Morphological and molecular characterisation, and phylogenetic position of X. browni sp. n., X. penevi sp. n. and two known species of Xiphinema americanum-group (Nematoda, Longidoridae)

Stela Lazarova¹, Vlada Peneva¹, Shesh Kumari²

¹ Department of Animal Biodiversity and Resources, Institute of Biodiversity and Ecosystem Research, Bulgarian Academy of Sciences 2, Gagarin Street, 1113 Sofia, Bulgaria ² Division of Plant Health, Crop Research Institute, Drnovská 507, Ruzyně, 16106 Prague 6, Czech Republic

Corresponding author: Vlada Peneva (vpeneva@ecolab.bas.bg)

Academic editor: S. Subbotin | Received 4 February 2016 | Accepted 1 March 2016 | Published 28 March 2016

Citation: Lazarova S, Peneva V, Kumari S (2016) Morphological and molecular characterisation, and phylogenetic position of X. browni sp. n., X. penevi sp. n. and two known species of Xiphinema americanum-group (Nematoda, Longidoridae). ZooKeys 574: 1–42. doi: 10.3897/zookeys.574.8037

Abstract

Using ribosomal (18S, ITS1, ITS2, D2-D3 expansion segments of 28S rDNA) and mitochondrial (partial cox1 and nad4) DNA markers in a study of several populations of Xiphinema americanum-group from Europe and Morocco, two cryptic species X. browni sp. n. (formerly reported as X. pachtaicum) and X. penevi sp. n. were revealed. The species are described, illustrated and their phylogenetic relationships discussed. The first species is most similar to X. parasimile and is a member of X. simile species complex. The phylogenetic reconstructions inferred from three molecular markers (18S, D2-D3 28S rDNA and cox1) showed that X. penevi sp. n. is part of X. pachtaicum-subgroup and is closely related to X. incertum, X. pachtaicum, X. parapachydermum, X. plesiopachtaicum, X. astaregiense and X. pachydermum. Also, a separate “X. simile-subgroup”, outside the X. pachtaicum-subgroup and so far consisting only of the parthenogenetic species X. simile, X. parasimile, X. browni sp. n. and probably X. vallense was formed. New primers for amplification and sequencing of part of the nad4 mitochondrial gene were designed and used.

Keywords

Bayesian Inference, Bulgaria, Cytochrome c oxidase subunit 1, Czech Republic, Morocco, Nicotinamide dehydrogenase subunit 4, phylogeny, ribosomal DNA, Slovakia
Introduction

The *Xiphinema americanum*-group is a well defined natural complex of species (Lamberti et al. 2000, Coomans et al. 2001, He et al. 2005b) with high significance to agriculture caused by the ability of several species to transmit economically important plant viruses (McFarlane et al. 2002), although there are controversial opinions defining the group (Archidona-Yuste et al. 2016). Even for experienced nematologists species delimitation within this group is challenging because they have rather similar morphology and metrics, and the existing keys (Lamberti et al. 2000, 2004) do not always allow species differentiation and identification. During the last decade wide usage of DNA sequencing in *Xiphinema* taxonomy including this group revealed the existence of a number of cryptic species (Gutiérrez-Gutiérrez et al. 2010, 2012, Archidona-Yuste et al. 2016). This was the case with several populations from the Czech Republic and Slovakia (Kumari et al. 2005, 2010b) originally identified as *X. pachtaicum* (Tulaganov, 1938) and one population from Morocco provisionally also determined as *X. pachtaicum*. The objectives of the present study were: i) to characterise populations from the Czech Republic, Slovakia and Morocco both morphologically and genetically; ii) to sequence populations of *X. pachtaicum* and *X. parasimile* Barsi & Lamberti, 2004 from Bulgaria for comparison; iii) to clarify phylogenetic relationships of identified species using ribosomal and mitochondrial DNA.

Material and methods

Sampling, nematode isolation and processing

The *Xiphinema* specimens examined originated from various localities in the Czech Republic (Kurdějov, Mohyla miru and Sokolnice, grapevines), Slovakia (Moča, grapevine), Bulgaria (Balgarene village, pear tree, Vinogradets vicinity, vineyard) and Morocco (Ifrane, holm oak tree). Details of the soil sampling, nematode isolation and processing for Czech and Slovakian populations are given in Kumari et al. (2005, 2010b). A decanting and sieving technique was used for extracting nematodes from soil samples from Bulgaria and Morocco. *Xiphinema* specimens recovered were heat killed at 55°C for two minutes, fixed in a 4% formalin, 1% glycerol solution, processed to anhydrous glycerol (Seinhorst 1959), and mounted on glass microscope slides. Drawings were prepared using an Olympus BX51 compound microscope with differential interference contrast (DIC). Photographs were taken using an Axio Imager.M2-Carl Zeiss compound microscope with a digital camera (ProgRes C7) and specialised software (CapturePro Software 2.8). Measurements were made using an Olympus BX41 light microscope, a digitising tablet (CalComp Drawing Board III, GTCO CalCom Peripherals, Scottsdale, AZ, USA), and computer Digitrak 1.0f programme (Philip Smith, Scottish Crop Research Institute, Dundee, UK).
DNA extraction, amplification and sequencing

Individual nematodes from Bulgaria, Morocco (DESS-preserved), Czech Republic and Slovakia (1M NaCl-preserved) were mounted on temporary slides containing glass beads and after taking measurements and photomicrographs the slides were dismantled, individual nematodes removed, and added in 0.25 M NaOH to digest overnight and thereafter heated to 99°C for 3 min. Afterwards 10 μl of 0.25 M HCl, and 5 μl each of 0.5 M Tris-HCl (pH 8) and 2% Triton X-100 were added and the mixture was incubated for another 3 min at 99°C (Stanton et al. 1998). Finally, the DNA suspension was cooled and the DNA was either used directly for PCR or stored at -20°C until template was needed for PCR reactions. Genomic DNA which was prepared by Kumari et al. (2010b) was also used in this study.

Six regions (18S, ITS1, ITS2, D2-D3 expansion segments of 28S, cox1 and nad4) of ribosomal and mitochondrial DNA were amplified and sequenced. Primer sequences and references to the primers are given in Table 1. The 18S gene of the Czech population was amplified by using primers SSU_F_04+SSU_R_09 (first fragment), SSU_F_22+SSU_R_13 (second fragment) and SSU_F_23+SSU_R_81 (third fragment). The 18S gene of other populations was amplified by using primer combination 988F+1912R (first fragment) and 1813F+2646R (second fragment).

Initially partial nad4 gene was amplified with the primers CDF+CDR but only one specimen was amplified using these primers. A pair of new primers (nadpachF+nadpachR) was designed using online software PRIMER 3 (http://frodo.wi.mit.edu/) from the sequences which were amplified by (CDF+CDR). For final analysis all specimens and populations of X browni sp. n. from the Czech Republic and Slovakia were amplified and sequenced by using nadpachF + nadpachR primers.

The PCR reaction was performed in 25 μl total volume containing 1 PCR bead (GE Healthcare, Buckinghamshire, UK), 20.5 μl double distilled sterile water, 2.0 μl of each primer (10pmol/μl) (synthesized by Generi Biotech, Hradec Králové, Czech Republic), and 0.5 μl of DNA added as a template for PCR. A negative control (sterilized water) was included in all PCR experiments. The cycling profile for all ribosomal DNA and mtDNA markers was as described by Kumari and Subbotin (2012) and by He et al. (2005a), respectively. All PCR reactions were performed in a DNA Engine PTC–1148 thermal cycler (Bio-Rad). Aliquots of PCR were analysed by gel electrophoresis and the remaining products were purified using High Pure Product Purification kit (Roche Diagnostics GmbH, Mannheim, Germany) and sequenced in both directions using each primer pair one forward and one reverse (Macrogen, Netherlands). Sequencher™ 4.8 (Genes codes. Corp., Ann Arbor, MI, USA) was used to assemble and view each sequence and check for base-calling errors. Accession numbers of all sequences are given in Table 2.

Sequence and phylogenetic analyses

A BLAST (Basic Local Alignment Search Tool) search at NCBI (National Center for Biotechnology Information) was performed using the obtained sequences as queries.
Table 1. Primers used to amplify ribosomal and mitochondrial DNA.

| Gene | Primer name | Direction | Primer sequence 5′ - 3′ | Reference |
|------|-------------|-----------|-------------------------|-----------|
| 18S  | SSU_F_04    | forward   | GCT TGT CTC AAA GAT TAA GCC | Blaxter et al. (1998) |
| 18S  | SSU_R_09    | reverse   | AGC TGG AAT TAC CGC GGC TG  | Blaxter et al. (1998) |
| 18S  | SSU_F_22    | forward   | TCC AAG GAA GGC AGC AGG C  | Blaxter et al. (1998) |
| 18S  | SSU_R_13    | reverse   | GGG CAT CAC AGA CCT GTT A  | Blaxter et al. (1998) |
| 18S  | SSU_F_23    | forward   | ATT CCG ATA AGC GAG A      | Blaxter et al. (1998) |
| 18S  | SSU_R_81    | reverse   | TGA TCC WKC YGC AGG TTC AC | Blaxter et al. (1998) |
| 18S  | 988F        | forward   | CTC AAA GAT TAA GCC ATG C   | Holterman et al. (2006) |
| 18S  | 1912R       | reverse   | TTT ACG TGC AGA ACT AGG G   | Holterman et al. (2006) |
| ITS1 | pXb101      | forward   | TTG ATT ACG TCC CTG CCC TTT | Vrain et al. (1992)  |
| ITS1 | ChR         | reverse   | ACG AGC CGA GTG ATC CAC CG  | Cherry et al. (1997) |
| ITS2 | WDF         | forward   | AGA CAC AAA GAG CAT CGA CT  | Kumari et al. (2009) |
| ITS2 | pXb481      | reverse   | TTT CAC TCG CCG TTA CTA AGG | Vrain et al. (1992)  |
| D2-D3| D2A         | forward   | ACA AGT ACC GTG AGG GAA AGT TG | Nunn (1992) |
| cox1 | COIF        | forward   | GAT TTT TTG GKC ATC CCG ARG | Nunn (1992) |
| cox1 | XIPHR2      | reverse   | GTA CAT AAT GAA AAT GTG CCA | Lazarova et al. (2006) |
| nad4 | CDF         | forward   | AAA AAG ATG GTA TTG GAG     | Kumari and Cesare (2013) |
| nad4 | CDR         | reverse   | GCA CAT GTA GAA GCT AGT     | Kumari and Cesare (2013) |
| nad4 | nadpachF    | forward   | ATA GAA GCA TTA CCA ACT A   | This study |
| nad4 | nadpachR    | reverse   | TAG TAC CAG AGG ATC AAT A    | This study |

to confirm their nematode origin and to identify the most closely related nematode sequences. Sequences revealing high similarity to those obtained here were included in the phylogenetic analyses of both ribosomal and mitochondrial gene regions (Neilson et al. 2004, Oliveira et al. 2004, He et al. 2005b, Gozel et al. 2006, Holterman et al. 2006; Lazarova et al. 2006, Kumari et al. 2009, Gutiérrez-Gutiérrez et al. 2010, Kumari et al. 2010a, Kumari et al. 2010b, De Luca and Agostinelli 2011, Gutiérrez-Gutiérrez et al. 2011a, Gutiérrez-Gutiérrez et al. 2011b, Meza et al. 2011, Sakai et al. 2011, Gutiérrez-Gutiérrez et al. 2012, Kumari and Subbotin 2012, Sakai et al. 2012, Kumari and Cesare 2013, Tzortzakakis et al. 2014, Getaneh et al. 2015, etc). Sequence numbers are presented in the trees. The multiple sequence alignments (MSA) of all datasets were performed using the GUIDANCE2 Server available at http://guidance.tau.ac.il/ (Sela et al. 2015). All three alignment algorithms (MAFFT, PRANK and ClustalW) were tested and the MSAs having highest alignment confidence scores were used for ITS phylogenetic reconstructions. Subsequently, the MSAs were manually optimised and trimmed using MEGA 6 (Tamura et al. 2013). The phylogenetic reconstructions were performed using the Bayesian Inference (BI) algorithm implemented in MrBayes 3.2.5. (Huelsenbeck and Ronquist 2001; Ronquist et al. 2012) using the General Time Reversible model plus Gamma distribution rates (GTR + G). The Bayesian MCMC tree searches were run using default heating parameters for 2 000 000 generations with a sample frequency of 1000 generations. The first 25% of the chains discarded as burning and the remaining 75% trees kept to summarise the tree topology, branch lengths,
Table 2. NCBI accession numbers of representative individual specimens for ribosomal and mitochondrial DNA.

| Species          | Xiphinema browni sp. n | X. pachtaicum | X. parasimile | X. penevi sp. n. |
|------------------|-------------------------|---------------|---------------|-----------------|
| Country          | Czech Republic          | Slovakia      | Bulgaria      | Morocco         |
| Locality         | Kurdějov                | Mohyla míru   | Moča          | Ifran           |
|                  | NSB1                    | NSB2          | NSB3          | NSB4            |
| Isolate          | NSB1                    | NSB2          | NSB3          | NSB4            | NSB5            | NSB6            | NSB7            |
| 18S              | KU250135                | KU250136      | KU250137      | KU250138        | KU250139        | KU250140        | KU250141        |
| 18S+ITS1         | KU250142                | KU250143      | KU250144      | NA              | NA              | NA              | NA              |
| 5.8S+ITS2+28S    | KU250145                | KU250146      | KU250147      | KU250148        | KU250149        | NA              | NA              | KU250150        |
| D2/D3            | KU250151                | KU250152      | KU250153      | KU250154        | KU250155        | KU250156        | KU250157        |
| cox1             | GU222424*               | *             | *             | KU250158        | NA              | KU250159        | NA              |
| nad4             | KU250160                | KU250161      | KU250162      | NA              | NA              | NA              |

* Kumari et al. (2010); NA = not acquired

and posterior probabilities (PP) of branch support. Convergence diagnostic values were calculated every 1000 generations with a predefined stop value equal to 0.01. A single strict consensus tree was visualised using FigTree v1.4.2 graphical viewer. Posterior probabilities values of ≥0.80 were considered as credible support values for nodes.

Taxonomy

**Xiphinema browni sp. n.**

http://zoobank.org/E385F7F7-2C78-4D54-BC57-0D24EDD43CB8

Figures 1–8, 15–18

**Xiphinema pachtaicum** (Tulaganov, 1938) Kirjanova, 1951 apud Kumari et al. 2005, syn. n.

Measurements. See Tables 3–5.

**Description.** Females. Body slender C to open spiral shaped. Cuticle with fine transverse striae. Thickness of the cuticle at postlabial region 1–1.5 μm, 1.5 rarely 2 μm at mid-body and 2 μm at post-anal region. Labial region set-off from the rest of the body by a constriction, expanded, rounded laterally, 5.0±1.1 (4–7) μm high. Amphidelal fovea hardly visible, funnel-shaped, its opening c. 5 μm (50%) wide visible posterior the constriction level. Distance between first and second guide ring in specimens with retracted odonostyle 5–10 μm long. Odontophore with moderately developed basal flanges 6.1±0.6 (5.5–7) μm wide. A small vestigium observed occasionally in slender part of pharynx. Pharyngeal characters presented at Table 4. Dorsal pharyngeal gland nucleus 2 μm diam. Ventrosobrateral nuclei barely visible. Rectum 20.8 ± 1.5 (18–23) μm, n=7, or c 1.3 times anal body diameter. Reproductive system amphidelphic, symbiont bacteria present in the ovaries. Separate uteri and ovejector present (Table 5), oviduct 90.5±13.0 (68–101) μm; vagina bell-shaped 39.5% of the corresponding body
width (33–50%, n=14), vulva post-equatorial. Numerous sperm observed in one female from Kurdějov (Figs 2B, 4B). Tail conical, dorsally convex, ventrally straight or slightly concave with narrowly rounded to pointed terminus. Two pairs of caudal pores.

**Male.** Very rare. One specimen found in Sokolnice population. Male similar to the female with posterior region more strongly curved. Lip region and tail shape as in females, differences were observed within body width and tail length, which reflected a and c' values. Spicules robust, slightly curved, lateral guiding piece 7 μm long. Adanal pair preceeded by a row of 5 irregularly spaced supplements, the two anteriormost weakly developed. Tail conoid, ventrally straight, dorsally convex with pointed terminus, caudal pores not visible. The slide of the only male specimen, described by Kumari et al. (2005), was subsequently damaged.

**Juveniles.** The scatter diagram based on functional and replacement odontostyle, and body length revealed the presence of four juvenile stages (Fig. 8). Tail shape and length similar in all stages and females with c' slightly decreasing in successive stages (Kumari 2005, Fig. 3, Table 3).

**Type locality and plant association.** Kurdějov, Břeclav County, South Moravia, Czech Republic, associated with grapevine. Other localities: Mohyla míru, Brno-Venkov County, South Moravia, the Czech Republic, in the rhizosphere of apple trees; Sokolnice, Brno-Venkov County, South Moravia, the Czech Republic, in the rhizosphere of grapevine; Moča, Komárno County, Nitra, Slovak Republic, in the rhizosphere of grapevine.

**Type material.** The holotype, 9 paratype females and juveniles from all stages are deposited in the nematode collection of the Institute of Biodiversity and Ecosystem Research, Sofia, Bulgaria. Other paratypes deposited as follows: 15 females in the Crop Research Institute, Prague, the Czech Republic; 5 females in the USDA Nematode Collection, Beltsville, Maryland, USA; 5 females in the Nematode Collection of the Institute of Plant Protection, Bari, Italy; 5 females in the Wageningen Nematode Collection (WANECO), Wageningen, the Netherlands. The ribosomal and mtDNA sequences (18S rDNA, ITS1, ITS2, D2-D3, cox1, nad4) of *X. browni* sp. n. are deposited in GenBank (for accession numbers see Table 2).

**Sequence and phylogenetic analyses.** There was no sequence variation between populations for 18S and D2-D3, ITS1 and ITS2 rDNA regions of *X. browni* sp. n. Of all four populations studied cox1 region of three population from the Czech Republic (Kurdějov, Mohyla Míru, Sokolnice) were sequenced by Kumari et al. (2010b) and all populations were identical therefore only one population was submitted to GenBank (accession number GU222424). The Slovakian population was sequenced in this study and it was identical to previously published sequence of Kurdějov the population identified as *X. pachtaicum* (GU222424, Kumari et al. 2010b). All four sequenced populations were also identical for nad4 part. BLAST at NCBI using 18S and D2-D3 region sequences as queries revealed highest similarity (99 and 87%) to the corresponding sequences of *X. simile* Lamberti, Choleva & Agostinelli, 1983 from Serbia (AM086681) and two Spanish populations of *X. opisthohysterum* Siddiqi, 1961 (JQ990040 and KP268967), respectively. The es-
Morphological and molecular characterisation, and phylogenetic position of *X.* browni...

Figure 1. *Xiphinema browni* sp. n. Female: Variations in: A–C Anterior end D–F Pharyngeal bulb G–I Tail shape A, D, G Kurđejov (type population) B, F, I Mohyla míru C, E, H Sokolnice. Scale bars: 25 μm
Figure 2. *Xiphinema browni* sp. n. Female: Variations in genital system: A, B Anterior genital branch C Posterior genital branch D–F Region of vagina and uteri A, B, D Kurdějov (type population) C, F Mohyla miru E Sokolnice. Scale bars: 25 μm.

Estimated divergences (p-distance) between the 18S rDNA sequences of the new species and the closest species, *X. parasimile* from Bulgaria (this study) and *X. simile* from Serbia (AM086681) were 0.3 (6 nt) and 1.2% (21 nt), respectively. Again, the new D2-D3
Figure 3. *Xiphinema browni* sp. n. A–C Entire body (A, C females B male); Female: D–F Anterior ends G–I Pharyngeal bulbus J–M Tail shape variation A, F, I, M Mohyla míru B, E, H, L Sokolnice C, D, G, J, K Kurdějov (type population). Scale bars: (A–C) 400 µm; (D–M) 30 µm.
Figure 4. *Xiphinema browni* sp. n. Female: **A–D** Genital system (**B** uterus full with sperm) **E–G** Labial region (**E** Amphid **F** Female **G** Male) **H** Ovary with endosymbionts **I, K–N** Variations in vagina **J** Lateral field **A, B, H, I, J** Kurďéjov **C, G** Sokolnice **D, K, M** Mohyla míru **E, F, L, N** Moča. Scale bars: 30 μm (**A–D, H–J**); 12 μm (**E–G, K–N**).
Figure 5. Xiphinema browni sp. n., Sokolnice. Male: A Anterior end B Pharyngeal bulbus C Posterior end D Spicules. Scale bars: 25 μm
Figure 6. *Xiphinema browni* sp. n., Kurdějov. Juveniles and female: **A–E** Anterior ends of first- to fourth-stage juveniles and female **F–J** Tails of first- to fourth-juvenile stages and female. Scale bar: 25 μm
Morphological and molecular characterisation, and phylogenetic position of *X. browni*...

**Figure 7.** *Xiphinema browni* sp. n. Kurdějov. Juveniles and female: A–E Anterior ends of first- to fourth-stage juveniles and female F–J Tails of first to fourth juvenile stages and female (G1 and G2 – second-stage juvenile). Scale bar: 30 μm.

**Figure 8.** Scatter plot of odontostyle (■) and replacement odontostyle (□) against body length of *Xiphinema browni* sp. n. juveniles and females from Kurdějov population.
| Localities          | Xiphinema browni sp. n | X. pachtaicum |
|---------------------|------------------------|--------------|
| Plant host          | Kurdějov grapevine     | Sokolnice grapevine | Mohyla miru apple | Moča grapevine |
|                     | n                      | n            | 50 females | 20 females | 12 females | 4 females | Balgarene pear | 6 females |
| L.                  | 1904                   | 2031±123     | 1886±89   | 1849      | 1972±90   | 1715±142   | 1735±232  |
|                     | (1798–2408)            | (1751–2099)  | (1785–2079) | (1603–1922) | (1522–2015) |               |           |
| a                   | 57.8                   | 60.5±4.4     | 73.9      | 60.1±3.14 | 69.5±6.57 | 58.7±4.9   |               |           |
|                     | (56.9–81.3)            | (52.3–69.9)  | (55.6–64.5) | (63.6–76.3) | (53.3–65.7) |               |           |
| b                   | 6.7                    | 6.9±0.38     | 6.9       | 7.0±0.32  | 8.2 ± 6.8 | 5.9±0.5    |               |           |
|                     | (6.1–8.7)              | (6.4–7.9)    | (6.4–7.4) |           |           |           |               |           |
| c                   | 64.8                   | 65.8±5.71    | 54.4      | 64.9±3.41 | 61.6±8.72 | 58.2±8.3   |               |           |
|                     | (54.7–83.0)            | (56–79.6)    | (58.5–70.3) | (53.4–73.9) | (50.9–66.3) |               |           |
| c'                  | 1.9                    | 1.8±0.14     | 1.89      | 1.8±0.08  | 1.8±0.17  | 1.7±0.1    |               |           |
|                     | (1.53–2.07)            | (1.61–2.13)  |           |           |           |           |               |           |
| V/Spicule length    | 56.1                   | 55±1.30      | 29.0      | 55.4±1.15 | 55.5±1.16 | 58.6±1.4   |               |           |
|                     |                        | (52.3–58.5)  | (49–57)   | (53.8–51) | (53.8–56.4) | (57.0–60.4) |           |
| Odontostyle         | 84                     | 79±2.6       | 76        | 82±3.39   | 77±4.69   | 84.2±3.7   |               |           |
|                     | (78–86)                | (74–83)      | (73–85)   | (72–81)   | (78–88.5) |               |               |           |
| Odontophore         | 43                     | 41±0.91      | 38        | 43.1±1.88 | 38±3.30   | 48.9±2.1   |               |           |
|                     | (38–48)                | (39–43)      | (39–46)   |           |           |           |               |           |
| Oral aperture to    | 72                     | 68±2.35      | 67        | 71±1.68   | 66±5.06   | 76.8±3.4   |               |           |
| guide ring          |                        | (52.3–58.5)  | (49–57)   | (53.8–51) | (53.8–56.4) | (57.0–60.4) |           |
| Tail length         | 29                     | 29±2.24      | 34        | 30±0.82   | 28±1.63   | 29.8±0.9   |               |           |
|                     | (25–33)                | (24–32)      | (29–32)   | (26–30)   | (28–30)   |               |               |           |
| Length of hyaline   | 8                      | 8±1.28       | 10        | 8±0.68    | 8±1.41    | 8.7±1.0    |               |           |
| part                | (6–12)                 | (6–10)       | (7–9)     | (7–10)    | (8–10)    |               |               |           |
| Body diam. at:      | 8                      | 8±0.58       | 9         | 9±0.43    | 8±0.50    | 8.8±0.2    |               |           |
| - lip region        | (8–10)                 | (8–9)        | (8.5–10)  | (7–8)     | (8.5–9)   |               |               |           |
| - guiding ring      | 22                     | 19±0.49      | 19        | 22±1.44   | 19±1.41   | 21.5±1.0   |               |           |
|                     | (19–21)                | (19–20)      | (19.5–24) | (18–21)   | (20.5–23) |               |               |           |
| - base of pharynx   | 29                     | 26±1.58      | 23        | 28±1.69   | 23.1, 23.8| 26.5±1.2   |               |           |
|                     | (22–32)                | (19–20)      | (25–30)   |           |           |               |               |           |
| - mid body          | 33                     | 31±2.58      | 25        | 34±2.66   | 25±1.89   | 28.9±2.1   |               |           |
|                     | (25–38)                | (26–37)      | (29–38.5) | (22–26)   | (26–32)   |               |               |           |
| - anus              | 16                     | 16±0.92      | 18        | 17±0.89   | 16±0.96   | 17.1±1.0   |               |           |
|                     | (14–19)                | (15–18)      | (16–19)   | (15–17)   | (16–19)   |               |               |           |
| - beginning of      | 7.5                    | 7±0.62       | 8         | 8±1.50    | 8.7±0.0   |               |               |           |
| hyaline part        | (5–10)                 | (6–8)        | (6–9)     |           | (9–9)     |               |               |           |
sequence of *X. parasimile* from Bulgaria was most similar (p-distance = 4.6%), followed by the Serbian populations of *X. parasimile* (p-distance = 7.6–7.9%, calculated for D2 region only) and various populations of *X. simile* (14.1–14.7%). The partial *cox1* sequences of *X. browni* sp. n revealed highest similarity to *X. simile* from Slovakia (AM086708). Surprisingly, these two species showed very high similarity 99% (2 nts difference) in *cox1* sequences and higher dissimilarity in 18S rDNA (p-distance = 1.2%, 21 nts). Other authors (Gutiérrez-Gutiérrez et al. 2012) have also reported similar observation namely, 100% identity in *cox1* part of two different species *X. duriense* Lamberti, Lemos, Agostinelli & D’Addabo, 1993 (JQ990053) and *X. opistohysterum* (JQ990054) and clear separation in D2-D3 28S sequences (or 96 % identity). Further, the *cox1* sequences of *X. browni* sp. n. and the closest species *X. parasimile, X. simile* (GU222425, Czech Republic) and *X. pachtaicum* (HM921369, Spain) were translated to amino acids and aligned (Fig. 9). The estimated p-distances between *X. browni* sp. n. and the three species were 10.1%, 21.7% and 23.3%, respectively.

In all three phylogeny reconstructions (18S, D2-D3 and *cox1*) *X. parasimile* from Bulgaria was a sister species of *X. browni* sp. n. and both species were part of a well supported clade with other European populations of *X. simile* (Figs 10–12). The recently described species *X. vallense* Archidona-Yuste, Navas-Cortes, Cantalapiedra-Navarrete, Palomares-Rius & Castillo, 2016 presented only with D2-D3 and ITS1 rDNA sequences seems also to be evolutionary very closely related (Figs 11 and 13), however amplifying additional sequences for other molecular markers (e.g. 18S and *cox1*) could help to better clarify its relationships. The position of the new species in the phylogeny trees based on ITS1 and ITS2 sequences was unstable (Figs 13 and 14). The analyses resulted in various tree topologies when using different alignment algorithms and reconstruction methods (ML and BI) and because of the absence of homologous sequences from closely related species. In most cases *X. browni* sp. n. was part of a clade of European *X. americanum*-group species considered as group II in a previous publication (Archidona-Yuste et al. 2016). Due to insufficient number of *nad4* sequences of species belonging to the *X. americanum*-group at NCBI no phylogenetic reconstructions are presented.

**Diagnosis and relationships.** *Xiphinema browni* sp. n. is characterised by a unique combination of traits: slender and medium sized body (1.6–2.41 mm) and odontostyle (73–85 μm), lip region expanded, laterally rounded, separated from the rest of body by a constriction, post-equatorial vulva position (V=52–58 %), symbiotic bacteria present, female tail conical dorsally convex, with narrow rounded to pointed tip, 24–35 μm long, (c=53.4–86.8; c’=1.5–2.1), and specific ribosomal and mtDNA sequences (Table 2). The alpha-numeric codes based on average values (ranges given in parentheses) using the polytomous key by Lamberti et al. (2004) are: A3 (2), B3 (2), C3 (4), D2 (1/3), E2 (3), F2 (1/3), G2, H1, I2 (1/3).

Species having similar morphometrics to *X. browni* sp. n. based on type populations are presented in Table 6. Recently described species *X. parasimile, X. parabrevicolle* Gutiérrez-Gutiérrez, Cantalapiedra-Navarrete, Decraemer, Vovlas, Prior, Palomares-Rius & Castillo, 2012, *X. parapachydermum* Gutiérrez-Gutiérrez, Cantalapie-
Table 4. Pharyngeal characters of females of *Xiphinema americanum* group species studied from different localities.

| Character         | *Xiphinema browni* sp. n. | *Xiphinema pachtaicum* | *Xiphinema penevi* sp. n. | *Xiphinema parasimile* |
|-------------------|---------------------------|------------------------|---------------------------|------------------------|
| Pharynx length (μm) | 278.0±17.9 (236–309) | 272.0±12.1 (247–297) | 271.4±20.1 (234–294) | 274.4±29.2 (291–317) |
| Bulbus length (μm)  | 60±3.48 (53–69) | 59±3.02 (56–67) | 60±2.88 (56–63) | 68.4±2.7 (65–72) |
| Bulbus width (μm)   | 13±1.30 (9–16) | 13±1.21 (11–16) | 14±1.14 (12–15) | 15, 15, 16 (12–14) |
| Bulbus length/Pharynx length (%) | 21.7±1.7 (21.8±1.6) | 21.8±1.6 (22.3±1.3) | 25.6±07 (26.0±07) | 21.8±0.7 (20.7–22.6) |
| DN* (%)           | 17.5±1.9 (15.3–21.1) | 13.1±2.3 (12.7–17.3) | 12.5, 13.0, 9.3, 10.3, 11.4±1.4 (9.9–12.9) | 16.7±3.3 (13.6–18.6) |
| DO* (%)           | 10.9±1.7 (8.8–13.8) | 7.9±3.8 (5.5–15.6) | 11.9±1.8 (8.8–13.3) | 12.1±1.6 (9.9–13.2) |
| SVN1* (%)         | 53.9±1.6 (51.8–55.0) | 55.6, 54.4 (55.6, 54.4) | 60.3 (56.7±2.0) (53.8–58.8) | 55.3–59.7 |
| SVN2* (%)         | 53.2 | 58.7±2.9 (55.4–61.0) | 57.3–60.1 |
| SVO (%)           | 74.2±1.9 (71.4–75.4) | 74.9±3.3 (67.6–76.4) | 68.5, 71.1, 72.0, 74.4 (71.8) | 75.4±2.4 (73.5–79.4) |
| Glandularium ** (μm) | 48.5±1.9 (46–52) | 50.6±2.3 (48–51) | 53, 48, 46, 68, 70, 70 (57–65) | 61.9±3.1 (49.9±1.4) |

Terminology adopted by Loof and Coomans (1972)*; and Andrássy (1998)**.

---

dra-Navarrete, Decraemer, Vovlas, Prior, Palomares-Rius & Castillo, 2012, *X. paratenicuicuts* Gutiérrez-Gutiérrez, Cantalapiedra-Navarrete, Decraemer, Vovlas, Prior, Palomares-Rius & Castillo, 2012, *X. plesiopachtaicum* Archidona-Yuste, Navas-Cortes, Cantalapiedra-Navarrete, Palomares-Rius & Castillo, 2016 and *X. vallense* (Barsi and Lamberti 2004, Gutiérrez-Gutiérrez et al. 2012, Archidona-Yuste et al. 2016) have also been compared. Six of these species have non-European distribution (Table 6) whereas the others were described from and/or found mainly in Europe. *Xiphinema simile* was also included in the table comparing morphometrical data because of the close relationships based on sequence and phylogenetic analyses and its wide distribution in many European countries.

Based both on morphology and molecular data *X. browni* sp. n. is most similar with *X. parasimile*, *X. simile* and *X. vallense*. Morphologically, it can be distinguished from:
Morphological and molecular characterisation, and phylogenetic position of *X. browni*...

Table 5. Measurements of uteri (including ovejector), ovejector and vaginal parts. All measurements in micrometres presented as mean ± standard deviation (range).

| Characters | Anterior uterus | Posterior uterus | Ovejector | Vagina length | Pars distalis vaginae | Pars proximalis vaginae |
|------------|----------------|-----------------|-----------|---------------|-----------------------|------------------------|
|            | Localities     |                 |           |               |                       |                        |
|            | Kurdějov       | 42.1±5.7 (35–54) | 38.9±5.0 (31–43) | 26            | 12.9±1.4 (11–15)     | 6.7±1.1 (5.5–8.5)      | 13.7±0.6 (13–14)       |
|            | n=8            |                 |           |               |                       |                        |                        |
|            | Sokolnice      | 45.5±3.7 (38–46) | 46.0±4.0 (40–49) | 30.5          | 5, 6, 6               | 8.5, 10, 10            |
|            | n=4            |                 |           |               |                       |                        |                        |
|            | Mohyla míru    | 39, 40, 50      | 39, 41.5, 44 | 26, 33        | 12.5±1.0 (11–14)     | 5, 6                   | 10, 10                 |
|            |                |                 |           |               |                       |                        |                        |
|            | Ifrane         | 52.2±9.0 (36–68) | 52.3±4.3 (46–58) | 26            | 8.9±0.3 (8–9)         | 10.6±1.2 (8–13)        |
|            |                |                 |           |               |                       |                        |                        |
|            | X. penevi      | 40, 48          | 42, 49, 50 | 37            | 13, 14, 15           | 9, 9                   | 12, 12                 |
|            |                |                 |           |               |                       |                        |                        |
|            | X. pachnaicum  | Vinogradets     | 33.1±0.4 (30–38) | 31.2±0.7 (24–39) | 29.4±4.3 (26–33.5) | 14.5±1.7 (13–15) | 7.4±0.5 (7–8) | 7.4±0.5 (7–9) |
|            |                | n=13            | n=13      | n=10          | n=17                  | n=15                  | n=19                  |
|            | X. parasimile  | Ralja, Trešna paratypes | 40.0±11.3 (27–46) | -             | -                     | 14.5±1.05 (13–16)     | 7.8±0.8 (7–8.5) | 8.75±0.3 (8–9) |
|            |                | n=3             |           |               |                       |                        |                        |
|            | X. simile      | Srebarma, Bulgaria | 18.8±2.8 (14–21) | 18.5±2.4 (15–20) | 36.3±6.4 (29–41) | 14.8±1.3 (13–16) | 5.8±0.4 (5.5–6) | 9.5±0.9 (8–11) |
|            |                | n=6             | n=6       | n=3           | n=5                   | n=8                   | n=7                   |
|            | Kalimok-Brashlen | 21.8±1.9 (16.5–24) | 21.5±1.8 (19–24) | 43.1±3.1 (36.5–48) | 16.8±0.8 (15–18) | 6.4±0.65 (5.5–7) | 8.6±0.5 (8–10) |
|            |                | n=14            | n=14      | n=12          | n=15                  | n=17                  | n=17                  |
|            | Orlyane        | 21.75±2.2 (17–24) | 22.1±2.3 (19–26) | 43.8±4.2 (36–50) | 16.9±1.1 (15–18) | 6.05±0.6 (5.5–7) | 9.05±0.8 (8–10) |
|            |                | n=7             | n=7       | n=8           | n=11                  | n=10                  | n=10                  |
|            | Kamen bryag    | 23.0±4.8 (18–30) | 24.2±4.15 (19–30) | 47.2±8.9 (37–60) | 15.9±1.8 (13–17) | 6.4±0.6 (6–7) | 9.8±0.8 (9–10) |
|            |                | n=5             | n=5       | n=5           | n=5                   | n=5                   | n=5                   |

Data for *X. parasimile* and *X. simile*, Lazarova et al. (2008).

*X. parasimile* by its different lip region shape (expanded vs not expanded), somewhat longer odontostyle av. 79–83 (73–85) μm vs av. 70 (64–74) μm in the type population, avs. 69–70 (63–74) in Bulgarian populations and avs. 68–70 (67–72) μm in females from Romania (Barsi and Lamberti 2004, Lazarova et al. 2008, Bontă et al. 2012);
Figure 9. Cox1 amino acid sequence alignment of *Xiphinema browni* sp. n. and the closest species *X. parasimile, X. simile* and *X. pachtaicum*.

*X. simile* by its longer odontostyle av. 79–83 (73–85) vs av. 66 (62–69) in type population, avs. 68.5–70 (66–72.5) in other Bulgarian populations, 67.5 (65–70) μm in a population from Bosna and Herzegovina, and avs. 67–68 (61–73) μm in females from the Czech Republic (Lamberti et al. 1983, Barsi and Lamberti 2004, Kumari 2006, Lazarova et al. 2008). However, it should be noted that females from Serbia and Crete (odontostyle 71.5 (66–74) μm and 75–77 μm, respectively) have slightly overlapping values between *X. browni* sp. n. and *X. simile* for this character (Barsi and Lamberti 2004, Tzortzakakis et al. 2014). Further *X. browni* sp. n. differs from *X. simile* in the length and structure of uteri (in the new species separate uteri and ovejector present vs separate uteri not present), different tail shape (conoid vs bluntly conoid), and in the shorter bulbus (53–69 vs 76–92 μm) (Lazarova et al. 2008) (Table 4). Finally, *X. browni* sp. n. develops though 4 vs 3 juvenile stages in *X. simile*; *X. vallense* by the position of amphideal fovea aperture (posterior constriction level vs on the lips); higher lip region (4–7 μm vs 2–3.5 μm); presence of symbiont bacteria in ovaries vs ovaries without symbionts; somewhat higher c’ values (c’=1.8 (1.53–2.07) vs c’=1.6 (1.4–1.7); the different tail shape (dorsoventral depression at hyaline region level not present vs present); shorter spicules in males (29 μm vs 38 μm).

Additionally, *X. browni* sp. n. can be differentiated from:

*X. pachtaicum* by the different vagina shape (bell-shaped vs funnel shaped, (Figs 16, 18) and shorter *pars distalis vaginae*, shorter pharyngeal bulb (53–69 vs 75–80 μm), more posterior location of the dorsal nucleus (DN=13–21% vs 9–10 %) (Table 4), different tail shape in both sexes (conical vs subdigitate). Illustrations of selected features of the closest species *X. pachtaicum, X. parasimile* and *X. penevi* sp. n. are presented in Figs 15–18 for comparison.
Figure 10. Hypothesis of the phylogenetic relationships of Xiphinema browni sp. n., X. parasimile, X. pachtaicum and X. penevi sp. n. based on 18S rDNA inferred from a Bayesian analysis using GTR+G model and Prionchulus punctatus (Cobb, 1917) Andrassy, 1958, Alaimus sp. and Tripylina sp. as an outgroup. Posterior probabilities higher than 0.8 are presented. The sequence of X. browni from Moča was not included due to the shorter length.

X. paratenuicutis in having symbionts in its ovaries vs absent, males rare vs abundant, higher values for c’ (1.8 (1.5–2.1) vs 1.4 (1.2–1.6), different location of dorsal nucleus (DN after beginning of the stronger cuticular lining of the bulbus vs before, see Fig. 1 D1-F and Fig. 2E in Gutiérrez- Gutiérrez et al. 2012);
Figure 11. Hypothesis of the phylogenetic relationships of *Xiphinema browni* sp. n., *X. parasimile*, *X. pachtaicum* and *X. penevi* sp. n. based on 28S rDNA inferred from a Bayesian analysis using GTR+G model and *Longidorus helveticus* Lamberti, Kunz, Grunder, Molinari, De Luca, Agostinelli & Radicci, 2001 and *L. poessneckensis* Altherr, 1974 as an outgroup. Posterior probabilities higher than 0.8 are presented.
Morphological and molecular characterisation, and phylogenetic position of *X. browni*...

**Figure 12.** Hypothesis of the phylogenetic relationships of *Xiphinema browni* sp. n. and *X. parasimile* based on cox1 inferred from a Bayesian analysis using GTR+G model and *X. italiae* Mayl, 1953 and *X. diversicaudatum* (Micoletzky, 1927), Thorne, 1939 as an outgroup. Posterior probabilities higher than 0.8 are presented.

*X. plesiopachtaicum* by the position of the amphidial aperture (posterior vs at the constriction level); somewhat shorter bulbus (avs. 59–60 (53–69) vs av. 73 (60–86); shorter uteri (av. 81 vs av. 138 μm); higher c’ values (c’=1.8 (1.53–2.07) vs c’=1.4 (1.3–1.7); and differently shaped vagina (bell-shaped vs funnel shaped).

For comparison between *X. browni* sp. n. and *X. penevi* sp. n. see below.

**Etymology.** The species is named after Prof Derek JF Brown, an outstanding nematologist, for his significant contributions to the knowledge of plant parasitic nematodes and the development of nematology in Bulgaria.
**Figure 13.** Hypothesis of the phylogenetic relationships of *Xiphinema browni* sp. n. based on ITS1 inferred from a Bayesian analysis using GTR+G model and *X. barense* Lamberti, Roca, Agostinelli, Bleve-Zacheo, 1986, *X. italae* and *X. diversicaudatum* as an outgroup. Posterior probabilities higher than 0.8 are presented.

*Xiphinema penevi* sp. n.
http://zoobank.org/C98CE5B3-9BAE-423C-B887-9BFFFD489798

Figures 15–23

**Measurements.** See Tables 4, 5, 7.

**Description.** *Females.* Body open spiral to C shaped. Thickness of the cuticle at postlabial region 1 μm, 1–1.5 μm at mid-body and 2–2.5 μm at post-anal region, outer cuticle layer not reaching the tail end. Labial region flat anteriorly, laterally rounded, set off from the rest of the body by constriction, 2.5–4 μm high. Amphideal fovea hardly visible, its opening 4 μm in a paratype specimen (40–47 % of the corresponding body width); Distance between first and second guide ring in specimens with retracted odonostyle, 2.5–5 μm long. Odontophore with well developed flanges, 6–9 μm wide, often a small vestigium located in odonthophore area. Pharyngeal characters presented at Table 4. Dorsal nucleus 2.5–3 μm diam., ventrosublateral nuclei well vis-
Morphological and molecular characterisation, and phylogenetic position of *X. browni*

Table 5

| Type material                  |
|--------------------------------|
| The holotype, 7 paratype females and juveniles from all stages are deposited in the nematode collection of the Institute of Biodiversity and Ecosys-

**Figure 14.** Hypothesis of the phylogenetic relationships of *Xiphinema browni* sp. n., *X. parasimile*, *X. pachtaicum* and *X. penevi* sp. n. based on ITS2 inferred from a Bayesian analysis using GTR+G model and *X. italic*ae, *X. diversicaudatum* and *X. vuittenezi* Luc, Lima, Weischer & Flegg, 1964 as an outgroup. Posterior probabilities higher than 0.8 are presented.

---

**Male.** Not found.

**Juveniles.** The scatter diagram based on functional and replacement odontostyle, and body length revealed the presence of four juvenile stages (Fig. 23). As in most species of the *X. americanum*-group there is a gradually decreasing of c' values with successive stages which reflects increasing body width while the tail length is more or less similar in juveniles and adults.

**Type locality and plant association.** Ifrane, Morocco, *Quercus ilex* L. forest.

**Type material.** The holotype, 7 paratype females and juveniles from all stages are deposited in the nematode collection of the Institute of Biodiversity and Ecosys-

**Prerectum indistinct, rectum 21.6±1.8 (19–24) μm, n=8, c 1.3 of corresponding body width. Reproductive system amphidelphic, symbiotic bacteria present in the ovaries. Uteri short, ovejector not developed, only in one specimen a structure resembling ovejector was observed (Table 5); vagina c. 2/3 of the corresponding body width, *pars proximalis vaginae* with well developed wall. Tail conoid, dorsally convex, ventrally slightly concave, gradually narrowing to a pointed tip, two distinct pairs of caudal pores.

**Male.** Not found.

**Juveniles.** The scatter diagram based on functional and replacement odontostyle, and body length revealed the presence of four juvenile stages (Fig. 23). As in most species of the *X. americanum*-group there is a gradually decreasing of c' values with successive stages which reflects increasing body width while the tail length is more or less similar in juveniles and adults.

**Type locality and plant association.** Ifrane, Morocco, *Quercus ilex* L. forest.

**Type material.** The holotype, 7 paratype females and juveniles from all stages are deposited in the nematode collection of the Institute of Biodiversity and Ecosys-

---

**Figure 14.** Hypothesis of the phylogenetic relationships of *Xiphinema browni* sp. n., *X. parasimile*, *X. pachtaicum* and *X. penevi* sp. n. based on ITS2 inferred from a Bayesian analysis using GTR+G model and *X. italic*ae, *X. diversicaudatum* and *X. vuittenezi* Luc, Lima, Weischer & Flegg, 1964 as an outgroup. Posterior probabilities higher than 0.8 are presented.
|                | Non-European species | European species |
|----------------|----------------------|------------------|
| **Species**    |                      |                  |
| *X. penevi* sp. n. | 1.69 (1.5–1.85) | 1.9 (1.7–2.3) |
| *X. bricolensis* | 1.9 (1.7–2.2) | 2 (1.8–2.2) |
| *X. californicum* | 1.6–1.8 | 1.6 (1.4–1.9) |
| *X. citricolum* | 1.6 (1.4–1.7) | 1.6 (1.5–1.7) |
| *X. intermedium* | 1.6 (1.5–1.7) | 1.8 (1.6–1.9) |
| *X. oxycaudatum* | 1.8 (1.6–1.9) | 1.9 (1.5–2.1) |
| *X. tenuicutis* | 61 (57.2–65.0) | 56 (52–62) |
| *X. plesiopachtaicum* | 57.7 (50.8–61.5) | 57 (49–65) |
| **Morphometric data** |                      |                  |
| **Body L**     | 1.69 (1.5–1.85) | 1.9 (1.7–2.3) |
| **a**          | 61 (57.2–65.0) | 56 (52–62) |
| **c**          | 57.7 (50.8–61.5) | 57 (49–65) |
| **c’**         | 1.8 (1.6–1.9) | 1.5 (1.3–1.6) |
| **Vulva [%]**  | 57 (51–61.5) | 52 (50–55) |
| **Odontostyle L** | 77 (72–79) | 87 (85–94) |
| **Tail L**     | 29 (26–32) | 36 (31–41) |
| **Length to GR** | 68 (66–71) | 68 (61–76) |
| **J**          | 9 (8–10) | (6–7) |
| **Lips width** | 8 (8–9) | 11 |
| **Juvenile stages** | 4 | ? |
| **Males (number of VM supplements)** | rare or absent | rare or absent 11 |
| **X. browni** sp. n. | 2.03 (1.8–2.40) | 2.6 (2.5–2.8) |
| **X. microstilum** | 2.6 (2.5–2.8) | 1.88 |
| **X. pachaicum** | 1.89 (1.75–2.26) | 2.01 (1.7–2.2) |
| **X. parasimile** | 2.01 (1.7–2.2) | 1.9 (1.7–2.1) |
| **X. parapachydermum** | 1.78 (1.41–2.0) | 2.0 (1.8–2.2) |
| **X. paratenuicutis** | 1.78 (1.41–2.0) | 2.0 (1.8–2.2) |
| **X. simile** | 69.3 (56.9–81.3) | 86 (77–93) |
| **X. vallense** | 72.3 | 74 (63–88) |
| **Body L**     | 70.5 (61.0–76.1) | 61.1 (51.9–69.7) |
| **a**          | 71 (63–77) | 64 (51.3–67.1) |
| **c**          | 68.9 (61.6–79.1) | 68.8 (58.8–79.9) |
| **c’**         | 67 (61–70) | 60.3 (46.3–75.5) |
| **Vulva [%]**  | 55 (52.3–58.5) | 57.5 (55–59.5) |
| **Odontostyle L** | 79 (75–83) | 74 (68–77) |
| **Tail L**     | 29 (25–33) | 35 (31–39) |
| **Length to GR** | 71 (65–75) | 63 (57–68) |
| **J**          | 7 (6–12) | 10 (7–12) |
| **Lips width** | 8 (8–10) | 9 (9–10) |
| **Juvenile stages** | 4 | 4 |
| **Males (number of supplements)** | rare or absent | frequent 4–5 |

**Notes:**
- VM = Vulval Matrix
- Supplementary data provided for each species, including additional morphometric data.
Morphological and molecular characterisation, and phylogenetic position of X. browni...

Morphological and molecular characterisation, and phylogenetic position of X. browni...

25

tem Research, Sofia, Bulgaria. Other paratypes deposited as follows: 2 females in the USDA Nematode Collection, Beltsville, Maryland, USA; 2 females in the Nematode Collection of the Institute of Plant Protection, Bari, Italy; 1 female in the Wageningen Nematode Collection (WANECO), Wageningen, the Netherlands. Three ribosomal sequences (18S, ITS2 and D2-D3) of X. penevi sp. n. are deposited in GenBank (for accession numbers see Table 2).

**Sequence and phylogenetic analyses.** Sequences for three gene regions were obtained (18S, D2-D3 and ITS2). BLAST at NCBI using any of these sequences as queries revealed highest similarity to X. pachtaicum (99% for 18S, 6 nt difference), two populations of X. incertum Lamberti, Choleva & Agostinelli, 1983 from Spain (99% for D2-D3, 1 and 3 nt difference) and X. pachtaicum (90% for ITS2). In both 18S and D2-D3 phylogeny reconstructions X. penevi sp. n. was part of well supported clades with other species of X. pachtaicum-subgroup (X. pachtaicum, X. parapachydermum for 18S and X. incertum, X. pachtaicum, X. parapachydermum, X. plesiopachtaicum, X. pachydermum Sturhan, 1983 for D2-D3). In the phylogeny reconstruction based on ITS2 sequences, the species grouped with two other X. pachtaicum populations.

**Diagnosis and relationships.** Xiphinema penevi sp. n. is characterised by specific combination of traits: slender body of medium size (1.54–1.85 mm), lip region rounded laterally, flattened anteriorly, separated from the body by a constriction, odontostyle 72–79 μm long, post-equatorial vulva position (V=56–58%), symbiont bacteria present in ovaria, female tail 26–32 μm long (c=50.8–61.2 and c’=1.7–1.9), conoid dorsally convex ventrally slightly concave with pointed tip, and specific ribosomal sequences (18S and ITS2). The alpha-numeric codes based on average values (ranges given in parentheses) using the polytomous key by Lamberti et al. (2004) are: A2, B3, C3 (4), D1 (2), E2, F2 (1), G2, H1, I2 (1). Subsequently described species X. parasimile, X. parabrevicolle, X. parapachydermum, X. paratenenuicu is (Barsi and Lamberti 2004, Gutiérrez-Gutiérrez et al. 2012) X. plesiopachtaicum, X. vallense and X. astarengenae (Archidona-Yuste et al. 2016) and X. browni sp. n. have been also compared. Species having most similar morphometrics with X. penevi sp. n. were: X. pachtaicum, X. plesiopachtaicum, X. browni sp. n., X. vallense and X. parasimile. Due to the close relationships based on phylogenetic analyses X. incertum, X. parapachydermum Sturhan, 1983 and X. parapachydermum were also compared. Xiphinema penevi sp. n. can be differentiated morphologically from:

X. pachtaicum by its shorter odontostyle av. 77 (72–79) vs 83 μm in holotype, av. 84 (78–88.5) in the present study, 89 (85–97) in females from Ethiopia, and distance of oral aperture to guide ring (68 (66–71) vs 78 in holotype, 77 (73–80) in the present study; shorter pharyngeal bulb (65–72 vs 75–80 μm) in the present study; different tail shape (conoid with gradually pointed tip vs conoid, subdigitate), outer cuticular layer not reaching vs reaching tail tip. (Lamberti and Siddiqi 1977, Getaneh et al. 2015); X. plesiopachtaicum by the position of the amphideal fovea aperture (posterior vs at constriction level); its somewhat shorter odontostyle (72–79 vs 77–89 μm) and uteri.
Figure 15. Xiphinema browni sp. n., X. penevi sp. n. X. pachtaicum and X. parasimile. Female: A–C Anterior ends D–F Tail shapes G–J Pharyngeal bulbs A, D, G X. browni sp. n. B, E, I X. penevi sp. n. C, F, J X. pachtaicum H X. parasimile. Scale bars: 25 μm

(104 vs 138 μm); different position of the dorsal nucleus (DN in front of or at the level of DO (beginning of cuticular lining of the bulb) vs DN below the level of DO); different tail shape (ventrally slightly concave vs straight), smaller values for c and larger for c’ ratios (c=50.8–61.5 vs c=62.5–88.7; c’=1.7–1.9 vs c’=1.3–1.7); X. vallense by the position of amphideal fovea (posterior constriction vs on the lips); its shorter body (L=1.69 (1.5–1.85) vs 2.01 (1.83–2.22), different position of dorsal
Figure 16. *Xiphinema browni* sp. n., *X. penevi* sp. n. and *X. pachtaicum*. Female genital system comparison: A, E Posterior genital branch B–D Anterior genital branch A, B *X. browni* sp. n. C *X. penevi* sp. n. D, E *X. pachtaicum*. Scale bars: 25 μm
Figure 17. *Xiphinema browni* sp. n., *X. parasimile*, *X. pachtaicum* and *X. penevi* sp. n. Female and male: A–D Anterior ends E–H Labial region I–K Pharyngeal bulbs L, M, R Male tails N–Q Female tails A, E, I, M, N *X. browni* sp. n. B, F, K, O, R *X. parasimile* C, G, J, L, P *X. pachtaicum* D, H, Q *X. penevi* sp. n. Scale bars: 30 μm (A–D, I–R); 12 μm (E–H).
Morphological and molecular characterisation, and phylogenetic position of *X. browni*...
Figure 19. *Xiphinema penevi* sp. n. Female. **A** Anterior end **B** Amphideal fovea outline **C** Variations in genital system: **C1** Anterior uterus and partim posterior genital branch **C2, C3** Region of vagina and uteri **D** Pharyngeal bulb **E** Tail. Scale bars: 25 μm.

nucleus (DN in front or at the level of DO vs DN below the level of DO); different tail shape (ventrally slightly concave vs straight) smaller values for c and larger values for c’ ratios (c=50.8–61.5 vs c=58.2–86.3; c’=1.7–1.9 vs c’=1.4–1.7), longer hyaline part (8–10 μm vs 6.5–8.5 μm);

*X. browni* sp. n. by its somewhat shorter body (L=1.69 (1.5–1.85) vs 2.03 (1.8–2.40) mm and longer bulbus (65–72 vs 53–69) μm; lower (2.5–4 vs 4–7 μm) and differently shaped lip region (not expanded vs expanded); different location of the dorsal nucleus (DN=9.9–12.9 % vs DN=12.7–21.1 %); different vagina shape (funnel- vs bell-like Figs 16, 18);

*X. parasimile* by its somewhat shorter body (L=1.69 (1.5–1.85) vs 1.99 (1.75–2.26) mm in type population and avs. 1.78 -1.82 (1.56–2.04) in females from Bulgaria), different lip region shape (laterally rounded vs not rounded), the different location of dorsal nucleus (DN 9.9–12.9 % vs 13.6–18.6 %), longer bulbus (65–72
Figure 20. *Xiphinema penevi* sp. n. Female. Variations in: A–D Anterior ends (B-holotype) F, J–L Vagina region G–I Tail shapes. Scale bars: 30 μm (A, B, F–I); 12 μm (C, D, J–L).
Figure 21. *Xiphinema penevi* sp. n. *Juveniles: A–E* Anterior ends of first- to fourth-stage juveniles and female *F–J* Tails of first to fourth juvenile stages and female (*F1* and *F2* tail of first stage juveniles). Scale bar: 25 μm.
Figure 22. Xiphinema penevi sp. n. Juveniles and female: A–D Neck region of first- to fourth-stage juveniles, E Anterior end of female F–J Tails of first to fourth juvenile stages and female. Scale bar: 30 μm.

vs 55.5–63 μm) (Table 4); different vagina shape (funnel vs bell-like), structure of uteri (ovejector not present vs ovejector and separate uteri present) and length of uterus (36–68 vs 27–46 μm in type population and 27–39 μm in population from Bulgaria (Table 5); shorter tail (av. 29 (26–32) vs 33 (30.3–37.1) in the type population and 30–32 (27–35) in females from Bulgaria, c’=1.8 (1.6–1.9) vs 2.02 (1.79–2.28) in the type population and 2.0 (1.7–2.3) in females from Bulgaria) (Barsi and Lamberti 2004, Lazarova et al. 2008); X. incertum by its different tail shape (elongate conoid vs bluntly conoid, ventrally slightly concave vs straight) and larger c’ values (c’=1.8 (1.6–1.9) vs c’=1.5 (1.4–1.7) in type material and 1.2 (0.9–1.3) in specimens from Spain, larger a values (a=61 (57–2-65) vs a=57 (56–58) in type population and a=49.7 (44.6–52.5) in the population from Spain and different vagina shape compared with females from Spain, this character not described for the type population (Lamberti et al. 1983, Gutiérres-Gutiérres et al. 2012); X. pachydermum by its shorter body (L=1.69 (1.5–1.85) mm vs 2.24 (2.08–2.44) mm), different location of dorsal nucleus (DN=10–13 % vs DN=15–20%), presence of symbiotic bacteria in ovaria vs not present; males occurrence (not present vs abundant);
Table 7. Morphometrics of *Xiphinema penevi* sp. n. (females and juveniles) from *Q. ilex* Morocco. All measurements except ratios in micrometres given as mean ± standard deviation (range).

| Characters                                      | Females          | Juveniles       |
|------------------------------------------------|------------------|-----------------|
|                                                | Holotype 12      | Paratypes 2     | J1 2   | J2 8   | J3 5   | J4 5   |
| n                                              | 1726             | 664, 602        | 777.6±37 | 1049.0±54 | 1318±38 |
| L                                              | (1532–1846)      | (702–816)       | (988–1126) | (1292–1384) |
| a                                               | 61.0±2.6 (57.2–65.0) | 40.6, 38.0      | 43.1±2.4 | 48.2±1.5 | 54.4±3.6 |
| b                                               | 6.1±1.1 (5.0–7.0) | 3.8, 3.6        | 4.0±0.2  | 4.5±0.1  | 5, 6   |
| c                                               | 57.7±3.9 (50.8–61.5) | 20.9, 22.4      | 24.7±2.5 | 33.4, 34.1 | 40.8±2.6 |
| c'                                              | 1.8±0.1 (1.6–1.9) | 2.9, 2.6        | 2.7±0.3  | 2.2, 2.3  | 2.1±0.2 |
| V (%)                                           | 57.1±0.6 (55.9–58.1) |                | 54.5±1.1 | 63.3±2.0 |        |
| G1(%)                                           | 11.2±0.5 (10.9–12.1) |                | 66.2±1.4 | 75.0±1.6 |        |
| G2(%)                                           | 12.3±3.2 (9.2–19.5) |                | 37.6±1.9 | 43.9±1.6 |        |
| Odontostyle                                     | 75 (72–79)       | 36.5, 37        | 43.8±1.0 | 54.5±1.1 | 63.3±2.0 |
| Replacement odontostyle                         | 43, 46           |                | 56.5±1.8 | 66.2±1.4 | 75.0±1.6 |
| Odontophore                                     | 50 (44–50)       | 28              | 33.6±1.2 | 37.6±1.9 | 43.9±1.6 |
| Oral aperture to guide ring                     | 71 (66–71)       | 30.5, 33        | 38.5±1.2 | 49.0±1.4 | 55.6±3.9 |
| Tail length                                     | 31 (26–32)       | 32, 27          | 32.0±2.7 | 32, 33  | 31.9±2.0 |
| Length of hyaline part                          | 9 (8–10)         | 4, 4            | 4.3±0.7  | 6, 6   | 6.9±0.7  |
| - lip region                                    | 9 (8–9)          | 7, 7            | 7.1±0.5  | 7.3±0.7  | 7.8±0.2  |
| - at guiding ring                               | 21 (20–21)       | 12, 13          | 14.5±0.6 | 16.2±0.7 | 18.6±0.9 |
| - base of pharynx                              | 25 (22–26)       | 15, 15          | 16.8±1.2 | 19.3±0.8 | 22.3±1.0 |
| - at mid body/at vulva                          | 28 (25–31)       | 16, 16          | 18.1±1.5 | 21.8±1.3 | 24.0±1.7 |
| - at anus                                       | 16 (15–17)       | 11, 10          | 11.9±0.8 | 25.2±22.1 | 15.4±0.5 |
| - at beginning of hyaline part                  | 7 (7–8)          | 5, 4            | 4.2±0.3  | 5, 5   | 6.3±0.5  |
Morphological and molecular characterisation, and phylogenetic position of X. browni...

Figure 23. Scatter plot of odontostyle (■) and replacement odontostyle (□) against body length of Xiphinema penevi sp. n. juveniles and females from Morocco.

X. parapachydermum by its different tail tip (not so acute and not with dorso-ventral depression) and in having symbionts in its ovaries vs absent, males occurrence (not present vs abundant).

Etymology. The new species is named after Dr Lyubomir Penev, an internationally recognised publisher and authority in entomology and ecology as acknowledgement of his invaluable help and support provided to one of the authors (VP) in her research activities.

Xiphinema pachtaicum (Tulaganov, 1938) Kirjanova, 1951
Figures 15–18

Measurements. Tables 3–6.

Note. Xiphinema pachtaicum has been recorded from Bulgaria and data on its morphology are available in previous studies (Lamberti et al. 1983; Peneva and Choleva 1992); here we present additional mornhometric data only for the population from Balgarene together with illustrations, LM micrographs and sequence data (Table 2). It is common and associated with a wide spectrum of cultivated and wild plants (Lamberti and Siddiqi 1977).
**Xiphinema parasimile** Barsi & Lamberti, 2004
Figures 15, 17, 18

Morphometric data and detailed description of *X. parasimile* from Bulgaria are reported previously (Lazarova et al. 2008). For the Vinogradets population two ribosomal and one mitochondrial DNA sequences were obtained (Table 2). *Xiphinema parasimile* has a limited distribution in Bulgaria (Lazarova et al. 2008).

**Sequence and phylogenetic analyses**

Three rDNA sequences were obtained for the Bulgarian *X. pachtaicum* population (18S, D2-D3 and ITS2) with BLAST showing identity or very high similarity to other *X. pachtaicum* populations available at NCBI (100% for 18S, 99/100% for D2-D3 and 98% for ITS2). Further, the DNA sequences of *X. parasimile* from Vinogradets (18S, D2-D3 and cox1) showed highest similarity to *X. simile* from Serbia (99% for 18S), various other populations of *X. simile* and *X. opisthobysterum* (88%, D2-D3) and 78% two cox1 sequences – *X. pachtaicum* from the Czech Republic (GU222424) and *X. simile* from Slovakia (AM086708). The first one is the previously published sequence of *X. browni* sp. n. identified as *X. pachtaicum* (Kumari et al. 2010b). The D2 28S rDNA region was further compared to the Serbian population of *X. parasimile* (D2 part of sequences AM490214, AM490217, Barsi and De Luca 2008) and the alignment showing the different nucleotides is presented (Fig. 24). The p-distance calculated for D2 part only was 1.8–2.1% that might indicate that *X. parasimile* population from Bulgaria could represent a cryptic species.

Based on the phylogenetic analyses performed (Figs 11–15) both new species described are members of two well-supported species complexes – *X. simile* and *X. pachtaicum*. The first subgroup includes *X. simile*, *X. parasimile*, *X. browni* sp. n. and probably *X. vallense*. All occur in Europe and *X. simile* has also been reported from Central Africa (Liškova and Brown 1996, Coomans and Heyns 1997, Barsi and Lamberti 2004, Kumari 2006, Repasi et al. 2008, Lazarova et al. 2008, Bontă et al. 2012). Whether some of these records represent *X. simile* or closely related species requires new investigations using morphological discrimination and molecular markers. So far, *X. parasimile* has been recorded from the Balkan region (Barsi and Lamberti 2004, Lazarova et al. 2008, Bontă et al. 2012). *Xiphiinema browni* sp. n. (previously reported as *X. pachtaicum*) seems to occur in central European countries. The second group of closely related species consists of *X. pachtaicum*, *X. penevi* sp. n., *X. incertum*, *X. parapachydermum*, *X. plesiopachtaicum*, *X. astaregiense* and *X. pachydermum*. Again, one of these species (*X. pachtaicum*) has a much wider distribution in Europe, Asia and Africa (Lamberti and Siddiqi 1977, Fadaei et al. 2003, Getaneh et al. 2015). *Xiphiinema incertum* has been reported from Bulgaria, Serbia, Croatia and Spain, all other species have limited distributions – *X. plesiopachtaicum*, *X. pachydermum*, *X. parapachydermum*, *X. astaregiense*, reported only from Spain, the latter three species being amphimictic, and *X. penevi* sp. n. so far found only in north-western Africa (Sturhan 1983, Lamberti et al. 1983, Barsi and Lamberti 2002, Gutiérrez- Gutiérrez, 2012).
Based on a hierarchical cluster analysis of morphometrics Lamberti and Ciancio (1993) distinguished five species subgroups, among them the X. pachtaicum-subgroup (IV) consisted of 8 species with five being described from Europe (X. fortutium Roca, Lamberti & Agostinelli, 1987, X. incertum, X. madeirensis, X. pachydermum and X. simile), one from North America (X. utahense Lamberti & Bleve-Zacheo, 1979), and one from Asia (X. opisthohystera). Our analyses using ribosomal and mitochondrial DNA sequences currently available in GenBank and the two new species described in this study supports the delimitation of the "X. pachtaicum-subgroup", however it also includes X. incertum, X. pachtaicum, X. pachydermum and the recently described species X. parapachydermum, X. astaregiense, X. plesiopachtaicum and X. penevi sp. n. Phylogenetic reconstructions showed that X. madeirensis, X. opisthobystera, X. simile and X. utahense are not part of this group, for X. fortutium no sequences are available. These results are in line with the findings of other recent studies on the X. americanum-group (Gutiérrez-Gutiérrez et al. 2012, Archidona-Yuste et al. 2016). Xiphinema simile (presented by two types of sequences for populations from Serbia and the Czech Republic in 18S rDNA and cox1 trees), X. parasimile and X. browni sp. n. formed a separate subgroup outside the X. pachtaicum-subgroup, so far consisting only of parthenogenetic species. Therefore we proposed this clade to be referred as the X. simile-subgroup. The recently described species X. vallense seems also evolutionary very closely related to this subgroup because of its high morphometric and DNA similarity, however amplifying additional sequences for other molecular markers (e.g. 18S and cox1) could help to clarify its relationships.

Figure 24. Sequence alignment of D2 28S rDNA region of Xiphinema parasimile from Bulgaria (KU250156) and Serbia (AM490214 and AM490217).
Acknowledgements

The authors are grateful to Dr Marta Lišková (Slovak Academy of Sciences) for providing nematodes and Dr Milka Elshishka (IBER-BAS) for her help. We are obliged also to Prof Derek JF Brown for linguistic improvement of the manuscript. The work was supported by the Ministry of Agriculture of the Czech Republic, Project number MZe–RO0414 and ANIDIV-2 project supported by the BAS.

References

Andrássy I (1998) The genus Boreolaimus gen. n. and its six species (Dorylaimida: Qudsianematidae), nematodes from the European Arctic. Fundamental and Applied Nematology 21: 553–567.

Archidona-Yuste A, Navas-Cortes JA, Cantalapiedra-Navarrete C, Palomares-Rius JE, Castillo P (2016) Cryptic diversity and species delimitation in the Xiphinema americanum-group complex (Nematoda: Longidoridae) as inferred from morphometrics and molecular markers. Zoological Journal of the Linnean Society 176: 231–265. doi: 10.1111/zoj.12316

Barsi L, De Luca F (2008) Morphological and molecular characterisation of two putative Xiphinema americanum-group species, X. parasimile and X. simile (Nematoda: Dorylaimida) from Serbia. Nematology 10: 15–25. doi: 10.1163/156854108783360212

Barsi L, Lamberti F (2004) Xiphinema parasimile sp. n. from Serbia and X. simile, first record from Bosnia and Herzegovina (Nematoda, Dorylaimida). Nematologica Mediterranea 32: 101–109.

Blaxter ML, De Ley P, Garey JR, Liu LX, Scheldeman P, Vierstraete A, Vanfleteren JR, Mackey LY, Dorris M, Frisse LM, Vida JT, Thomas WK (1998) A molecular evolutionary framework for the phylum Nematoda. Nature 392: 71–75. doi: 10.1038/32160

Bontă (Groza) M, Peneva V, Lazarova S, Roşca I (2012) Diversity of Xiphinema species (Nematoda: Dorylaimida) associated with different crops in Romania. Scientific Papers. Series A. Agronomy LV: 387–390.

Cherry T, Szalanski AL, Todd TC, Powers TO (1997) The internal transcribed spacer region of Belonolaimus (Nemat, Belonolaimidae). Journal of Nematology 29: 23–29.

Coomans A, Huys R, Heyns J, Luc M (2001) Character analysis, phylogeny, and biogeography of the genus Xiphinema Cobb, 1913 (Nematoda: Longidoridae). Annales du Musée Royal de l’Afrique Centrale (Zoologie), Tervuren. Belgique 287: 1–289. doi: 10.1163/005025997X00021
Morphological and molecular characterisation, and phylogenetic position of X. browni...
Huelsenbeck JP, Ronquist F (2001) MRBAYES: Bayesian inference of phylogenetic trees. Bioinformatics 17: 754–755. doi: 10.1093/bioinformatics/17.8.754

Kumari S (2006) *Xiphinema simile* (Nematoda: Longidoridae) in the Czech Republic and a note on other *Xiphinema* species. Helminthologia 43: 43–50. doi: 10.2478/s11687-006-0009-x

Kumari S, Cesare AD (2013) Nicotinamide dehydrogenase subunit 4 analysis of *Xiphinema diversicaudatum* and *Xiphinema simile* (Nematoda: Longidoridae). European Journal of Plant Pathology 136: 803–810. doi: 10.1007/s10658-013-0208-5

Kumari S, Decraemer W, De Luca F (2010a) Molecular characterization of *Xiphinema brevicollum* (Nematoda: Longidoridae) from the Czech Republic. European Journal of Plant Pathology 128: 243–250. doi: 10.1007/s10658-010-9651-8

Kumari S, Decraemer W, De Luca F, Tiefenbrunner W (2010b) Cytochrome c oxidase subunit 1 analysis of *Xiphinema diversicaudatum*, *X. pachtaicum*, *X. simile* and *X. vuittenezi* (Nematoda, Dorylaimida). European Journal of Plant Pathology 127: 493–499. doi: 10.1007/s10658-010-9614-0

Kumari S, Decraemer W, Traversa D, Lišková M (2009) Molecular and morphological delineation of *Longidorus poessneckensis* Altherr, 1974 (Nematoda: Dorylaimida). European Journal of Plant Pathology 123: 125–137. doi: 10.1007/s10658-008-9348-4

Kumari S, Polák J, Choutka R (2005) Plant-parasitic nematodes of the genus *Xiphinema* (Nematoda: Longidoridae) in the vineyards of the Czech Republic. Nematology 7(1): 81–93. doi: 10.1163/1568541054192117

Kumari S, Subbotin SA (2012) Characterization of *Longidorus helveticus* (Nematoda: Longidoridae) from the Czech Republic. European Journal of Plant Pathology 133: 923–933. doi: 10.1007/s10658-012-9959-7

Lamberti F, Choleva B, Agostinelli A (1983) Longidoridae from Bulgaria (Nematoda, Dorylaimida) with description of three new species of *Longidorus* and two new species of *Xiphinema*. Nematologia Mediterranea 11: 49–72.

Lamberti F, Ciancio A (1993) Diversity of *Xiphinema americanum*-group species and hierarchical cluster analysis of morphometrics. Journal of Nematology 25: 332–343.

Lamberti F, Molinari S, Moens M, Brown DJF (2000) *The Xiphinema americanum* group. I. Putative species, their geographical occurrence and distribution, and regional polytomous identification keys for the group. Russian Journal of Nematology 8: 65–84.

Lamberti F, Siddiqi MR (1977) *Xiphinema pachtacium* (=*X. mediterraneum*). CIH Descriptions of Plant-parasitic Nematodes No. 94. CAB International, Wallingford, UK.

Lamberti FS, Hockland A, Agostinelli, Moens M, Brown DJF (2004) *The Xiphinema americanum* group. III. Keys to species identification. Nematologia Mediterranea 32: 53–56.

Lazarova S, De Luca F, Peneva VK (2008) On two closely related species of the *Xiphinema americanum*-group: *X. simile* Lamberti, Choleva et Agostinelli, 1983 and *X. parasimile* Barsi et Lamberti, 2004 (Longidoridae), with a description of the male of *X. parasimile*. ZooKeys 3: 29–50. doi: 10.3897/zookeys.3.26

Lazarova SS, Malloch G, Oliveira CMG, Hübschen J, Neilson R (2006) Ribosomal and mitochondrial DNA analyses of *Xiphinema americanum*-group populations. Journal of Nematology 38: 404–410.
Lišková M, Brown DJF (1996) Taxonomic validity and ecological relations of *Xiphinema pachtaicum* and *X. simile* (Nematoda: Dorylaimida), two members of the *X. americanum* group occurring in Slovakia. Helminthologia 33: 137–142.

Loof PAA, Coomans A (1972) The oesophageal gland nuclei of Longidoridae (Dorylaimida). Nematologica 18: 213–233. doi: 10.1163/187529272X00458

McFarlane SA, Neilson R, Brown DJF (2002) Nematodes. In: Plumb RT (Ed.) Advances in botanical research. Academic Press, San Diego, CA, USA, 169–198. doi: 10.1016/s0065-2296(02)36063-4

Meza P, Aballay E, Hinrichsen P (2011) Molecular and morphological characterisation of species within the *Xiphinema americanum*-group (Dorylaimida: Longidoridae) from the central valley of Chile. Nematology 13: 295–306. doi: 10.1163/138855410X00458

Neilson R, Ye W, Oliveira CMG, Hübschen J, Robbins RT, Brown DJF, Szalanski AL (2004) Phylogenetic relationships of selected species of Longidoridae (Nematoda: Longidoridae) from North America inferred from 18S rDNA gene sequence data. Helminthologia 41: 209–215.

Nunn GB (1992) Nematode molecular evolution: an investigation of evolutionary patterns among nematodes based upon DNA sequences. Ph.D. Dissertation, University of Nottingham, Nottingham, UK

Oliveira CMG, Hubschen J, Brown DJF, Ferraz LCCB, Wright F, Neilson R (2004) Phylogenetic relationships among *Xiphinema* and *Xiphidorus* nematode species from Brazil inferred from 18S rDNA sequences. Journal of Nematology 36: 153–159.

Peneva V, Choleva B (1992) Nematodes of the family Longidoridae from forest nurseries in Bulgaria. II. Genus *Xiphinema* Cobb, 1913. Khelmintologiya 32: 47–66.

Repasi V, Agostinelli A, Nagy P, Coiro MI, Hecker K, Lamberti F (2008) Distribution and morphometrical characterization of *Xiphinema pachtaicum, X. simile* and *X. brevicollum* from Hungary. Helminthologia 45: 96–102. doi: 10.2478/s11687-008-0018-z

Roca F, Lamberti F, Agostinelli A (1987) *Xiphinema fortuitum* a new longidorid nematode from Italy. Nematologica Mediterranea 15: 219–223.

Ronquist F, Teslenko M, van der Mark P, Ayres DL, Darling A, Höhna S, Larget B, Liu L, Suchard MA, Huelsenbeck JP (2012) MrBayes 3.2: Efficient Bayesian Phylogenetic Inference and Model Choice Across a Large Model Space. Systematic Biology 61: 539–542. doi: 10.1093/sysbio/sys029

Sakai H, Takeda A, Mizukubo T (2011) First report of *Xiphinema brevicolle* Lordello et Costa, 1961 (Nematoda, Longidoridae) in Japan. ZooKeys 135: 21–40. doi: 10.3897/zookeys.135.1716

Sakai H, Takeda A, Mizukubo T (2012) Intra-specific variation of *Xiphinema brevicolle* Lordello et Costa, 1961 (Nematoda: Longidoridae) in Japan. Nematological Research 42: 1–7. doi: 10.3725/jijn.42.1

Seinhorst JW (1959) A rapid method for the transfer of nematodes from fixative to anhydrous Glycerin. Nematologica 4: 67–69. doi: 10.1163/187529272X00381

Sela I, Ashkenazy H, Katoh K, Papko T (2015) GUIDANCE2: accurate detection of unreliable alignment regions accounting for the uncertainty of multiple parameters. Nucleic Acids Research 43 (Web Server issue): W7–W14. doi: 10.1093/nat/gkq443
Stanton JM, McNicol CD, Steele V (1998) Non–manual lysis of second–stage *Meloidogyne* juveniles for identification of pure and mixed samples based on the polymerase chain reaction. Australasian Plant Pathology 27: 112–115. doi: 10.1071/AP98014

Sturhan D (1983) Description of two new *Xiphinema* species from Portugal with notes on *X. pachtaicum* and *X. opisthohysterum* (Nematoda, Longidoridae). Nematologica 29(3): 270–283. doi: 10.1163/187529283X00032

Tamura K, Stecher G, Peterson D, Filipski A, Kumar S (2013) MEGA6: Molecular Evolutionary Genetics Analysis Version 6.0. Molecular Biology and Evolution 30: 2725–2729. doi: 10.1093/molbev/msr121

Tzortzakakis EA, Archidona-Yuste A, Cantalapiedra-Navarrete C, Nasiou E, Lazanaki MS, Kabourakis EA, Palomares-Rius JE, Castillo P (2014) Integrative diagnosis and molecular phylogeny of dagger and needle nematodes of olives and grapevines in the island of Crete, Greece, with description of *Xiphinema cretense* sp. n. (Nematoda, Longidoridae). European Journal of Plant Pathology 140: 563–590. doi: 10.1007/s10658-014-0488-4

Vrain TC, Wakarchuk DA, Levesque AC, Hamilton RI (1992) Intraspecific rDNA restriction fragment length polymorphism in the *Xiphinema americanum* group. Fundamental and Applied Nematology 15: 563–573.