500-600	8	Fs: 9-10	Fs: 8-10	-	Fs: 50-54
D. cobbi	M:500	8	M:9	M:12	51	M:44
Allen & Noffsinger, 1978	F:500	0	F:8	F:14	51	F:53
D. cornutus	Ms:495-610	10.11	Ms:16	Ms:13		Ms:26-28
Decraemer, 1988	Ms:480	10-11	Ms: 18	Ms: 16	55-56	Ms:49
D. demani	Ms:500-800	0	Ms: 5-7	Ms: 8-12	15 50	Ms: 24-39
Allen & Noffsinger, 1978	Fs:500-800	8	Fs: 6-8	Fs: 10-13	45-53	Fs: 41-51
D. eira	M:500	8	M: 12	M: 8	48	M: 48
(Inglis, 1968)	F: 600	0	Fs: 12	Fs: 8	40	F: 41
D. falcatus (Irwin-	M:800	12	M:12	M:17	71	M:32
Smith, 1918)	F:900	12	F:21	F:23	/1	F:48
D. filipjevi	Ms:500-700	8	Ms:8-11	Ms:9-11	27 40	Ms:40-50
Allen & Noffsinger, 1978	Fs:600-700	8	Fs:12-14	Fs:9-11	37-40	Fs:46-55
D. gerlachi	M:600	10	M: 13	M: 18	20	M: 28
Allen & Noffsinger, 1978	F:700	13	F: 24	F: 21	39	F: 28
D. gilbertae	M:581	8	M:10	M: 9	59	

Verschelde & Vincx, 1993	F:639		F:13	F: 10		M:20 F: 42
<i>D. grootarti</i> Decraemer, 1988	M:650 Fs:675-755	8	M:10 Fs:12-13	M:14 Fs:7-9	68	M:46 Fs:43-61
<i>D. kreisi</i> Allen & Noffsinger, 1978	M:400 F: 400	8	M:5 F: 12	M:11 F 9	36	M: 40 F: 69
<i>D. laingensis</i> Decraemer, 1988	M:460 F:440	8	M:8-9 F:5	M:8 F:5	39	M:24 F:43
D. mawsoni Allen & Noffsinger, 1978	Ms:500-600 F: 700	8	Ms:13 F:15	Ms:13 F:16	52-54	Ms:28-35 F: 58
<i>D.miguelitus</i> sp. nov.	Ms:630-735 Fs:748-788	4	Ms:10-12 Fs:13	Ms:10 Fs:13	47-54	Ms:40-41 Fs:56-59
<i>D. minutus</i> Decraemer, 1988	M:290	6	M:5	M: 2-3	18	M: 24
<i>D. ngakei</i> Leduc & Zhao, 2016	M:576 Fs:586-615	8	M:11 Fs:13	M:10 Fs:10-12	50	M:28 Fs:37-50
D. papuensis Decraemer, 1988	M: 310 Ms:350-400	8	M: 10 Ms:9-11	M: 11 Ms:9-11	29	M: 75 Ms:46-56
<i>D. pusillus</i> Decraemer, 1988	M:310	8	M:10	M: 6	26	M: 28%
D. solidus (Gerlach, 1952)	M:700 Fs:600-800	8	M:7 Fs:8-11	M:11 Fs: 8-11	46	M:29 Fs:46
D. spinosus Decraemer, 1988	M:340	8	M:8	M:10	45	M: 49%
D. stekhoveni Allen & Noffsinger, 1988	Ms:500-600 Fs:500-600	14	Ms:16-23 Fs:20-25	Ms:16-23 Fs:21-29	40-50	Ms:22-34 Fs:37-47
<i>D. timmi</i> Allen & Noffsinger, 1978	Ms:500-700 Fs:500-600	8	Ms:7-10 Fs:9-12	Ms:19-23 Fs: 7-11	41-51	Ms:29-36 Fs:43-52

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D. trispinosus	Ms:700		Ms:10	Ms:6-7		Ms:26-27
(Allen & Noffsinger, 1978)	Fs:600-800	8	Fs:12-13	Fs:8-13	59-64	Fs:52-61
D. trukensis	Ms:593-642	o	Ms:10	Ms:8-10	34-42	Ms:43-48
Min et al., 2016	Fs:663-778	8	Fs:13-15	Fs:9-11	34-42	Fs:45-58
D. wieseri	M:600	o	M:17	M:13	46	M:26
Allen & Noffsinger, 1978	F: 500	8 F: 500		F:12	40	F:45

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Table 3(on next page)

Table 3. Descriptive comparison for all valid species of Dracograllus genus including *Dracograllus miguelitus* sp. nov

Ann. Ornam., annules ornamentation; Fov. M., amphideal fovea male; Fov., amphideal fovea female; Parav. set., paravalvular setae; "-": not provided in the original description or not applicable.

1 Table 3. Descriptive comparison for all valid species of the *Dracograllus* genus including *Dracograllus miguelitus* sp. nov.. Ann.

2 ornam., annules ornamentation; Fov. M., amphideal fovea male; Fov., amphideal fovea female; Parav. set., paravalvular setae; "-": not

3 provided in the original description or not applicable.

Specie	Ann. Ornam.	Fov. M.	Fov. F.	Parav. set.	Anal flap	Diff. Diagnosis
D. antillensis	Spine-like	Large, conspicuously 'U' - shape with ventral arm often slightly longer than dorsal	-	_	Absent	Spicules 35-40 μm long, arcuated and cephalated. Gubernaculum 11-15 μm long, with corpus and lateral wind. Four long somatic setae between the eighteenth SlATs.
D. chitwoodi	Spine-like	-	Elongated loop- shape	Absent	Present. short	Fewer SIATs and SvATs, absence of PS, and setae without collar at the base.
D. cobbi	Without	Elongated loop- shape	Elongated loop- shape	One pair anterior to the vulva (6-7 μm long)	Absent	Great number of SIATs in males, shorter caudal glands and anterior position of the vulva.

D. cornutus	Without	Short loop-shape	Short loop-shape	Absent	Absent	Similar to Dracotoramonema Allen & Noffsinger, 1978, but cornifor setae and length of SIATs less conspicuous than in <i>Dracotoramonema</i> <i>trispinosum</i>
D. demani	Without	Elongated loop- shape	Elongated loop- shape	Two setae (7-9 μm long)	Absent	PS in ventro-sublateral rows, but only anterior to the SIATs.
D. eira	Without	Elongated loop- shape	Elongated loop- shape	Absent	Absent	All CATs anterior to the amphid, and 1 SIAT on the non-annulated tail region. Males with SIATs posterior to the anus.
D. falcatum	Without	Elongated loop- shape	Elongated loop- shape	Two pairs, one anterior and one posterior to the vulva	Absent	Rostrum without Ceph Acan-set and with 12 CATs
D. filipjevi	Dot-like	Elongated loop- shape	Elongated loop- shape	Two pairs, one anterior and one posterior to the vulva. (5-6 µm	Absent	Scattered minute spiny on cuticle, Absence of PS.

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long)

D. gerlachi	Dot-like punctations, more evident at mid- body	Elongated loop- shape	Elongated loop- shape	Two pairs, one anterior and one posterior	Absent	13 CATs on rostrum and great swollen esophageal region
D. gilbertae	Broad interannual space, ornamented with a slit	Large, ventrally whorled, open loop-shape	Large, closed loop-shaped	Absent	Absent	Large amphideal fovea, long and slender PATs. Slender tail with ventral post cloacalpostcloacal swelling. Spicules long and well cuticularized gubernaculum.
D. grootaerti	Spine-like	Long, inverted U- shaped, with longer ventral arm extending to the first annule	As in male, but shorter	Absent	Absent	Long body, with spiny ornamentaded annulated cuticle. Two of the SIATs in females on the tail region.
D. kreisi	Dot-like punctations	Elongated loop- shape	Elongated loop- shape	Absent	Absent	Absence of PS in ventro-sublateral row and shorter spicules in males.
D. laingensis	Spine-like	Long, inverted U- shaped	Elongated unispiral	Absent	Absent	Long swollen pharyngeal regions, and stiff posteriorly directed

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setae anterior to PATs.

D. mawsoni	Without	Elongated loop- shape	Slightly smaller, with more open loop than in male	Two pairs, one anterior and one posterior to the vulva. (4-7 µm long)	Absent	Great number of SIATs in males, and females with 1 SIAT posterior to anus
D. miguelitus sp. nov.	Without	Elongated loop- shaped, ventrally coiled, ventral arm slightly longer	Inverted U- shaped with branches more equal in size and more closed than in males	Two pairs, one anterior and one posterior to the vulva (6 μm long). Single seta emerging from the vulvar aperture	Absent	4 CATs on the rostrum, PS longer in males than in females.
D. minutus	Spine-like	Very large, loop- shape, ventrally whirled	-	-	Absent	Smaller body size within the genus, only six CATs on rostrum, short spicules. Largest fovea within the genus.
D. ngakei	Without	Loop-shaped, with two arms of equal length	Loop-shaped, with two arms of equal length	Absent	Absent	11 SvATs per row in male, all anterior to anus. Females with 12 SvATs with one of themn posterior to anus.

D. papuensis	Finely annulated	Long, inverted U- shaped, ventrally coiled, ventral arm slightly longer	Large, loop- shaped. dorsal arm slightly longer than ventral one	Minute setae: two ventral posterior and one anterior to the vulva	Absent	Shorter swollen pharyngeal region, spicule and c-value.
D. pusillus	Dot-like punctations at ring edges in the pharyngeal region	Long, inverted U- shaped. ventrally coiled, ventral arm slightly longer	-	-	Absent	Short and stout body with minute spiny ornamentations, short spicule. Long non- annulated tail tip.
D. solidus	Without	Elongated loop- shape	Elongated unispiral	Absent	Absent	11 long setae intermingled with SIATs in males, unispiral amphid and 2 SIATs posterior to anus in females.
D. spinosus	Without	Large, oblique loop-shape by position of sublateral CAT, ventrally whirled; Ventral arm slightly longer than dorsal arm	-	-	Absent	Spiny ornamentation at the insertion base of several somatic setae in the posterior body region. Females and juveniles not found.
D. stekhoveni	Without	Elongated loop-	Elongated loop-	Two pairs, one	Absent	1 pair of sub-lateral

		shape	shape	anterior and one posterior to the vulva. (3-5 µm long)		cephalic acant setae on rostrum.
D. timmi	Spine-like projections	Elongated loop- shape, some specimens ventral arm curved anteriorly toward dorsal arm almost forming unispiral	Elongated loop- shape	Two pairs, one anterior and one posterior to the vulva. (5-7 µm long)	Absent	Faint annular ridges with spine-like projections appearing as 2 rows of fine punctations.
D. trispinosum	Dot-like punctations	Very large, loop- shape	Elongated unispiral	-	Absent	Males with 3 large Corn-set, a single ventral mid-body setae and 1 preanal pair.
D. trukensis	Ridges with spiny protrusion, spiny ornamentation	Large. Elongated, open loop-shaped, longer ventral arm extending to the first body cuticular annule	Large, elongated and closed loop- shaped, shorter than in male	Absent	Absent	Numerous minute spiny ornamentation on male and female cuticle. Shorter spicule in males.
D. wieseri	Granules and vacuoles	Elongate loop- shape	Elongate loop- shape	Absent	Present. Short	6 long setae intermingled with SIATs in males, and SIAT 1 in females posterior to anus.

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Table 4(on next page)

Table 4.Distribution and ecological characteristics of Dracograllus species

References marked with an asterisk (*) indicate the original description and type locality, while those without an asterisk refer to additional localities.

Table 4. Distribution and ecological characteristics of *Dracograllus* species. References marked with an asterisk (*) indicate the
 original description and type locality, while those without an asterisk refer to additional localities.

Species / reference	Ocean	Geographic distribution	Habitat	Habitat type, sampling and conditions	Remarks
D. antillensis Decraemer & Gourbault, 1986*; Stock & Nadler, 1998	Atlantic	Guadeloupe Island: Anse de la Gourde, Grande Terre; Les Galets, Capesterre; Petite Anse, La Marie-Galante. Martinique Island: Anse l'Étang; Anse Figuiers.	Intertidal region	Sandy beach; interstitial waters with coarse and calcareous sediments.	-
D. chitwoodi Allen & Noffsinger, 1978*	Atlantic	Coral Key, Florida, USA	Subtidal region	Sandy beach; sediment associated with calcareous algae (<i>Halimeda sp.</i>).	Males only measured, without complete description. No third o fourth-stage juvenile observed.
D. cobbi Allen & Noffsinger, 1978*; Decraemer, 1988.	Atlantic	Coral Key, Florida, USA; Anse de la Gourpe, Guadeloupe.	Intertidal region	Sandy beach; sediment associated with calcareous algae (<i>Halimeda sp.</i>).	Females from Guadeloupe lack paravalvular setae and show other differences compared to the origina description. See Decraemer, 1998.

D. cornutus Decraemer, 1988*	Pacific	Laing Island, Papua New Guinea and River Mouth, NT, Australia	Subtidal region	Sandy beach; sediment sampling	-
D. demani Allen & Noffsinger 1978*; Decraemer 1988; Verschelde & Vincx 1993; Shahina et al., 2019	Atlantic, Pacific and Indian	Marseille, France; Laing Island, Duangit Reef, Papua New Guinea; Malindi, Kenya; Pakistan.	Subtidal region, down to 42 m depth	Sandy beach; coarse sand with algae and coarse coral sand	Specimens from Papua New Guinea differ from the type locality by having a shorter general body length, shorter PATs, and shorter spicules.
<i>D. eira</i> Inglis 1968*; Decraemer 1988; Verschelde & Vincx 1993	Pacific and Indian	St. Vincent's Bay, New Caledonia; Laing Island, Papua New Guinea; Malindi, Kenya	Subtidal and intertidal zone	Sediments associated with polychaete tubes and large pieces of dead coral.	-
<i>D. falcatus</i> Irwin-Smith, 1918*; Allen & Noffsinger, 1978	Pacific	Cremorne, Port Jackson, New South Wales, Australia; Long Reef and Vaucluse, Australia	Subtidal region, from 1.2 - 1.5 m depth	Sandy beach; sediment sampling with seaweed and shells	-
D. <i>filipjevi</i> Allen & Noffsinger, 1978* ; Rho et al., 2006	Pacific	Oarai, Ibaraki-ken, Honshu Island, Japan; Daebo-ri, Guryongpo, Korea	Subtidal region	Washings of holdfasts of Kelps and also in shallow littoral calcareous algae	-

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D. gerlachi Allen & Noffsinger, 1978* ; Jesús- Navarrete, 2021	Pacific and Atlantic	Ibusuki, Kyushu Island, Japan and Laguna de Términos, Gulf of Mexico	Subtidal region	Sandy beach; sediment sampling with brown algae growing on rocks
<i>D. gilbertae</i> Verschelde & Vincx, 1993*	Indian	Gazi, Kenya	Subtidal region	Sandy beach; core of 3.5 cm diameter into the sediment down to 20 cm depth, close to mangrove plants (<i>Sonneratia</i> sp.)
D. grootaerti Decraemer, 1988*	Pacific	Madang Province, Hansa Bay, Duangit Reef, Laing Island, Papua New Guinea	Subtidal region at 42 m depth.	Sandy beach; sediment sampling with polychaete tubes, and coral sand
D. kreisi Allen & Noffsinger, 1978*	Atlantic	Coco Solo, on Galeta Beach, Panama	Subtidal region	Sediment associated with calcareous algae (<i>Halimeda sp.</i>).
D. laingensis Decraemer, 1988*	Pacific	Laing Island, Papua New Guinea	Subtidal region at 42 m depth.	Sediment sampling with polychaete tubes, and coral sand
D. mawsoni Allen & Noffsinger,	Pacific	Long Nose Point, Port Jackson, New South Wales, Australia	Subtidal region	Sandy beach; sediment sampling with bottom debris

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1978*

D. minutus Decraemer, 1988*	Pacific	Laing Island, Papua New Guinea	Subtidal region	Sediment sampling with polychaetes tubes of sand and mucus	No female or juvenile known
D. miguelitus sp. nov. Johnson et al., 2024*	Atlantic	Lucky Strike vent field, Mid Atlantic Ridge	Deep-sea	Hydrothermal inactive vent structure;	Only present in the inactive vent structure at LS, without individuals in active or periphery samples.
D. ngakei Leduc & Zhao, 2016*	Pacific Ocean	Hataitai Beach, Wellington, New Zealand	Mid-intertidal region	Sandy beach; sediment sampling (0 to 10 cm sediment depth) with coarse sand and gravel	SSU Molecular sequences available in original description.
D. papuensis Decraemer, 1988*	Pacific	Laing Island, Papua New Guinea	Subtidal region	Sediment sampling with dead coral debris, also with polychaete tubes of sand and mucus	Only one male found, without non-annulated tail tip length known.
<i>D. pusillus</i> Decraemer, 1988*	Pacific	Laing Island, Papua New Guinea	Subtidal region	Sediment sampling, with dead coral debris	-

<i>D. solidus</i> Gerlach, 1952*	Atlantic and Indian Ocean	Banyuls, France; Bay of Biscay, Mediterranean sea; Mascarene Islands.	Subtidal region	Sandy beach; sediment sampling	Also recorded in Mascarene Islands, no juvenile known.
D. spinosus Decraemer, 1988*	Pacific	Laing Island, Papua New Guinea	Subtidal region	Sediment sampling with polychaete tubes, sand and mucus	Males and juveniles not found.
D. stekhoveni Allen & Noffsinger, 1988*	Pacific	Solano, Colombia; Port Jackson, Australia; Isla Taboga, Panama	Subtidal region	Sandy beach; sediment sampling with corals	Juveniles specimens third stage without PS.
D. timmi Allen & Noffsinger, 1978*	Pacific	Bora Bora Island, Society Islands	Subtidal region	Sandy beach; sediment sampling of coarse sand	Second and third-stage juveniles not found
D. trispinosus Allen & Noffsinger, 1978*	Atlantic	Southwest of the Pomegues Ratonneau jetty, near Marseille, France	Subtidal region	Sandy beach; sediment sampling at 20 m depth.	-
D. trukensis Min et al., 2016*	Pacific	Weno, Chuuk, Micronesia	Subtidal region	Sandy beach; sediment sampling with seagrass bed (<i>Zostera</i> sp., from 1 to 2 m depth)	-
D. wieseri Allen & Noffsinger,	Pacific	Juan Fernandez Islands, Chile	Subtidal region during high tide	Sandy beach; sediment sampling with green algae	-

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Table 5(on next page)

Global distribution, habitats, and environmental characteristics of unidentified *Dracograllus* sp.

1 Table 5. Global distribution, habitats, and environmental characteristics of unidentified *Dracograllus* sp.

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Ocean	Site	Coords.	Habitat	Substratum type	Morphospecies	Reference
Atlantic ocean	Great Meteor Seamount	30°00'N, 28°30'W.	Plateau of the seamount, with 1,465 km2	Calcareous biogenic sands	Dracograllus sp.4; Dracograllus sp.5; Dracograllus sp.6; Dracograllus sp.7	Gad, 2009
	Condor Seamount	38°32.949'N, 29°02.879'W.	Summit of the seamount, at 206 m depth.	Large rocky outcrops, gravels, and coarse bioclastic deposits	<i>Dracograllus</i> sp.	Zeppilli et al., 2014; Zeppilli et al., 2013
	Snake Pit vent field	23°22.0'N, 44°57.0'W.	Sampling located 70 m from one black smoker. Depth between 3.480 m and 3.570 m	Reddish sediments covering the corer, with several polychaete tubes and individuals	<i>Dracograllus</i> sp.	Spedicato et al., 2020

Anse Laborde, Guadeloup e island	16°29.2'N, 61°30.3'W	Intertidal beach zone with high hydrodynamic activity.	Composed of detrital fragments, mostly carbonates	<i>Dracograllus</i> sp. 1	Decraemer & Bourbaults, 1986 ; Renauld- Mornant & Gourbault
Raisins, clairs, Guadeloup e	16.24892°N, 61.28345°W	Sandy beach on the characterized by a low sandy ridge (2 to 3 meters in height) facing frequent waves and subject to significant coastal erosion	Sediments consist of a low sandy ridge, with a "beach-rock" (sandstone) layer along the coastline.	Dracograllus sp. 2 and Dracograllus sp. 3	Decraemer & Bourbaults, 1986; Renauld- Mornant & Gourbault, 1981
La Marie Galante. Guadeloup e	15°55'59.99"N, 61°15'60.00"W 15.912°N, 61.269°W	Sandy Beach with a topography that includes a low sand ridge parallel to the shoreline and sparse vegetation	Sandy beach composed of sediments ranging from fine volcanic sands to coarse organogenic sands.	<i>Dracograllus</i> sp. 4	Decraemer & Bourbaults, 1986 ; Renauld- Mornant & Gourbault, 1981

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	Guanabara Bay, Rio de Janeiro, Brazil.	22°24'S - 22°57'S, 42°33'W - 43°19'W	Sandy Beach, intertidal zone.	Substratum composed of sand, from medium to very coarse sediments. Highly impacted beach region subject to anthropogenic pressures.	<i>Dracograllus</i> sp.	Maria et al., 2008
Pacific ocean	Munseon Island, Jjeudo, Korea	33°13′66″N, 126°34′18″E	Subtidal zone, 37 m deep.	Sampling based on washings of shallow subtidal detritus and coarse sediments	<i>Dracograllus</i> sp. 1	Rho et al;, 2011
	Geomundo Island, Jeonranam d, Korea	34°05′57″N, 127°14′84″E	Intertidal zone, associated with invertebrates	Substratum with associated invertebrates	Dracograllus sp. 2	Rho et al;, 2011
	Volcanic Island of Moorea, French Polynesia	(17°30'S - 149°50'W)	Flat beaches surrounded by a large coral reef	Sediments with coarse coral sand	<i>Dracograllus</i> sp. 1 e <i>Dracograllus</i> sp. 2	Gourbaulet al., 1995

Indian ocean	Gazi Kenya	-4.4222°S, 39.5050°E	Sandy beach, intertidal zone	Sample taken in mangrove region, with <i>Ceriops sp.</i> tree	Dracograllus spec.	Verschelde & Vincx, 1993
	Chidiyatap u, South Andaman Island, India	11°29'30"N - 11°30'34"N, 92°35'10"E - 92°42'30"E	Rocky coastal area	Sediments associated with several seagrasses patches (<i>Thalassia</i> hempirichi, Halodule uninervis and Halophila ovalis)	Dracograllus sp.	Naufal & Padmavati, 2016
	Marina Park, Andaman Islands	11°40'15.39"N, 92°45'39.16"E	Sublitoral sediments	Substratum composed of silty- sand and clayey-sand	Dracograllus sp.	Arunima et al., 2023
	Huvadhoo Atoll, Maldives	08°33'20.88"N, 73°81'4.76"E	Central atoll region	Sediments with coarse and gravelly sand, at 61 m deep.	<i>Dracograllus</i> sp.	Semprucci et al., 2014

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Souther n ocean	Halley Bay, Weddel Sea	74°S - 75°S, 25°W - 29°W	Shelf break region, 500 m deep	Sediment poorly to extremely poorly sorted, with significant variations in grain size, with presence of pellite and gravel	<i>Dracograllus</i> sp.	Vanhove et al., 1999	