Calcaridorylaimus castaneae sp. n. (Nematoda, Dorylaimidae) from Bulgaria with an identification key to the species of the genus

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¶ http://zoobank.org/415009FF-DE2E-4A02-83D4-66125561AFA7  
‖ http://zoobank.org/4C865CC6-DED3-4DCE-AAA1-71A51922752B  
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Abstract

An unknown species belonging to the genus Calcaridorylaimus Andrássy, 1986 was collected from the litter of broadleaf forests dominated by Castanea sativa Mill. and mixed with Quercus daleshampii Ten. and Fagus sylvatica L. on Belasitsa Mountain, south-western Bulgaria. Calcaridorylaimus castaneae sp. n. is characterised by its long body (1.4–2.1 mm), lip region practically not offset, vulva transverse, short odontostyle (14.5–16 μm) and tail (75.5–110.5 μm, c=14.7–23.6; c'=2.9–4.4) in females and 38–46 μm long spicules with small spur before their distant end in males. It is most similar to C. andrassyi Ahmad & Shaheen, 2004, but differs in having transverse vs pore-like vulva and shorter spicules (38–46 μm vs 52–57 μm). An identification key to the species of the genus Calcaridorylaimus is proposed. Phylogenetic analyses were performed on 18S and D2-D3 expansion domains of 28S rRNA genes by Neighbor-Joining, Maximum Likelihood and Bayesian Inference methods. The phylograms inferred from 18S sequences showed closest relationships...
of the new species with some species belonging to the genus *Mesodorylaimus*. However, insufficient molecular data for members of both genera do not allow the phylogenetic relationships of *Calcaridorylaimus* and the new species described herein to be elucidated.

**Keywords**
Taxonomy, morphology, 18S and D2-D3 rRNA genes, compendium

**Introduction**

During an ecological study of chestnut forests on Belasitsa Mountain (2003-2005) an undescribed species belonging to the genus *Calcaridorylaimus* Andrásy, 1986 was recovered. The genus *Calcaridorylaimus* is represented by nine species worldwide: *C. calcarifer* Andrásy, 1986, *C. promissus* Andrásy, 1986, *C. ruwenzorii* (De Coninck, 1935) Andrásy, 1986, *C. signatus* (Loof, 1975) Andrásy, 1986, *C. similimus* Andrásy, 1986, *C. sirgeli* Heyns & Meyer, 1995, *C. arcticus* Gagarin, 1997, *C. andrassyi* Ahmad & Shaheen, 2004 and *C. beatus* Andrásy, 2011. The genus is distributed mainly in the southern hemisphere: three species occur in Africa, two in South America and one each in Antarctic, Central America, and Europe, and *C. promissus* was recorded from Australia and Alaska, North America (Andrássy 1986, 2003) (Fig. 1). The most characteristic features of these species are the shapes and structures of the spicules which are provided with a small spur before the distal tip. The new species is described based on both morphological and molecular data.

**Materials and methods**

**Sample collection**

The litter samples were collected in 2003 by the last author (VP) from three sites on Belasitsa Mountain representing different types of broadleaf forests dominated by *Castanea sativa* Mill. mixed with *Quercus daleshampii* Ten. and *Fagus sylvatica* L. (Forest Management Plan database, sub-compartments 104g, 140b and 146a). Subsequently, on 17.10.2012 new litter samples were collected by Dr Michaela Ilieva from one of these sites, sub-compartment 140b, in order to obtain fresh material for molecular studies. Nematodes were recovered from the litter using the Baermann funnel method. They were killed by heat (65 °C), fixed in TAF (Triethanolamine-formalin, Courtney et al. 1955), and processed to anhydrous glycerine (Seinhorst 1959). Drawings were prepared using an Amplival 30-G048b and a drawing tube RA-6V42. Photographs were taken using an Axio Imager M2-Carl Zeiss compound microscope equipped 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 programme Digittrak 1.0f (Philip Smith, Scottish Crop Research Institute, Dundee, UK).

DNA extraction, amplification and sequencing

Genomic DNA was extracted from two female and two male worms using a standard nematode digestion protocol (Holterman et al. 2006). Two overlapping fragments of 18S rRNA genes (~1600 bp) were amplified from each specimen using primer sets 988F (5’-CTC AAA GAT TAA GCC ATG C-3’) and 1912R (5’-TTT ACG GTC AGA ACT AGG G -3’) for the first fragment, and 1813F (5’-CTG CGT GAG AGG TGA AAT-3’, 2646R 5’-GCT ACC TTG TTA CGA CTT TT-3’) for the second fragment (Holterman et al. 2006). The D2/D3 expansion segments of the 28S rRNA gene (~900 bp) were additionally amplified from all specimens using the primers D2A (5’-ACA AGT ACC GTG AGG GAA AGT TG-3’) and D3B (5’-TCG GAA GGA ACC AGC TAC TA-3’) (De Ley et al. 1999). Each PCR reaction was performed under the following conditions: initial denaturation 94 °C for 5 min; 40 cycles (denaturation 94 °C for 30 sec; primer annealing 50 °C for 30 sec; extension 72 °C for 1 min), and final extension 72 °C for 10 min. PCR products were visualized on 1% agarose gel with GreenSafe (NZYtech) under visible and UV light. Fragment size was determined using GeneRuler™ 100 bp Ladder Plus (Ferments, Thermo Scientific). The amplified products were sequenced by Eurofins MWG Operon.
Sequence and phylogenetic analysis

The sequences of the new species have been deposited in GenBank with the accession numbers KF717497 and KF717498 for the 18S and the D2-D3 rRNA genes, respectively. A BLAST (Basic Local Alignment Search Tool) search at NCBI (National Center for Biotechnology Information) was performed using the obtained sequences as queries to confirm their nematode origin and to identify the most closely related nematode sequences. The sequences revealing a similarity up to 97% and 85% with nematodes from various Dorylaimida families were included in the phylogenetic analyses of 18S and D2-D3 regions, respectively (Griffiths et al. 2006; Holterman et al. 2006; Meldal et al. 2007; Lesaulnier et al. 2008; Pedram et al. 2010; Pedram et al. 2011; Álvarez-Ortega and Peña-Santiago 2012a; 2012b; Donn et al. 2012; Álvarez-Ortega et al. 2013). The Multiple Sequence Alignments (MSA) of both datasets were performed using the Clustal Omega tool (Sievers et al. 2011) via the EBI webserver: http://www.ebi.ac.uk/Tools/msa/clustalw2/. Subsequently, the MSAs were manually optimised and trimmed using MEGA 5 (Tamura et al. 2011). *Eudorylaimus* sp. (family Qudsianematidae) was used as an outgroup taxon for both 18S and D2-D3 rDNA sequence datasets (accession numbers AY284800 and AY593037, respectively; Holterman et al. 2008). The phylogenetic reconstructions of three datasets D2-D3, complete and partial 18S rDNA were performed using Neighbor Joining (NJ), Maximum Likelihood (ML) and the Bayesian Inference (BI) algorithms and implemented in MEGA 5.0 and MrBayes v. 3.2.1 (Huelsenbeck and Ronquist 2001, Tamura et al. 2011, Ronquist et al. 2012). The NJ phylogenetic inferences were performed under the following settings: Maximum Composite Likelihood method for computing evolutionary distances; Gamma distributed rates among sites, estimated values set up to 0.3429 (D2-D3) and 0.05 (18S rDNA); 2000 bootstrap replications. A total of 755 and 1593 positions in the final datasets were used for both analyses, respectively. General Time Reversible model (GTR) plus Gamma distribution rates (G) and 1000 bootstrap replications were used as ML analyses settings for all datasets. The Bayesian MCMC tree searches were conducted using MrBayes 3.2.1. Each analysis was run for 10, 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, and posterior probabilities (PP) of branch support. The evolutionary models for nucleotide substitutions were set up as for ML analyses. 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 v1.4.0 graphical viewer (http://tree.bio.ed.ac.uk/software/figtree/). Posterior probabilities values of ≥0.80 (BI) and bootstrap values of ≥70 (NJ and ML) were considered as credible support values for nodes.
Taxonomy

*Morphology*

**Calcaridorylaimus castaneae** sp. n.

http://zoobank.org/9BF1D302-1986-47C5-B0D8-2F5DC75BD6D2

http://species-id.net/wiki/Calcaridorylaimus_castaneae

Figs 2–6

**Measurements.** See Table 1.

**Description. Female.** Body slender, more or less curved ventrally. Cuticle ca 2 μm at anterior part of neck, 3 μm thick at midbody, 4.5–5.5 μm at postanal region; outer layer with fine transverse striae. Lateral chord ca 1/4 of body width. Three dorsal, two ventral and three lateral pores are observed in the spear area. Lip region practically not offset, lip region width ca 20% of its height. Lips partly fused, labial papillae slightly protruding. Body at proximal end of pharynx 3–4 times the width of the lip region diameter. Amphidial aperture 5–6 μm wide or about half the lip region width. Odontostyle 1.2–1.3 times the lip region diameter, aperture occupying 35–40% of its length. Odontophore simple 1.3–1.7 times odontostyle length. Guiding ring at 8.5–9 μm from anterior end. Nerve ring surrounding the pharynx at 36–39% of neck length from head end. Hemizonid and conspicuous excretory pore observed in the nerve ring region. Pharyngeal characters (five females and five males): pharynx beginning to widen at 56–60% and attaining its full width at 61–64% of neck length from anterior end. DO (for terminology see Loof and Coomans 1970) lying near the point where the pharynx attains its full width; DO–DN 8–13 μm. The two S 1 N lying at a small distance behind the middle of the distance DN–S 2 N, the anterior one (S 1 N 1 ) smaller, ca 2 μm diam; S 1 N 2 comparatively large and distinct, 3–4 μm diam. DN nucleolus 4–4.5 μm diam.; S 2 N 2–3 μm diam. (DN>S 1 N 2 >S 2 N). Locations (%):

DO=60–64    S 1 N 1 =78–81    S 2 N=91–92    K=73–84
DN=62–66    S 1 N 2 =82–85    S 2 O=93–95    K’=78–85
DO–DN=2.2–3.8    S 1 N 1 –S 1 N 2 =2.9–4.8.

Cardia conoid, variable in length, microvilli visible only here. Genital system didelphic-amphidelphic, ovaries reflexed, reaching rarely the vulva level. Oviducts and ovaries very long compared to uteri. Uteri not differentiated, short, anterior 96-125 μm and posterior 95-135 μm long, in one female with no sperm inside the genital system uteri shorter, 83 μm each, sphincter between uterus and pars dilatata oviductus well developed. Sperm present in uteri and pars dilatata oviductus. Synchronous uterine eggs 1–3, measuring 72–78 × 32–44 μm. Vulva transverse. Vagina extending to 60–70% inwards: pars proximalis 9–13 μm wide, 14.5–19.5 μm long, pars refringens more or less rounded trapezoid, 10–12.5 μm wide, 4–6 μm deep; pars distalis approx. 1.5 μm (terminology following De Ley et al. 1993). Peculiar tongue-like valve present at intestine-prerectum junction. Rectum 1.3–1.5, prerectum 3–4 times anal body width long, respectively. Tail first conoid, then more or less uniformly tapering to a
Table 1. Morphometrics of *Calcaridorylaimus castaneae* sp. n., females and males, from Belasitsa Mountain. All measurements, unless indicated otherwise, in μm and in the form: mean ± SD (range).

<table>
<thead>
<tr>
<th>Characters</th>
<th>Holotype</th>
<th>Paratypes</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Females</td>
<td>Males</td>
</tr>
<tr>
<td>n</td>
<td>40</td>
<td>20</td>
</tr>
<tr>
<td>L (mm)</td>
<td>1.7 ± 0.11 (1.5–2.0)</td>
<td>1.6 ± 0.08 (1.4–1.8)</td>
</tr>
<tr>
<td>a</td>
<td>35.7</td>
<td>36.6 ± 2.4 (32.5–42.7)</td>
</tr>
<tr>
<td>b</td>
<td>4.9</td>
<td>5.0 ± 0.3 (4.5–5.7)</td>
</tr>
<tr>
<td>c</td>
<td>19.4</td>
<td>18.8 ± 0.4 (16.0–22.0)</td>
</tr>
<tr>
<td>c'</td>
<td>3.9</td>
<td>3.6 ± 0.4 (2.9–4.4)</td>
</tr>
<tr>
<td>V/T %</td>
<td>50.7</td>
<td>51.6 ± 1.7 (47.9–54.8)</td>
</tr>
<tr>
<td>G1%</td>
<td>17</td>
<td>18.4 ± 1.7 (14–22.6)</td>
</tr>
<tr>
<td>G2%</td>
<td>16</td>
<td>18.3 ± 2.3 (14.1–24.4)</td>
</tr>
<tr>
<td>Odontostyle</td>
<td>14.5</td>
<td>15.0 ± 0.4 (14.5–15)</td>
</tr>
<tr>
<td>Odontophore</td>
<td>26</td>
<td>24.8 ± 2.2 (20–29)</td>
</tr>
<tr>
<td>Spear</td>
<td>41</td>
<td>39.8 ± 2.2 (35–43)</td>
</tr>
<tr>
<td>Cardia length</td>
<td>19</td>
<td>19 ± 1.8 (16–24)</td>
</tr>
<tr>
<td>Body width at: lip region</td>
<td>12</td>
<td>12.3 ± 0.3 (12–12.5)</td>
</tr>
<tr>
<td>mid-body</td>
<td>48</td>
<td>46.8 ± 4.7 (39.5–55)</td>
</tr>
<tr>
<td>anus</td>
<td>22</td>
<td>24.9 ± 1.8 (22–30)</td>
</tr>
<tr>
<td>Lateral chord</td>
<td>12</td>
<td>12.7 ± 1.8 (10.5–18)</td>
</tr>
<tr>
<td>Prerectum</td>
<td>89</td>
<td>89 ± 11 (68–117)</td>
</tr>
<tr>
<td>Rectum</td>
<td>41</td>
<td>37 ± 2.4 (30–41)</td>
</tr>
<tr>
<td>Tail</td>
<td>88</td>
<td>90.9 ± 7.1 (76–110)</td>
</tr>
<tr>
<td>Spicules</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>T-distance anterior end of anterior testis-cloaca (mm)</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>
Calcaridorylaimus castaneae sp. n. (Nematoda, Dorylaimidae) from Bulgaria...

Figure 2. Calcaridorylaimus castaneae sp. n. Female: A Pharyngeal gland nuclei B Anterior region C Pharyngeal region D Genital system E Vulval region F Sperm cells in uterus. Scale bars: A, B, E, F – 30 μm; C, D – 50 μm.
Figure 3. Calcaridorylaimus castaneae sp. n. Female: A Posterior region. Male: B, C Extruded spicules with supplements D Posterior region. Scale bar: A–D – 30 μm.
Calcaridorylaimus castaneae *sp. n.* (Nematoda, Dorylaimidae) from Bulgaria...

**Figure 4.** *Calcaridorylaimus castaneae* sp. n. Female: A Anterior end C Amphid F–I Tail shapes Male B Anterior end D Posterior end with extruded spicules, arrow indicating the spur E Posterior end. Scale bars: A, B, D–I – 20 μm; C – 6 μm.
Calcaridorylaimus castaneae sp. n. (Nematoda, Dorylaimidae) from Bulgaria...

narrowly rounded terminus. Posterior part of tail usually slightly curved dorsally. Two pairs of caudal pores, one subventral, other subdorsal.

Male. General morphology similar to that of female, body curved ventrally in J-shape when fixed. Genital system dicrochic, testes opposed, well developed. Spicules dorylaimoid, with double contour on dorsal arm, 1.3–1.6 times the corresponding body diameter long; ventral arm smaller than dorsal. A spur present dorsally before the distal tip, distinctly visible in extruded spicules. Lateral guiding pieces 9-11 μm long or ca. 23 % spicule length. In addition to adcloacal pair seven to twelve (mostly nine or ten), regularly spaced ventromedian supplements present (9 supplements in 8 specimens; 10 suppl. – in 7, and 7, 8
and 12 each in one specimen). Prerectum 4.5–7.5 times the corresponding body diameter long, extending 0.7–1.8 body widths anterior to the supplement series. Tail dorsally co-noid and broadly rounded. One subdorsal and one subterminal pair of caudal pores.

**Differential diagnosis and relationships.** *Calcaridorylaimus castaneae* sp. n. differs from all species in the genus by a combination of the following characters: long body (1.4–2.1 mm), lip region practically not offset, short odontostyle (14.5–16 μm in females and 14–16 μm in males) and short female tail (75.5–110.5 μm; c=14.7–23.6; c'=2.9–4.4); vulva transverse, 38–46 μm long spicules with spur before its distal end. The new species is most similar to *C. andrassyi* from which it can be differentiated by having transverse vulva vs pore-like and shorter spicules (38–46 μm vs 52–57 μm). Further, *C. castaneae* differs from *C. ruwenzorii* by having shorter odontostyle (14.5-16 μm vs 19.5-25 μm), and tail (75.5-110.5 μm vs 160 μm), higher c (c=14.7-23.6 vs c=10) and lower c’ (c’=2.9–4.4 vs 7) values in females. It can be differentiated from *C. arcticus*, *C. beatus*, *C. calcarifer* and *C. signatus* by having different vulva shape (transverse vs longitudinal) and shorter spicules (38–46 μm vs 57–67 μm; 48–55 μm; 52–54 μm and 72 μm). From *C. sirgeli* it differs by having transverse vulva vs pore-like, higher c (c=14.7–23.6 vs c=9.7–11.3) and lower c’ (c’=2.9–4.4 vs c’=5.3–6.7) values. Finally, *C. castaneae* differs from *C. promissus* and *C. similimus* by having longer odontostyle (14.5–16 μm vs 13 μm and 11 μm) and shorter tail (75.5–110.5 μm vs 158–178 μm and 175 μm).


**Type locality and plant association.** Belasitsa Mountain, south-western Bulgaria, litter from old broadleaf forest (100-140 years) dominated by *C. sativa*, mixed with *Q. daleshampii* and *F. sylvatica*. Site is located in the vicinity of Belasitsa hut, N41°22’12”; E23°11’12” (sub-compartment 140b). Second locality: young sweet chestnut forest (30-40 years old) near Belasitsa village (sub-compartment 104g).

**Type material.** Holotype and 80 paratype females and 47 males deposited in the nematode collection of the Institute of Biodiversity and Ecosystem Research, Sofia, Bulgaria. Other paratypes deposited as follows: four females and two males in the Nematode Collection of the Foodland Environment Research Agency, Sand Hutton, UK (former Rothamsted Nematode Collection); three females and three males in the USDA Nematode Collection, Beltsville, Maryland, USA; two females and two males in the Riverside Nematode Collection, University of California, Riverside, USA; four females, and four males in the Wageningen Nematode Collection (WANECO), Wageningen, the Netherlands; four females and three males in the Nematode Collection of the Zoology Museum of the Ghent University, Belgium.

**Etymology.** The scientific name is derived from the generic name of dominant tree species, the sweet chestnut tree (*Castanea*) in the forest where this nematode was found.
### Table 2. Main morphological and morphometrical data of *Calcaridorylaimus* species, habitat type and distribution.

<table>
<thead>
<tr>
<th>Species</th>
<th>Body length (mm)</th>
<th>Body width (mm)</th>
<th>Lip region width (mm)</th>
<th>Tail length (mm)</th>
<th>Vulva shape</th>
<th>Spicule length (mm)</th>
<th>Habitat</th>
<th>Distribution</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>C. andrassyi</em></td>
<td>1.18–1.5</td>
<td>0.8–1.2</td>
<td>0.6–0.7</td>
<td>4.5–5</td>
<td>13–14</td>
<td>44.5–49</td>
<td>aquatic freshwater</td>
<td>forest</td>
<td>Andrássy 2003</td>
</tr>
<tr>
<td><em>C. arcticus</em></td>
<td>1.6–2.3</td>
<td>1.0–1.25</td>
<td>0.6–0.7</td>
<td>5–6.5</td>
<td>12–16</td>
<td>48–55</td>
<td>Russian Arctic</td>
<td>coastal forest</td>
<td>Ahmad and Shaheen 2004</td>
</tr>
<tr>
<td><em>C. beatus</em></td>
<td>1.28–1.3</td>
<td>0.9–1.0</td>
<td>0.6–0.8</td>
<td>14.0–16.3</td>
<td>10–12</td>
<td>115–167</td>
<td>Moist grassy soil</td>
<td>forest</td>
<td>Andrássy 2011</td>
</tr>
<tr>
<td><em>C. calcarifer</em></td>
<td>1.18–1.3</td>
<td>0.9–1.0</td>
<td>0.6–0.8</td>
<td>14.0–16.3</td>
<td>10–12</td>
<td>115–167</td>
<td>Republic of Congo</td>
<td>forest soil</td>
<td>Andrássy 1986</td>
</tr>
<tr>
<td><em>C. castaneae</em></td>
<td>1.4–2.1</td>
<td>1.4–1.8</td>
<td>0.7–0.9</td>
<td>16.0–23.6</td>
<td>10–12</td>
<td>53.7–89.7</td>
<td>Broadleaf forest</td>
<td>litter</td>
<td>Present study</td>
</tr>
<tr>
<td><em>C. dieckii</em></td>
<td>1.28–1.6</td>
<td>1.0–1.25</td>
<td>0.6–0.7</td>
<td>14.0–16.3</td>
<td>10–12</td>
<td>115–167</td>
<td>Republic of Congo</td>
<td>forest</td>
<td>Andrássy 2003</td>
</tr>
<tr>
<td><em>C. signatus</em></td>
<td>1.3–1.7</td>
<td>0.9–1.0</td>
<td>0.6–0.8</td>
<td>14.0–16.3</td>
<td>10–12</td>
<td>115–167</td>
<td>South Africa</td>
<td>forest</td>
<td>Heyns and Meyer 1975</td>
</tr>
</tbody>
</table>

Shape of vulva: **L** – longitudinal; **P** – pore like; **T** – transverse; when average values are present ranges in parentheses; * from the drawing; **C. promissus** from Alaska (Andrássy 2003) was not included in a subsequent paper by the same author (Andrássy 2011); since most of the characters deviate substantially from the original description, probably belongs to another species.
**Key to species of Calcaridorylaimus**

1. Odontostyle 19.5–25 μm long ................................................... *C. ruwenzorii*
   - Odontostyle shorter, ≤17 μm ................................................... 2

2. \( c' = 11 \) ................................................... *C. simillimus*
   - \( c' < 9 \) ................................................... 3

3. \( c' = 2.2–2.4 \) ................................................... *C. beatus*
   - \( c' ≥ 2.9 \) ................................................... 4

4. \( c = 11 \) ................................................... 5
   - \( c > 11 \) ................................................... 7

5. \( V = 45–47, \) supplements 10–13 ................................................... *C. promissus*
   - \( V > 49, \) supplements 6–9 ................................................... 6

6. Vulva longitudinal, spicules 52–54 μm ................................................... *C. calcarifer*
   - Vulva pore-like, spicules 39–45 μm ................................................... *C. sirgeli*

7. Lip region 14–17 μm wide, vulva longitudinal ................................................... 8
   - Lip region 11–13 μm wide, vulva not longitudinal ................................................... 9

8. Tail 120–171 μm long, \( L = 1.6–2.27, \) spicules 57–67 ................................................... *C. arcticus*
   - Tail 100 μm long, \( L = 1.3–1.7, \) spicules 72 ................................................... *C. signatus*

9. Vulva pore-like, spicules 52–57 ................................................... *C. andrassyi*
   - Vulva transverse, spicules 38–46 ................................................... *C. castaneae*

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**Phylogenetic relationships of Calcaridorylaimus castaneae based on partial 28S and 18S rDNA sequences**

The sequences of D2-D3 expansion domains of 28S and 18S ribosomal DNA from two female and two male nematodes have been processed. No inter-individual variability within both domains have been observed, thus only two consensus sequences for each of the genes (787 and 1699 bp long) have been submitted to GenBank (accession numbers KF717498 and KF717497). A BLAST search for D2-D3 region showed highest similarity (88%) to the sequence of *Labronema vulvapapillatum* clone 2 (AY592997, Holtermann et al. 2008) while the 18S rDNA showed 99% similarity (4-6 nucleotide differences) to the sequences of *Mesodorylaimus bastiani* (AJ966488) from France (Meldal et al. 2007) and two sequences of unidentified species from environmental samples from Scotland (AJ875133 and JN049666) (Griffiths et al. 2006, Donn et al. 2012).

The phylogenetic analyses based on 18S rDNA and D2-D3 of 28S rDNA sequences from various dorylaimid species with the highest matches of the BLAST search (up to 97% and 85%, respectively) were aligned along with our sequence. The phylograms obtained by NJ, ML and BI methods showed similar topology and differed only in the positions of poorly supported clades. The BI trees (Figs 7 and 8) with posterior probabilities higher than 0.8 and NJ-ML trees with bootstrap values above 70% are presented (Figs 9 and 10). The new species has clustered in a well-supported
Figure 7. Phylogenetic relationships of Calcaridorylaimus castaneae sp. n. and its closest species for 18S rRNA gene. Bayesian Inference strict consensus tree acquired under GTR+G model. Posterior probabilities higher than 0.8 are presented.

The genus Calcaridorylaimus was erected by Andrássy (1986) to accommodate a few species having different shapes and structures of spicules from those of Mesodorylaimus, with the males and females being practically indistinguishable. The phylograms inferred from
Figure 8. Phylogenetic relationships of *Calcaridorylaimus castaneae* sp. n. and its closest species for D2-D3 expansion segments of the 28S rRNA gene. Bayesian Inference strict consensus tree acquired under GTR+G model. Posterior probabilities higher than 0.8 are presented.
Figure 9. Phylogenetic relationships of *Calcaridorylaimus castaneae* sp. n. and its closest species for 18S rRNA gene. Tree acquired with Neighbor Joining and Maximum Likelihood (GTR+G model) methods. Bootstrap values higher than 70% are presented.
Figure 10. Phylogenetic relationships of *Calcaridorylaimus castaneae* sp. n. and its closest species for D2-D3 expansion segments of the 28S rRNA gene. Tree acquired with Neighbor Joining and Maximum Likelihood (GTR+G model) methods. Bootstrap values higher than 70% are presented.

18S sequences showed the closest relationships of *C. castaneae* with some members of the latter genus; however, the insufficiency of molecular data complementary to detailed morphological studies of species belonging to both genera does not allow the elucidation of evolutionary relationships among them and the position of the new species herein described.
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References


Calcaridorylaimus castaneae *sp. n.* (*Nematoda, Dorylaimidae*) *from Bulgaria...*