A new rare nematode *Nothocriconemoides hangzhouensis* n. sp. (Nematoda: Criconematidae) from Hangzhou, China

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Abstract

The Family Criconematidae is commonly referred as ring nematodes that include some members with economic importance as plant parasites. During a recent nematode inventory survey at Zhejiang Province, China, a new species of genus *Nothocriconemoides* was detected in the rhizosphere of elm tree. *Nothocriconemoides hangzhouensis* n. sp. can be characterized by the female body having annuli with fine longitudinal striations and 2 to 3 anastomoses at the posterior half of the body. The first cephalic annulus is rounded and expanded enclosing the lip region, and the second annulus is narrow, offset, collar like. En face view shows a central elevated labial disk bearing four distinct equal-sized submedian lobes and “I” shaped oral aperture. Excretory pore is located 3–4 annuli posterior to esophageal bulb. Vagina is straight and vulva closed. The ventral side of postvulval annuli is inverted, in majority of individuals. Anus is indistinct and located on the next annuli posterior to vulva. Tail is short, conoid, with forked or branched terminus. Juveniles are devoid of collar-shaped annuli in the lip region. The cephalic region has two rounded annuli where the first annulus shows slight depression in the middle. Body annuli are finely crenated. Anus is indistinct and located 3 to 4 annuli from tail terminus. Tail is short ending in a single lobed terminus. Phylogenetic studies based on analysis of the D2–D3 expansion segments of the 28S rRNA, ITS rRNA, partial 18S rRNA, and cox1 gene revealed that the new species formed a separate clade from other criconematid species, thereby supporting its status as a new species of the genus. The new species showed close relationships with *Discocriconemella sinensis*. Additionally, this is the first record of genus *Nothocriconemoides* from China.

Keywords

DNA sequencing, Elm tree, Morphology, Morphometrics, Nematode, New record, Species, Phylogeny, Scanning electron microscopy.
collar like. So far, the genus contains only two species i.e. *Nothocriconemoides crenulatus* (Ivanova, 1984) and *Nothocriconemoides lineolatus* (Maas et al., 1971) that were described from Tadzhikistan and Suriname, respectively. Both species were found associated with forest soils; however, no association has been reported from soils of cultivated areas (Geraert, 2010).

During our nematode inventory survey, a population of *Nothocriconemoides* was detected in the rhizosphere of elm tree. As *Nothocriconemoides* was never reported from China, the present work was undertaken to identify the species status. The morpho-molecular characterization and SEM data of this population were compared with the existing species of the genus. Careful examination revealed that the species under investigation presents unique characteristics and is a new member of the genus *Nothocriconemoides*. Therefore, the paper describes a new *Nothocriconemoides* species with the following objectives: to provide a morphological and molecular characterization of the new species; to elucidate important morphological details through SEM observations; and to study the phylogenetic relationships of these species with other related criconematid species.

**Materials and methods**

**Nematode samplings, extraction and morphological study**

Nematodes were extracted from soil and root samples using the modified Cobb sieving and flotation-centrifugation method (Jenkins, 1964). For morphometric studies, nematodes were killed and fixed in hot formalin (4% with 1% glycerol) and processed in glycerin (Seinhorst, 1959). The measurements and light micrographs of nematodes were made with a Nikon Eclipse Ni-U 931845 compound microscope. For the SEM examination, the nematodes were fixed in a mixture of 2.5% paraformaldehyde and 2.5% glutaraldehyde, washed three times in 0.1 M cacodylate buffer, post-fixation in 1% osmium tetroxide, dehydrated in a series of ethanol solutions and critical point-dried with CO₂. After mounting on stubs, the samples were coated with gold with 6 to 10-nanometer thickness and the micrographs were made with 3 to 5kV operating system (Maria et al., 2018a).

**Molecular analyses**

DNA was extracted by transferring individual nematodes into the Eppendorf tube containing 16μL ddH₂O. Nematodes were crushed using a sterilized pipette tip, the tubes were centrifuged at 12,000 rpm for 1 min and frozen at −68°C for at least 30 min. Tubes were heated to 85°C for 2 min, and then, 2μL proteinase K and PCR buffer solution were added. The tubes were incubated at 56°C for 1 to 2 hr and, then, at 95°C for 10 min. After incubation, these tubes were cooled to 4°C and used for conducting PCR (Zheng et al., 2003). Several sets of primers (synthesized by Invitrogen, Shanghai, China) were used in the PCR analyses to amplify the partial 18S, ITS region, D2–D3 of 28S of rDNA and partial coxl fragments. Primers for amplification of partial 18S were 18s900–18s1713 (Olson et al., 2017). Primers for amplification of ITS were TW81-AB28 (Joyce et al., 1994). The primers for amplification of D2–D3 of 28S were D2A and D3B (De Ley et al., 1999). And, finally, the primers used for coxl amplification were COI-F5 (5’-AATWTTWGGTGGAGACCTTCTTGAAAC-3’) and COI-R9 (5’-CTTAAACAAAAATGAGWACACACWACATAAGTATC-3’) (Powers et al. 2014). PCR conditions were as described by Ye et al. (2007) and Powers et al. (2010). PCR products were evaluated on 1% agarose gels stained with ethidium bromide. PCR products of sufficiently high quality were sent for sequencing by Invitrogen (Shanghai, China).

**Phylogenetic analysis**

Newly obtained sequences of *Nothocriconemoides hangzhouensis* n. sp. (D2-D3 expansion segments of 28S, ITS, partial 18S rRNA, and partial coxl) and the available sequences of other criconematid nematodes obtained from NCBI were used for phylogenetic analyses. Outgroup taxa for the dataset were chosen according to previous published data (Afshar et al., 2019; Maria et al. 2019). Multiple alignments of the different sequences were made using the FFT-NS-2 algorithm of MAFFT v. 7.205 (Katoh and Standley 2013). Sequence alignments were manually visualized using BioEdit (Hall 1999) and edited by Gblocks ver. 0.91b (Castresana 2000) in the Castresana Laboratory server (http:// molevol.cmima.csic.es/castresana/Gblocks_server. html) using options for a less stringent selection (minimum number of sequences for a conserved or a flanking position: 50% of the number of sequences +1; maximum number of contiguous non-conserved positions: 8; minimum length of a block: 5; allowed gap positions: with half). Phylogenetic analyses of the sequence datasets were based on Bayesian inference (BI) using MrBayes 3.1.2 (Ronquist et al., 2012). The best-fit model of DNA evolution was obtained using JModelTest V.2.1.7 (Darriba et al. 2012) with the Akaike Information Criterion (AIC). The best-fit model,
the base frequency, the proportion of invariable sites, and the gamma distribution shape parameters and substitution rates in the AIC were then given to MrBayes for the phylogenetic analyses. An unlinked general time-reversible model with invariable sites and a gamma-shaped distribution (GTR + I + G) was used for the D2-D3 expansion segments of 28S rRNA, ITS, partial 18S, and partial coxl. These BI analyses were run separately per dataset using four chains for $2 \times 10^6$ generations for all of the molecular markers. A combined analysis of the three genes was not undertaken due to some sequences not being available for all species. The Markov chains were sampled at intervals of 100 generations. Two runs were conducted for each analysis. After discarding burn-in samples and evaluating convergence, the remaining samples were retained for further analyses. The topologies were used to generate a 50% majority-rule consensus tree. Bayesian posterior probabilities (BPP) are given on appropriate clades. Trees from all analyses were visualized using FigTree software V.1.42 (http://tree.bio.ed.ac.uk/software/figtree/).

Results and description
Systematics

*Nothocriconemoides hangzhouensis* n. sp. (Figs 1–4; Table 1).

Description

**Females**

Body is slightly curved ventrally after heat-killing. Body annuli are wide (13-17 µm thick in the middle of the body) with fine longitudinal striations that look like annulus bearing rough cuticular margins. Anastomoses are 2 to 3, located at the posterior half of the body. The first cephalic annulus is rounded and expanded enclosing the lip region. The second annulus narrow, offset, collar like. *En face* view shows a central elevated labial disk bearing four equal-sized submedian lobes and “I”-shaped oral aperture. Stylet is robust with anchor-shaped basal knobs, and DGO indistinct. Esophageal lumen is looped in median esophageal bulb having a medium-sized valvular apparatus. Isthmus is narrow, short, encircled by nerve ring, and basal esophageal bulb distinct. Excretory pore is 3 to 4 annuli posterior to esophageal bulb. Monodelphic gonad is outstretched, and spermatheca is spherical, filled with sperm. Vagina is straight, vulva is closed, and vulval lips do not project above body contour. The ventral side of postvulval annuli is inverted, in majority of individuals. Anus is indistinct and located on the next annuli posterior to vulva. Tail is short and conoid, longitudinal striations are more prominent on the terminal annulus that gives the appearance of forked or branched terminus.

**Male**

Not found.

**Juveniles ($n=5$)**

Except for the cephalic region, they are similar to females; cephalic region of juveniles are devoid of collar-shaped annuli. Two rounded annuli are present...
Nothocrinemonoides hangzhouensis n. sp. from China: Maria et al.

Figure 2: Light photomicrographs of Nothocrinemonoides hangzhouensis n. sp. Female A: Entire body; B-E: Cephalic regions; F-H: Esophageal regions, arrow pointing on the excretory pore (exp); I-K: Cuticle markings; L-O: Tail regions, arrows pointing on vulva (v) and anus (a). (Scale bars=A=50 μm, B-I=10 μm).
and the first annulus show slight depression in the middle. Body annuli are narrower (4.5-5.5), slightly higher in number $R = (48-51)$ and finely crenated. Stylet is (32.5-36.7) $\mu$m long and esophageal components are similar as those of females but less developed. Anus is indistinct and located 3 to 4 annuli from tail terminus. Tail is short and ends in a single lobed terminus.

**Type host and locality**

This population was found in the rhizosphere of *Ulmus* sp. from Zijingang Campus, Zhejiang University, Hangzhou, Zhejiang Province, P.R. China, on February 2019. The geographical position of the sampling site is E: 120°4’54” N: 30°17’5.”

**Type material**

Holotype female and 13 female paratypes (slide numbers ZJU-30-01-ZJU-30-03) were deposited in the nematode collection of Zhejiang University, Hangzhou, China. Four females and two juveniles paratypes on two slides (Slide numbers T-7353, T-7356) and ten additional females on two slides were (T-7354-55) deposited at USDA nematode collection, Beltsville, Maryland, USA. The Zoobank code is as follows: LSID urn:lsid:zoobank.org:pub:E977B880-9EF3-4E2C-BB29-4BD970D63F0A

**Etymology**

The species epithet refers to the City name where the species was detected.

**Diagnosis and relationships**

The new species can be characterized by wider body annuli that have fine longitudinal striations that look like annulus bearing rough cuticular margins, and 2 to 3 anastomoses at the posterior half of the body. The first cephalic annulus is rounded and expanded enclosing the lip region, and the
Table 1. Morphometric data for *Nothocriconemoides hangzhouensis* n. sp.

|                | Holotype    | Paratype    |
|----------------|-------------|-------------|
| n              | 494.0       | 17          |
| Body Length    | 487.1±43.8  | (419.6-572.3)|
| R              | 37.2±1.2    | (35.0-39.0) |
| Rst            | 6.5±0.5     | (6.0-7.0)   |
| Rex            | 14.7±0.6    | (13.0-15.0) |
| RV             | 2.9±0.2     | (2.0-3.0)   |
| Ryan           | 0.0±0.0     | (0.0-0.0)   |
| Ran            | 2.9±0.2     | (2.0-3.0)   |
| a              | 7.9±0.7     | (6.3-9.4)   |
| b              | 4.1±0.3     | (3.5-4.5)   |
| c              | 17.3±1.7    | (14.2-20.0) |
| c'             | 0.9±0.1     | (0.7-1.1)   |
| V              | 92.6±0.9    | (90.5-94.0) |
| VL/VB          | 1.0±0.1     | (0.9-1.2)   |
| Lip height     | 9.5±0.7     | (7.8-10.6)  |
| Lip diam.      | 20.2±1.4    | (17.2-22.1) |
| Stylet length  | 71.0±3.0    | (64.4-75.5) |
| Stylet percentage | 14.7±1.2  | (13.1-17.4) |
| Pharynx length | 117.5±5.0   | (111.8-129.6)|
| Body width     | 61.8±5.3    | (52.0-69.4) |
| Vulval body diam.| 35.7±2.5   | (31.8-38.5) |
| Anal body diam. | 32.1±3.1   | (26.4-37.4) |
| Vulva to tail terminus | 36.0±4.0 | (29.8-41.7) |
| Tail length    | 28.3±1.9    | (23.3-30.5) |
| Annuli width   | 14.5±1.2    | (13.1-16.9) |

Notes: All measurements are in µm and in the form of mean ± SD (range).

The genus only contains two species; It can be differentiated from *N. crenulatus* by having shorter stylet 64.4 to 75.5 vs 87 to 96µm long, less number of body annuli R=35.0 to 39.0 vs 57 to 64, less number of annuli between vulva and tail terminus RV=2.0 to 3.0 vs 6 to 8, location of anus (next annuli to vagina vs 3 to 4 annuli posterior to vagina), and tail terminus morphology (terminal annulus forked or branched vs button shaped).

The new species differs from *N. lineolatus* by less number of body annuli R=35.0 to 39.0 vs 57 to 64, less number of annuli between vulva and tail terminus RV=2.0 to 3.0 vs 7 to 9, location of anus (next annuli to vagina vs 3 to 5 annuli posterior to vagina), anastomosis (2-3 vs 1), lip annuli (2 vs 4), submedian lobes (separate as four vs connected as 2 subdorsal and sublateral lobes), position of excretory pore.
Figure 4: Scanning electron microscopy of *Nothocriconemoides hangzhouensis* n. sp. Female. A: Entire body; B-D: En face view; E: Cuticle markings; F-H: tail regions arrows pointing on vulva (v) and anus (a) (Scale bars, A = 100 μm; B, C = 10 μm; D, H = 20 μm; G-F = 30 μm).

(Molecular profiles and phylogenetic status)

The new species was molecularly characterized using partial 18S, D2-D3 of 28S, ITS and coxl sequences and obtained sequences were deposited.
Nothocriconemoides hangzhouensis n. sp. from China: Maria et al.

Figure 5: Phylogenetic relationships of *Nothocriconemoides hangzhouensis* n. sp. with other criconematids species as inferred from Bayesian analysis using the 18S rRNA gene sequence dataset with the GTR + I + G model (−lnL=7,315.8130; AIC = 14,859.6260; freqA=0.2371; freqC=0.2413; freqG=0.2833; freqT=0.2384; R(a)=1.5166; R(b)=2.2509; R(c)=0.9364; R(d)=0.7246; R(e)=6.0997; R(f)=1.0000; Pinva=0.6630; and Shape=0.6070). Posterior probability more than 70% is given for appropriate clades. Newly obtained sequences are indicated in bold.
Figure 6: Phylogenetic relationships of Nothocriconemoides hangzhouensis n. sp. with other criconematid species as inferred from Bayesian analysis using the D2-D3 of 28 S rRNA gene sequence dataset with the GTR + I + G model (−lnL=8,382.1334; AIC=16,972.2669; freqA=0.1451; freqC=0.2354; freqG=0.3515; freqT=0.2681; R(a)=0.8404; R(b)=2.5613; R(c)=1.6924; R(d)=0.4616; R(e)=4.7092; R(f)=1.0000; PinvA=0.2730; and Shape=0.8370). Posterior probability more than 70% is given for appropriate clades. Newly obtained sequences are indicated in bold.
**Discussion**

Since 1960, the taxonomy of criconematids has been revised independently and more or less simultaneously by several nematologists, which caused confusions and conflicting definitions of genera as well as contradictions in proposed species synonyms (Subbotin et al., 2005). One such example is *Mesocriconema xenoplax* (Raski, 1952; Loof, 1988), it has been called *Macroposthonia xenoplax* (Siddiqi, 2000; Wouts, 2006), *Criconemella xenoplax* (Xiang et al., 2010), or *Criconemoides xenoplax* (Decraemer and Geraert, 2006; Decraemer and Hunt, 2006; Cid Del Prado Vera and Talavera, 2012).

Criconematids have been widely accepted as a monophyletic group based on the esophageal structure, monodelphic ovary and sexual dimorphism. However, there is no concrete phylogenetic evidence of criconematids subfamily, genus and subgenus grouping (Powers et al., 2017). Geraert (2010) placed *Nothocriconemoides* in the subfamily Macroposthoniinae but phylogenetically the new species grouped with various species of *Bakernema*, *Criconemoides*, *Mesocriconema*, *Neobakernema* (subfamily Macroposthoniinae) *Criconema*, *Lobocriconema*, *Neolobocriconema* (Criconematinae) and *Discocriconemella sinensis* (Discocricinomella). *Nothocriconemoides hangzhouensis* n. sp. can be differentiated from *Discocriconemella sinensis* and other criconematids based on the presence of fine longitudinal striae on the cuticle, and
Figure 8: Phylogenetic relationships of *Nothocriconemoides hangzhouensis* n. sp. with other criconematids species as inferred from Bayesian analysis using the coxI gene sequence dataset with the GTR + I + G model (−lnL=13,473.0592; AIC=27,198.1184; freqA=0.3715; freqC=0.0509; freqG=0.0477; freqT=0.5299; R(a)=0.7544; R(b)=36.5547; R(c)=1.6680; R(d)=51.5187; R(e)=20.3538; R(f)=1.0000; Pinva=0.2510; and Shape=0.3470). Posterior probability more than 70% is given for appropriate clades. Newly obtained sequences are indicated in bold.
from Lobocricerena, Neolobocricerena based on the absence of rows of scales on juveniles.

Additionally, Nothocriconemoides hangzhouensis n. sp. appeared as a sister species of Discocricerena sinensis in our phylogenetic analysis. Discocricerena species are characterized by the presence of cephalic disc (Geraert, 2010), and currently the genus contains 29 species but only D. hengsungica (Choi and Geraert, 1975), D. limitanea and D. sinensis are molecularly characterized; interestingly, none of these species display monophyletic behavior (Maria et al., 2018b, 2019). Nothocriconemoides hangzhouensis n. sp. has a large labial annulus resembling the Discocricerena type 1 cephalic disc (round to oval with uninterrupted margins) of sensu Vovlas (1992). It is also noted that Discocricerena species having type 3 (disc intend medially and laterally giving a four-lobed appearance) cephalic disc is not easy to differentiate from Mesocriconema (Geraert, 2010). Several authors have expressed their concerns that Discocricerena species showed considerable variation in distinguishing characters (Orton Williams, 1981; Vovlas, 1992; Siddiqi, 2000). It is likely that a large labial disc is a homoplastic character that independently appears in several criconematids lineages. To this point, we only assume that close phylogenetic relationship between Nothocriconemoides hangzhouensis n. sp. and D. sinensis is mainly because of a similar arrangement of the labial annulus, except the submedian lobes. We agreed with Jahanshahi-Afshar et al. (2019) that majority of criconematids genera and species have yet to be sequenced, and with the inclusion of additional/new sequences of criconematids, the phylogenetic studies could provide better insights than now.

Nothocriconemoides hangzhouensis n. sp. is the first-named species of this genus to be molecularly characterized. In our coxl tree, an unknown Nothocriconemoides sp. (KJ788064) from Costa Rica is arranged distantly from Nothocriconemoides hangzhouensis n. sp. When the information attached to this unidentified species is examined (at https://nematode.unl.edu/sp-16137.htm), it is observed that this population does not fit with the generic definition of Nothocriconemoides i.e. the second cephalic annulus of female is offset collar like, lips have four distinct submedian lobes and vulva is closed. Uncertainties concerning the correct identity of some GenBank sequences and lack of sufficient ultra-morphological characterization presenting challenges in the taxonomy of criconematids. The phylogeny of the majority of criconematids taxa is not well resolved, and to this point, we only suggest that molecular identification can be an efficient way of identifying species; however, linking the correct molecular information to the detected species is an important aspect. The generic status of new species is assigned primarily on the basis of morphological characters of females and juveniles. This is the first report of the genus Nothocriconemoides from China.

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