

# On the species status of the root-knot nematode *Meloidogyne mayaguensis* Rammah & Hirschmann, 1988

Gerrit Karssen<sup>1</sup>, Jinling Liao<sup>2</sup>, Zhuo Kan<sup>2</sup>,  
Evelyn YJ van Heese<sup>1</sup>, Loes JMF den Nijs<sup>1</sup>

**1** Plant Protection Service, Wageningen Nematode Collection, P.O. Box 9102, 6700 HC Wageningen, The Netherlands **2** Plant Nematode Lab, College of Environment and Natural Resource, South China Agricultural University, Guangzhou, 510642, PR China

Corresponding author: Gerrit Karssen (g.karssen3@chello.nl)

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## Abstract

Holo- and paratypes of the root-knot nematodes *Meloidogyne mayaguensis* Rammah & Hirschmann, 1988 and *M. enterolobii* Yang & Eisenback, 1983 were morphometrically and morphologically compared. All observed female, male and second-stage juvenile morphometrical and morphological characters are identical for the two studied species. Additionally, contradictions between the original species descriptions were unravelled.

The present study of holo- and paratypes confirms the taxonomical status of *Meloidogyne mayaguensis* as a junior synonym for *M. enterolobii*.

## Keywords

Junior synonym, *Meloidogyne*, *M. enterolobii*, *M. mayaguensis*, Nematoda, root-knot nematode, synonymisation

## Introduction

In 1983 Yang and Eisenback described the root-knot nematode *Meloidogyne enterolobii* from roots of pacara earpod trees (*Enterolobium contortisiliquum* (Vell.) Morong), on Hainan Island in China. The authors reported severe damage on these pacara earpod trees. In 1988 Rammah and Hirschmann described the root-knot nematode *M. maya-*

*guensis* from eggplant (*Solanum melongena* L.) roots, from Puerto Rico. *Meloidogyne mayaguensis* was described by the authors as: “superficially resembles *M. enterolobii*”, and reported at the same time “several distinct morphologically features and a unique malate dehydrogenase pattern (N3c)”.

It was Fargette and Braaksma (1990) and Fargette et al. (1996) who reported for the first time on the resistance-breaking behaviour of *M. mayaguensis* in Africa and concluded that it is present in both continents of Africa and America. The authors reported (1996) on *M. enterolobii*: “*M. enterolobii* from China has been described as having the same esterase phenotype as *M. mayaguensis*. However it is not known whether their DNA are closely related”. In 2000 Carneiro et al. published esterase and malate dehydrogenase patterns for a Brazilian population of *M. mayaguensis*, and detected a different (N1a) malate dehydrogenase pattern. Additionally Blok et al. (2002) published mtDNA results from different *M. mayaguensis* populations, including type material from Puerto Rico.

In their comprehensive studies on the characterisation of *Meloidogyne* species from China, with isozymes and mtDNA, Meng et al. (2004) and Xu et al. (2004) included two *M. enterolobii* populations from Hainan Island, isolated from the fruit tree Guava (*Psidium guajava* L.). They proved for the first time that *M. enterolobii* esterase (VS1-S1) and malate dehydrogenase (N1a) patterns and mtDNA results are identical to reported *M. mayaguensis* data, and concluded carefully: “the mtDNA sequence evidence presented here, suggests that *M. mayaguensis* could be conspecific with *M. enterolobii*”.

In 2005–2006 we compared the available holo- and paratypes of *M. enterolobii* and *M. mayaguensis*. Meanwhile our Chinese co-authors collected live *M. enterolobii* material on Hainan Island at the type locality from the type host and we kindly received live *M. mayaguensis* type material from Dr. V. Blok (originating from Dr. M. Fargette). The preliminary isozyme and morphological results were presented by the first author during a Pest Risk Analysis meeting on *M. enterolobii* at EPPO in Paris (Anonymous, 2008). Additionally this type material of both species was compared at DNA level to *Meloidogyne* sp. from Switzerland and we identified the Swiss population as *M. enterolobii* (Kiewnick et al. 2008).

Finally, as again at DNA level no differences were found, the two species were synonymised: “The species *M. enterolobii* (syn. *M. mayaguensis*)” and “...of *M. mayaguensis* (junior synonym of *M. enterolobii*)” (Kiewnick et al. 2009).

Although taxonomical not strictly necessary, we present herein a morphological and morphometrical comparison between the holo- and paratype slides of *M. mayaguensis* and *M. enterolobii*. Additionally we discuss anomalies between the descriptions of *M. mayaguensis* and *M. enterolobii*.

## Material and methods

Holo- and paratype slides (Table 1) originating from USDA Nematode Collection (USDANC), Beltsville, USA were kindly provided by Dr. Z. Handoo. The type slides

**Table 1.** *Meloidogyne mayaguensis* and *M. enterolobii* holo-, allo- and paratype slides studied, including USDANC codes.

	<i>M. mayaguensis</i>		<i>M. enterolobii</i>	
Holotype	1 female	T-428t	1 female	T-360t
Allotype*	1 male	T-429t	1 male	T-361t
Paratype	10 perineal patterns	T-3849p	8 perineal patterns	T-3147p
Paratype	6 males	T-3843p	10 males	T-3149p
Paratypes	25 J2's	T-3846/7p	25 J2's	T-3152p

\*According to the ICZN rules (4<sup>th</sup> edition) the allotype concept is no longer valid, and treated herein as a paratype.

are in good condition and includes female holotypes, male allotypes, perineal patterns and second-stage juvenile paratypes. These slides were observed by compound light microscopy (Olympus BH-2 and Zeiss Axio Imager), including Differential Interference Contrast and photographed by Leica DMC-50 digital camera. For the overall morphological and morphometrical comparison between the types we focussed on the most differential and supplementary *Meloidogyne* characters, as described by Jepson (1987) and as previously applied by Karssen (2002). Live type material of both species was propagated and maintained on tomato at the greenhouse of the PPS the Netherlands. This material was studied morphologically (females, males and second-stage juveniles) and used for isozyme electrophoresis (Mdh; EC 1.1.1.37 and Est; EC 3.1.1.1). For details on the preparation of slides and applied electrophoresis method we respectively refer to Karssen (1996) and Karssen et al. (1995).

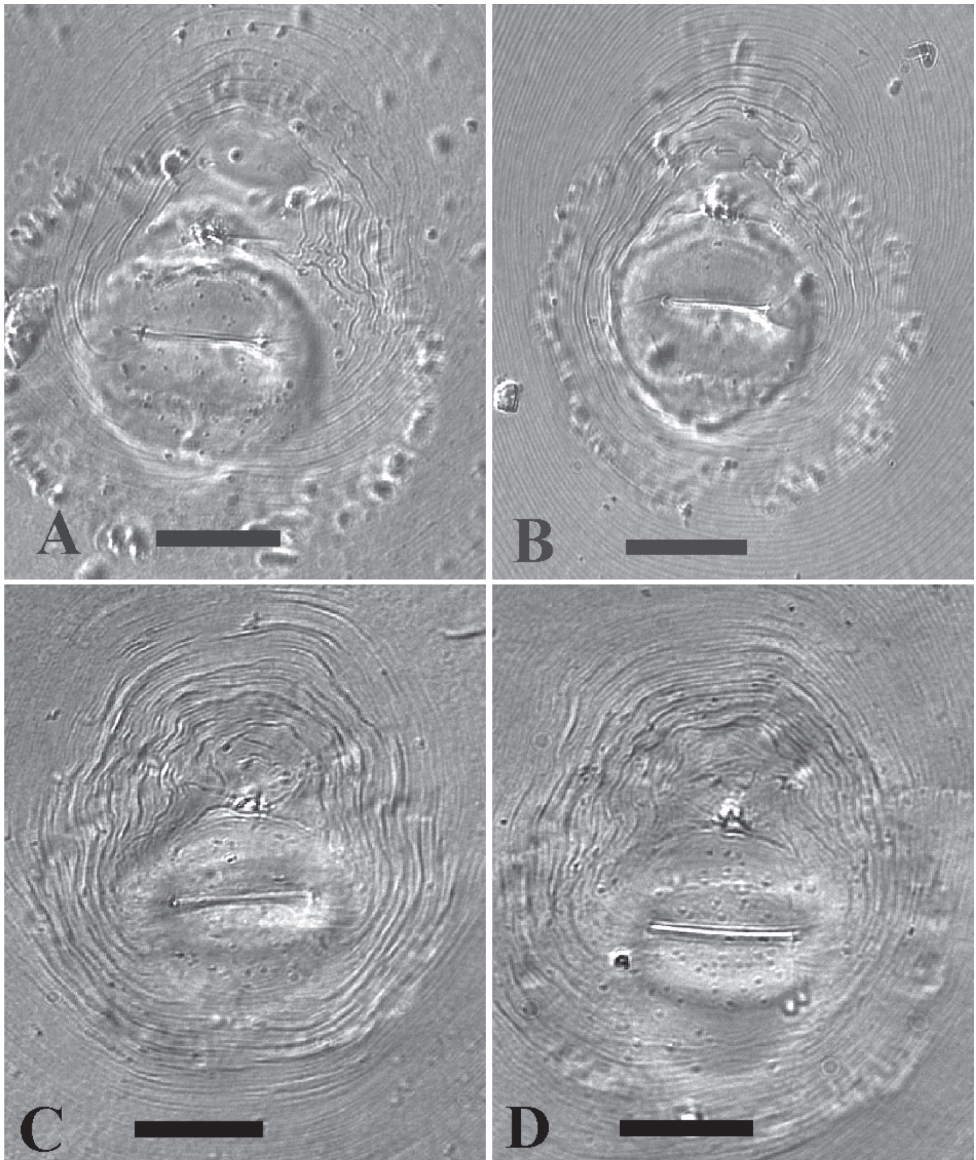
## Results and discussion

See Figure 1 and 2 for LM photographs of female and second-stage juvenile morphological characteristics.

See Table 2–5 for respectively female, male and second-stage juvenile morphological and morphometrical observations.

### Females

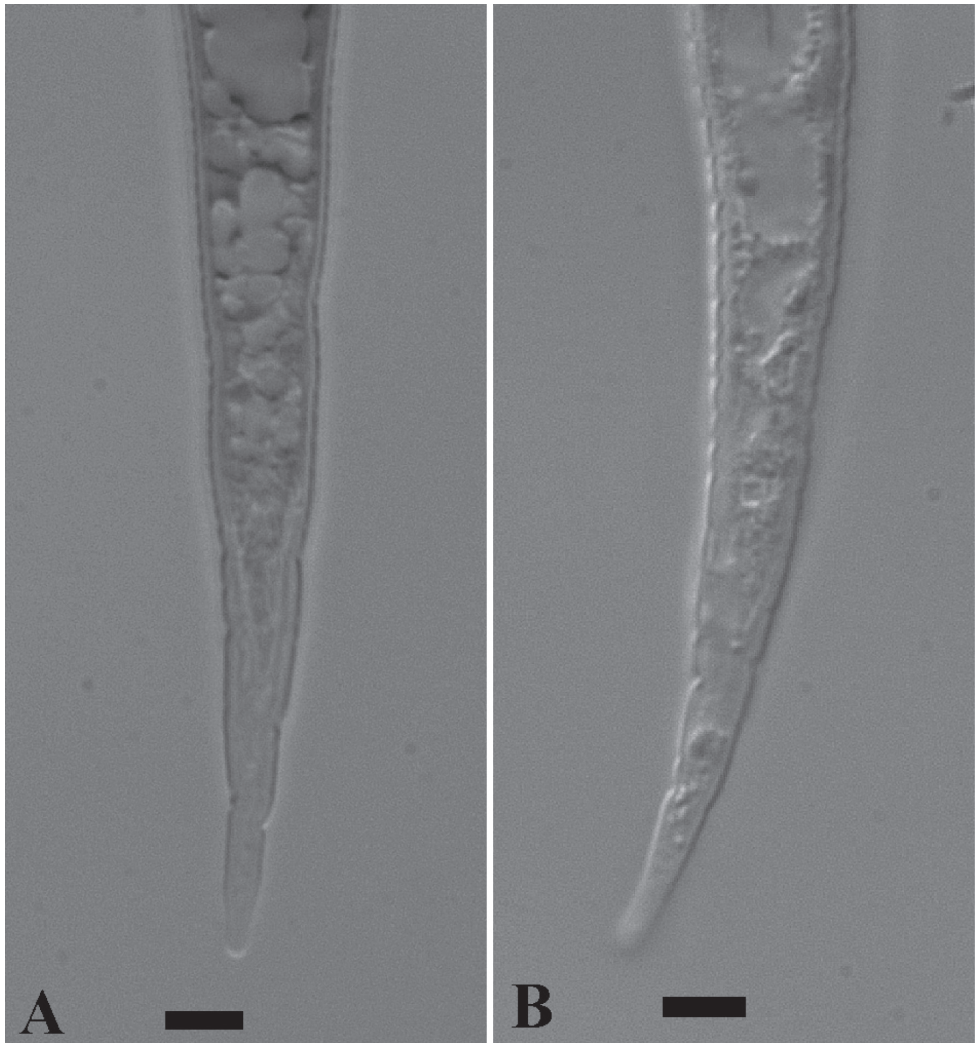
The important morphological characters, like female stylet knob and perineal pattern shape do not differ between the species, as can already be observed by comparing the original illustrations between *M. mayaguensis* and *M. enterolobii* (see original descriptions respectively Fig. 2 A–D & Fig. 3 A–D). This perineal pattern type is not species specific within the genus *Meloidogyne* and can best be marked as typical for many species within the *M. incognita*-group, including the observed variation within the dorsal part. Additionally we observed a relatively large tail remnant area, free of any striae, just above the covered anus (Fig. 1 A–D). Also the observed stylet knob position variation,



**Figure 1.** LM photographs of perineal patterns of *M. mayaguensis* (A, B) and *M. enterolobii* (C, D). Bar = 25  $\mu$ m.

slightly sloping backward to set off from the shaft, is a common *Meloidogyne* feature. Strangely this variation is also clearly visible in the SEM photographs of excised female stylets of *M. mayaguensis* (see original description, Fig. 3 A-C), but not described. With the light microscope one can observe a weak longitudinal indentation, for both species, in the female stylet knobs at the anterior side. The reported differences “not divided so conspicuously as those of *M. enterolobii*” as mentioned in the *M. mayaguensis* descrip-





**Figure 2.** LM photographs of second-stage juvenile tails of *M. mayaguensis* (A) and *M. enterolobii* (B). Bar = 5  $\mu$ m.

tion (see diagnosis original description), was not confirmed by our observations. Also the described position of one of the *M. mayaguensis* stylet knobs “the dorsal knob is slightly sloping posteriad in lateral view” was not observed by us.

### Males

The male head shape for *M. mayaguensis* is described as “not set off”, while a slightly set off head region was observed as described for *M. enterolobii*. Comparing the original SEM pictures of the head for *M. mayaguensis* and *M. enterolobii* (see original descrip-

**Table 2.** Morphological observations of primary female, male and second-stage juvenile characters of *Meloidogyne mayaguensis* and *M. enterolobii* holo- and paratypes compared to described data.

Species	<i>M. mayaguensis</i>	<i>M. enterolobii</i>	
Character	described		observed
<b>Female</b>			
Stylet knobs	knobs reniform or transversely elongated, distinctly indented, merging gradually with shaft	knobs set off from shaft, and divided longitudinally by groove so that each knob appears as two	oval, anteriorly often indented, slightly sloping backward to set off
Perineal pattern	round to dorso-ventrally ovoid, dorsal arch rounded, striae fine, single lateral line may occur	oval shaped, with coarse and smooth striae, dorsal arch moderately high to high, often rounded, nearly square in some, lateral lines not distinct	oval shaped, striae mostly fine, dorsal arch rounded to square, weak lateral line(s) sometimes present
<b>Male</b>			
Head shape	head not set off, shallowly rounded to truncate, head region high without annulations	head cap high and rounded, head region only slightly set off from body	head cap high and rounded, head region slightly set off, not annulated
Stylet knobs	knobs large, set off from shaft, rounded, sloping backward, dorsal knob base concave	knobs large, rounded, distinctly set off, in some specimens each knob divided longitudinally	knobs large, ovoid to rounded, slightly sloping backwards
<b>Second-stage juvenile</b>			
Stylet knobs	knobs small, rounded, set off from shaft, distinctly sloping backward	knobs large, rounded, set off from shaft	knobs ovoid to rounded, slightly sloping backwards
Tail shape	slender, gradually tapering to bluntly rounded tip	very thin, tip broad, bluntly rounded	slender, posterior part nearly straight and parallel, tapering to rounded tip
Hyaline tail part	distinctly set off, often containing small fat droplet at tip	clearly defined, a few fat droplets may occur in terminus	anterior part not clearly delimited

tions respectively Fig. 6 A–D & 5 A,B) shows clearly not any differences in head morphology. Also the male stylet knobs have been SEM studied for the original descriptions (Fig. 3 E, F & Fig. 6 B) of both species. Large oval to rounded shaped knobs, slightly sloping backwards are clearly visible. This was also observed by LM for both species, however described as “rounded and set off” for *M. enterolobii* and “set off from the shaft, rounded, sloping backward” for *M. mayaguensis*. The later description of the knobs is rather odd, i.e. set off and sloping backward at the same time! The same results were described and observed for the second-stage juvenile knobs for both species.

### Second-stage juveniles

The second-stage juvenile stylet knob size is described as small for *M. mayaguensis* and large for *M. enterolobii*. We indeed observed a larger size variation for *M. enterolobii*

**Table 3.** Morphometrical (in  $\mu\text{m}$ ) observations (mean, SD & range) of female *Meloidogyne mayaguensis* and *M. enterolobii* holo- (single female) and paratypes (perineal patterns) compared to described data.

Species	<i>M. mayaguensis</i>		<i>M. enterolobii</i>	
	description	observed	description	observed
<b>Holotype (N)</b>	1	1	1	1
Body length	720	674	667	693
Body width	570	576	415	462
Neck length	190	168	265	262
Neck width	160	169	--	--
DGO	6.2	6.4	3.7	4.8
Excretory pore to head end	46.4	45.8	44.8	64.0
Stylet length	15.1	15.7	13.4	14.7
Stylet knob height	2.2	2.0	2.7	2.3
Stylet knob width	4.4	4.5	4.3	4.5
<b>Paratypes (N)</b>	35	10	20	8
Interphasmidial dist.	23.2 $\pm$ 2.5 (18.1–29.6)	28.8 $\pm$ 3.7 (24.3–33.3)	30.7 $\pm$ 4.8 (22.2–42.0)	33.5 $\pm$ 7.6 (22.4–41.9)
Vulval slit length	26.1 $\pm$ 1.9 (20.9–30.4)	27.0 $\pm$ 1.4 (25.0–29.4)	28.7 $\pm$ 2.0 (25.3–32.4)	28.0 $\pm$ 1.0 (25.9–29.1)
Vulva-anus distance	18.4 $\pm$ 1.5 (12.7–21.1)	21.4 $\pm$ 3.1 (17.0–27.1)	22.2 $\pm$ 1.8 (19.7–26.6)	23.4 $\pm$ 1.6 (21.1–26.2)
DGO	4.8 $\pm$ 0.8 (3.5–6.7)	–	4.9 $\pm$ 0.8 (3.7–6.2)	–
Excretory pore to head end	48.2 $\pm$ 13.6 (25.9–86.6)	–	62.9 $\pm$ 10.5 (42.3–80.6)	–
Stylet length	15.8 $\pm$ 0.8 (13.8–16.8)	–	15.1 $\pm$ 1.4 (13.2–18.0)	–

stylet knob width (2.5 – 4.0  $\mu\text{m}$ ) compared to *M. mayaguensis* (2.2 – 2.9  $\mu\text{m}$ ). However when observing live second-stage juveniles, the same large stylet knob width variation was observed for both species.

As for the males, the published SEM second-stage juvenile head shape is absolute identical for *M. mayaguensis* and *M. enterolobii* (see original descriptions respectively Fig. 7 A–D & Fig. 8 A, B). The tail is distinctly tapering and in the posterior tail (roughly the hyaline tail part) nearly straight and running parallel for both second-stage juvenile paratypes. Also, for both species the hyaline tail part is described as “distinctly set off” or “clearly defined”. We observed for both species however not a clearly anterior delimited hyaline tail part, in fact the body content runs deep into the hyaline tail part (Fig 2 A, B), as comparable to *M. hapla* (Karssen, 2002). The second-stage juvenile drawings for both species descriptions (Fig. 4 E, F & Fig. 7 E–F) show a clearly delimited anterior hyaline tail part, while the original photographs (Fig. 5 F, G & Fig. 9 B) do not show this at all. The fact that both descriptions did not include the hyaline tail measurements (a standard procedure), suggest strongly that the hyaline tail part is not clearly defined. Also in live second-stage juveniles we did not observe a clearly defined hyaline tail part (Table 2).

**Table 4.** Morphometrical (in  $\mu\text{m}$ ) observations (mean, SD & range) of male *Meloidogyne mayaguensis* and *M. enterolobii* paratypes compared to described data.

Species	<i>M. mayaguensis</i>		<i>M. enterolobii</i>	
	description	observed	description	observed
N	30	7	20	11
Body length	1503 $\pm$ 142 (1175–1742)	1431 $\pm$ 63 (1337–1496)	1600 $\pm$ 160 (1349–1913)	1230 $\pm$ 316 (865–1667)
Greatest body width	37.8 $\pm$ 3.1 (32.2–44.4)	34.5 $\pm$ 1.9 (32.0–37.4)	42.3 $\pm$ 3.6 (37.0–48.3)	32.0 $\pm$ 6.0 (23.7–39.2)
Stylet length	22.9 $\pm$ 0.8 (20.7–24.6)	22.1 $\pm$ 0.7 (20.8–23.0)	23.4 $\pm$ 1.0 (21.2–25.5)	21.5 $\pm$ 1.7 (19.2–23.4)
Stylet knob height	3.0 $\pm$ 0.3 (2.4–3.7)	3.2 $\pm$ 0.3 (2.6–3.4)	3.3 $\pm$ 0.3 (2.6–3.9)	2.5 $\pm$ 0.3 (2.1–3.2)
Stylet knob width	5.0 $\pm$ 0.3 (4.3–5.6)	5.3 $\pm$ 0.5 (4.5–5.8)	5.4 $\pm$ 0.3 (4.5–5.8)	4.5 $\pm$ 0.6 (3.5–5.0)
DGO	4.1 $\pm$ 0.4 (3.3–5.0)	4.1 $\pm$ 0.7 (3.2–5.1)	4.7 $\pm$ 0.4 (3.7–5.3)	4.7 $\pm$ 0.6 (3.7–5.8)
Excretory pore to head end	166.4 $\pm$ 8.8 (147.2–180.8)	158.6 $\pm$ 14.9 (132.5–177.9)	178.2 $\pm$ 11.2 (159.7–206.2)	155.8 $\pm$ 22.3 (129.9–199.7)
Spicule length	28.3 $\pm$ 1.5 (24.4–31.3)	29.0 $\pm$ 2.4 (25.6–32.3)	30.4 $\pm$ 1.2 (27.3–32.1)	28.0 $\pm$ 1.1 (26.2–29.4)
Gubernaculum length	7.1 $\pm$ 0.6 (6.1–9.3)	7.5 $\pm$ 1.0 (6.4–9.0)	6.2 $\pm$ 1.0 (4.8–8.0)	6.5 $\pm$ 0.8 (6.1–8.0)
Tail length	14.3 $\pm$ 1.1 (11.3–16.3)	13.0 $\pm$ 1.1 (10.9–14.7)	12.5 $\pm$ 2.2 (8.6–20.2)	11.9 $\pm$ 1.2 (10.2–13.4)
A	39.9 $\pm$ 3.9 (31.1–49.6)	41.6 $\pm$ 2.9 (37.2–44.7)	37.9 $\pm$ 3.2 (34.1–45.5)	38.1 $\pm$ 4.0 (30.0–43.4)
C	105.7 $\pm$ 10.0 (85.8–124.3)	110.5 $\pm$ 10.8 (98.5–133.7)	131.6 $\pm$ 24.2 (72.0–173.4)	103.2 $\pm$ 23.7 (71.4–135.9)

## Morphometrics

The morphometrical characters between the types of *M. mayaguensis* and *M. enterolobii* (Table 3–5), are comparable for the described and observed data, i.e. all mean data are the same or at least within the calculated range. Body length and body width data are generally slightly smaller when comparing observed to described data, this is a well known effect due to a slight shrinking of the nematode body within permanent slides. For *M. enterolobii* males we noticed however an unusual difference in greatest body width between the described 42.3  $\mu\text{m}$  (37–48  $\mu\text{m}$ ) and observed 32.0  $\mu\text{m}$  (24–39)  $\mu\text{m}$  data. The differences can not only be explained due to a shrinking effect, particularly as the observed greatest body width data agrees with the observed data for *M. mayaguensis*. Also for the *M. enterolobii* female holotype unexplainable differences were noticed between described and observed data for the DGO (3.7  $\mu\text{m}$  versus 4.8  $\mu\text{m}$ ) and stylet length (13.4  $\mu\text{m}$  versus 14.7  $\mu\text{m}$ ).

The described and discussed *M. mayaguensis* differences (see diagnosis original description) within the female perineal pattern for the interphasmidial distance, vulval



**Table 5.** Morphometrical (in  $\mu\text{m}$ ) observations (mean, SD & range) of second-stage juvenile *Meloidogyne mayaguensis* and *M. enterolobii* paratypes compared to described data.

Species	<i>M. mayaguensis</i>		<i>M. enterolobii</i>	
	description	observed	description	observed
N	35	25	30	25
Body length	454 $\pm$ 28 (390–528)	420 $\pm$ 21 (386–456)	437 $\pm$ 17 (405–473)	408 $\pm$ 18 (380–442)
Greatest body width	14.7 $\pm$ 0.5 (13.8–15.8)	13.9 $\pm$ 0.7 (13.1–15.4)	15.3 $\pm$ 0.9 (13.9–17.8)	14.8 $\pm$ 2.1 (11.0–18.0)
Body width at anus	10.9 $\pm$ 0.5 (10.2–12.2)	9.8 $\pm$ 0.6 (9.0–11.2)	–	9.8 $\pm$ 0.9 (8.0–11.0)
Stylet length	11.6 $\pm$ 0.3 (11.1–12.2)	11.5 $\pm$ 0.4 (10.9–12.1)	11.7 $\pm$ 0.5 (10.8–13.0)	11.3 $\pm$ 0.7 (10.5–13.0)
Stylet base to head end	15.2 $\pm$ 0.3 (14.8–15.8)	15.4 $\pm$ 0.3 (14.7–16.0)	–	15.0 $\pm$ 0.7 (14.0–16.0)
Stylet knob height	–	1.5 $\pm$ 0.1 (1.2–1.7)	1.6 $\pm$ 0.1 (1.3–1.8)	1.8 $\pm$ 0.3 (1.5–2.0)
Stylet knob width	–	2.5 $\pm$ 0.2 (2.2–2.9)	2.9 $\pm$ 0.3 (2.4–3.4)	3.0 $\pm$ 0.4 (2.5–4.0)
DGO	3.9 $\pm$ 0.2 (3.3–4.3)	3.7 $\pm$ 0.4 (3.2–4.2)	3.4 $\pm$ 0.3 (2.8–4.3)	3.8 $\pm$ 0.3 (3.0–4.5)
Excretory pore to head end	87.6 $\pm$ 3.3 (79.9–97.9)	88.3 $\pm$ 3.0 (83.5–95.3)	91.7 $\pm$ 3.3 (84.0–98.6)	80.8 $\pm$ 4.4 (70.0–88.0)
Tail length	54.4 $\pm$ 3.6 (49.2–62.9)	54.2 $\pm$ 2.7 (48.7–58.5)	56.4 $\pm$ 4.5 (41.5–63.4)	52.1 $\pm$ 3.4 (45.0–57.0)
a	30.9 $\pm$ 1.9 (26.4–34.7)	30.1 $\pm$ 1.6 (26.9–32.8)	28.6 $\pm$ 1.9 (24.0–32.5)	28.0 $\pm$ 3.7 (23.3–34.6)
c	8.3 $\pm$ 0.4 (7.0–9.2)	7.8 $\pm$ 0.3 (7.1–8.4)	7.8 $\pm$ 0.7 (6.8–10.1)	7.9 $\pm$ 0.6 (7.0–9.0)
Excretory pore (%)	19.4 $\pm$ 1.0 (17.8–22.3)	21.1 $\pm$ 0.9 (19.2–22.7)	–	19.8 $\pm$ 1.1 (17.6–21.9)

slit length and vulva-anus distance is not confirmed by our observations. All these measurements are within the observed range. Perineal pattern measurements are generally highly variable and a logical reason for Jepson (1987) not to list this type of data when discussing differential characters for the genus *Meloidogyne*.

### Reproduction and cytogenetics

The two species descriptions report also on the mode of reproduction and number of chromosomes, both reproduce by mitotic parthenogenesis (= apomixes) and have a somatic chromosome number of  $2n = 44\text{--}45$  for *M. mayaguensis* and  $2n = 44\text{--}46$  for *M. enterolobii*. In conclusion, both species have the same mode of reproduction and somatic chromosome number.

## Host plants

Additionally, both species descriptions report in their introduction part some hosts, i.e. they both previously applied the North Carolina differential host test (Hartman and Sasser, 1985). Both species showed the same positive host response for tobacco, pepper, watermelon and tomato and no host response on peanut. Beside this, *M. mayaguensis* did not infest cotton, while *M. enterolobii* moderately infested cotton. As the details of the previously applied host tests have not been described in the material and method part of the species descriptions, we can not explain the reported host response differences on cotton for *M. mayaguensis* and *M. enterolobii*. Interesting is the *M. mayaguensis* study by Brito et al. (2004) with four isolates from Florida (USA). All four isolates, maintained on tomato, reproduced also on cotton, tobacco, pepper and watermelon but not on peanut, i.e. identical to the published results for *M. enterolobii*.

## Isozymes

The observed esterase (VS1-S1 type) and malate dehydrogenase (N1a type) isozyme patterns are identical for both species and agrees with previous results (Carneiro et al. 2000; Xu et al. 2004).

## Conclusion

In conclusion, the holo- and paratype material of *Meloidogyne mayaguensis* and *M. enterolobii* is morphological and morphometrical identical and it confirms the taxonomical status of *M. mayaguensis* as a junior synonym for *M. enterolobii*.

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