1 Comparative molecular and morphological characterisations in the 2 3 nematode genus Rotylenchus: Rotylenchus paravitis n. sp., an example of cryptic speciation 4 5 Carolina Cantalapiedra-Navarrete¹, Juan A. Navas-Cortés¹, Gracia Liébanas², Nicola Voylas³, 6 Sergei A. Subbotin^{4,5}, Juan E. Palomares-Rius¹, and Pablo Castillo^{1,*} 7 8 9 ¹Institute for Sustainable Agriculture (IAS), Spanish National Research Council (CSIC), 10 Alameda del Obispo s/n, Apdo. 4084, 14080 Córdoba, Campus de Excelencia Internacional 11 Agroalimentario, ceiA3, Spain ² Department of Animal Biology, Vegetal Biology and Ecology, University of Jaén, Campus 12 13 'Las Lagunillas' s/n, Edificio B3, 23071 Jaén, Spain ³ Istituto per la Protezione delle Piante, UOS-Bari, Consiglio Nazionale delle Richerche 14 (C.N.R.), Via Amendola 122/D, 70126 Bari, Italy 15 ⁴ Plant Pest Diagnostic Center, California Department of Food and Agriculture, 3294 16 17 Meadowview Road, Sacramento, CA 95832-1448, USA ⁵ Center of Parasitology of A.N. Severtsov Institute of Ecology and Evolution of the Russian 18 19 Academy of Sciences, Leninskii Prospect 33, Moscow, 117071, Russia 20 21 22 Received: ; revised:.....; accepted for publication: 23 24 25 *Corresponding author

26

E-mail address: p.castillo@csic.es

Abstract

The nematode Rotylenchus paravitis n. sp. infesting roots of commercial sunflowers in
southern Spain is described. The new species is characterised by a truncate lip region with 7-9
annuli and continuous with the body contour, lateral fields areolated at pharyngeal region
only, body without longitudinal striations, stylet length of 44-50 μm , vulva position at 43-
54%, tail rounded to hemispherical with 12-18 annuli. A comparative phenetic study based on
a multivariate principal component analysis was developed to determine potential species
discrimination. The degree of variation for most characters among specimens of Rotylenchus
paravitis n. sp. and R. vitis was comparable to that observed among specimens belonging to
each of the two studied populations of R. robustus from Spain and USA. Molecular
comparison of the partial 18S, D2-D3 expansion segments of 28S-rRNA, ITS1-rRNA, partial
COI and hsp90 from R. paravitis n. sp. and R. vitis, and other species in the genus, clearly
supports the proposal of R. paravitis n. sp. as a new species. Consequently, R. paravitis n. sp.
should be considered as an example of cryptic speciation within the genus Rotylenchus. PCR-
ITS-RFLP was provided for diagnostics of <i>R. paravitis</i> n. sp. and PCR with specific primers
were also developed for diagnostics of this new species, R. vitis and R. robustus. The results
of the phylogenetic analysis based on the sequences of the D2-D3 expansion regions of the
28S, ITS1-rRNA genes, and the partial COI, have proven to be a powerful tool for providing
accurate species identification and assessing phylogenetic relationships within the genus
Rotylenchus. Phylogenetic testing of D2-D3 expansion segments of 28S-rRNA gene
sequences did not refute the monophyly of the genera Rotylenchus, Helicotylenchus,
Hoplolaimus, based on tree topologies and the Shimodaira-Hasegawa test even with the split
in several clades for some of the genera.

Keywords: Nematoda, new species, phylogeny, spiral nematodes, taxonomy.

2

1. Introduction

3

4

5

6

7

8

9

10

11

12

13

14

15

16

17

18

19

20

21

22

23

24

25

26

The genus Rotylenchus Filipiev, 1936 belongs to Hoplolaimidae Filipiev, 1934, a family which also contains agricultural and economically important genera such as Helicotylenchus Steiner 1945, Hoplolaimus Von Daday 1905, Rotylenchulus Lindford & Oliveira 1940 and Scutellonema Andrássy 1958. This genus tends to be greatly conserved in gross morphology which makes species identification a very difficult task. More than 95 valid species have been recognised in this genus, which confirms the previously mentioned difficulty for identification (Castillo and Vovlas, 2005; Atighi et al., 2011; Cantalapiedra-Navarrete et al., 2012). All the known *Rotylenchus* spp. are obligate plant parasites of a wide range of wild and cultivated plants and are closely associated with plant roots. They are migratory ectoparasites and browse on the surface of roots. As migratory ectoparasites do not enter the plant root, the damage they cause is usually limited to necrosis of those cells penetrated by stylet. The large number of species within the genus *Rotylenchus* complicates the identification process and has required the construction of tabular and dichotomous keys, based on a combination of major and supplementary characters, to enable pragmatic morphological identification (Castillo and Vovlas 2005). Recently, DNA-based approaches have been successfully used for the molecular diagnostics of *Rotylenchus* (Vovlas et al., 2008; Atighi et al., 2011; Cantalapiedra-Navarrete et al., 2012). Phylogenetic studies within Rotylenchus have been carried out based on the D2-D3 expansion regions of the 28S and ITS rRNA gene, providing initial insight toward resolving phylogenetic relationships among Rotylenchus, and demonstrating paraphyly of the genus in the majority rule consensus trees within Hoplolaimidae (Subbotin et al., 2007; Vovlas et al. 2008; Cantalapiedra-Navarrete et al., 2012). Molecular analysis of D2-D3 sequences also revealed several Rotylenchus species

1 with almost identical sequences, i.e. Rotylenchus goodeyi Loof and Oostenbrink, 1958, 2 Rotylenchus incultus Sher, 1965, and Rotylenchus laurentinus Scognamiglio and Talamé, 3 1973; or Rotylenchus robustus (de Man, 1876) Filipjev, 1936 and Rotylenchus uniformis 4 (Thorne, 1949) Loof and Oostenbrink, 1958 (Vovlas et al., 2008). Mitochondrial DNA 5 (mtDNA), particularly the protein-coding mitochondrial gene, cytochrome c oxidase subunit 1 6 (COI), has proven to be a powerful tool for providing accurate species identification and 7 assessing phylogenetic patterns across the animal kingdom, including plant-parasitic and free-8 living nematodes (Hugall et al., 1994; Derycke et al., 2010; Gutiérrez-Gutiérrez et al., 2011). 9 Similarly, the heat-shock protein *hsp90* gene has also been considered to be a useful 10 molecular marker for species identification or phylogenetic analysis of plant-parasitic 11 nematode species (Skantar and Carta, 2004; Madani et al., 2011). Consequently, both markers 12 may be valuable tools for diagnostics as well as for clarifying phylogenetic relationships in 13 those Rotylenchus species with almost identical D2-D3, although no information of these 14 markers is available in this genus and so they need to be developed. Also, molecular 15 techniques have recently shown that many presumed monospecific species are in fact siblings 16 or cryptic species (Subbotin et al., 2003; Vovlas et al., 2008; Gutierrez-Gutiérrez et al., 2010). 17 Consequently, the nematode species concept should be based on principles of polyphasic 18 taxonomy, which assembles and assimilates all available data and information (phenotypic, 19 morphometric, genotypic and phylogenetic) used for delimiting taxa at all levels (Subbotin 20 and Moens, 2006; Vovlas et al., 2008). 21 Nematode surveys in agricultural and natural environments in Southern Spain revealed 22 low to moderate soil infestations by two amphimictic populations of *Rotylenchus* species. 23 Preliminary morphological observations indicated that these species appeared to be 24 morphologically similar to Rotylenchus robustus (de Man, 1876) Filipiev, 1936, and 25 Rotylenchus vitis Cantalapiedra-Navarrete, Liébanas, Archidona-Yuste, Palomares-Rius and 26 Castillo, 2012. Detailed observations using light and scanning electron microscopy (SEM).

and molecular characterisation analyses, indicated that these specimens should be assigned to
 R. robustus and to a new species showing close morphological and morphometric
 resemblance with *R. vitis*, but clearly differentiated by molecular analyses, being considered
 as an example of cryptic species within the genus *Rotylenchus* (i.e. species genetically distinct
 but sharing common morphological diagnostic characters). In addition, several *Rotylenchus*

out a morphological characterization combined with molecular analyses which may clarify the

populations from Australia and USA were collected from natural and cultivated areas to carry

phylogeny of the genus.

Therefore, the objectives of this study were: *i*) to conduct a comparative phenetic study of the Spanish species resembling *R. vitis* with holotype and paratypes of *R. vitis* of the nematode collection from IAS-CSIC, Córdoba, Spain, as well as two populations of *R. robustus* from Spain and USA, using the most useful diagnostic morphological and morphometric characters for *Rotylenchus* species based on a multivariate principal component analysis; *ii*) to verify the taxonomic status of this species close to *R. vitis*, which is described herein as *Rotylenchus paravitis* n. sp., as well as other species from Australia, Spain and USA, conducting detailed morphometric and molecular studies of these *Rotylenchus* species; *iii*) to determine the molecular phylogenetic affinities of *R. paravitis* n. sp. and other known *Rotylenchus* species identified in Australia, Spain and USA with closely related species using the rRNA gene sequences (D2-D3 of 28S and ITS1-rRNA), the partial sequences of mitochondrial gene *COI*, and heat shock protein (*hsp90*) gen; and *iv*) to provide PCR-ITS-RFLP for *R. paravitis* n. sp. and develop PCR with species specific primers for diagnostics of *R. paravitis* n. sp., *R. vitis* and *R. robustus*.

2. Material and methods

2.1. Nematode populations

2

3

4

5

6

7

8

9

10

11

12

13

14

15

16

17

18

19

20

21

22

23

24

25

26

Nematodes of R. paravitis n. sp. used in this study were obtained from the rhizosphere of sunflower plants in Jerez de la Frontera (Cádiz province), southern Spain (36°46'28.85"N latitude, 6°15'26.27" W longitude) at an altitude of 16 m a.s.l., and were collected with a shovel from the upper 30 cm of soil on May 2011 by J. Martin-Barbarroja and G. León-Ropero (IAS-CSIC). In addition, Spanish and American populations of R. robustus from stone pine in Lucena del Puerto (Huelva province, southern Spain) and grasses in Tomales. California (USA), respectively, a population of *Rotylenchus brevicaudatus* Colbran, 1962 from grasses in Brisbane (Australia), and American populations of Rotylenchus buxophilus Golden, 1956, and Rotylenchus pumilus (Perry et al., 1959) Sher, 1961, were collected and studied morphometrically and molecularly. In addition, previously morphologically wellcharacterized Rotylenchus populations (Vovlas et al., 2008) from Italy and Spain (including Rotylenchus cazorlaensis Castillo and Gómez-Barcina, 1987, Rotylenchus eximius Siddigi 1964, Rotylenchus jaeni Vovlas, Subbotin, Troccoli and Castillo, 2008, R. laurentinus, and Rotylenchus magnus Zancada, 1985) were collected to carry out molecular analyses of the partial COI sequences which may clarify the phylogeny of the genus. For molecular analyses, a list of studied *Rotylenchus* species and populations is given in Table 1. The nematodes were extracted from rhizosphere soil samples by the centrifugal-flotation method (Coolen, 1979). Specimens to be observed under light microscopy (LM) were killed by gentle heat, fixed in a solution of 4% formaldehyde + 1% propionic acid and processed to pure glycerin using De Grisse's (1969) method. Specimens were examined using a Zeiss III compound microscope with Nomarski differential interference contrast at up to ×1000 magnification. Measurements were done using a *camera lucida* attached to this microscope. For line drawing, handmade pictures were scanned and imported to CorelDraw software version 12 and redrawn. LM micrographs were based on live specimens for the Spanish population (Fig. 3), and glycerine-mounted specimens for the American populations (Fig. 5-7).

For SEM studies, fixed specimens were dehydrated in a graded ethanol series, critical point dried, sputter-coated with gold and observed with a JEOL JSM-5800 microscope (Abolafia et al., 2002).

2.2. Multivariate principal component analysis

A multivariate principal component analysis was performed on *R. vitis, R. paravitis* n. sp., as well as two populations of *R. robustus* from Spain and USA in order to determine the morphometric discrimination among species. The analyses were based upon the following characters: body length (L), lip width, lip height, number of lip annuli, stylet length, stylet-conus length, knobs width, dorsal gland orifice (D.G.O.), anterior end to beginning of median pharyngeal bulb distance, anterior end to centre of median pharyngeal bulb distance, total pharyngeal length, anterior end to pharyngo-intestinal junction distance, anterior end, maximum body width, pharyngeal overlapping, cuticle tail tip width, vulva position (V), anterior and posterior gonads, female tail length, anal body width, number of female tail annuli, phasmid to terminus distance, phasmid to anus distance, number of annuli between phasmid and anus, and the de Man ratios a, b, b', c, and c', G₁, G₂ and O (Siddiqi, 2000; Table 2).

Principal component analysis was performed with the PRINCOMP procedure of SAS (Statistical Analysis System, version 9.2; SAS Institute Inc., Cary, NC, USA). This analysis produced a set of variables (principal components) that were linear combinations of the original variables. The new variables (principal components) were independent of each other and ranked according to the amount of variation accounted for.

2.3. DNA extraction, PCR and sequencing

1 Nematode DNA from R. paravitis n. sp. and other studied Rotylenchus species was 2 extracted from single individuals using proteinase K as described by Castillo et al. (2003). 3 Detailed protocols for PCR and sequencing were applied as described by Castillo et al. 4 (2003). The primers used for amplification of D2-D3 regions of 28S, ITS1-rRNA, the partial 5 18S, the COI and hsp90 gene, as well as those for species specific PCR are listed in Table 3. 6 These sequences were used for molecular species characterization and phylogenetic analyses. 7 PCR products were purified after amplification with Geneclean turbo (O-BIOgene SA, 8 Illkirch Cedex, France) or QIAquick (Qiagen, USA) gel extraction kits, quantified using a 9 Nanodrop spectrophotometer (Nanodrop Technologies, Wilmington, DE, USA) and used for 10 direct sequencing in both directions using the primers referred above. The resulting products 11 were purified and run on a DNA multicapillary sequencer (Model 3130XL genetic analyser; 12 Applied Biosystems, Foster City, CA, USA), using the BigDye Terminator Sequencing Kit 13 v.3.1 (Applied Biosystems, Foster City, CA, USA), at the SCAI, University of Córdoba 14 sequencing facilities (Córdoba, Spain). The newly obtained sequences were submitted to the 15 GenBank database under accession numbers indicated on the phylogenetic trees and Table 1.

16

2.4. Phylogenetic analysis

18

19

20

21

22

23

24

25

26

17

D2-D3 expansion segments of 28S, ITS1-rRNA, partial *COI*, the partial 18S, and partial *hsp90* gen newly sequenced and sequences obtained from GenBank were used for phylogenetic reconstruction. Outgroup taxa for D2-D3 expansion regions of 28S-rRNA dataset was chosen according to previous published data (Vovlas et al., 2008). The newly obtained and published sequences for each gene were aligned using ClustalW (Thompson et al., 1997) with default parameters. Sequence alignments were manually edited using BioEdit (Hall, 1999). Phylogenetic analyses of the sequence data sets were performed under maximum likelihood (ML) using PAUP * 4b10 (Swofford, 2003) and Bayesian inference (BI)

1 using MrBayes 3.1.2 (Huelsenbeck and Ronquist, 2001). The best fitting model of DNA 2 evolution was obtained based on the Akaike Information Criterion (AIC) using iModelTest v. 3 0.1.1 (Posada, 2008) with. The Akaike-supported model, the base frequency, the proportion of 4 invariable sites, and the gamma distribution shape parameters and substitution rates in the 5 AIC were then used in phylogenetic analyses. BI analysis under TVM + I + G model for D2-6 D3 expansion segment of 28S-rRNA; TVM + G for ITS1 region and GTR + I + G for COI, 7 TIM1 + I + G for partial 18S, and K80 + G for hsp90 gene were run with four chains for 5×10^{-1} 10^6 , 1×10^6 , 1×10^6 , 1×10^6 , and 2×10^6 generations, respectively. The Markov chains were 8 9 sampled at intervals of 100 generations. Two runs were performed for each analysis. After 10 discarding burn-in samples and evaluating convergence, the remaining samples were retained 11 for further analyses. The topologies were used to generate a 50% majority rule consensus tree. 12 Posterior probabilities (PP) are given on appropriate clades. Trees were visualised using 13 TreeView (Page, 1996). In ML analysis the estimation of the support for each node was 14 obtained by bootstrap analysis with 100 fast-step replicates. In order to test alternative tree 15 topologies by constraining hypothetical monophyletic groups, we performed the Shimodaira-16 Hasegawa test (SH-test) as implemented in PAUP (Swofford, 2003) and using the RELL 17 option, based on D2-D3 expansion segments of 28S and partial 18S from 37 and 15 selected 18 taxa, respectively: D2-D3 expansion segments of 28S for the genera Rotylenchus, 19 Helicotylenchus, Scutellonema, Aorolaimus and Hoplolaimus; partial 18S for Rotylenchus, 20 Helicotylenchus and Scutellonema. Aglenchus agricola (AY780979, FJ969113), Coslenchus 21 costatus (DQ328719, AY284581) and Basiria gracillis (DQ328717, EU130839) were used as 22 outgroups in both datasets (Vovlas et al., 2008; Subbotin et al., 2007). Species in datasets 23 including D2-D3 expansion segments of 28S were selected from the clades formed in the 24 previous study of phylogeny of Hoplolaimidae performed by Cantalapiedra-Navarrete et al. 25 (2012). Tests constraining hypothetical groups were performed using ML (Table 8).

2.5. RFLP-ITS-rRNA

Three to seven µl of purified PCR product of the D2-D3 of 28S-rRNA gene for *R. paravitis* n. sp. was digested by one of following restriction enzymes: *AvaI*, *RsaI*, *BseNI*, *MvaI* or *HpaII*, in the buffer stipulated by the manufacturer. The digested DNA was run on a 1.4% TAE buffered agarose gel, stained with ethidium bromide, visualized on UV transilluminator and photographed. The exact lengths of each restriction fragment from the PCR products were obtained by a virtual digestion of the sequences using WebCutter 2.0 (www.firstmarket.com/cutter/cut2.html).

2.6. PCR with species specific primers

Species specific primers were developed for several species: *R. paravitis* n. sp., *R. vitis* parasitizing grapevine in Spain and the type species of the genus and agricultural important species *R. robustus*. Specific primers were designed based on unique fragments for each species using sequence alignment of ITS-rRNA gene. Several *Rotylenchus* samples were used to test species specific primers. PCR mixture was prepared as described in Tanha Maafi et al. (2003). The universal forward TW81 primer was used in PCR with combinations of the specific reverse *R. paravitis*, *R. vitis* or *R. robustus* primers (Table 3) for diagnostics of *R. paravitis* n. sp., *R. vitis* or *R. robustus*, respectively. The PCR amplification profile consisted of 4 min at 94°C; 30 cycles of 1 min at 94°C, 45 s at 57°C and 45 s at 72°C, followed by a final step of 10 min at 72°C. Four μl of the PCR product was run on a 1.4% TAE buffered agarose gel, stained and photographed.

3. Results

3.1. Comparative multivariate analysis of Rotylenchus paravitis n. sp., Rotylenchus vitis,

and two populations of Rotylenchus robustus

In the principal component analysis, the first four principal components accounted for 67.5% of the total variance of morphological and morphometric characters of the four *Rotylenchus* populations included in the analysis (Table 2). Table 2 includes the eigenvalues for the first four principal components (PC) that were used to interpret the significance of the PCs. Principal component 1separates *R. vitis* and *R. paravitis* n. sp. specimens from the two populations of *R. robustus*, characterized by smaller values of stylet-conus length, D.G.O, and O ratio, and higher values for body size (length and width), knobs width, cuticle tail tip width, phasmid to terminus and anus distances, number of annuli between phasmid and anus, and b ratio (Table 2, Fig. 1), but did not separate *R. vitis* from *R. paravitis* n. sp. Similarly, the stylet conus and orifice of dorsal gland of *R. robustus* specimens were lower than that observed for *R. vitis* and *R. paravitis* n. sp. with no differences between these two species or between the two *R. robustus* populations (Fig. 1A-C).

Principal component 2 is dominated by high negative weights for anterior body end to

Principal component 2 is dominated by high negative weights for anterior body end to centre and beginning of median pharyngeal bulb distances, total pharyngeal length and anterior end to pharyngo-intestinal junction distance (Table 2, Fig. 1A), relating this factor with pharyngeal dimensions. These characters showed a large but similar degree of variation among specimens of the three *Rotylenchus* species in the study (Fig. 1A).

When projected on the plane of PCs 1 and 3, female tail shape and size and c' ratio decreased from bottom to top. According to their relative position to y axis in Figure 1B, specimens of *R. paravitis* n. sp. were characterized by lower values for these two characters when compared to that of *R. vitis*, although some specimens of the two species showed also similar values (Fig. 1B). Similarly, when projected on the plane of PCs 1 and 4, specimens of *R. vitis* tended to have longer gonads than that of *R. paravitis* n. sp. (Fig. 1 B, C).

1 Concerning R. robustus populations, a similar range of variation for characters related 2 to PC 3 (Fig. 1B) or PC 4 (Fig. 1C) were observed among specimens of the two R. robustus 3 populations. 4 5 3.2. Systematics 6 Family Hoplolaimidae Filipjev, 1934 7 Rotylenchus Filipiev, 1936 8 **3.2. 1.** *Rotylenchus paravitis* n. sp. 9 Figs. 2-4; Table 4. 10 11

3.2.1.1. Holotype and paratypes

Holotype female, extracted from in a clay soil from the rhizosphere of sunflower (Helianthus annuus L.) in Jerez de la Frontera, Cádiz province, southern Spain, by J. Martín-Barbarroja and G. León Ropero, mounted in pure glycerine and deposited in the nematode collection at Institute for Sustainable Agriculture (IAS) of Spanish Council for Scientific Research (CSIC), (collection number IAS-J174-02). Female and male paratypes extracted from the same soil as holotype, and deposited in the Nematode Collection at the Institute for Sustainable Agriculture, CSIC, Córdoba, Spain. Another two female paratypes deposited at each of the following collections: Istituto per la Protezione delle Piante (IPP) of Consiglio Nazionale delle Ricerche (C.N.R.), Sezione di Bari, Bari, Italy; USDA Nematode Collection, Beltsville, MD, USA (collection number G-18577) and WaNeCo, Plant Protection Service, Wageningen, The Netherlands. Specific D2-D3, ITS1, 18S-rRNA, COI and hsp90 sequences are deposited in GenBank with accessions numbers JX015422, JX015434, JX015429, JX015415, JX015416, and JX015393, respectively.

26

12

13

14

15

16

17

18

19

20

21

22

23

24

3.2.1.2. Diagnosis

Rotylenchus paravitis n. sp. is a gonochoristic species assigned to the species group having more than six lip annuli, a truncate lip region, female tail usually hemispherical to broadly rounded and stylet more than 35 μm long (Castillo and Vovlas, 2005). It is characterised by a truncate lip region clearly narrowing in the first half, with 7-9 annuli, continuous with body contour, lateral fields areolated at pharyngeal region only, body without longitudinal striations, stylet length of 44-50 μm, vulva position at 43-54%, tail rounded to hemispherical, with 12-18 annuli, and a specific D2-D3, ITS1-rRNA, partial 18S, *COI* and *hsp90* sequences. Intra-specific variability of D2D3 was evaluated by sequencing two specimens showing high similarity (99%), differing in 1/682 nucleotides and showing no indels between them.

3.2.1.3. Description

Female: Body large, habitus slightly curved ventrally to open C-shaped. Lateral fields with four smooth equidistant lines, beginning anteriorly at $12\text{-}14^{\text{th}}$ annulus as three lines forming two bands; after 20-23 annuli, central line dividing to form a third band. The three bands are 10.6 ± 0.8 (9.5-12.0) μm wide at mid-body, approximately one-fifth as wide as body diam. Regular areolation of lateral fields (external bands) observed in pharyngeal region. Cuticle 2.0-3.0 μm thick, clearly annulated, annuli 1.5 μm wide at mid-body. Body without longitudinal striations in any region. Labial region truncate, continuous with body, clearly narrowing in the first half, bearing 7-9 narrow annuli which are not divided longitudinally. Labial disc distinct under SEM, but not in lateral view under LM. Oral aperture oval $(1.0\text{-}1.2 \times 0.8\text{-}1.0 \,\mu\text{m})$, labial disc oval $(4.5\text{-}5.0 \times 3.5\text{-}4.0 \,\mu\text{m})$ bordered by amphidial apertures. Oral disc clearly separated from first annulus of lip region, which is

1 undivided. Labial framework well developed, $5.3 \pm 0.4 \mu m$ (5-6) long, posterior margin at level of 7th or 9th body annulus. Stylet robust, 3.1-3.8 times longer than labial region diam. 2 Basal knobs slightly flattened posteriorly, 8.0-10.5 um wide, at level of 25th-32th annulus 3 4 posterior to labial region. Dorsal pharyngeal gland opening 4.0-7.0 µm posterior to stylet 5 base. Procorpus of pharynx cylindrical, with slight depression just anterior to median bulb, 6 51.4 ± 5.0 (44-61) µm long. Median pharyngeal bulb well developed, broadly oval, $21-29 \times$ 7 19-24 µm, occupying 10-12 annuli, valvular apparatus 5.0-6.5 µm long, located at 46.1 ± 3.8 8 (41-51)% of pharyngeal length. Isthmus 23.4 \pm 2.8 (19-28) μ m long, encircled by nerve ring 9 at mid-point. Pharyngeal glands sacciform, with three nuclei, overlapping intestine dorsally 10 for 24-52 annuli. Nerve ring enveloping isthmus at middle, at 127 ± 10.8 (113-153) µm from 11 anterior end. Excretory pore usually located near pharyngo-intestinal junction level, but 12 varying from anterior to posterior of pharyngeal-intestinal valve. Hemizonid distinct, located 13 at anterior to excretory pore, extending for ca 1.5-2.0 body annuli width, just anterior to 14 excretory pore, rarely posterior. Reproductive system with both genital branches equally 15 developed; anterior branch 315 ± 64.1 (217-416) µm long, posterior branch 300 ± 54.9 (203-16 369) µm long. Vulva slightly posterior to mid-body, undistinct epiptygma under LM but 17 clearly distinguishable under SEM (Fig. 4D, E). Vagina with internal walls slightly 18 sclerotised, 20.5 (18-24) µm long. Ovaries with a single row of oocytes, spermathecae not 19 functional, without sperm. Phasmids pore-like, 19.1 ± 2.6 (12-24) annuli anterior to level of 20 anus or 24.9 ± 3.9 (15.5-28.0) µm anterior to anus. Tail short, usually hemispherical to 21 broadly rounded, with 12-18 annuli, terminus striated and cuticle at tail tip 8.2 ± 0.6 (7.5-22 10.0) um wide. 23 24 *Male*: Very rare, only one specimen detected. Habitus slightly curved ventrally.

25 Morphology similar to that of females, except for the following characters: lip region slightly 26 more elevated in outline, 10.0 um wide and 5.0 um high; stylet shorter than that of female:

1 stylet knobs less developed (5.5 µm wide). Testis single, anteriorly outstretched, 494 µm 2 long. Tail conoid, 1.7 times shorter than that of female. Phasmid distinct, situated anteriorly at 3 anus level. Bursa 54 µm long, completely surrounding tail which is tapering with a rounded-4 pointed tip. Gubernaculum prominent, with distinct titillae. 5 6 **3.2.1.4.** Type locality 7 8 Rotylenchus paravitis n. sp. was found in a clay soil from the rhizosphere of sunflower 9 (Helianthus annuus L.) in Jerez de la Frontera, Cádiz province, southern Spain. 10 11 **3.2.1.5. Etymology** 12 13 The species epithet refers to Gr. prep. para, alongside of, resembling; N.L. masc. n. 14 vitis, a morphologically close species (Rotylenchus vitis). 15 **3.2. 2. Rotylenchus brevicaudatus** Colbran, 1962 16 17 Fig. 5; Table 5. 18 Female: Habitus spiral. Lip region continuous with body contour, marked by 4 annuli. 19 20 Lateral fields with four smooth equidistant lines, the three bands are 5.1 ± 0.3 (5.0-5.5) µm 21 wide at mid-body, approximately one-fifth as wide as body diam. Regular areolation of lateral 22 fields (external bands) observed in pharyngeal region. Cuticle 1.5 µm thick, clearly annulated, 23 one annulus 1.5 µm wide at mid-body. Body without longitudinal striations in any region.

posterior to stylet base. Procorpus of pharynx cylindrical, with slight depression just anterior

Stylet moderately robust, 2.3-2.5 times longer than labial region diam. Basal knobs rounded

slightly backwards directed, 4.0-4.5 µm wide. Dorsal pharyngeal gland opening 5.0-5.5 µm

24

25

Comparative molecular and morphological characterisations in *Rotylenchus*

- 1 to median bulb, 34.8 ± 1.0 (34-36) µm long. Median pharyngeal bulb well developed, broadly
- 2 oval, $10.0-11.0 \times 9.0-10.0$ μm, occupying 6-8 annuli, valvular apparatus 2.0-2.5 μm long,
- 3 located at $55.6 \pm 4.1 (51-59)\%$ of pharyngeal length. Isthmus $22.5 \pm 1.3 (21-24) \mu m long$,
- 4 encircled by nerve ring at mid-point. Pharyngeal glands short, with three nuclei, slightly
- 5 overlapping intestine dorsally for 4-6 annuli. Nerve ring enveloping isthmus at middle, at $81 \pm$
- 6 3.6 (76-84) μm from anterior end. Excretory pore usually located near pharyngo-intestinal
- 7 junction level. Hemizonid distinct, 2-3 annuli long, located 1-3 annuli anterior to excretory
- 8 pore, extending for ca 1.0-1.5 body annuli width. Reproductive system with both genital
- 9 branches equally developed; anterior branch 91 \pm 18.9 (72-114) µm long, posterior branch 86
- ± 19.3 (67-111) µm long. Vulva clearly posterior to mid-body, without a distinct epiptygma.
- 11 Vagina with internal walls slightly sclerotized, 10.5 (10.0-11.0) μm long. Ovaries with a
- single row of oocytes. Phasmids pore-like, 10.0 ± 2.8 (8-14) annuli anterior to level of anus or
- 13 13.9 ± 1.9 (12.0-16.5) µm anterior to anus. Tail short, conoid-rounded, terminus
- hemispherical, coarsely striated and cuticle at tail tip 2.6 ± 0.3 (2.5-3.0) µm wide.

15

- Male: Habitus a closed C shape. Lip region similar than that of female. Bursa crenate,
- enveloping tail, 43.7 ± 2.4 (41-48) µm. Spicules slightly cephalated and ventrally arcuate.
- 18 Gubernaculum protrusible. Testis single, outstretched.

19

20

- **3.2. 3. Rotylenchus buxophilus** Golden, 1956
- 21 Fig. 6; Table 5.

- Female: Habitus from close C to spiral. Lip region hemispherical, separated from
- body contour by a slight constriction, marked by 4-5 annuli. Lateral fields with four smooth
- equidistant lines, the three bands are 7.8 ± 0.6 (7.0-8.5) µm wide at mid-body, approximately
- one-fifth as wide as body diam. Regular areolation of lateral fields (external bands) observed

Comparative molecular and morphological characterisations in *Rotylenchus*

- 1 in pharyngeal region. Cuticle 1.5 µm thick, clearly annulated, annuli 1.0-1.5 µm wide at mid-
- 2 body. Body without longitudinal striations in any region. Stylet robust, 2.8-3.1 times longer
- 3 than labial region diam. Basal knobs rounded slightly laterally directed, 6.0-7.5 μm wide.
- 4 Dorsal pharyngeal gland opening 4.0-5.0 μm posterior to stylet base. Procorpus of pharynx
- 5 cylindrical, with slight depression just anterior to median bulb, 42.9 ± 1.7 (40-46) µm long.
- 6 Median pharyngeal bulb well developed, broadly oval, 15.0-16.0 × 12.0-13.0 μm, occupying
- 7 9-11 annuli, valvular apparatus 2.0-2.5 μ m long, located at 50.9 \pm 1.5 (48-53)% of pharyngeal
- 8 length. Isthmus 28.5 ± 1.9 (26-32) µm long, encircled by nerve ring at mid-point. Pharyngeal
- 9 glands forming a compact lobe, with three nuclei, overlapping intestine dorsally. Nerve ring
- enveloping isthmus at middle, at 105 ± 11.1 (98-126) µm from anterior end. Excretory pore
- usually located near pharyngo-intestinal junction level. Hemizonid distinct, located 2-3 annuli
- long, 1-3 annuli anterior to excretory pore. Reproductive system with both genital branches
- almost equally developed; anterior branch 210 ± 29.3 (171-250) µm long, posterior branch
- 14 193 ± 19.3 (158-242) µm long. Vulva clearly posterior to mid-body, without a distinct
- epiptygma. Vagina with internal walls slightly sclerotized, 13.5 (13.0-14.0) µm long. Ovaries
- with a single row of oocytes. Phasmids pore-like, 8.2 ± 2.2 (6-11) annuli anterior to level of
- anus or 10.3 ± 4.4 (6.0-16.0) µm anterior to anus. Tail dorsally convex-conoid-rounded,
- terminus pointed, striated and cuticle at tail tip 3.2 ± 0.6 (2.5-4.0) µm wide.

19

20

- **3.2. 4.** *Rotylenchus pumilus* (Perry et al., 1959) Sher, 1961
- 21 Fig. 7; Table 5.

- Female: Habitus usually forming a spiral, slightly tapering towards extremities. Lip
- region hemispherical, continuous with body contour or slightly separated by a depression,
- 25 marked by 4-5 annuli. Lateral fields with four smooth equidistant lines, the three bands are
- 8.1 ± 0.2 (8.0-8.5) µm wide at mid-body, approximately one-fourth as wide as body diam.

1 Regular areolation of lateral fields (external bands) observed in pharyngeal region. Cuticle 2 1.5-2.0 um thick, clearly annulated, annuli 1.5 um wide at mid-body. Body without 3 longitudinal striations in any region. Stylet robust, 2.8-3.2 times longer than labial region 4 diam. Basal knobs rounded slightly convex anteriorly, 6.0-6.5 µm wide. Dorsal pharyngeal 5 gland opening 4.0-5.0 µm posterior to stylet base. Procorpus of pharynx cylindrical, with 6 slight depression just anterior to median bulb, 36.8 ± 2.1 (34-40) µm long. Median pharyngeal 7 bulb well developed, broadly oval, 13.0-15.0 × 10.5-12.0 µm, occupying 8-10 annuli, valvular 8 apparatus 3.0 µm long, located at 53.9 ± 1.0 (53-56)% of pharyngeal length. Isthmus $26.3 \pm$ 9 1.0 (25-28) um long, encircled by nerve ring at mid-point. Pharvngeal glands short, with three 10 nuclei, slightly overlapping intestine dorsally for 2-4 annuli. Nerve ring enveloping isthmus at 11 middle, at 85 ± 2.9 (81-89) µm from anterior end. Excretory pore usually located near middle 12 of isthmus level. Hemizonid distinct, located at anterior to excretory pore, extending for ca 1.0-1.5 body annuli width, just anterior to excretory pore. Reproductive system with both 13 14 genital branches equally developed; anterior branch $158 \pm 13.0 \, (144-182) \, \mu m \, long$, posterior

branch 143 ± 14.9 (122-160) µm long. Vulva clearly posterior to mid-body, without a distinct

epiptygma. Vagina with internal walls slightly sclerotized, 11.8 (11.0-12.5) um long. Ovaries

with a single row of oocytes. Phasmids pore-like, 5.0 ± 1.1 (4-6) annuli anterior to level of

anus or 5.4 ± 2.2 (3.0-9.0) µm anterior to anus. Tail short, of variable shape, conoid-rounded,

20

21

15

16

17

18

19

3.2. 5. Rotylenchus robustus (de Man, 1876) Filipjev, 1936

terminus striated and cuticle at tail tip 2.6 ± 0.2 (2.5-3.0) µm wide.

Fig. 8; Table 6.

23

24

25

26

Female: Body large, habitus usually forming a spiral. Lateral fields with four smooth equidistant lines, 11.1 ± 0.8 (10.0-12.0) µm wide at mid-body, approximately one-fourth as wide as body diam. Regular areolation of lateral fields (external bands) observed in

1 pharyngeal region, an irregularly areolated along-body (Fig. 8). Cuticle 1.5-2.0 µm thick, 2 clearly annulated, annuli 1.5-2.0 µm wide at mid-body. Labial region hemispherical, set off 3 from body by a constriction, bearing 6-8 annuli, irregularly divide longitudinally, particularly 4 the 2-3 basal annuli. Labial disc distinct under SEM, rounded. Centrally located on the oral 5 disc is the oval opening of the prestoma without any labial sensillae surrounding (Fig. 8). The 6 oral disc is clearly separated from the first annulus of the lip region, which is divided into six 7 sectors, with lateral sectors, bordering the amphidial apertures, smaller than the subventral 8 and subdorsal sectors. Each amphidial opening appears as a half ellipse, i.e. a wide slit with a 9 curved distal margin between the oral disc and the lateral sectors of the first lip annulus. Labial framework well developed, $4.7 \pm 0.4 \mu m$ (4-5) long, posterior margin at level of 6^{th} or 10 8th body annulus. Stylet robust, 3.6-4.1 times longer than labial region diam. Basal knobs 11 rounded, 7.5-9.0 µm wide, at level of 22th-26th annulus posterior to labial region. Pharyngeal 12 13 glands sacciform, with three nuclei, overlapping intestine dorsally for 22-46 annuli. Excretory 14 pore usually located near pharyngo-intestinal junction level. Reproductive system with both 15 genital branches equally developed. Vulva slightly posterior to mid-body, with a distinct 16 epiptygma. Ovaries with a single row of oocytes, spermathecae rounded, with sperm (1.5-2.0 µm wide). Phasmids pore-like, 2.8 ± 1.1 (1-4) annuli anterior to level of anus or 4.5 ± 1.8 17 18 (3.0-8.0) µm anterior to anus. Tail short, hemispherical, with 10-21 annuli, terminus striated 19 and cuticle at tail tip 6.8 ± 0.4 (6.0-7.0) µm wide.

20

21

22

23

24

Male: Common, habitus an open C shape. Lip region more distinctly set off and more elevated than in female. Testis single, outstretched. Bursa crenate 76.2 ± 4.8 (68.0-82.0) µm long, completely surrounding tail which is tapering with a rounded-pointed tip. Spicules ventrally arcuate. Gubernaculum protrusible, with prominent titillae distally.

25

3.3. Molecular characterisation and phylogenetic position of Rotylenchus paravitis n. sp.

within the genus and other Hoplolaimidae

3

4

5

6

7

8

9

10

11

12

13

14

15

16

17

18

19

20

21

22

23

24

25

26

1

2

Similarity values from D2-D3 expansion segments of 28S and ITS1-rRNA among species of *Rotylenchus* are presented in Table 7. Interspecific variations for the D2-D3 sequence among Rotylenchus species retrieved from GenBank and R. paravitis n. sp. varied from 34 to 57 nucleotides (5-12%). Phylogenetic analysis (BI and ML) of Rotylenchoidinae and Hoplolaiminae based on D2-D3 expansion segments of 28S-rRNA of a multiple alignment including 82 sequences of 574 bp in length showed 5 moderately or highly supported lineages in the genus Rotylenchus (Fig. 9): (i) R. incultus, R. goodeyi, R. laurentinus, R. buxophilus and R. pumilus; (ii) R. montanus, Rotylenchus sp. SAS-2006, and R. brevicaudatus; ; (iii) R. robustus, R. uniformis, R. magnus and R. jaeni, R. cazorlaensis; and (iv) R. eximius, R. vitis and R. paravitis n. sp. (Fig. 9). Helicotylenchus species formed three highly supported clades, Scutellonema formed a single highly supported clade, and Hoplolaimus and Aorolaimus clustered together in a separate clade together with R. unisexus (Fig. 9). Rotylenchus paravitis n. sp. formed a poorly supported clade with R. eximius and R. vitis, but occupied a monophyletic position among the other genera of Hoplolaimidae (Helicotylenchus, Hoplolaimus and Scutellonema) included in the analysis (Fig. 9). D2-D3 expansion segments of 28S-rRNA did not refute the monophyly of the genera Rotylenchus (P = 0.071), Helicotylenchus (P = 0.572), Hoplolaimus (P = 1.00), even regarding the tree topology in several clades within some of the genera (Table 8). The exclusion of some species strongly associated to Helicotylenchus (viz. R. conicaudatus, HQ700698) and other not so strongly associated to *Hoplolaimus* and *Aorolaimus* (R. unisexus, EU280799) did not refute the monophyly of *Rotylenchus* (P = 0.635) (Table 8). Similar results were obtained for the analysis based on the partial 18S. However, the absence of some sequences in genus Hoplolaimus or some Rotylenchus species (viz. R. conicaudatus and R.

1 unisexus), may be the cause of these results. In both cases the constriction of the genera 2 Rotylenchus, Helicotylenchus, Hoplolaimus and Scutellonema did not refute to the monophyly 3 of these genera (P = 0.064 for D2-D3 and P = 0.298 for partial 18S). 4 ITS1-rRNA from R. paravitis n. sp. sequence differed to the aligned sequences of 5 Rotylenchus species in a range from 84 to 155 nucleotides (18-34%). Since only seven partial 6 18S-rRNA sequences from *Rotylenchus* species are deposited in GenBank, phylogenetic 7 analysis with this gene was carried out including 18S sequences from six *Helicotylenchus* 8 species. The alignment for 28 ITS1-rRNA sequences of *Rotylenchus* samples was 800 bp 9 long. After discarding ambiguously aligned regions from the alignment, the ITS1 dataset 10 included 448 bp. The 50% majority rule consensus phylogenetic tree generated from the 11 ITS1-rRNA alignment by BI analysis under the TVM + G model is presented in Fig. 10. The 12 tree topologies between ML and BI were congruent and showed a similar clustering topology 13 to that obtained for D2-D3, including 3 clades with similar species in each one. Rotylenchus 14 paravitis n. sp. which does not form supported clades with any of the other Rotylenchus 15 species (Fig. 10). On the other hand, R. robustus formed a well supported clade with R. 16 magnus and R. jaeni. Rotylenchus vitis formed a moderately supported clade with R. iranicus, 17 but not with the most similar species R. paravitis n. sp., which occupied a basal position in a 18 clade with R. jaeni, R. magnus, R. robustus, R. cazorlaensis and R. eximius (Fig. 10). 19 Similarly, the alignment for partial 18S-rRNA of 8 Rotylenchus samples and other 20 Rotylenchoidinae and Hoplolaiminae in the GenBank with 1713 positions in length showed a 21 highly supported clade with R. uniformis, R. robustus, and R. jaeni, a moderately supported 22 clade formed by R. paravitis n. sp. and R. vitis, whereas R. goodeyi, R. eximius, and 23 Rotylenchus sp. JH-2004 were clearly separated from the former, and Helicotylenchus species 24 formed two well supported clades (Fig. 11). Similarity values from the partial 18S-rRNA 25 sequence of R. paravitis n. sp. with those deposited in GenBank were high and ranged from 26 95% for Rotylenchus sp. (AY284608) to 98% for R. vitis (JN032583). Similarly, the

1 alignment for partial 18S-rRNA of 8 Rotylenchus samples and others Rotylenchoidinae and 2 Hoplolaiminae in the GenBank with 1713 positions in length showed a highly supported clade 3 with R. uniformis, R. robustus, and R. jaeni, a moderately supported clade formed by R. 4 paravitis n. sp. and R. vitis, whereas R. goodeyi, R. eximius, and Rotylenchus sp. JH-2004 5 were clearly separated from the former, and *Helicotylenchus* species formed two well 6 supported clades (Fig. 11). 7 There was not any sequence of *COI* from *Rotylenchus* species deposited in GenBank, 8 so only the partial COI sequences of this study have been used. Similarity values of COI 9 sequences of R. paravitis n. sp. with those of other Rotylenchus species ranged from 90% for 10 R. eximius (JX015401, JX015402), 87% for R. vitis (JX015417, JX015418), 82% for R. 11 cazorlaensis (JX015399, JX015400), 77% for R. robustus (JX015411- JX015414), to 76% for 12 R. magnus (JX015408-JX015410). There was no intraspecific variation among COI of 13 different populations from the same species, except for COI for R. robustus which showed 14 92% similarity (321/359 bp) between the two Spanish isolates and 84% similarity (298/354 15 bp) with USA isolate. The *COI* alignment consisted of 22 sequences with 409 bp in length. 16 The 50% majority rule consensus phylogenetic tree generated from the COI alignment by BI 17 analysis under the GTR+I+G model is presented in Fig. 11. The tree topologies between ML 18 and BI were congruent. The COI tree showed the same clade that appears in D2-D3 or ITS1 19 trees, separating R. paravitis n. sp. from R. vitis and R. eximius, and a major clade including 20 R. cazorlaensis, R. buxophilus, R. incultus, R. laurentinus, R. jaeni, R. magnus, and a 21 moderately supported sub-clade with R. robustus clearly separated by its geographical origin 22 (Spanish or American isolates) (Fig. 12). 23 The alignment generated from hsp90 sequences from R. paravitis n. sp. and R. vitis 24 showed the presence of insertions in the hsp90 sequence of R. paravitis n. sp. After discarding ambiguously aligned regions from the alignment, the size was 150 bp with a similarity of 81% 25 26 (122/150 and 5 gaps, 3%) between them. Finally, the hsp90 alignment consisted of 35

- 1 sequences with 244 bp in length. The 50% majority rule consensus phylogenetic tree
- 2 generated from the *hsp90* alignment by BI analysis under the K80 + G model is presented in
- 3 Fig. 13. The *hsp90* tree showed that *Rotylenchus* spp. clustered together with *H*.
- 4 pseudorobustus, and R. paravitis n. sp. was clearly separated from R. vitis and other
- 5 Rotylenchus spp. (Fig. 13).

3.4. Molecular diagnostics of some *Rotylenchus* species

8

7

- 9 PCR-D2-D3-28S-RFLP profile for R. paravitis n. sp. is given in Fig. 14. The
- 10 following restriction profiles are obtained for this species: unrestricted PCR 792 bp; AvaI –
- 11 395, 267, 130 bp, *Rsa*I 328, 259, 161, 38, 6 bp; *BseN*I 684, 108 bp; *Mva*I 560, 232 bp;
- 12 *Hpa*II –215, 206, 164, 145, 62 bp.
- Results of PCR with species specific primers are given in Fig. 15. The combinations of
- 14 the universal TW81 primer with the species specific R. paravitis, R. vitis or R. robustus
- primers yielded amplicons of approximately 131, 258, or 438 bp in lengths for corresponding
- species, respectively. No additional bands were observed in any tested samples.

17

18

4. Discussion

20

21

19

4.1. Multivariate analysis of morphological characters

- Results of multivariate analyses identified size, lip annuli, stylet knobs, cuticle at tail
- 23 tip, and position of phasmid, the stylet conus and orifice of dorsal gland as key characters to
- 24 differentiate R. vitis and R. paravitis n. sp. from those of R. robustus. However, no characters
- could be found to clearly discriminate between specimens of R. vitis and R. paravitis n. sp.
- since their values overlapped for the two species. Moreover, their degree of variation was
- 27 comparable to that observed among specimens belonging to each of the two populations of R.

- 1 *robustus*. Consequently, on the basis of this morphometric multivariate analysis as well as
- 2 morphological crypticism, *R. paravitis* n. sp. and *R. vitis*, should be considered cryptic
- 3 species, since both taxa are cryptic to human perception largely due to the lack of conspicuous
- 4 differences in morphometric appearance (Palomares-Rius et al., 2010).

6

- 4.2. Comparative morphology and morphometry of Rotylenchus paravitis n. sp. and
- 7 other *Rotylenchus* spp. studied

8

9 Delimiting closely related nematode species is a particularly difficult issue. For this 10 reason, Castillo & Vovlas (2005) established a specific matrix code for Rotylenchus spp. 11 according to the following: group A: 1 = lip region (l.r.) annulation absent or smooth; 2 = l.r.12 with 2-3 annuli; 3 = 1.r. with 4 annuli; 4 = 1.r. with 5 annuli; 5 = 1.r. with 6 annuli; 6 = 1.r. with 13 7-8 annuli; 7 = 1.r, with 9-10 annuli, group B: 1 = 1.r, hemispherical; 2 = 1.r, rounded; 3 = 1.r. conoid; 4 = 1.r. truncate. group C: 1 = only in pharyngeal region (ph. reg.); 2 = in ph. reg. and 14 15 irregularly at mid-body; 3 = in ph. reg. and incompletely at mid-body; 4 = in ph. reg. and near 16 phasmids; 5 = whole body length except tail region; 6 = whole body length included tail region; 7 = incompletely along whole body. group D: 1 = punctuated along body annuli; 2 = 17 18 longitudinally striated in ph. reg.; 3 = longitudinally striated over whole body; 4 = without body striations. group E: $1 = < 30 \mu m$; $2 = by 30 to 35.9 \mu m$; $3 = by 36 to 40.9 \mu m$; 4 = > 4119 20 μ m. group F: 1 = < 2 μ m; 2 = by 2 to 6.9 μ m; 3 = by 7 to 12 μ m; 4 = > 12 μ m. group G: 1 = < 21 $5 \mu \text{m}$; $2 = \text{by } 6 \text{ to } 20.9 \mu \text{m}$; $3 = \text{by } 21 \text{ to } 30.9 \mu \text{m}$; $4 = \text{by } 31 \text{ to } 40.9 \mu \text{m}$; $5 = > 41 \mu \text{m}$. group 22 H: 1 = hemispherical; 2 = rounded; 3 = conoid; 4 = pointed; 5 = with ventral projection. group I: 1 = < 50 %; 2 = by 50 to 70 %; 3 = > 70 %. group J: 1 = present; 2 = absent. group K: 1 = >23 24 5 annuli anterior to anus; 2 = from 5 anterior to 5 posterior to anus; 3 = > 5 annuli posterior to anus. Rotylenchus paravitis sp. n. is very close to R. vitis in general morphology and 25 26 morphometry, since most of the morphometric characters are within the same range, including

1 de Man ratios, DGO, O, spicules and gubernaculum. Morphologically R. paravitis n. sp. can 2 be also distinguished from the most similar species by a number of particular characteristics 3 resulting from its specific matrix code (A6,7, B4, C1, D4, E4, F2, G5, H1,2, I1,2, J1, K1 4 sensu Castillo & Vovlas, 2005). It is also close to R. cazorlaensis, Rotylenchus capitatus 5 Eroshenko, 1981, Rotylenchus elegans (Khan and Khan, 1982) Fortuner, 1987, Rotylenchus 6 fabalus Baydulova, 1984, R. iranicus Atighi, Pourjam, Pedram, Cantalapiedra-Navarrete, 7 Palomares-Rius and Castillo, 2011, R. labiodiscus Wouts & Sturhan, 1999, R. montanus 8 Vovlas, Subbotin, Troccoli, Liébanas and Castillo, 2008, and R. troncapitatus Scotto La 9 Massese and Germani, 1998. It differs from R. cazorlaensis by lip annuli (7-9 narrow annuli 10 not divided longitudinally vs 6-7 annuli, irregularly divided into blocks), first lip annulus 11 (undivided vs divided into six sectors), distance of orifice of dorsal pharyngeal gland to stylet 12 base (4-7 vs 8.5-11.5μm), stylet length (44-50 vs 46.5-56.5 μm), phasmid position (12-24 13 annuli anterior to level of anus vs 0-4 annuli posterior to anus); spicule length (33 vs 42-48 14 μm), and gubernaculum length (13 vs 19.5-25 μm). It differs from R. capitatus by body and 15 stylet length (1383-1856, 44-50 vs 680-850, 26-29 µm, respectively), lip region shape 16 (truncate lip region with 7-9 annuli, continuous with body contour vs truncate with 7-8 annuli, 17 slightly set off from body), and female tail shape (hemispherical vs conoid). It differs from R. 18 elegans by body and stylet length (1383-1856, 44-50 vs 500-600, 22-25 μm, respectively), 19 and position of phasmids (12-24 annuli anterior to level of anus vs at 5-8 annuli anterior to 20 anus). It differs from R. fabalus by lip region shape (truncate lip region with 7-9 annuli, 21 continuous with body contour vs conoid, without annulation, continuous with body contour), 22 tail tip (regularly annulated vs smooth), phasmid position (12-24 annuli anterior to level of 23 anus vs 0-2 annuli anterior to anus), and presence vs absence of males. It differs from R. 24 iranicus by body length (1383-1856 vs 954-1237 um), lip region shape (truncate lip region 25 with 7-9 annuli, continuous with body contour vs hemispherical, with 5-6, rarely 7 annuli and 26 set off from body), pharyngeal glands overlapping (53-57 vs 3-16 um long), female tail shape

1 (hemispherical with regularly annulated tip vs short, rounded, slightly conoid in some 2 specimens, and typical smooth end). It differs from R. labiodiscus by body and stylet length 3 (1383-1856, 44-50 vs 820-980, 33-37 µm, respectively), and position of phasmids (12-24) 4 annuli anterior to level of anus vs located from five annuli anterior to one annulus posterior to 5 anus level). It differs from R. montanus by lip region shape (truncate lip region with 7-9 6 annuli, continuous with body contour vs hemispherical with 6-7 annuli), body and stylet 7 length (1383-1856, 44-50 vs 913-1135, 33-37 um, respectively. It differs from R. 8 troncapitatus by body length (1383-1856 vs 940-1180 µm), lip region shape (truncate lip 9 region with 7-9 annuli, continuous with body contour vs truncate, with 7-10 annuli and set off from body), and phasmid position (12-24 annuli anterior to level of anus vs varying from four 10 11 annuli anterior to four posterior to anus). 12 The *R. brevicaudatus* population from grasses in Brisbane (Australia) was 13 morphologically and morphometrically similar with that analyzed in the original description 14 and subsequent reports, as showed by de Man ratios, except for minor differences which may 15 be considered as intraspecific (Colbran, 1962; Van den Berg and Heyns, 1974). This 16 population was characterised by a slightly shorter body and stylet length than those from the 17 original population from Lawnton, Queensland (533-587 vs 700-800 μm, 21-24 vs 22-27 μm, 18 respectively); but almost identical to a population from South Africa (21-24 vs 19-25 µm, 19 533-587 vs 500-800 µm, respectively) (Colbran, 1962; Van den Berg and Heyns, 1974). 20 Similarly, the alpha-numeric codes for *R. brevicaudatus* to be applied to the polytomic 21 identification key for *Rotylenchus* species by Castillo and Vovlas (2005) are coincident with 22 previous descriptions A3-B2-C1-D4-E1-F2-G3-H1-I2-J1-K1. 23 The R. buxophilus population from Napa County (California, USA) agrees fairly well 24 with the original description and the population from Iran, the morphometric of which do not 25 exceed the intraspecific variation reported herein, as showed by de Man ratios and other

diagnostic characters (Golden, 1956; Sher, 1965; Geraert and Barooti, 1996).

1	The <i>R. pumilus</i> population from <i>Urtica</i> sp. in San Jose park, California (USA)
2	completely fit the original description and data from a French population, except for minor
3	differences which may be considered as intraspecific (Sher, 1961; Germani and Scotto La
4	Massese, 2002). This population was characterised by a slightly larger body and stylet length
5	than those from the original population from Wisconsin, USA (773-906 \textit{vs} 600-700 μm , 27-
6	$31\ vs\ 23\text{-}28\ \mu m$, respectively) (Sher, 1961); but almost identical in stylet length to a
7	population from France (27-31 vs 26-29 μm, respectively) (Germani and Scotto La Massese,
8	2002). Similarly, the alpha-numeric codes for <i>R. pumilus</i> to be applied to the polytomic
9	identification key for Rotylenchus species by Castillo and Vovlas (2005) are coincident with
10	previous descriptions A4-B1-C1-D4-E1-F2-G2-H1-I2-J1-K2.
11	The R. robustus populations from stone pine and grasses in southern Spain and
12	California (USA), respectively, closely agree with the original description and previous
13	reports, showing minor differences which could be considered as intraspecific variations
14	(Sher, 1965; Castillo and Vovlas, 2005). Both populations were clearly differentiated from R
15	uniformis (Thorne, 1949) Loof and Oostenbrink, 1958 by a higher number of lip annuli (6-8
16	vs 5); lateral fields areolated in pharyngeal region and irregularly areolated at mid-body vs
17	areolated only in pharyngeal region; and female tail hemispherical vs rounded (Castillo and
18	Vovlas, 2005). The alpha-numeric codes for <i>R. robustus</i> to be applied to the polytomic
19	identification key for Rotylenchus species by Castillo and Vovlas (2005) are coincident with
20	previous descriptions A6-B1-C2-D4-E4-F2-G3-H1-I2-J1-K2. SEM studies for the Spanish
21	and American populations of R. robustus showed a similar pattern of lip region, with
22	longitudinal striations in lip annuli, giving a tiled surface appearance (Fig. 8), and were
23	coincident with previous studies (De Grisse et al., 1974; Abrantes et al., 1987).

25

4.3. Molecular phylogenetic relationships

1 The phylogenetic relationships inferred in this study based on the D2-D3 of 28S-2 rRNA and the ITS1 of rRNA gene sequences mostly agrees with well-differentiated lineages 3 obtained in previous studies (Vovlas et al., 2008; Athigi et al., 2011; Cantalapiedra-Navarrete 4 et al., 2012). Nevertheless, small differences may be attributed to additional sequences added 5 in this study. In particular, the position of R. agnetis in the present tree differed after including 6 R. brevicaudatus, which formed a moderately supported clade with R. montanus and with 7 Rotylenchus sp. SAS-2006. Phylogeny based on the D2-D3 of 28S-rRNA clearly showed the 8 paraphyly of Hoplolaimidae, which agrees with previous studies (Subbotin et al., 2007; 9 Vovlas et al., 2008; Atighi et al., 2011; Cantalapiedra-Navarrete et al., 2012). This molecular 10 marker separated Rotylenchus from other genera within Hoplolaimidae (i.e. Helicotylenchus, 11 Hoplolaimus, Scutellonema and Aorolaimus), which agree with the separation by 12 morphological characters, including position of pharyngeal overlapping (dorsal vs ventral), 13 type of phasmid (scutellum-type or pore-like) and stylet knobs (tulip-shaped or rounded). 14 Also, in *Rotylenchus* some lineages derived from molecular markers were congruent with 15 morphological and morphometrical traits for D2-D3 with the major number of species 16 studied, i.e., lineage (iii)included species with hemispherical lip regions of 7-8 annuli, long 17 bodies and stylets, but the first one possesses three pharyngeal gland nuclei; and lineage (iv) 18 grouped two closely related species (R. vitis and R. paravitis n. sp.) and R. eximius sharing 19 only a long stylet and a broadly rounded tail with those species. The ITS1 data set clearly 20 indicated similar relationships and lineages composition with previous studies (Vovlas et al., 21 2008; Atighi et al., 2011; Cantalapiedra-Navarrete et al., 2012), except for some species, such 22 as R. vitis which formed a moderately supported clade with R. iranicus, instead of clustering 23 with R. paravitis and R. eximius which clustered separately or with very low clade support 24 (Figs. 9, 10). However, these lineages differ between D2D3 and ITS.

The phylogenetic relationships inferred in this study based on the partial *COI* data set was provided for the first time in the genus *Rotylenchus*. Phylogenetic relationships based on

25

- 1 the partial *COI* showed similar phylogenies than those detected in D2-D3 and ITS1 regions 2 for the majority of species. Finally, phylogeny of the available partial 18S sequences suggests 3 a monophyletic relationship within the genus Rotylenchus, but additional studies are needed 4 for clarifying this aspect. Molecular markers based on D2-D3 expansion segments of 28S-5 rRNA and partial 18S did not refute the monophyly of *Rotylenchus* and other genera in the 6 family Hoplolaimidae (*Helicotylenchus* in both markers and *Hoplolaimus* only in D2-D3). 7 However, additional sequences of *Hoplolaimus*, *Scutellonema* and *Aorolaimus* will help in the 8 study of these genera from a phylogenetic point of view. Our results showed that D2-D3 of 9 28S-rRNA, the ITS1 of rRNA, the partial COI, the partial 18S, and partial hsp90 gen are 10 useful markers for species delimitation. Nevertheless, if the speciation event has taken place 11 recently, the D2-D3 of 28S-rRNA might not be served a good marker to separate Rotylenchus 12 spp., as it has been shown for R. laurentinus and R. incultus (Vovlas et al., 2008). 13 Consequently, phylogenetic analyses based on the different markers did not result in a general 14 consensus of species grouping, since lineages were maintained for some species (i.e. species 15 with hemispherical lip regions of 7-8 annuli, long bodies and stylets), but not in others (i. e. 16 position of R. vitis, R. breviannulatus were quite variable). Diagnostic PCR-ITS-RFLP profiles with five restriction enzymes, as well as species-17 18
 - Diagnostic PCR-ITS-RFLP profiles with five restriction enzymes, as well as species-specific primer proved to be useful tools for identification of *Rotylenchus* species.

 Nevertheless, specificity of species specificity of *R. robustus* primers still requires testing with *R. uniformis* samples, and species identity for nematodes previously named as *R. uniformis* by Subbotin et al. (2007) should be confirmed by molecularly comparing with the type materials of this species. There is some controversy on the synonymy of *R. uniformis* with *R. robustus*, since some authors consider both as valid species (Seinhorst and Kuniyasu, 1969, Castillo and Vovlas, 2005) based on differences in six characters including body length, ratio a, dimensions and number of lips annuli, stylet length, and posterior extensions of labial

20

21

22

23

24

1	framework, while other authors consider both species as synonyms (Loof and Oostenbrink,
2	1958, Seinhorst, 1991).
3	In summary, molecular characterisation and phylogenetic analysis of D2-D3 region
4	ITS1 of rRNA, partial 18S-rRNA, COI, and hsp90 sequences and morphological and
5	morphometric analyses clearly support the proposal of R. paravitis n. sp. as a new species.
6	And also, the recognition of this cryptic species within the genus <i>Rotylenchus</i> shows that the
7	biodiversity of these nematodes is still not fully understood and need some additional studies.
8	Interestingly, R. paravitis n. sp. and R. vitis showed clearly different sequences and positions
9	in the phylogenetic analysis, in spite of showing scarce or no differences in morphology or
10	morphometry. Also, RFLP profiles as well as slight minor phenotypic traits suggest speciation
11	between the two taxa. Understanding the range of R. vitis and R. paravitis n. sp. by new
12	locality reports could help to understand the speciation process in these nematodes.
13	
14	
15	Acknowledgements
16	
17	This research was supported by grant AGL2009-06955 from 'Ministerio de Ciencia e
18	Innovación' of Spain, grant AGR-136 from 'Consejería de Economía, Innvovación y Ciencia'
19	from Junta de Andalucía, and the European Social Fund. The authors thank J. Martín-
20	Barbarroja, and G. León-Ropero from IAS-CSIC for excellent technical assistance, M.
21	Pedram (Tarbiat Modares University, Tehran, Iran) for his help with line drawing, and B.
22	Landa (IAS-CSIC) for critical reading of the manuscript.
23	
24	References

- 1 Abolafia, J., Liebanas, G., Peña-Santiago, R., 2002. Nematodes of the order Rhabditida from
- 2 Andalucía Oriental, Spain. The subgenus *Pseudacrobeles* Steiner, 1938, with description
- of a new species. J. Nematode Morph. Syst. 4, 137-154.
- 4 Abrantes, I. M. d. O., Vovlas, N., Santos, M.S.N., 1987. Morphological studies on six
- 5 tylenchid nematode species associated with olive in Portugal. Ciencia Biol. Ecol. Syst. 7,
- 6 1-9.
- 7 Andrássy, I., 1958. Hoplolaimus tylenchiformis Daday, 1905 (Syn. H. coronatus Cobb, 1923)
- 8 und die Gattungen der Unterfamilie Hoplolaiminae Filipiev, 1936. Nematologica 3, 44-
- 9 56.
- 10 Atighi, M.R., Pourjam, E., Pedram, M., Cantalapiedra-Navarrete, C., Palomares-Rius, J.E.,
- 11 Castillo, P., 2011. Molecular and morphological characterisations of two new
- 12 Rotylenchus species from Iran. Nematology 13, 951-964.
- Baydulova, L.A., 1984. [A new species from the genus *Rotylenchus* (Nematoda:
- Hoplolaimidae) from the northern pre-Caspian region.] Byulletin' Vsesoyuznogo
- 15 Instituta Gel' mintologii im. K. l. Skryabina 36, 67-68.
- 16 Boutsika, K., Brown, D.J.F., Phillips, M.S., Blok, V.C., 2004. Molecular characterisation of
- the ribosomal DNA of *Paratrichodorus macrostylus*, *P. pachydermus*, *Trichodorus*
- primitivus and T. similis (Nematoda: Trichodoridae). Nematology 6, 641-654.
- 19 Cantalapiedra-Navarrete, C., Liébanas, G., Archidona-Yuste, A., Palomares-Rius, J.E.,
- Castillo, P., 2012. Molecular and morphological characterisation of *Rotylenchus vitis* n.
- sp. (Nematoda: Hoplolaimidae) infecting grapevine in southern Spain. Nematology 14,
- 22 235-247.
- 23 Castillo, P., Gómez Barcina, A., 1987. *Rotylenchus cazorlaensis* sp. n. and new record of *R*.
- fallorobustus Sher, 1965 (Nematoda: Tylenchida) from south-eastern Spain.
- 25 Nematologica 33, 393-400.

- 1 Castillo, P., Vovlas, N., 2005. *Bionomics and identification of the genus* Rotylenchus
- 2 (Nematoda: Hoplolaimidae). Nematology Monographs and Perspectives, vol. 3 (series
- 3 editors: Hunt, D.J. and Perry, R.N.). Leiden, The Netherlands, Brill Academic
- 4 Publishers, 377 pp.
- 5 Castillo, P., Vovlas, N., Subbotin, S., Troccoli, A., 2003. A new root-knot nematode,
- 6 *Meloidogyne baetica* n. sp. (Nematoda: Heteroderidae), parasitizing wild olive in
- 7 Southern Spain. Phytopathology 93, 1093-1102.
- 8 Colbran, R.C., 1962. Studies of plant and soil nematodes. 5. Four new species of
- 9 Tylenchoidea from Queensland Pineapple Fields. Queens. J. Agric. Sci. 19, 231-239.
- 10 Coolen, W.A., 1979. Methods for extraction of *Meloidogyne* spp. and other nematodes from
- roots and soil. In: Lamberti, F., Taylor, C.E. (Eds), Root-knot nematodes (Meloidogyne
- species). Systematics, biology and control. Academic Press, New York, pp. 317-329.
- Curran, J., Driver, F., Ballard, J.W.O., Milner, R.J., 1994. Phylogeny of *Metarhizium*:
- analysis of ribosomal DNA sequence data. Mycol. Res. 98, 547-555.
- Daday, E. von, 1905. Untersuchungen über die Süsswasser-Mikrofauna Paraguays. Miteinem
- Anhang von W. Michaelsen. Zoologica, Stuttg. 18, 1-349 and 362-374.
- 17 De Grisse, A.T., 1969. Redescription ou modifications de guelques techniques utilisées dans
- 18 l'étude des nématodes phytoparasitaires. Meded. Fac. Landbouw. Rijksuniv. Gent 34,
- 19 351-369.
- 20 De Grisse, A.T., Lippens, P.L., Coomans, A., 1974. The cephalic sensory system of
- 21 Rotylenchus robustus and a comparison with some other Tylenchids. Nematologica 20,
- 22 88-95.
- Derycke, S., Vanaverbeke, J., Rigaux, A., Backeljau, T., Moens, T., 2010. Exploring the use
- of cytochrome oxidase c subunit 1 (*COI*) for DNA barcoding of free-living marine
- 25 nematodes. PLoS ONE 5(10), e13716. doi:10.1371/journal.pone.0013716.

- 1 Eroshenko, A.S., 1981. Phytopathogenic nematodes of forest undergrowth of the families
- 2 Tylenchorhynchidae and Hoplolaimidae (Nematoda). In: Eroshenko, A.C., Belogurov,
- 3 O.I. (Eds). Freeliving and plant-parasitic nematodes in the Far-East. Vladivostok.
- 4 Dal'nev. Nauch. Tsentr Akad. Nauk SSR, pp. 22-27 and 85-92.
- 5 Filipjev, I.N., 1934. The classification of free-living nematodes and their relations to parasitic
- 6 nematodes. Smithson. Misc. Collect. 89, 1-63.
- 7 Filipjev, I.N., 1936. On the classification of the Tylenchinae. Proc. Helminth. Soc. Wash. 3,
- 8 80-82.
- 9 Fortuner, R., 1987. A reappraisal of Tylenchina (Nemata). 8. The family Hoplolaimidae
- 10 Filip'jev, 1934. Rev. Nématol. 10, 219-232.
- 11 Geraert, E., Barooti, S., 1996. Four *Rotylenchus* from Iran, with a key to the species.
- 12 Nematologica 42, 503-520.
- 13 Germani, G., Scotto La Massese, C., 2002. Description of four new species and two
- populations of *Rotylenchus* (Nematoda: Hoplolaimidae). Nematol. Mediterr. 30, 203-208.
- 15 Golden, A.M., 1956. Taxonomy of the spiral nematodes (*Rotylenchus* and *Helicotylenchus*),
- and the developmental stages and host-parasite relationships of R. buxophilus, n. sp.,
- attacking boxwood. Bulletin A-85, University of Maryland, MD, USA, Agricultural
- Experimental Station, 28 pp.
- 19 Gutiérrez-Gutiérrez, C., Palomares-Rius, J.E., Cantalapiedra-Navarrete, C., Landa, B.B.,
- Esmenjaud, D., Castillo, P., 2010. Molecular analysis and comparative morphology to
- resolve a complex of cryptic *Xiphinema* species. Zool. Scr. 39, 483-498.
- Gutiérrez-Gutiérrez, C., Castillo, P., Cantalapiedra-Navarrete, C., Landa, B.B., Derycke, S.,
- Palomares-Rius, J.E., 2011. Genetic structure of *Xiphinema pachtaicum* and *X. index*
- populations based on mitochondrial DNA variation. Phytopathology 101, 1168-1175.
- 25 Hall, T.A., 1999. BioEdit: a user-friendly biological sequence alignment editor and analysis
- program for windows 95/98/NT. Nucl. Acids Symp. Ser. 41, 95-98.

- 1 Huelsenbeck, J.P., Ronquist, F., 2001. MrBAYES: Bayesian inference of phylogenetic trees.
- 2 Bioinformatics 17, 754-755.
- 3 Hugall, A., Moritz, C., Stanton, J., Wolstenholme, D.R., 1994. Low, but strongly structured
- 4 mitochondrial DNA diversity in root knot nematodes (*Meloidogyne*). Genetics 136, 903-
- 5 912.
- 6 Khan, M.L., Khan, S.H., 1982. Three new species of genus *Orientylus* Jairajpuri & Siddiqi,
- 7 1977 associated with fruit trees in India (Rotylenchoidinae: Nematoda). Ind. J. Nematol.
- 8 12, 111-117.
- 9 Linford, M.B., Oliveira, J.M., 1940. Rotylenchulus reniformis, nov. gen., n. sp., a nematode
- parasite of roots. Proc. Helminth. Soc. Wash. 7, 35-42.
- 11 Loof, P.A.A., Oostenbrink, M., 1958. Die identitat van *Tylenchus robustus* de Man.
- Nematologica 3, 34-43.
- 13 Madani, M., Ward, L., De Boer, S., 2011. *Hsp90* gene, an additional target for discrimination
- between the potato cyst nematodes, *Globodera rostochiensis*, *G. pallida*, and the related
- species G. tabacum tabacum. Eur. J. Plant Pathol. 130, 271-285.
- 16 Man, J.G. de, 1876. Onderzoekingen over vrij in de aarde levende Nematoden. Tijdschr.
- 17 Nederl. Dierk. Ver. 2, 78-196.
- Page, R.D.M., 1996. TREEVIEW: an application to display phylogenetic trees on personal
- 19 computers. Comp. Appl. Biosc. 12, 357-358.
- Palomares-Rius, J.E., Castillo, P., Liébanas, G., Vovlas, N., Landa, B.B., Navas, -Cortés, J.A.,
- Subbotin, S.A. 2010. Description of *Pratylenchus hispaniensis* n. sp. from Spain and
- considerations on the phylogenetic relationship among selected genera in the family
- Pratylenchidae. Nematology 12, 429-451.
- Perry, V.G., Darling, H.M., Thorne, G., 1959. Anatomy, taxonomy, and control of certain
- spiral nematodes attacking blue grass in Wisconsin. Univ. Wisconsin Res. Bull. 207, 1-
- 26 24.

- 1 Posada, D., 2008. jModelTest: Phylogenetic model averaging. Mol. Biol. Evol. 25, 1253-
- 2 1256.
- 3 Scognamiglio, A., Talamé, M., 1972. Rotylenchus laurentinus n. sp. (Nematoda:
- 4 Hoplolaimidae). Boll. Lab. Entomol. Agraria Filippo Silvestri 30, 1-7.
- 5 Scotto La Massèse, C., Germani, G., 1998. Description of two new species of *Rotylenchus*
- 6 and two populations of *R. agnetis*. Nematologica 44, 37-44.
- 7 Seinhorst, J.W. 1991. The identity of *Tylenchus robustus* de Man, 1876, with a remark on
- 8 distortions in de Man's (1876) figures. Nematologica 37, 119-122.
- 9 Seinhorst, J.W., Kuniyasu, K., 1969. *Rotylenchus uniformis* (Thorne) on carrots. Neth. J.
- 10 Plant Pathol. 75, 205-223.
- 11 Sher, S.A., 1961. Revision of the Hoplolaiminae (Nematoda). I. Classification of nominal
- genera and nominal species. Nematologica 6, 155-169.
- 13 Sher, S.A., 1965. Revision of the Hoplolaiminae (Nematoda). V. *Rotylenchus* Filipjev, 1936.
- 14 Nematologica 6, 173-198.
- 15 Siddiqi, M.R. (2000). Tylenchida parasites of plants and insects. 2nd edition. Wallingford,
- 16 UK, CABI Publishing, 833 pp.
- 17 Skantar, A.M., Carta, L.K., 2004. Molecular characterization and phylogenetic evaluation of
- the *hsp90* gene from selected nematodes. J. Nematol. 36, 466-480.
- 19 Steiner, G., 1945. *Helicotylenchus*, a new genus of plant-parasitic nematodes and its
- relationship to *Rotylenchus* Filipjev. Proc. Helminth. Soc. Wash. 12, 34–38.
- Subbotin, S.A., Sturhan, D., Chizhov, V.N., Vovlas, N., Baldwin, J.G., 2006. Phylogenetic
- analysis of Tylenchida Thorne, 1949 as inferred from D2 and D3 expansion fragments
- of the 28S rRNA gene sequences. Nematology 8, 455-474.
- Subbotin, S.A., Sturhan, D., Rumpenhorst, H.J, Moens, M., 2003. Molecular and
- 25 morphological characterisation of the *Heterodera avenae* complex species (Tylenchida:
- Heteroderidae). Nematology 5, 515-538.

- 1 Subbotin, S.A., Moens, M., 2006. *Molecular Taxonomy and Phylogeny*. In R. Perry & M.
- 2 Moens (Eds) *Plant Nematology*. Wallingford, UK, CABI, pp. 33-58.
- 3 Subbotin, S. A., Sturhan, D., Vovlas, N., Castillo, P., Tanyi Tambe, J., Moens, M., Baldwin,
- 4 J.G., 2007. Application of the secondary structure model of rRNA for phylogeny: D2-
- 5 D3 expansion segments of the LSU gene of plant-parasitic nematodes from the family
- 6 Hoplolaimidae Filipjev, 1934. Mol. Phylogenet. Evol. 43, 881-890.
- 7 Swofford, D.L., 2003. PAUP*: Phylogenetic analysis using parsimony (*and other methods),
- 8 version 4.0b 10. Sinauer Associates, Sunderland, Massachusetts.
- 9 Tanha Maafi, Z., Subbotin, S.A., Moens, M., 2003. Molecular identification of cyst-forming
- nematodes (Heteroderidae) from Iran and a phylogeny based on ITS-rDNA sequences.
- 11 Nematology 5, 99-111.
- 12 Thompson, J.D., Gibson, T.J., Plewniak, F., Jeanmougin, F., Higgins, D.G., 1997. The
- 13 CLUSTAL X windows interface: flexible strategies for multiple sequence alignment
- aided by quality analysis tools. Nucl. Acids Res. 25, 4876-4882.
- 15 Thorne, G. 1949. On the classification of the *Tylenchida*, new order (Nematoda, Phasmidia).
- 16 Proc. Helminth. Soc. Wash. 16, 37-73.
- 17 Van Den Berg, E., Heyns, J., 1974. South African Hoplolaiminae. 3. The genus *Rotylenchus*
- Filipjev, 1936. Phytophylactica 6, 165-184.
- 19 Vovlas, N., Subbotin, S. A., Troccoli, A., Liébanas, G., Castillo, P., 2008. Description of
- 20 Rotylenchus montanus sp. n. and recognition of R. jaeni comb. n. as a separate species
- with approaches to molecular phylogeny of the genus *Rotylenchus* (Nematoda,
- 22 Tylenchida). Zool. Scr. 37, 521-537.
- Wouts, W. M., Sturhan, D., 1999. Descriptions of *Rotylenchus* species from New Zealand,
- including R. labiodiscus sp. n. (Nematoda: Hoplolaimidae. New Zeal. J. Zool. 26, 395-
- **25** 404.

- 1 Zancada, M.C., 1985. Rotylenchus magnus sp. n. and R. mesorobustus sp. n. (Nematoda:
- Tylenchida) from Spain. Nematologica 31, 134-142.

Comparative molecular and morphological characterisations in Rotylenchus

Table 1. *Rotylenchus* species and populations used and sequenced in the present study*.

Species	Locality	Host	D2-D3	ITS1/ITS1- 5.8S-ITS2	18S	COI
R. agnetis	Potenza, Italy	Ruscus aculeatus (Butcher's broom)	EU280795	-	-	-
R. brevicaudatus	Brisbane, Australia	Grasses	JX015419 JX015420	JX015430 JX015431	-	-
R. brevicaudatus	Republic of China	-	-	-	DQ309587	-
R. buxophilus	Napa County, California, USA	Unknown	JX015421	JX015432 JX015433	-	JX015398
R. buxophilus	Arkansas, USA	-	FJ485646, FJ485647	-	-	-
R. cazorlaensis	Cazorla, Spain	Quercus faginea (Portuguese oak)	EU280793	EU373668, EU373669	EU373668	JX015399 JX015400
R. cazorlaensis	Cazorla, Spain	Quercus rotundifolia (oak)	EU280792	EU373670, EU373671	-	-
R. conicaudatus	Mazandaran, Iran	Grasses	HQ700698	HQ700700	_	_
R. eximius	Brindisi, Italy	Pistacia lentiscus (lentisc)	EU280794	EU373663	-	JX015401 JX015402
R. eximius**	Huelva, Spain	Olea europaea subsp. silvestris (wild olive)	DQ328741	EU373664	JX015427	
R. goodeyi	Vejer, Spain	Olea europaea subsp. silvestris (wild olive)	DQ328756	-	-	-
R. incultus	Niebla, Spain	Vitis vinifera (grapevine)	EU280796	EU373673	-	JX015403
R. incultus	Bollullos, Spain	Vitis vinifera (grapevine)	EU280797	EU373672	-	-
R. iranicus	Mazandaran, Iran	Fagus orientalis (beech tree)	HQ700698	HQ700699	-	-
R. jaeni**	Santa Elena, Spain	Quercus suber (cork tree)	EU280791	EU373661, EU373662	JX015428	JX015404
R. laurentinus	Torre Canne, Italy	Pistacia lentiscus (lentisc)	DQ328757	-	-	JX015405 JX015406 JX015407
R. laurentinus	Zahara, Spain	Pistacia lentiscus (lentisc)	EU280798	EU373666	EU373667	-
R. magnus	Arévalo, Spain	Ilex aquifolium (holly)	EU280789	EU373660, EU373676	-	JX015408 JX015409 JX015410
R. magnus	Lubia, Spain	Quercus robur (pedunculate oak)	EU280790	EU373659, EU373665		-
R. montanus	Trentino, Italy	Malus domestica (apple)	DQ328743	EU280800, EU280801	-	-
R. paravitis n. sp.**	Jerez, Spain	Helianthus annuus (sunflower)	JX015422	JX015434	JX015429	JX015415 JX015416
R. pumilus	Santa Clara County, California, USA	Urtica sp.	JX015423	JX015435 JX015436	-	-
R. robustus** R. robustus	Lucena del Puerto, Spain Bonares, Spain	Pinus pinea (stone pine) Pinus pinea (stone pine)	JX015424	JX015437	-	JX015413 JX015414
R. robustus	Marin County, California, USA	Grasses	JX015425	JX015438	-	-
R. robustus	Tomales. California, USA	Grasses	JX015426	JX015439 JX015440	-	JX015411 JX015412
R. robustus	Michigan, USA	Unknown	EU280788	-	-	-
R. uniformis	Bruges, Belgium	Unknown	DQ328735, DQ328736	-	-	-
R. uniformis	Ghent, Belgium	Unknown	DQ328738	-	-	-
R. uniformis	Poppel, Belgium	Unknown	DQ328739, DQ328740	-	-	-
R. uniformis	Elst, the Netherlands	Unknown	DQ328737	-	-	-
R. unisexus	Seville, Spain	Citrus aurantium (citrus)	EU280799	EU373674, EU373675	-	-
R. vitis**	Montemayor, Spain	Vitis vinifera (grapevine)	JN032581	JN032582	JN032583	JX015417 JX015418
Rotylenchus sp.	Moscow, Russia	Unknown	DQ328742	EU280802		

^{4 *}Newly sequenced samples are indicated by bold font.

^{5 **} hsp90 gene sequence= **JX015395** for *R. eximius*, **JX015396** for *R. jaeni*, **JX015393** for *R.*

⁶ paravitis n. sp., JX015397 for R. robustus (Lucena del Puerto), and JX015394 for R. vitis.

1 **Table 2.** Eigenvector and eigenvalues of principal components derived from nematode

2 morphometric characters for *Rotylenchus paravitis* n. sp., *R. vitis* and two populations of *R*.

3 robustus^a.

	Principal component						
Character ^b	PC1	PC2	PC3	PC4			
Body length (L)	<u>-0.788</u>	-0.154	-0.077	0.467			
Lip width	-0.098	-0.256	0.448	-0.003			
Lip height	0.237	-0.113	0.088	0.217			
Number of lip annuli	<u>-0.660</u>	-0.234	-0.034	0.209			
Stylet length	0.307	-0.406	0.390	0.120			
Stylet-conus length	0.676	-0.202	0.178	0.096			
Knobs width	<u>-0.670</u>	-0.030	0.284	-0.214			
Dorsal gland orifice (D.G.O.)	0.727	-0.462	-0.223	0.115			
O	<u>0.700</u>	-0.395	-0.328	0.099			
Anterior end to beginning of median pharyngeal bulb distance	-0.079	<u>-0.779</u>	0.066	0.285			
Anterior end to centre of median pharyngeal bulb distance	-0.062	<u>-0.837</u>	0.037	0.377			
Total pharyngeal length	-0.332	<u>-0.727</u>	0.246	0.029			
Anterior end to pharyngo-intestinal junction distance	-0.231	<u>-0.634</u>	0.232	0.161			
Excretory pore to anterior end (EP)	0.309	-0.592	0.240	0.435			
Maximum body width	<u>-0.745</u>	-0.342	0.249	0.059			
Pharyngeal overlapping	-0.593	-0.344	-0.038	-0.133			
Cuticle tail tip width	<u>-0.755</u>	0.042	0.251	-0.071			
Vulva position	0.538	-0.195	-0.003	-0.323			
Anterior gonad	-0.561	-0.375	-0.232	-0.567			
Posterior gonad	-0.571	-0.382	-0.243	-0.527			
G_1	-0.190	-0.315	-0.195	<u>-0.846</u>			
G_2	-0.194	-0.334	-0.214	<u>-0.827</u>			
Female tail length	0.002	-0.361	<u>-0.831</u>	0.271			
Anal body width	-0.592	-0.441	0.113	0.208			
Number of female tail annuli	-0.478	-0.256	-0.497	-0.020			
Phasmid to terminus distance	<u>-0.839</u>	0.094	-0.322	0.121			
Phasmid to anus distance	<u>-0.827</u>	0.182	-0.214	0.097			
Number of annuli between phasmid and anus	<u>-0.909</u>	0.190	-0.124	0.045			
a	-0.121	0.197	-0.379	0.525			
b	<u>-0.777</u>	0.298	-0.105	0.274			
b'	-0.572	0.309	-0.258	0.456			
c	-0.485	0.207	0.745	0.044			

Comparative molecular and morphological characterisations in Rotylenchus

c'	0.332	-0.134	<u>-0.864</u>	0.156
Eigenvalues	9.973	4.823	3.776	3.699
% of total variance	30.22	14.61	11.44	11.21
Cumulative % of total variance	30.22	44.83	56.28	67.49

1

^a Based on 21 female specimens of *Rotylenchus vitis* and *R. paravitis* n. sp., respectively, and

^{3 12} and nine female specimens of two populations of R. robustus from Spain and USA,

⁴ respectively. Values of morphometric and morphological characters dominating Principal

⁵ components 1 to 4 (eigenvector >0.63) are underlined.

^{6 &}lt;sup>b</sup> Morphological and diagnostic characters according to Castillo and Vovlas (2005).

Table 3. Primers sets used in the present.

Primer code	Sequences 5'-3'	Amplified gene	References		
TW81	GTTTCCGTAGGTGAACCTGC	ITS-rRNA	Curran et al. (1994)		
AB28	ATATGCTTAAGTTCAGCGGGT	110-11111	Curran et al. (1994)		
D2A	ACAAGTACCGTGAGGGAAAG	D2-D3 of 28S	Subbotin et al. (2006)		
D3B	TCGGAAGGAACCAGCTACTA	rRNA	Subbotin et al. (2000)		
TW81	GTTTCCGTAGGTGAACCTGC				
5.8SM5	GGCGCAATGTGCATTCGA	ITS1-rRNA	Vovlas et al. (2008)		
A	AAAGATTAAGCCATGCATG				
13R	GGGCATCACAGACCTGTTA	18S	Boutsika et al. (2004)		
18P-SSU_R_81	TGATCCWKCYGCAGGTTCAC				
JB3	TTTTTTGGGCATCCTGAGGTT	COI	Derycke et al. (2010)		
JB4	TAAAGAAAGAACATAATGAA	COI	Derycke et al. (2010)		
U831	AAYAARACMAAGCCNTYTGG	hsp90	Skantar and Carta		
L1110	TCRCARTTVTCCATGATRAAV	пзрэо	(2004)		
TW81	GTTTCCGTAGGTGAACCTGC	ITS-rRNA	This study		
R_paravitis	GCTCCATCACGCAGCAGAC	115-IKIVA	This study		
TW81	GTTTCCGTAGGTGAACCTGC	ITS-rRNA	This study		
R_vitis	CTTACGTGTGTGCCAAATAGT	II D-IIIIIA	Tills study		
TW81	GTTTCCGTAGGTGAACCTGC	ITS-rRNA	This study		
R_robustus	GACGTGGACATCATACAGTC	II S-IKIVA	Tills study		

4 5

Table 4. Morphometrics of *Rotylenchus paravitis* n. sp. from the rhizosphere of sunflower (*Helianthus annuus* L.) from Jerez de la Frontera, Cádiz province, southern Spain. All measurements are in μ m and in the form: mean \pm s.d. (range)^a.

	F	Female	Male
Character	Holotype	Paratypes	Paratype
n	_	20	1
L	1644	1599 ± 144.1	1167
a	35.0	(1383-1856) 32.1 ± 1.7	37.6
b	14.7	(29.3-34.6) 15.3 ± 0.9 (13.6-17.0)	17.2
b'	7.7	7.6 ± 0.5	8.7
c	67.1	(6.7-8.5) 79.7 ± 15.2	55.6
c'	0.7	(55.3-104.9) 0.6 ± 0.1	1.2
V or T	52.0	(0.4-0.8) 51.2 ± 2.7	42
G_1	18	(43-54) 17.0 ± 5.0	_
G_2	19	(12.6-29.8) 16.5 ± 4.2	_
Stylet	47.5	$(12.1-26.7)$ 46.5 ± 1.7	33.5
Stylet conus	21.5	(44.0-50.0) 21.1 ± 1.0	15.5
DGO	4.0	(19.0-22.5) 4.4 ± 0.8	4.0
O	8.4	(4.0-7.0) 9.5 ± 1.6	11.9
Anterior end to centre of median	112.0	(8.2-14.6) 106 ± 6.0	68
hulh Anterior end to excretory pore	151.0	$(97-116)$ 148 ± 11.3	95
Pharynx length	213.0	(136-166) 210 ± 14.6	134
Pharyngeal overlap	60.0	(195-244) 57.7 ± 4.5	47
Max. body diam.	47.0	(53-67) 50.5 ± 4.7	31
Anal body diam.	34.0	(44-61) 36.3 ± 3.5	18
Tail	24.5	(30.0-43.0) 20.8 ± 3.5	21
Tail annuli	14.0	(16.0-28.0) 15.4 ± 1.7	_
Phasmid to terminus	58.0	$(12-18)$ 46.7 ± 4.6	_
Spicules	_	(40.0-54.0) -	33
Gubernaculum	_	_	13

^a Abbreviations are defined in Siddiqi (2000)

Table 5. Morphometrics of *Rotylenchus brevicaudatus* Colbran, 1962 from grasses (Brisbane, Australia), *Rotylenchus buxophilus* Golden, 1956 from Napa Valley (California, USA), and *Rotylenchus pumilus* (Perry et al., 1959) Sher, 1961 from *Urtica* sp.San Jose park, California, USA. All measurements are in μm and in the form: mean ± s.d. (range) ^a.

	R. brevic	audatus	R. buxophilus	R. pumilus		
Character	Females	Males	Females	Females		
n	4	6	10	6		
L	571.8 ± 56.0 (502-618)	565.8 ± 20.1 (533-587)	948 ± 95.0 (829-1142)	825 ± 50.4 (773-906)		
a	23.9 ± 1.8 (21.8-25.7)	23.8 ± 0.7 (23.1-24.9)	30.8 ± 2.6 (25.4-28.9)	26.9 ± 1.5 (25.4-28.9)		
b	8.9 ± 0.6 (8.2-9.5)	8.8 ± 0.1 (8.5-8.9)	7.3 ± 0.8 (6.4-8.9)	11.1 ± 0.7 $(10.3-12.2)$		
b'	5.3 ± 0.0 (5.3-5.3)	4.8 ± 0.2 (4.5-5.0)	6.1 ± 0.6 (5.4-7.3)	6.5 ± 0.4 (5.9-7.0)		
c	54.8 ± 11.0 (41.8-68.7)	34.9 ± 2.0 (32.6-38.1)	37.5 ± 2.9 (33.2-42.0)	46.6 ± 2.9 (42.9-50.2)		
c'	0.6 ± 0.1 (0.5-0.7)	1.1 ± 0.01 (1.1-1.2)	1.2 ± 0.1 (1.1-1.5)	0.9 ± 0.1 (0.8-1.0)		
V or T	56.5 ± 1.3 (55.0-58.0)	46.2 ± 2.9 (42.3-49.7)	55.4 ± 1.8 (54.0-57.0)	56.7 ± 1.8 (54.0-59.0)		
G_1	16.0 ± 3.1 (11.7-18.5)	-	21.5 ± 0.7 $(15.0-25.9)$	19.1 ± 0.7 (18.3-20.1)		
G_2	15.0 ± 3.0 (10.8-18.0)	-	19.6 ± 2.7 $(16.1-24.0)$	17.3 ± 1.5 (14.6-18.8)		
Stylet	22.4 ± 1.3 (21.0-24.0)	20.8 ± 0.8 (20.0-22.0)	35.5 ± 1.2 (33.0-37.0)	29.3 ± 1.4 (27.0-31.0)		
Stylet conus	11.1 ± 0.5 (10.5-11.5)	10.6 ± 0.5 $(10.0-11.0)$	17.6 ± 0.7 $(16.5-18.5)$	13.3 ± 0.8 $(12.0-14.0)$		
DGO	5.3 ± 0.3 (5.0-5.5)	5.3 ± 0.4 (5.0-6.0)	4.2 ± 0.3 (4.0-5.0)	4.3 ± 0.5 (4.0-5.0)		
O	23.9 ± 0.8 (22.7-24.4)	25.6 ± 1.9 (22.7-28.6)	11.8 ± 0.9 $(11.0-13.9)$	14.8 ± 1.6 (12.9-16.7)		
Anterior end to centre of median bulb	64 ± 2.5 (61-67)	64 ± 2.6 (60-67)	85 ± 2.1 (82-88)	74 ± 1.8 (72-77)		
Anterior end to excretory pore	89 ± 3.6 (84-92)	83 ± 3.9 (77-87)	127 ± 2.5 (124-132)	105 ± 9.5 (97-117)		
Pharynx length	108 ± 10.9 (95-117)	118 ± 3.8 (114-123)	156 ± 6.8 (147-168)	128 ± 2.8 (124-131)		
Pharyngeal overlap	20.8 ± 2.2 (18.0-23.0)	21.7 ± 2.7 (18.0-26.0)	17.3 ± 2.5 (14.0-20.0)	6.8 ± 3.1 (5.0-13.0)		
Max. body diam.	23.9 ± 0.6 (23.0-24.5)	23.8 ± 1.0 (23.0-25.0)	30.8 ± 1.6 (29.0-33.0)	30.7 ± 1.6 (29.0-33.0)		
Anal body diam.	17.5 ± 0.4 $(17.0-18.0)$	14.7 ± 0.8 (14.0-16.0)	21.1 ± 0.7 (20.0-22.5)	20.3 ± 1.3 (18.5-22.0)		
Tail	10.6 ± 1.4 (9.0-12.0)	16.3 ± 1.2 (15.0-18.0)	25.4 ± 2.7 (22.5-32.0)	17.8 ± 1.5 (16.0-20.0)		
Tail annuli	7.3 ± 1.0 (6-8)	-	16.6 ± 1.9 (14-19)	10.7 ± 0.8 (10-12)		
Phasmid to terminus	30.5 ± 2.1 (28.0-33.0)	-	34.5 ± 5.0 (30.0-40.0)	23.8 ± 4.6 (21.0-33.0)		
Spicules	-	22.0 ± 0.7 (21.0-23.0)	_	_		
Gubernaculum	-	9.6 ± 0.9 (9.0-11.0)	-	-		

^a Abbreviations are defined in Siddiqi (2000)

Table 6. Morphometrics of *Rotylenchus robustus* (de Man, 1876) Filipjev, 1936 from Tomales (California, USA) and from Lucena del Puerto (Huelva province, southern Spain). All measurements are in μ m and in the form: mean \pm s.d. (range) ^a.

_	R. robustus								
Population	Tomales, Cali	fornia,USA	Lucena del Puerto, Spain						
Character	Females	Males	Females	Males					
n L	12 1429 ± 83.6 $(1207-1533)$	9 1311 ± 129.7 (1117-1555)	9 1423 ± 143.4 (1178-1583)	$6 \\ 1203 \pm 57.3 \\ (1117-1272)$					
a	33.0 ± 2.7 (29.4-37.6)	37.3 ± 2.4 (33.5-40.3)	34.6 ± 2.8 (30.2-37.4)	38.8 ± 1.0 (37.2-39.8)					
b	12.9 ± 0.9 (11.4-14.3)	12.6 ± 0.7 (11.6-14.1)	13.5 ± 1.2 (11.2-14.9)	13.7 ± 0.6 (13.1-14.6)					
b'	7.1 ± 0.7 (6.3-8.3)	6.7 ± 0.5 (6.2-7.5)	7.2 ± 0.8 (5.8-8.5)	6.8 ± 0.3 (6.5-7.2)					
c	54.6 ± 5.6 (42.7-63.8)	39.3 ± 4.0 (34.3-45.7)	65.2 ± 8.3 (57.8-80.4)	40.2 ± 3.2 (36.0-45.4)					
c'	0.8 ± 0.1 (0.7-0.9)	1.3 ± 0.1 (1.2-1.5)	0.7 ± 0.05 $(0.6-0.8)$	1.4 ± 0.1 (1.2-1.5)					
V or T	54.0 ± 1.2 (53.0-57.0)	39.8 ± 8.4 (28.9-51.6)	53.8 ± 0.7 (53.0-55.0)	41.6 ± 8.0 (30.6-51.7)					
G_1	18.3 ± 3.7 (14.0-23.9)	-	17.1 ± 3.0 (13.2-21.6)	-					
G_2	18.0 ± 4.3 (12.0-23.6)	-	16.3 ± 2.9 (12.0-21.1)	_					
Stylet	46.8 ± 2.1 (43.0-50.0)	41.7 ± 1.7 (40.0-45.0)	49.6 ± 1.7 (46.0-51.0)	43.5 ± 2.1 $(41.0-47.0)$					
Stylet conus	23.2 ± 1.6 (21.0-26.0)	21.4 ± 0.7 (20.5-22.5)	25.3 ± 0.8 (24.0-26.5)	21.8 ± 0.9 (21.0-23.0)					
DGO	7.4 ± 0.9 (6.5-9.0)	7.1 ± 0.8 (6.0-9.0)	6.7 ± 0.4 $(6.0-7.0)$	5.3 ± 0.4 (5.0-6.0)					
0	15.8 ± 1.8 (13.3-19.6)	16.9 ± 1.9 (14.4-20.9)	13.5 ± 0.7 (12.0-14.6)	12.2 ± 0.4 (11.6-12.8)					
Anterior end to centre of median bulb	113 ± 7.9 (100-126)	104 ± 6.3 (91-110)	105 ± 3.3 (99-109)	88 ± 4.7 (82-95)					
Anterior end to excretory pore	150 ± 15.7 (134-176)	162 ± 13.7 (135-178)	162 ± 13.8 (143-179)	138 ± 4.5 (131-143)					
Pharynx length	203 ± 19.4 (178-234)	194 ± 12.1 (181-213)	198 ± 10.5 (181-210)	177 ± 11.0 (165-195)					
Pharyngeal overlap	47.9 ± 9.0 (39.0-67.0)	50.2 ± 3.6 (45.0-55.0)	46.1 ± 4.2 (41.0-52.0)	44.2 ± 3.4 $(41.0-49.0)$					
Max. body diam.	43.5 ± 3.6 (38.0-49.0)	35.2 ± 3.4 (31.0-41.0)	41.1 ± 1.3 (39.0-43.0)	31.0 ± 1.1 $(30.0-32.0)$					
Anal body diam.	34.1 ± 2.4 (30.0-37.0)	25.1 ± 3.7 (19.0-30.0)	31.4 ± 2.2 (29.0-36.0)	21.6 ± 1.1 (20.5-23.5)					
Tail	26.4 ± 2.4 (23.0-32.0)	33.5 ± 3.3 (26.5-38.0)	22.0 ± 2.7	30.0 ± 1.7 $(28.0-32.0)$					
Tail annuli	14.3 ± 2.2 $(13-21)$	-	$(19.0-26.0)$ 14.0 ± 2.2 $(10-17)$	-					
Phasmid to terminus	34.2 ± 4.1 (28.0-43.0)	-	26.6 ± 2.2 (22.0-30.0)	-					
Spicules	-	37.1 ± 2.8 (33.0-41.0)	-	35.7 ± 2.0 (33.0-38.0)					
Gubernaculum	-	18.6 ± 1.6 (16.0-21.0)	_	17.7 ± 1.0 (16.0-19.0)					

^a Abbreviations are defined in Siddiqi (2000)

Table 7. Similarity values (%) of rRNA sequences among *Rotylenchus* species. Above diagonal D2-D3 expansion segments of 28S rRNA and below diagonal ITS1 region*.

										Ro	tylen	chus s	spp.									
Rotylenchus spp.	01	02	03	04	05	06	07	08	09	10	11	12	13	14	15	16	17	18	19	20	21	22
01. R. paravitis n. sp.*		92	-	89	88	92	92	91	93	90	90	89	93	90	93	90	90	93	92	90	94	90
02. R. agnetis	-		_	88	89	92	91	88	90	90	90	90	92	90	92	92	90	92	92	89	90	91
03. R. brevicaudatus	74	-		_	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
04. R. brevicaudatus	74	-	99		86	89	88	88	87	93	93	86	89	93	88	88	95	89	88	87	88	89
05. R. buxophilus	-	-	-	-		97	90	87	87	93	93	86	89	93	89	89	95	89	89	88	87	88
06. R. buxophilus	81	-	-	81	-		93	91	91	96	96	90	92	97	92	91	98	92	92	91	91	91
07. R. cazorlaensis	77	-	74	74	-	61		90	91	91	92	89	95	92	95	90	91	94	94	91	90	90
08. R. conicaudatus	62	-	62	62	-	69	59		90	87	88	88	90	88	90	88	90	91	91	89	89	88
09. R. eximius	74	-	71	70	-	60	73	56		89	89	89	92	89	92	89	89	92	92	90	92	90
10. R. goodeyi	-	-	-	-	-	-	-	-	-		99	88	90	99	90	89	95	90	90	89	88	89
11. R. incultus	66	-	65	65	-	83	66	66	61	-		88	90	99	90	90	95	90	90	90	88	89
12. R. iranicus	67	-	67	67	-	71	65	67	63	-	68		89	88	90	88	88	90	90	87	87	88
13. R. jaeni	77	-	64	73	-	63	81	59	71	-	72	68		90	98	90	90	96	96	90	92	90
14. R. laurentinus	65	-	65	65	-	83	65	66	61	-	68	72	68		90	90	95	90	90	90	89	89
15. R. magnus	77	-	75	74	-	63	85	59	73	-	65	68	92	67		90	90	97	97	90	92	90
16. R. montanus	65	-	65	64	-	64	65	61	62	-	67	65	65	66	65		90	91	90	88	89	99
17. R. pumilus	81	-	-	81	-	97	62	68	59	-	83	71	62	83	62	64		91	90	89	90	90
18. R. robustus	73	-	70	69	-	59	80	59	70	-		67	81	65	85	65	58		99	91	92	90
19. R. uniformis	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-		90	92	90
20. R. unisexus	69	-	64	64	-	68	66	65	65	-	72	72	66	67	66	65	-	62	-		89	88
21. <i>R. vitis</i>	72	-	62	61	-	68	65	62	63	-	64	64	66	63	67	72	68	66	-	65		89
22. Rotylenchus sp.	66	-	65	64	-	87	65	61	63	-	65	65	65	66	65	98	86	81	-	66	72	

^{*}Newly obtained sequences are in bold letters. (-) Not available.

Accessions numbers (D2-D3, ITS1, respectively): **1= JX015422, JX015434**; 2= EU280795; 3= DQ309587; **4= JX015419, JX015430**; 5= FJ485647; **6= JX015421, JX015432**; 7= EU280792, EU373671; 8= HQ700698; HQ700700; 9= EU280794, EU373664; 10= DQ328756; 11= EU280797; EU373673; 12= HQ700697; HQ700699; 13= EU280791, EU373662; 14= EU280798, EU373667; 15= EU280789, EU373665; 16= DQ328743, EU280800; **17= JX015423, JX015435**; **18= JX015424, JX015437**; 19= DQ328737; 20= EU280799; EU373675; 21= JN032581, JN032582; 22= DQ328742, EU280802.

Table 8. Results of the SH-tests for alternative hypotheses using ML trees.

		D2-D3			18S	
Topologies and hypothesis tested	-LnL	Difference of -LnL	P *	-LnL	Difference of -LnL	P *
ML tree	6727.44	best		5810.80	best	
All Rotylenchus spp. constrained into one group	6752.96	25.52	0.071	5814.55	3.75	0.298
All Helicotylenchus spp. constrained into one group	6733.37	5.93	0.572	5810.80	0.00	1.00
All <i>Hoplolaimus</i> spp. constrained into one group**	6727.44	0.00	1.00	Non-available	-	-
Grouping separately Rotylenchus-Helicotylenchus-Hoplolaimus-Scutellonema**	6753.99	26.55	0.064	5814.55	3.75	0.298
All $Rotylenchus$ spp. constrained, with exception of R . $conicaudatus + R$. $unisexus$	6729.92	2.48	0.635	Non-available	-	-

^{*}P < 0.05 indicates the significant differences between the two inferred tree topology. **Non available partial 18S sequences for *Hoplolaimus* or some specific *Rotylenchus* sequences in GenBank.

1 Figure legends

2

- 3 Fig. 1. Principal component (PC) analysis of 33 morphometric characters used to characterise
- 4 21 specimens of *Rotylenchus vitis* and *R. paravitis* n. sp., respectively, and 12 and nine
- 5 specimens of two populations of *R. robustus* from Spain and USA, respectively. Projection of
- 6 morphometric characters on the plane of PC 1 and 2 (A), 1 and 3 (B), and 1 and 4 (C).

7

- 8 Fig. 2. Rotylenchus paravitis n. sp. (A) Female pharyngeal region. (B) Detail of pharyngeal
- 9 gland. (C) Female anterior body region. (D, E) Male and female habitus. (F) Vulval region.
- 10 (G, H) Female tail regions. (I) Male tail region.

11

- **Fig. 3.** Light micrographs of *Rotylenchus paravitis* n. sp. (A) Female habitus. (B) Female
- pharyngeal region. (C-F) Female anterior body region. (G, H) Detail of pharyngeal gland. (I)
- 14 Vulval region. (J) Detail of spermatheca. (K) Detail of ovary. (L-O) Female tail regions. (P-
- O) Male tail region. Abbreviations: a = anus; n = gland nucleus; ph = phasmid. (Scale bars: A
- 16 = 500 μm; B = 50 μm; C-F = 25 μm; G-I = 50 μm; J-O = 25 μm; P-Q = 50 μm).

17

- **Fig. 4.** Scanning electron microscope photographs of *Rotylenchus paravitis* n. sp. (A, B)
- 19 Female lip region, lateral view, showing anterior narrowing (arrowed) and beginning of
- 20 lateral fields (lf). (C) En face view showing oral aperture (oa), amphid (ap) and labial disc
- 21 (ld). (D, E) Vulval region showing non-areolated lateral fields and few developed epiptygma
- 22 (ept). (F) Female tail region showing anus (a) and phasmid (ph). (Scale bars: A, D-F = $20 \mu m$;
- 23 B, $C = 10 \mu m$).

- 25 **Fig. 5.** Light micrographs of *Rotylenchus brevicaudatus* Colbran, 1962 (A-C) Female anterior
- body region. (D) Detail of pharyngeal region. (E) Lateral fields at mid-body. (F-K) Female

- 1 tail regions. (L) Male tail region. Abbreviations: a = anus; ep = excretory; ph = phasmid.
- 2 (Scale bars: A-D = 20 μ m; B = 50 μ m; E-L = 10 μ m).

- 4 Fig. 6. Light micrographs of *Rotylenchus buxophilus* Golden, 1956 (A) Female habitus. (B)
- 5 Female anterior body region. (C-E) Female lip regions. (F) Detail of pharyngeal region. (G)
- 6 Vulval region. (H) Lateral fields at mid-body. (I-M) Female tail regions. (N) Abnormal
- 7 female tail. Abbreviations: a = anus; ep = excretory; ph = phasmid. (Scale bars: $A = 100 \mu m$;
- 8 B = 20 μ m; C-N= 10 μ m).

9

- **Fig. 7.** Light micrographs of *Rotylenchus pumilus* (Perry et al., 1959) Sher, 1961. (A) Female
- pharyngeal regions. (B, C) Details of pharyngeal gland. (D-E) Female anterior body region.
- 12 (F) Detail of lateral fields at mid-body. (G) Vulval region. (H) Female habitus. (I, J, L)
- Female tail regions. (K) Detail of phasmid. Abbreviations: a = anus; n = gland nucleus; ph = anus
- 14 phasmid. (Scale bars: A, H = 50 μ m; B-E, G = 25 μ m; F, K = 10 μ m; I-J, L = 20 μ m).

15

- **Fig. 8.** Scanning electron microscope photographs of *Rotylenchus robustus* (de Man, 1876)
- 17 Filipjev, 1936 from Lucena del Puerto, Huelva province, southern Spain (A-D) and Tomales,
- 18 California, USA (E-H). (A, B, E) Female lip region, lateral views, showing basal lip annulus
- with longitudinal striations. (C, F, G) En face view showing oral aperture (oa), amphid (ap)
- and labial disc (ld). (D, H) Female tail region showing anus (a) and phasmid (ph). (Scale bars:
- 21 A, C, E-G = 10 μ m; B, D, H = 10 μ m).

- **Fig. 9.** The 50% majority rule consensus tree from Bayesian analysis generated from the D2-
- D3 of 28S-rRNA gene dataset with the TVM + I + G model. Posterior probabilities more than
- 25 65% are given for appropriate clades; bootstrap values greater than 50% are given on
- appropriate clades in ML analysis. Newly obtained sequences are in bold letters.

1	
2	Fig. 10. The 50% majority rule consensus tree from Bayesian analysis generated from the
3	ITS-rRNA gene dataset with TVM + G model. Posterior probabilities more than 65% are
4	given for appropriate clades; bootstrap values greater than 50% are given on appropriate
5	clades in ML analysis. Newly obtained sequences are in underline letters.
6	
7	Fig. 11. The 50% majority rule consensus tree from Bayesian analysis generated from the
8	18S-rRNA gene dataset with TVM1 + I + G model. Posterior probabilities more than 65% are
9	given for appropriate clades; bootstrap values greater than 50% are given on appropriate
10	clades in ML analysis. Newly obtained sequences are in underline letters.
11	
12	Fig. 12. The 50% majority rule consensus tree from Bayesian analysis generated from the
13	partial <i>COI</i> gene dataset with GTR + I + G model. Posterior probabilities more than 65% are
14	given for appropriate clades; bootstrap values greater than 50% are given on appropriate
15	clades in ML analysis. All sequences are newly obtained.
16	
17	Fig. 13. The 50% majority rule consensus tree from Bayesian analysis generated from the
18	hsp90 gene dataset with K80 + G model. Posterior probabilities more than 65% are given for
19	appropriate clades; bootstrap values greater than 50% are given on appropriate clades in ML
20	analysis. Newly obtained sequences are all.
21	
22	Fig. 14. PCR-D2-D3-28S-RFLP profile for <i>Rotylenchus paravitis</i> n. sp. Lines: M – 100bp
23	DNA ladder (Promega); U – unrestricted PCR product, 1- AvaI, 2- RsaI, 3 - BseNI, 4 – MvaI,
24	5 - HpaII.

Comparative molecular and morphological characterisations in Rotylenchus

- 1 Fig. 15. The gel with specific amplicons obtained in the results of PCR with species specific
- 2 primers. (A) PCR with the Rotylenchus robustus specific primer (TW81 + R robustus
- 3 primers). (B) PCR with the R. vitis specific primer (TW81 + R_vitis). (C) PCR with the R.
- 4 paravitis n. sp. specific primer (TW81 + R paravitis). Lanes: M 100 bp DNA ladder
- 5 (Promega); 1 R. robustus (CA, USA); 2 R. vitis; 3 R. paravitis n. sp.; 4 R.
- 6 brevicaudatus (Australia); 5 R. buxophilus (CA, USA); 6 R. pumilus (CA, USA), 7 -
- 7 Scutellonema brachyurus (USA); 8: control without DNA.