Morphological and molecular characterization of *Heterodera dunensis* n. sp. (Nematoda: Heteroderidae) from Gran Canaria, Canary Islands

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Abstract

*Heterodera dunensis* n. sp. from the coastal dunes of Gran Canaria, Canary Islands, is described. This new species belongs to the *Schachtii* group of *Heterodera* with ambifenestrate fenestration, presence of prominent bullae, and a strong underbridge of cysts. It is characterized by vermiform second-stage juveniles having a slightly offset, dome-shaped labial region with three annuli, four lateral lines, a relatively long stylet (27-31 µm), short tail (35-45 µm), and 46 to 51% of tail as hyaline portion. Males were not found in the type population.

Phylogenetic trees inferred from D2-D3 of 28S, partial ITS, and 18S of ribosomal DNA and COI of mitochondrial DNA sequences indicate a position in the ‘Schachtii clade’.

Keywords

18S, 28S, Canary Islands, COI, Cyst nematode, ITS, Gran Canaria, *Heterodera dunensis*, Plant-parasitic nematodes, *Schachtii*, Systematics, Taxonomy.

The cysts forming nematodes of the genus *Heterodera* Schmidt, 1871 (Nematoda: Heteroderidae) are an economically important plant-parasitic nematode (PPN) group with a worldwide distribution and a broad host range causing prominent damages to the host plants ranging from stunted and reduced growth to wilting, chlorosis, and reduced root system (Perry et al., 2018; Sikora et al., 2018). The vermiform second-stage juveniles (J2) of this PPN migrate in the root system of a host plant to feed on the vascular cylinder where they become obese sedentary females; subsequently, following fertilization and egg production, these females turn into protective cysts of more or less lemon shape, housing numerous embryonated eggs. These eggs can remain viable for years inside the cysts, until favorable environmental conditions initiate hatching of the cysts to continue further life cycle (Subbotin et al., 2010; Perry et al., 2018).

Within this genus, 85 nominal species, eight species inquirendae, and a nomen nudum have been listed in a recent update by Handoo and Subbotin (2018). Using morphological and molecular characteristics, the species of this genus have been divided into nine groups, i.e., *Afenestrata*, *Avenae*, *Bifenestra*, *Cardiolata*, *Cyperi*, *Goettingiana*, *Humuli*, *Sacchari*, and *Schachtii*. Morphological characterization of *Heterodera* species is mainly done based on vulva-slit length, vulval cone fenestration, presence or absence of bullae and underbridge in female cysts, and stylet length, lateral field differentiation, tail length, and hyaline tail length in J2 (Subbotin et al., 2010). Since the last two decades, employing molecular data such as ITS and 28S of ribosomal DNA and COI gene of mitochondrial DNA to characterize *Heterodera* species has been a common practice, including DNA barcoding, phylogeny, and even phylogeography (Ferris et al., 1999; Toumi et al., 2013a; Subbotin et al., 2017, 2018).

Herein, we characterize *Heterodera dunensis* n. sp. discovered in a recent exploratory survey of PPN from Canarian coastal dunes of Gran Canaria in May
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2019. The species characterization is done based on light microscopy (LM), scanning electron microscopy (SEM), and molecular information of ITS, 18S, and 28S of ribosomal DNA and COI of mitochondrial DNA.

Materials and methods

Nematode extraction and morphological analysis

A sandy soil sample was collected from around the root system of Tetraena fontanesii (Webb & Berthel.) Beier & Thulin, commonly known as Sea Grape or Canarian Bean-Caper. This halophilic succulent plant was growing on a dune (GPS coordinates: 27°44'19.11" N; 15°35'0.3" W), about 200 m away from the Maspalomas beach of Gran Canaria. Vermiform J2 was extracted from the sand using the modified Bearmann method (Whitehead and Hemming, 1965) and stored at 4°C during the course of analysis.

For collecting female cysts, sand dried at room temperature was mixed thoroughly in water using a spoon, and after letting the sand settle, floating female cysts were picked out using a fine brush.

Morphological study of J2 was done using both heat-relaxed and fixed specimens. Individual live nematodes were heat-relaxed in a drop of water on a glass slide and examined, photographed, and measured using an Olympus BX51 DIC Microscope (Olympus Optical, Tokyo, Japan), equipped with an Olympus C5060Wz camera and a drawing tube as described in the study of Singh et al. (2018). After recording morphological information, the specimens were recovered from the slide and their genomic DNA was extracted as described in the next section. The remaining J2 juveniles were concentrated in a drop of water in a glass embryo dish, followed by adding a few drops of freshly prepared Trump’s fixative [2% paraformaldehyde, 2.5% glutaraldehyde in 0.1 M Sorenson buffer (sodium phosphate buffer at pH = 7.3)]. The nematodes were then immediately heated in a microwave (700 watts) for about 4 sec, left to rest for 1 hr at room temperature and at 4°C for 24 hr, followed by gradually transferring to anhydrous glycerin to be heat-relaxed and fixed specimens. Individual live nematodes after morphological analysis were recovered from temporary slides. Each individual nematode was cut into pieces in distilled water using a blade, and the pieces were transferred to a PCR tube with 20 µl of worm lysis buffer [50 mM KCl, 10 mM Tris at pH=8.3, 2.5 mM MgCl₂, 0.45% NP 40 (Tergitol Sigma), and 0.45% Tween 20]. DNA extraction from cysts containing embryonated eggs was also done for which an individual cyst was crushed and transferred in the PCR tube containing worm lysis buffer as mentioned. The PCR tubes were then incubated at -20°C (10 min) followed by adding 1 µl of proteinase K (1.2 mg/ml), incubation at 65°C (1 hr) and 95°C (10 min), and ending by centrifuging the mixture at 14,000 rpm for 1 min (Singh et al., 2018). PCR amplifications of partial ITS and 18S regions of ribosomal DNA were done using the primer pairs, Vrain2F: 5′-CTTGTACACCGCGGCTGCT-3′/Vrain2R: 5′-TTTCACCTCGCGGTTACTAAGGGAATC-3′ (Vrain et al., 1992) and SSU18A: 5′-AAAGATTAACCATGCGAT-3′/SSU26R: 5′-CATGTTGCAAAATGCTTTTCG-3′ (Mayer et al., 2007) and with thermal profiles described in the study of Singh et al. (2018, 2019). For amplification of D2-D3 expansion segment of 28S of ribosomal DNA, two primer sets were used, the primer pair, 391: 5′-AGCGGAGAAAAGAAACTAA-3′/501: 5′-TCCGAGGAGGACCACTACTA-3′ was used as described in the study of Nadler et al. (2006) and D2A: 5′-AAGTACCAGTGAGGAAAAGTTG-3′/D3B: 5′-TCCTCGGAAGGAACCAGCTACTA-3′ (Nunn, 1992) with the thermal profile from Singh et al. (2019). For the amplification of the COI region of mitochondrial DNA, the primer pair, JB3: 5′-TTTTGGGCTCC TGGAGTTAT-3′/JB4.5: 5′-TAAAGAAAAAATAAATG AAAATG-3′ was used according to Bowles et al. (1992). The PCR products were enzymatically cleaned with alkaline phosphatase (1 U/ml) and exonuclease I (20 U/ml) for 15 min at 37°C followed by 15 min at 80°C and sent for sequencing at Macrogen (https://dna.macrogen.com), and contigs were made from the newly produced forward and backward sequences using Geneious Prime 2020.0.5 (https://www.geneious.com) and deposited in GenBank.
Phylogenetic analysis

The phylogenetic relationships of the new species with other related species were analyzed based on the D2-D3, ITS, 18S, and COI sequences. Phylogenetic programs implemented in Geneious Prime 2020.0.5 were used. The obtained consensus contigs were subjected to BLAST search to check for closely related species on GenBank, and all the collected sequences for each gene fragment were aligned using MUSCLE alignment of Geneious Prime 2020.0.5 using default parameters, followed by manually trimming off of the poorly aligned ends. The best nucleotide substitution model of each gene alignment (see Figures) was determined by jModelTest 2.1.10. Bayesian phylogenetic analysis (MrBayes 3.2.6) was carried out using the selected models, analyses were run under $1 \times 10^6$ generations (4 runs), and Markov chains were sampled every 100 generations, and 20% of the converged runs were regarded as burn-in (Huelsenbeck and Ronquist, 2001).

Results

Systematics

*Heterodera dunensis* n. sp.

Figures 1–6, Tables 1 and 2.

Description

Cyst: Body typical lemon shape to sometimes ovoid shape with protruding prominent neck and vulva. Neck regularly bent. Cysts wall light to medium brown in color with irregular zig-zag pattern on surface. Fenestration ambifenestrate. Vulva cone dome-shaped with sub terminal anus. No egg-sac observed. Vulva slit longer than fenestral length. Bullae prominent, medium brown in color, variable shape, in some cysts commonly finger-like or elongated, irregularly distributed at the periphery of vulva cone slightly above underbridge level. Underbridge furcated with central thickening, prominent in young cysts, breaks down in older cysts. Cysts containing 100-200 eggs.

J2: Body slender, tapering posteriorly. Labial region slightly offset, dome-shaped with two clear incisure under LM appearing as three lip annuli, second annule wider than the other two. *En face* showing oral disc fused with submedial sectors, well-separated lateral lip sectors and rectangular to square-shaped stoma opening. Lateral field with four longitudinal incisures forming three bands, outer two bands slightly wider than inner. All bands irregularly areolated, sometimes with incomplete areolation. Stylet robust, 27-31 µm long, with large rounded strongly anteriorly projecting knobs. Pharynx well-developed, ca one-third of body length with well-developed median bulb, valves and glands overlapping intestine ventrally. Nerve ring encircling isthmus. Hemizonid distinct, about two cuticular annuli long, just above secretory-excretory (SE) pore opening. SE pore at ca one-fourth of body length from anterior end. Tail 35-45 µm long, tapers gradually to a rounded terminus, hyaline region ca 50% of tail length. Phasmid opening small, roughly halfway between anus and start of hyaline tail part.

Male: Not found.

Diagnosis and relationships

*Heterodera dunensis* n. sp. is characterized by moderate-sized J2 of 0.43 to 0.52 mm long, lateral field with four lines, the inner band slightly smaller than the outer two bands, and all bands with irregular areolation throughout the length; a relatively long J2 stylet of 27 to 31 µm with anteriorly projected knobs, a relatively short tail of 35 to 45 µm in length, small rounded phasmids, and tail hyaline part usually ca 50% of the tail; cyst ovoid to regularly lemon-shaped, ambifenestrate, the presence of prominent finger-like bullae, and a strong underbridge.

This new species belongs to the *Schachtii* group that comprises sixteen *Heterodera* species, i.e., *Heterodera agrostis* Kazachenko, 1993; *Heterodera betae* Wouts, Rumpenhorst and Sturhan, 2001; *Heterodera cajani* Koshy, 1967; *Heterodera ciceri* Vovlas, Greco and Di Vito, 1985; *Heterodera daverti* Wouts and Sturhan, 1978; *Heterodera galeopsidis* Goffart, 1936; *Heterodera glycines* Ichinohe, 1952; *Heterodera lespedezae* Golden and Cobb, 1963; *Heterodera medicaginis* Kirjanova in Kirjanova and Krall, 1971; *Heterodera mediterranea* Vovlas, Inserra and Stone, 1981; *Heterodera rosii* Duggan and Brennan, 1966; *Heterodera schachtii* A. Schmidt, 1871; *Heterodera sonchophila* Kirjanova, Krall and Krall, 1976; *Heterodera spiraeae* Kazachenko, 1993; *Heterodera swarupi* Sharma, Siddiqi, Rahaman, Ali and Ansari, 1998; and *Heterodera trifolii* Goffart, 1932. They are all similar in having J2 with a lateral field with four lines, with more or less anteriorly projected stylet knobs; cysts presented with ambifenestrate fenestration, the presence of prominent bullae, and a strong underbridge. *Heterodera dunensis* n. sp. can be easily differentiated from other members of *Schachtii* group based on J2 with a long stylet, short tail, and
Figure 1: Light microscopy and scanning electron microscopy images of second-stage juveniles of *Heterodera dunensis* n. sp. A to D: *En face* view, E: Anterior part up to pharyngeal gland end, F to H: Labial region showing stylet and labial annuli, I: Total body, J to L: Anterior part showing median bulb, hemizonid, and secretory-excretory pore, M to Q: Tail region showing hyaline portion and anus.
a short hyaline region. It differs from H. agrostis, H. daverti, H. glycines, H. lespedezae, H. medicaginis, H. schachtii, H. spiraeae, and H. swarupi in having a distinctly longer J2 stylet of 29 µm (27-31 µm) vs stylet length always shorter than 27 µm.

This new species can also be easily separated from all the members of the group, except H. mediterranea by its shorter J2 tail length of 41 µm (35-45 µm) vs always above 45 µm on average (38-77 µm) and a shorter hyaline tail part of 21 µm (16-23 µm) vs always above 23 µm on average (20-45 µm) in the other species. Heterodera dunensis n. sp. is morphologically closest to H. mediterranea with several overlapping morphometrics, such as the dimension of cyst cone fenestration and length of vulva slit, the length of J2 tail and hyaline part, but differs from this species in the J2 body length (426-520 vs 360-430 µm), by a slightly longer J2 stylet (27-31 vs 25-27 µm), presence vs absence of finger-like bullae and central thickening of an underbridge in its respective cysts.

Heterodera dunensis n. sp. is also based on D2-D3, ITS, 18S, and COI sequences clearly different from all known species, see below.

**Molecular characterization**

**D2-D3 of 28S rDNA**

Three D2-D3 sequences (MT508987-MT508989) of 987-1039 bp were produced without intraspecific sequence variation. The closest available sequence on GenBank was MK292129 of H. glycines with 95.9% similarity (43 out of 1039 bp differences). The D2-D3 alignment of 750 bp long consisted of 75 Heterodera sequences of 33 species and a Cryphodera sinensis sequence (JX566455) as the outgroup. The resulting D2-D3 tree revealed an
Figure 3: Line illustrations of cysts and second-stage juveniles (J2) of *Heterodera dunensis* n. sp. A: Whole cysts, B, C: Vulva cones showing cone fenestration, vulva slit, bullae, underbridge, and anus, D, F: Anterior part of J2 showing lip region, pharynx, hemizonid, and secretory excretory pore, E: Stylet of J2, G to I: Tail region of J2 showing lateral field differentiation, anus, phasmid, and tail hyaline portion.

unresolved position of *H. dunensis* n. sp. in a clade (PP = 0.93) comprising eight members of Schachtii group, i.e. *H. glycines*, *H. medicaginis*, *H. schachtii*, *H. mediterranea*, *H. trifolii*, *H. betae*, *H. daverti*, and *H. cajani*.

**ITS of rDNA**

Three partial ITS sequences (MT508990-MT508992) of 747-1025 bp were produced with intraspecific sequence variation of only one bp. The closest available sequence was LC030416 of *H. trifolii* with 84.6% sequence similarity (150 out of 971 bp differences). The ITS alignment was 1514 bp long and consisted of 105 *Heterodera* sequences of 54 species and a *Cryptodera sinensis* sequence (JX566457) as the outgroup. In the inferred ITS tree, *H. dunensis* n. sp. occupies a well-supported sister relationship with *H. cajani* (PP = 0.99) within a maximally supported clade of other members of Schachtii group, namely *H. daverti*, *H. betae*, *H. trifolii*, *H. schachtii*, *H. ciceri*, *H. medicaginis*, *H. glycines*, and *H. mediterranea*.
Figure 4: Phylogenetic relationships of *Heterodera dunensis* n. sp. with 33 known *Heterodera* species. Bayesian 50% majority-rule consensus tree as inferred from the analysis of D2-D3 of 28S rDNA sequences under GTR + G model. Posterior probabilities of more than 0.5 are given for appropriate clades.
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Figure 5: Phylogenetic relationships of *Heterodera dunensis* n. sp. with 54 known *Heterodera* species. Bayesian 50% majority-rule consensus tree as inferred from the analysis of partial ITS of rDNA sequences under GTR + I + G model. Posterior probabilities of more than 0.5 are given for appropriate clades.
Figure 6: Phylogenetic relationships of Heterodera dunensis n. sp. with 29 known Heterodera species. Bayesian 50% majority-rule consensus tree as inferred from the analysis of COI of mtDNA sequences under GTR+I+G model. Posterior probabilities of more than 0.5 are given for appropriate clades.
Table 1. Morphometric data of *Heterodera dunensis* n. sp. from fixed specimens mounted in glycerin, except for cysts that were not fixed.

| Cysts |  |  |
|---|---|---|
| n | 21 |  |
| Length without neck (*L*) | 482 ± 75 (316-610) |  |
| Maximum cyst width (*W*) | 360 ± 72.1 (241-478) |  |
| Neck length | 110 ± 31.9 (55-180) |  |
| *L*/*W* | 1.4 ± 0.2 (1.2-1.9) |  |
| Vulval area |  |
| n | 13 |  |
| Fenestral length | 42.1 ± 4.3 (37-51) |  |
| Mean semifenestral width | 41.8 ± 5.2 (34-53) |  |
| Vulva bridge width | 8.1 ± 1.6 (5.2-11) |  |
| Vulva slit length | 52.1 ± 7.3 (35-63) |  |
| Underbridge length | 63.4 ± 9.8 (44-83) |  |
| Underbridge width | 18.2 ± 6.2 (8.8-29) |  |
| Vulva-anus distance | 58.4 ± 5.7 (50-68) |  |
| Juveniles |  |
| n | 21 |  |
| Body length (*L*) | 471 ± 25.3 (426-520) |  |
| *a*=*L*/MBD | 20.1 ± 1.4 (19-23) |  |

Note: All measurements are except percentage and ratio in μm and in the form: mean ± sd (range).

**18S of rDNA**

Three partial 18S sequences (MT509422-MT509424) of 806-845 bp were generated without intraspecific sequence variation. The sequences were closest to EU306357 of *H. koreana* with 98.9% similarity (9 out of 845 bp differences). The resulting phylogenetic tree revealed an unresolved position of several species, including *H. dunensis* n. sp., and is, therefore, not provided.

**COI of mtDNA**

Eight COI sequences (MT511092-MT511099) of 309-405 bp were generated without intraspecific sequence variation. The closest sequence available on GenBank was MN31179 of *H. medicaginis* with 87.6% similarity (46 out of 370 bp difference). A COI sequence alignment of 418 bp long was made consisting of 90 sequences from 29 *Heterodera* species, including the new species, five unidentified sequences, and a *Cryptodera sinensis* sequence (MF425738) as the outgroup. From the phylogenetic tree inferred, *H. dunensis* n. sp. formed a poorly supported sister relationship (PP = 0.74) with a clade consisting of *H. glycines*, *H. medicaginis*, *H. schachtii*, *H. trifolii*, *H. betae*, *H. daverti*, and *H. ciceri*.

**Etymology**

The species epithet refers to the coastal dunes, the type locality where this new species was found.

**Type host and locality**

*Heterodera dunensis* n. sp. was recovered from the rhizosphere of the halophilic host plant, *Tetraena fontanesii* (Webb & Berthel.) Beier & Thulin, growing on a dune, roughly 30 cm high, about 200 m inland of Maspalomas beach of Gran Canaria; GPS coordinates: 27°44′19.11″ N; 15°35′0.3″ W.
Table 2. Comparison of important characters of seventeen *Heterodera* species of the *Schachtii* group.

| Character | Cyst length (L) | Cyst width (W) | L/W ratio | Fenestral length | Fenestral width | Vulval slit length | J2 body length | J2 stylet length | J2 tail length | J2 tail hyaline length | Hyaline% of the tail |
|-----------|----------------|----------------|-----------|------------------|----------------|-------------------|---------------|-----------------|----------------|------------------------|---------------------|
| *H. agrostis* | 429-800 | 320-541 | 1.2-2.6 | 34-54 | 16-24 | 39-48 | 384-472 | 25-26 | 46-61 | 27-40 | 59-66 |
| *H. betae* | 830-878 | 455-518 | 1.6-1.9 | 44-55 | 38-43 | 48-57 | 547-607 | 29-31 | 70-74 | 38-42 | 54-57 |
| *H. cajani* | 448-670 | 209-422 | 1.4-2.1 | 27-65 | 25-40 | 43-55 | 420-519 | 22-27 | 42-52 | 23-31 | 55-60 |
| *H. ciceri* | 570-930 | 350-550 | 1.6-2.4 | 32-52 | 20-37 | 43-60 | 440-585 | 27-30 | 53-72 | 31-42 | 58-58 |
| *H. daverti* | 650-749 | 380-491 | 1.4-1.5 | 42-54 | 31-40 | 47-52 | 457-476 | 25-26 | 54-57 | 30-33 | 56-58 |
| *H. dunensis* n. sp. | 316-610 | 241-478 | 1.2-1.9 | 37-51 | 34-53 | 35-63 | 426-520 | 27-31 | 35-45 | 16-23 | 46-51 |
| *H. galeopsidis* | 576-797 | 408-556 | 1.4-1.5 | 41-50 | 31-38 | 39-50 | 485-553 | 26-28 | 61-75 | 35-40 | 57-53 |
| *H. glycines* | 474-709 | 327-535 | 1.3-1.7 | 34-58 | 16-41 | 38-50 | 386-471 | 21-23 | 39-51 | 22-30 | 56-59 |
| *H. lespedezae* | 678-719 | 371-522 | 1.4-1.8 | 43-59 | 35-41 | 45-47 | 457-481 | 24-25 | 54-56 | 26-30 | 48-54 |
| *H. medicaginis* | 568-728 | 364-570 | 1.4-1.5 | 39-55 | 30-40 | 39-55 | 417-512 | 24-26 | 41-60 | 22-33 | 54-55 |
| *H. mediterranea* | 430-690 | 240-570 | 1.2-1.6 | 38-45 | 37-42 | 42-48 | 360-430 | 25-27 | 38-45 | 19-26 | 50-58 |
| *H. rosii* | 537-1173 | 403-634 | 1.0-1.7 | 48-65 | 40-45 | 45-59 | 430-662 | 27-34 | 58-77 | 37-45 | 64-58 |
| *H. schachtii* | 768-815 | 512-529 | 1.5-1.6 | 35-38 | 25-31 | 41-44 | 436-489 | 25-26 | 45-49 | 24-27 | 53-55 |
| *H. sonchophila* | 732-1032 | 381-616 | 1.6 | 37-58 | 29-50 | 42-56 | 437-492 | 24-27 | 47-56 | 26-30 | 55-54 |
| *H. spiraeae* | 467-861 | 283-566 | 1.1-1.7 | 33-60 | 20-45 | 39-45 | 371-446 | 22-23 | 39-49 | 21-27 | 54-55 |
| *H. swarupi* | 520-700 | 320-475 | 1.6-2.1 | 45 | 35 | 41 | 400-440 | 21-23 | 39-54 | 20-29 | 51-54 |
| *H. trifolii* | 608-841 | 341-536 | 1.3-1.8 | 43-53 | 33-44 | 40-53 | 492-613 | 25-28 | 60-72 | 32-37 | 53-51 |

Note: The measurements of the new species are shown in bold. All measurements are in µm and presented as range.
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**Type material**

Holotype J2 and seven J2 paratypes in two slides, two cyst vulval cones, and two whole cysts in separate slides were deposited at the National Plant Protection Organization, Wageningen Nematode Collection, Wageningen, The Netherlands (WaNeCo). Six paratype J2 and three cyst vulval cones in two slides were submitted to the Ghent University Museum, Zoology Collections, Belgium. Additionally, five J2 paratypes and two cyst vulval cones were also deposited at the UGent Nematode Collection (slide UGnem-189-190) of the Nematology Research Unit, Department of Biology, Ghent University, Belgium.

**Discussion**

*Heterodera dunensis* n. sp. is easily distinguishable from other *Heterodera* species and from all other members of *Schachtii* group by both morphology and molecular data (D2-D3, ITS, 18S, and COI sequences). The obtained phylogenetic trees revealed a consistent phylogenetic position of the new species always forming a clade together with other members of the *Schachtii* group. The ITS tree provided slightly better-resolved phylogenetic relationships among different *Heterodera* species compared to the D2-D3 and the COI trees, while the 18S tree had an inferior resolution.

*Heterodera dunensis* n. sp. was present in a Canarian dune sample with a moderately large population of only J2 and cysts together with very few saprophytic nematodes and no other PPN species. A resampling in search of males from the type location and an attempt to culture the species at NPPO, Wageningen green house did not succeed. Coastal regions are a relatively common habitat for *Heterodera* spp. To our knowledge, nine *Heterodera* spp. have been reported from similar habitats around the world, namely *Heterodera arenaria* Cooper, 1955, parasitizing on marram grass (*Ammophila arenaria*) on mobile sand dunes from several coastal sites in the United Kingdom and the Netherlands (Robinson et al., 1996; Brzeski, 1998); *Heterodera aucklandica* Wouts and Sturhan, 1995 associated with meadow rice grass (*Microlaena stipoides*) in Auckland, New Zealand (Wouts and Sturhan, 1995); *Heterodera hordecalis* Andersson, 1975 on marram grass in the Netherlands (Van der Putten and Van der Stoel, 2006); *Heterodera leucelyma* Di Edwardo and Perry, 1964 from the coastal regions of Florida parasitizing on St Augustine grass (*Stenotaphrum secundatum*) (Di Edwardo and Perry, 1964); *Heterodera litoralis* Wouts and Sturhan, 1996 associated with a succulent plant, beaded glasswort (*Sarcocornia quingueflora*) in South Island, New Zealand (Wouts and Sturhan, 1996); *H. mediterranea* associated with a woody plant, lentisc (*Pistacia lentiscus*) on the Adriatic coast of Southern Italy (Vovlas et al., 1981); *Heterodera pratensis* Gäbler, Sturhan, Subbotin and Rumpenhorst, 2000 in a pasture near the coast of the Baltic Sea at Lindhöft, Germany (Gäbler et al., 2000); *Heterodera riparia* (Kazachenko, 1993) Subbotin, Sturhan, Rumpenhorst and Moens, 2003 associated with couch grass, *Elymus repens* (L.) from along the coast of Olga Bay, the coast of Okhot Sea, Kamchatka region of Russia (Subbotin et al., 2003), and *Heterodera salixophila* Kirjanova, 1969 parasitizing roots of the willow tree, *Salix purpurea* from the shores of Kurdish Bay, Baltic Sea, Kaliningrad region of Russia (Kirjanova, 1969). This new species was found associated with *Tetraena fontanesii* (Sea Grape or Canarian Caper), a succulent plant with a limited distribution, in Canary Islands and some parts of West Africa. To the best of our knowledge, no report of an association of PPN with this host has been made before. Sampling in similar habitats should reveal if *H. dunensis* n. sp. is endemic for the Canarian islands and to what extent it is host-specific.

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**References**

Andersson, S. 1975. *Heterodera hordecalis* n. sp. (Nematoda: Heteroderidae), a cyst nematode of cereals and grasses in southern Sweden. Nematological 20:445–4.

Bowles, J., Blair, D. and McManus, D. P. 1992. Genetic variants within the genus *Echinococcus* identified by mitochondrial DNA sequencing. Molecular and Biochemical Parasitology 54:165–73.

Brzeski, M. W. 1998. Nematodes of Tylenchina in Poland and temperate Europe Muzeum i Instytutu Zoologii, Polska Akademia Nauk (MiIZ PAN, Warszawa, Poland.

Copper, B. A. 1955. A preliminary key to British species of *Heterodera* for use in soil examination. In Kevan, D. K. McE. (Eds), Soil zoology. London: Butterworths, pp. 269–80.

Di Edwardo, A. A. and Perry, V. G. 1964. *Heterodera leucelyma* n. sp. (Nematoda: Heteroderidae), a severe pathogen of St Augustine grass in Florida. Agricultural Experiment Stations. Gainesville, and Bulletin: University of Florida, Vol. 687, p. 35.
Duggan, J. J. and Brennan, P. A. 1966. Heterodera rosi (Heteroderidae), a new species of cyst-forming nematode from curled dock (Rumex crispus L.). Irish Journal of Agricultural Research 5:113–20.

Ferris, V. R., Subbotin, S. A., Ireholm, A., Spiegel, Y., Faghihi, J. and Ferris, J. M. 1999. Ribosomal DNA sequence analysis of Heterodera filipjevi and H. latipons isolates from Russia with sequences from other nematode isolates. Russian Journal of Nematology 7:121–5.

Gäbler, C., Sturhan, D., Subbotin, S. A. and Rumpenhorst, H. J. 2000. Heterodera pratensis sp. n., a new cyst nematode of the H. avenae complex (Nematoda: Heteroderidae). Russian Journal of Nematology 8:115–26.

Goffart, H. 1932. Untersuchungen am Hafernematoden Heterodera schachtii Schm. unter besonderer Berücksichtigung der schlewigholsteinischen Verhältnisse. I. III. Beitrage zu: Rassenstudien an Heterodera schachtii Schm. Arbeiten aus der Biologischen Reichsanstalt, Berlin 20:1–26.

Goffart, H. 1936. Heterodera Schachtii Schmidt an gemeiner Hanfnessel (Galeopsis tetrahit L.) und an Kakteen. Zeitschrift für Parasitenkunde 8:528–32.

Golden, A. M. and Cobb, G. S. 1963. Heterodera lespedezae (Heteroderidae) a new species of cyst-forming nematode. Proceedings of the Helminthological Society of Washington 30:281–6.

Handoo, Z. A. and Subbotin, S. A. 2018. “Taxonomy, identification and principal species”, In Perry, R. N., Moens, M. and Jones, J. T. (Eds), “Taxonomy, identification and principal species”, In Perry, R. N., Moens, M. and Jones, J. T. (Eds), “Taxonomy, identification and principal species”, In Perry, R. N., Moens, M. and Jones, J. T. (Eds), “Taxonomy, identification and principal species”, In Perry, R. N., Moens, M. and Jones, J. T. (Eds), “Taxonomy, identification and principal species”, In Perry, R. N., Moens, M. and Jones, J. T. (Eds), “Taxonomy, identification and principal species”, In Perry, R. N., Moens, M. and Jones, J. T. (Eds), “Taxonomy, identification and principal species”, In Perry, R. N., Moens, M. and Jones, J. T. (Eds), “Taxonomy, identification and principal species”, In Perry, R. N., Moens, M. and Jones, J. T. (Eds), “Taxonomy, identification and principal species”, In Perry, R. N., Moens, M. and Jones, J. T. (Eds), “Taxonomy, identification and principal species”, In Perry, R. N., Moens, M. and Jones, J. T. (Eds). Bioinformatics 17:754–5.

Ichinohe, M. 1952. On the soybean nematode, Heterodera glycines n. sp., from Japan. Magazine of Applied Zoology 17:1–9.

Kazachenko, I. P. 1993. Cyst-forming nematodes of Far East and their control. Vladivostok, USSR: Dalnauka.

Kirjanova, E. S. 1969. On the structure of the subcystcrntilline layer of the nematode genus Heterodera (Nematoda: Heteroderidae) with a description of two new species. Parazitologiya 3:81–91.

Kirjanova, E. S. and Krall, E. L. 1971. Parasitic nematodes of plants and their control. Vol. II. Leningrad, USSR: Nauka.

Kirjanova, E. S., Krall, E. L. and Krall, H. 1976. The sowthistle cyst-nematode Heterodera sonchophila n. sp. (Nematoda: Heteroderidae) from Estonia. Eesti NSV Teaduste Akademia Toimetised, Biologiline Seeria 25:305–25.

Koshy, P. K. 1967. A new species of Heterodera from India. Indian Phytopathology 20:272–4.

Mayer, W. E., Herrmann, M. and Sommer, R. J. 2007. Phylogeny of the nematode genus Pristionchus and implications for biodiversity, biogeography and the evolution of hermaphroditism. BMC Evolutionary Biology 7:104.

Nadler, S. A., Bolotin, E. and Stock, S. P. 2006. Phylogenetic relationships of Steinernema Travassos, 1927 (Nematoda: Cephalobina: Steinernematidae) based on nuclear, mitochondrial and morphological data. Systematic Parasitology 63:159–79.

Nunn, G. B. 1992. Nematode molecular evolution. An investigation of evolutionary patterns among nematodes based upon DNA sequences. PhD thesis, University of Nottingham.

Perry, R. N., Moens, M. and Jones, J. T. 2018. Cyst Nematodes. Oxfordshire: CABI, Oxfordshire.

Robinson, A. J., Stone, A. R., Hooper, D. J. and Rowe, J. A. 1996. A redescription of Heterodera arenaria Cooper 1955, a cyst nematode from marram grass. Fundamental and Applied Nematology 19:109–17.

Schmidt, A. 1871. Über den Rümennematoden. Zeitschrift der Vereinte Rübenzuckerindustrie Zollverein 21:1–19.

Sharma, S. B., Siddiqi, M. R., Rahaman, P. F., Ali, S. S. and Ansari, M. A. 1998. Description of Heterodera swarupi sp. n. (Nematoda: Heteroderidae), a parasite of chickpea in India. International Journal of Nematology 8:111–16.

Sikora, R. A., Coyne, D., Hallmann, J. and Timper, P. 2018. Plant Parasitic Nematodes in Subtropical and Tropical Agriculture. Boston: CABI.

Singh, P. R., Couvreur, M., Decraemer, W. and Bert, W. 2019. Survey of slug-parasitic nematodes in East and West Flanders, Belgium and description of Angiostoma gandavensis n. sp. (Nematoda: Angiostomidae) from arionid slugs. Journal of Helminthology 94.

Singh, P. R., Nyiragatane, A., Janssen, T., Couvreur, M., Decraemer, W. and Bert, W. 2018. Morphological and molecular characterisation of Pratylenchus rwandae n. sp. (Tylenchida: Pratylenchidae) associated with maize in Rwanda. Nematology 20:781–94.

Subbotin, S. A., Mundo-Ocampo, M. and Baldwin, J. G. 2010. Systematics of cyst nematodes (Nematoda: Heteroderidae). Nematology Monographs and Perspectives 8B, Leiden: Brill.

Subbotin, S. A., Sturhan, D., Rumpenhorst, H. J. and Moens, M. 2003. Molecular and morphological characterisation of the Heterodera avenae species complex (Tylenchida: Heteroderidae). Nematology 5:515–38.

Subbotin, S. A., Toumi, F., Elekciójú, I. H., Waeyenberge, L. and Maafi, Z. T. 2018. DNA barcoding, phylogeny and phylogeography of the cyst nematode species of the Avenae group from the genus Heterodera (Tylenchida: Heteroderidae). Nematology 20:671–702.

Subbotin, S. A., Akanwari, J., Nguyen, C. N., Cid Del Prado Vera, I., Chitambar, J. J., Inserra, R. N. and Chizhov, V. N. 2017. Molecular characterisation and phylogenetic relationships of cystoid nematodes of the family Heteroderidae (Nematoda: Tylenchida). Nematology 19:1065–81.

Toumi, F., Waeyenberge, L., Viene, N., Dababat, A., Nicol, J. M., Ogbonnaya, F. and Moens, M. 2013a. Development of a species-specific PCR to detect the
heterodera dunensis n. sp. from canary islands: Singh et al.

cereal cyst nematode Heterodera latipons. Nematology 15:709–17.

Van der Putten, W. and Van der Stoel, C. 2006. Pathogenicity and host range of Heterodera arenaria in coastal foredunes. Nematology 8:255–63.

Vovlas, N., Greco, N. and Di Vito, M. 1985. Heterodera ciceri sp. n. (Nematoda: Heteroderidae) on Cicer arietinum L. from northern Syria. Nematologica Mediterranea 13:239–52.

Vovlas, N., Inserra, R. N. and Stone, A. R. 1981. Heterodera mediterranea n. sp. (Nematoda: Heteroderidae) on Pistacia lentiscus in southern Italy. Nematologica 27:129–38.

Vrain, T. C., Wakarchuk, D. A., Levesque, A. C. and Hamilton, R. I. 1992. Intraspecific rDNA restriction fragment length polymorphism in the Xiphinema americanum group. Fundamental and Applied Nematology 15:563–73.

Whitehead, A. G. and Hemming, J. R. 1965. A comparison of some quantitative methods of extracting small vermiform nematodes from soil. Annals of Applied Biology 55:25–38.

Wouts, W. M. and Sturhan, D. 1978. The identity of Heterodera trifolii Goffart, 1923 and the description of H. daverti n. sp. (Nematoda: Tylenchida). Nematologica 24:121–8.

Wouts, W. M. and Sturhan, D. 1995. Heterodera aucklandica sp. n. (Nematoda: Heteroderidae) from a New Zealand native grass, with notes on the species of the H. avenae group. New Zealand Journal of Zoology 22:199–207.

Wouts, W. M. and Sturhan, D. 1996. Heterodera litoralis sp. n. (Nematoda: Heteroderidae) from austral glasswort, Sarcocornia quinqueflora, in New Zealand. Nematologica 42:62–70.

Wouts, W. M., Rumpenhorst, H. J. and Sturhan, D. 2001. Heterodera betae sp. n., the yellow beet cyst nematode (Nematoda: Heteroderidae). Russian Journal of Nematology 9:33–42.