Infectivity of entomogenous nematodes (Steinernematidae and Heterorhabditidae) to *Pediculus humanus capitis* De Geer (Anoplura: Pediculidae)

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Summary – The susceptibility of human head lice, *Pediculus humanus capitis*, to three species of Argentinean entomogenous nematodes, *Steinernema rara*, *S. feltiae*, and *Heterorhabditis bacteriophora* (Oliva and Rio Negro strains), was studied. All species and strains, except *S. feltiae*, killed adult and nymph head lice. None of the species killed the eggs. *S. rara* and the Rio Negro strain of *H. bacteriophora* killed both adults and nymphs; on the contrary *H. bacteriophora* Oliva was more aggressive to adults. Apparently, penetration into the lice body takes place through the spiracles, which means that the body size of the infective juveniles is a limiting factor. This is the first report of parasitism in head lice by entomogenous nematodes.

Résumé – Pouvoir infestant de nématodes entomopathogènes (Steinernematidae et Heterorhabditidae) envers Pediculus humanus capitis De Geer (Anoplura: Pediculidae) – La sensibilité du pou de tête, Pediculus humanus capitis, aux nématodes entomoparasites Steinernema rara, S. feltiae, Heterorhabditis bacteriophora (souches Oliva et Rio Negro) a été évaluée. Tous ces nématodes (à l'exception de S. feltiae) tuent les adultes et les nymphes, mais non les œufs. La souche Rio Negro de H. bacteriophora et S. rara se montrent bien plus agressives envers les adultes et les nymphes que les autres nématodes. H. bacteriophora souche Oliva est plus agressif envers les adultes. La pénétration des larves infestantes a lieu apparemment par les spiracles et le diamètre du corps des larves infestantes constituerait un facteur limitant. Le parasitisme du pou de tête par des nématodes entomoparasites est relaté pour la première fois.

Key-words : infectivity, Heterorhabditidae, nematode, Pediculus humanus capitis, Steinernematidae.

Nematodes of the genera *Steinernema* and *Hetero-rhabditis* can be used for insect control, as an alternative to chemical insecticides in agriculture (Begley, 1990). Infective juveniles (IJ) of these nematodes are capable of killing a wide range of insects within 24-48 h (Poinar, 1979), and their pathogenicity is associated with lethal bacteria, a nematode toxin (Akhurst & Boemare, 1990), and the ability of the IJ to search, find and penetrate the host (Dadd, 1971, Glazer, 1992).

Besides agricultural applications, steinernematid and heterorhabditid nematodes have been used against insects and arthropods of medical and veterinary significance, including flies, mosquito larvae, and black flies (Begley, 1990), cat flea (Silverman *et al.*, 1982), ticks (Zhioua *et al.*, 1995), and spiders (Poinar, 1989).

It has been demonstrated that S. glaseri, S. carpocapsae ("Mexican" and "Pye" isolates), and H. bacteriophora (strain HP 88) are pathogenic against body lice (Pediculus humanus humanus L.) (Weiss et al., 1993). The present study examined the pathogenicity of four Argentinean isolates, *i.e.*, two Steinernema spp. and two isolates of H. bacteriophora. The nematodes used in this work were: an isolate of *S. feltiae* recently obtained from Los Chorrillos, Córdoba, the isolate "Noetinger" of *S. rara* from Córdoba, and isolates "OLI" and "RN" of *H. bacteriophora* from Córdoba and Rio Negro, respectively. The nematodes were reared on the greater wax moth *Galleria mellonella* following standard methods and were stored in water suspension at 6°C. The head lice were obtained from infested children. The stages considered were adults, nymphs and eggs. The head lice were put in contact with the nematodes (100 IJ per insect) in Petri dishes provided with two moistened filter papers, at 25°C. Mortality caused by the nematode was verified 36 h after exposure by dissecting the dead insects.

Results

The percentage of mortality caused by the different nematode isolates is summarized in Table 1. With the exception of *S. feltiae*, the nematodes tested were effective against adults and nymphs, but the eggs of head lice were not parasitized. The greatest mortality

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Fig. 1. Pediculus humanus capitis (De Geer) infected by entomogenous nematodes. A: Adult nematodes exiting from the abdomen through a rupture in the cuticle. Nematode inside the body; B: In a segment of the leg; C: In the head (Abbreviations: n= nematode; hn=head of the nematode).

was observed in adults (84 %) and nymphs (100 %) infected by *H. bacteriophora* "RN". Exposure of lice to *H. bacteriophora* "OLI" resulted in poor mortality in nymphs (25 %). On the contrary, the mortality caused to adults (65 %) was similar to that caused by *S. rara* to both stages (65, 62,51 %).

Dissection of infected adults and nymphs showed that a first generation of nematodes can develop in lice. The production of viable IJ was not observed. Mortality of nematodes occurs within 36 to 48 h, and a second generation was never found. The body of parasitized insects is more or less transparent and nematodes could be seen in all parts of the body, including thorax, abdomen, head, antennae, and legs (Fig. 1).

Discussion

The data presented here are the first report on the infectivity of entomogenous nematodes in the genera

Lice mortality (in %)				Nematodes (Infective juveniles)		
Adults	Nymphs	Eggs	Total	Species and strain	Length*	Max. diam.*
0	0	0	0	S. feltiae	947 (870-990)	34 (31-38)
62.5	5	0	64.5	S. rara	465 (400-510)	22.3 (19-25)
84	100	0	91	H. bacteriophora RN	530 (510-670)	24.4 (23-26)
65	25	0	46	H. bacteriophora OLI	540 (490-610)	23 (22-25)

Table 1. Mortality in Pediculus humanus capitis (De Geer) caused by different Steinernema spp. and Heterorhabditis strains.

* in µm

Steinernema and Heterorhabditis on P. h. capitis. Differences in infectivity of these nematode species and strains on insect hosts have already been reported (Doucet *et al.*, 1992).

In the present study also, significant differences were observed, from 100 % nymph mortality caused by *H. bacteriophora* "RN" to no infection with *S. feltiae.* Differences in pathogenicity are related to the specificity of nematodes and this can be attributed to several factors: *i*) ecological and mechanical factors (discovery of, and penetration in the insect body: Glazer, 1992; Dadd, 1971); and *ii*), physiological factors (capacity of the bacteria for growing and invading the host hemocel: Glazer, 1992).

Our results shows that *S. rara* and *H. bacteriophora* were able to infect head lice, while *S. feltiae* (the largest of the nematodes tested here) was unable to do so (Table 1).

It has been demonstrated that *Steinernema* IJ use natural openings to enter the haemocel (Poinar, 1979) and that *Heterorhabditis* individuals can also use their dorsal "tooth" for boring into soft cuticular areas (Bedding & Molyneux, 1982). In the case of lice, penetration *via* the oral and anus routes would not be possible because of the buccal apparatus structure and the intense activity of excretion that are characteristic of these insects, which would prevent the entry of IJ (Kaya, 1990). Therefore, spiracles are the most common point of entry. The spiracles of head lice can be open or closed; in our study, the pore diameters were never greater than 30 μ m. This can cause some size limitations for the penetration of nematodes through natural openings (Dadd, 1971).

The aggressiveness of entomogenous nematodes to body lice (Weiss *et al.*, 1993) and head lice (present study) has been demonstrated. Nevertheless, these human parasites cannot be controlled by nematodes because of their habits; however, control by nematode toxins (Akhurst & Boemare, 1990) is a topic worthy of further investigations.

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Screening bananas for root-knot (Meloidogyne spp.) and lesion nematode (Pratylenchus goodeyi) resistance for the Canary Islands

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Summary – Fifteen banana cultivars and accessions of interest to the Canary Islands were evaluated against *Meloidogyne java*nica, *M. incognita* and *Pratylenchus goodeyi*. Most were highly susceptible to both root-knot nematode species, although different degrees of susceptibility were observed. Nearly all of the plants tested were good hosts for *P. goodeyi*, with the exception of cv. Yangambi Km 5 that had a lower root lesion index and supported a significantly lower nematode reproduction. The five local commercial banana cultivars (Gruesa, Pequeña Enana, Williams, Johnson Negrita and Grande Naine) cultivated in the Canary Islands had the highest susceptibility to *M. incognita* and *P. goodeyi*.

Résumé – Tests de résistance du bananier aux nématodes galligènes Meloidogyne spp. et à Pratylenchus goodeyi pour les cultures des Canaries – Quinze cultivars et clones de bananier destinés aux Canaries ont été évalués vis-à-vis de Meloidogyne javanica, M. incognita et Pratylenchus goodeyi. La plupart se sont montrés hautement sensibles aux deux espèces de Meloidogyne, même si des niveaux différents de sensibilité ont pu être détectés. Tous les matériels végétaux testés sont hôtes de P. goodeyi, à l'exception du cv. Yangambi Km 5 qui a montré un indice de lésions racinaires et une reproduction du nématode significativement inférieurs. Les bananiers les plus sensibles à M. incognita et P. goodeyi sont les cinq cultivars commercialisés aux Canaries (Gruesa, Pequeña Enana, Williams, Johnson Negrita et Grande Naine).

Key-words : Meloidogyne javanica, M. incognita, Musa spp. Pratylenchus goodeyi, resistance.

Bananas cultivated in the Canary Island are Cavendish cultivars, which are all susceptible to root-knot and lesion nematodes. Several species of Meloidogyne are common in banana plantations in the Canary Islands, especially at altitudes of less than 300-400 m above sea level (Rodríguez, 1990). Root-knot nematodes have been shown to damage bananas in sandy and loamy soils where they cause yield reductions of over 20% (Rodríguez, 1975). In recent years, damage has become evident in intensive greenhouse operations, although loss estimates for bananas grown under these conditions are unknown. The root lesion nematode Pratylenchus goodeyi causes serious root damage similar to that of P. coffeae, which is also present but rarely observed in the Canary Islands (de Guiran & Vilardebó, 1962). P. goodeyi is widespread at altitudes above 300-500 m in the three major bananaproducing islands - Tenerife, Gran Canaria, and La Palma - and is considered the most important nematode pest attacking bananas in the Canary Islands (Rodríguez, 1975). It is estimated that about 80% of

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cultivated bananas are infected with *P. goodeyi* (Rodríguez, 1990). By comparison with nematicidetreated plots, this nematode causes 16% loss in Gran Canaria (Rodríguez, 1975). This species is also common in cooler subtropical regions and highland environments. It causes damage to bananas in East and Central Africa (Bridge, 1988; Gowen & Quénéhervé, 1990; Price & Bridge, 1995), and to plantains grown at altitudes of more than 900 m above sea level, which are widely cultivated in Cameroon (Bridge *et al.*, 1995).

Breeding Musa for resistance against nematodes has been one of the most neglected approaches as a pest control management alternative (Rowe, 1984; Pinochet, 1992; Sarah & Jones, 1993). Efforts made for detecting sources of resistance and incorporating them into commercial bananas have been mainly oriented against Radopholus similis, the nematode of major concern in the humid tropics (not present in the Canary Islands). Furthermore, bananas are attacked by several important pathogens, some of epidemic proportions, mainly Panama disease (Fusarium oxysporum f. sp. cubense), Black Sigatoka disease (Mycosphaerella fijiensis), and Moko disease (Pseudomonas solanacearum), that have devastated the banana industry during the last 40 years (Stover & Simmonds, 1991). Any attempt at breeding to obtain nematode resistant material against R. similis, P. coffeae or P. goodeyi should incorporate resistance to these diseases, taking into account type of plant material, market, and pest and disease priorities in each geographical region (Rowe & Richardson, 1975; Buddenhagen, 1996). Some of the Musa material assembled for this study have resistance against Fusarium wilt R4 (Panama disease), a disease of concern in the Canary Islands. Also, a few cultivars / accessions are known to have resistance against R. similis, and in one case (Yangambi Km 5) against P. goodeyi (Fogain & Gowen, 1998). Little information is available on the response of this material to root-knot nematodes (*Meloidogyne* spp.).

The purpose of the present study was to evaluate the host suitability of locally cultivated bananas against *Meloidogyne javanica*, *M. incognita*, and *P. goodeyi* in the Canary Islands. Another goal was to screen the currently available accessions in order to establish their potential use as sources of resistance to important pests and diseases of banana (including *P. goodeyi*).

Materials and methods

PLANT MATERIAL

Commercial plant material and germplasm of interest to the Canary Islands was assembled for screening against nematodes. Micropropagated (in vitro) plant material was provided from several sources. Commercial Musa AAA cultivars, including Grande Naine, Johnson, Negrita, Gruesa, and Brier Pequeña Enana, were provided by ICIA and by Cultivos Vegetales in vitro de Tenerife (CULTESA, Tenerife, Canary Islands, Spain). Accessions provided by the International Network of Improvement of Bananas and Plantains (INIBAP) Germplasm Collection were: FHIA 01 (Goldfinger), PV 03.44, PA 03.22, GCTCV119, GCTCV215, Saba, Yangambi Km 5, Pisang Jari Buaya, Pisang Lilin, Bluggoe and Williams. This group was genetically more diverse and it included diploid, triploid and tetraploid material with known sources of resistance to several pests and diseases for use in plant breeding (Table 1).

Plantlets were transferred to 200 cm³ minipots in a 1:1 (v:v) peat (Floratorf®, Floraguard GmbH, Germany) and perlite (Iberperlita®, Stavik S.A., Huesca, Spain) substrate and acclimatized for 3 weeks under controlled mist chamber conditions until the plants grew to a height of 8 to 12 cm. A minimal number of plants were lost during shipping and acclimatization.

NEMATODE INOCULUM

Two root-knot nematode species were originally collected from banana hosts. Meloidogyne incognita was isolated from banana cv. Pequeña Enana in Valleguerra, Tenerife, and M. javanica from banana cv. Williams, in Telde, Gran Canarias. Both isolates were increased on tomato cv. Roma from single-egg-mass cultures. Identification of isolates was made by perineal patterns (twenty females per population) and confirmed by the Random Amplified Polymorphic DNA technique (Cenis, 1993). The nematode inoculum was prepared by shreding infected tomato roots in a blender for 15 s at 14 500 rpm in a 0.12-0.15% NaOCl solution (Hussey & Barker, 1973). Eggs and juveniles (J2) were collected using a 20 µm-pore sieve (600 mesh) and rinsed with tap water. The inoculum was adjusted to deliver a suspension of 2000 nematodes (eggs and J2) per plant through four holes made in the substrate at 4-5 cm from the base of the plant.

A population of *P. goodeyi* was isolated from banana cv. Pequeña Enana in Tacoronte, Tenerife. Nematodes were extracted from infected root tissues and reared monoxenically for several generations on carrot disk cultures (Moody *et al.*, 1973) incubated at 21°C. The inoculum was recovered from carrot disk cultures by adding water to the cultures and collecting the nematodes with a pipette. The suspension was filtered through a 20 μ m pore-screen (600 mesh) and rinsed with tap water. The inoculum of *P. goodeyi* was adjusted to deliver 2000 nematodes per plant, as previously described for root-knot nematodes.

ROOT-KNOT NEMATODE EXPERIMENTS

Following the hardening phase, plants were individually transplanted to 3-liter containers filled with pasteurized 5:1:1 soil (83% sand, 14% silt, 3% clay), peat, and perlite substrate, with pH 7.3, less than 6% organic matter, and a cation-exchange-capacity of less than 13 meq/100 g soil. Two experiments, one per root-knot nematode species, were conducted using two sets of the same plant material. In each experiment, each cultivar/accession was replicated ten times in completely randomized design. Plants were kept in the greenhouse for 2 weeks before nematode inoculation. Plants were harvested 105 (M. javanica) and 125 (M. incognita) days after inoculation. Percentage of galled root system (Barker, 1985), final nematode population level in roots, and number of nematodes per gram of root were determined for each plant. For the extraction of nematodes in roots, the entire root system was weighed, cut into pieces with scissors, thoroughly mixed, and a 10% subsample was shreded in a blender at 14 500 rpm in a stronger solution of NaOCI (0.25-0.30%) for three periods of 15 s, separated by two 5 s-intervals. Eggs and J2 were then concentrated using 150, 25, and 20 µm-pore sieves (100,

Cultivar/ accession	Genome	General information/ outstanding features	Source
Grande Naine	(AAA)	Most grown dessert banana worldwide. In expansion, replacing other cultivars in the Canary Islands	Stover & Simmonds 1991; Galán & Cabrera, 1992
Brier Pequeña Enana	(AAA)	Locally cultivated in Tenerife, Canary Islands	Stover & Simmonds, 1991
Williams	(AAA)	Widely cultivated in the Canary Islands	Galán & Cabrera, 1992
Johnson Negrita	(AAA)	Locally cultivated in La Palma, Canary Islands	CULTESA (pers. comm.)
Gruesa	(AAA)	Locally cultivated in La Palma, Canary Islands	CULTESA (pers. comm.)
FHIA-01 (Goldfinger)	(AAAB)	Resistant to FOC R4*, Black Sigatoka disease**, and <i>Radopholus similis</i>	Rowe, 1984; Stover & Buddenhagen, 1986
PV 03.44 EMB-402	(AAAB)	Resistant to FOC R4	Shepherd et al., 1994
PA 03.22 EMB-404	(AAAB)	Resistant to FOC R4	Shepherd et al., 1994
GCTCV 119	(AAA)	Resistant to FOC R4. Somaclonal mutant from Taiwan	Hwang & Ko, 1987
GCTCV 215	(AAA)	Resistant to FOC R4. Somaclonal mutant from Taiwan	Hwang & Ko, 1987. Tang & Hwang, 1994
Saba	(BBB/ABB)	Resistant to Fusarium wilt	Stover & Simmonds 1991
Pisang Jari Buaya	(AA)	Source of resistance to R. similis	Wehunt <i>et al.</i> , 1978; Pinochet & Rowe, 1979
Yangambi Km 5	(AAA)	Resistant to R. similis and P. goodeyi	Price, 1994; Fogain & Gowen, 1998
Pisang Lilin	(AA)	Source of resistance to Black Sigatoka disease	Stover & Buddenhagen, 1986
Bluggoe	(ABB)	International reference cultivar	Anon., 1994

Table 1. Information on Musa plant material tested against Pratylenchus goodeyi, Meloidogyne incognita and M. javanica in this study.

* FOC R4 = Fusarium oxysporum f.sp. cubense Race 4; ** Black Sigatoka disease = Mycosphaerella fijiensis.

500, and 600 mesh, respectively). Root tissue and debris collected on the 100-mesh sieve were discarded.

LESION NEMATODE EXPERIMENT

This trial was conducted as described for the previous experiments, except that each plant material was replicated eight times instead of ten. Due to the slower nematode build-up in roots, the experiment with *P. goodeyi* was ended at 180 days after inoculation. Final nematode population levels in roots, number of nematodes per gram of root, and root lesion index were assessed for each plant at the end of the experiment. The last parameter, the root lesion index, measures the length of roots with lesions and is expressed as a percentage of the root system (Pinochet, 1988). Nematodes in roots were extracted as described for root-knot nematodes, but without NaOCl.

Plants were watered daily or as needed, and fertilized with Osmocote® Plus (15-10-12 + micronutrients, Sierra Grace España S. A., Tarragona, Spain). Experiments were conducted in a greenhouse where temperatures fluctuated between 22 and 33° C).

DATA ANALYSIS

Results from the three experiments were analyzed by a one-way analysis of variance. Data for final nematode population and nematodes per gram of root were log_{10} (x+1) transformed for analysis. Data for percentages of galled root system and root lesion index were arcsin transformed. Means were compared by Fisher's LSD test ($P \le 0.05$).

Results and discussion

All banana cultivars were susceptible to both rootknot nematode species, although different degrees of susceptibility were detected (Tables 2, 3). Root galling varied widely from 48% (Bluggoe inoculated with *M. javanica*) to 99% (Grande Naine inoculated with *M. incognita*) of the total root system. Higher final nematode populations were obtained with *M. incognita*, probably due to a slightly longer nematode exposure (125 days) than with *M. javanica* (105 days)

Cultivar/ accession	Percentage of galled roots*	Final nematode population in roots	Nematodes per gram of root
Bluggoe	48 a	124 240 a	4 680 <i>ab</i>
FHIA-01 (Goldfinger)	56 <i>ab</i>	141 450 ab	4 710 <i>ab</i>
Pisang Lilin	61 <i>abc</i>	261 380 cd	9 550 fgh
GCTCV 119	63 <i>abcd</i>	164 720 <i>ab</i>	6 390 bcd
Saba	63 bcde	134 780 <i>a</i>	4 370 <i>a</i>
Brier Pequeña Enana	64 bcde	135 970 ab	5 470 abc
GCTCV 215	66 bcde	172 080 abc	5 550 abc
Williams	69 bcdef	139 580 ab	5 780 abc
PA 03.22 (EMB 404)	69 bcdef	443 320 d	14 440 h
Yangambi Km 5	71 <i>cdef</i>	317 990 d	6 990 cde
PV 03.44 (EMB 402)	71 cdef	185 840 abc	6 370 <i>abc</i>
Grande Naine	79 defg	194 310 bc	8 550 def
Johnson Negrita	81 efgh	179 250 abc	10 670 gh
Pisang Jari Buaya	87 gh	188 070 abc	4 520 <i>ab</i>
Gruesa	90 h	144 320 <i>ab</i>	8 120 cdef

Table 2. Galling and reproduction of Meloidogyne javanica on Musa cultivars and accessions 105 days after inoculation with 2000 nematodes per plant.

Data are means of ten replications. Actual data are presented for nematode reproduction based on log_{10} (x+1) transformed values for analysis. Percentage of galling based on arcsin transformed values for analysis. Means in the same columns followed by the same letter do not differ significantly according Fisher's LSD test ($P \le 0.05$).

* Galling based on percentage of total root system galled: 0 = no galls to 100% = totally galled (Barker, 1985).

Table 3. Galling and reproduction of Meloidogyne incognita on Musa cultivars and accessions 125 days after inoculation with 2000 nematodes per plant.

Cultivar/ accession	Percentage of galled roots*	Final nematode population in roots	Nematodes per gram of root
Pisang Lilin	51 a	186 670 <i>a</i>	7 370 <i>a</i>
FHIA-01 (Goldfinger)	54 <i>ab</i>	499 200 bcd	8 050 <i>a</i>
GCTCV 215	63 <i>abc</i>	419 400 abc	7 500 a
Yangambi Km 5	68 <i>abcd</i>	712 800 bcde	7 990 a
GCTCV 119	71 <i>bcd</i>	528 000 bcde	8 670 <i>a</i>
Saba	72 bcd	307 200 ab	4 010 <i>a</i>
PA 03.22 (EMB 404)	74 cde	964 000 bcde	14 440 <i>ab</i>
Pisang Jari Buaya	75 cde	772 800 bcde	10 500 <i>ab</i>
PV 03.44 (EMB 402)	80 <i>de</i>	949 330 bcde	12 590 <i>ab</i>
Bluggoe	86 e	749 330 bcde	8 060 <i>a</i>
Williams	86 e	931 200 bcde	11 870 <i>ab</i>
Johnson Negrita	96 <i>f</i>	1 394 400 de	8 250 <i>a</i>
Brier Pequeña Enana	98 <i>f</i>	1 170 400 ef	10 120 <i>ab</i>
Gruesa	98 <i>f</i>	3 516 000 cde	22 660 b
Grande Naine	99 <i>f</i>	2 978 400 f	19 650 <i>ab</i>

Data are means of ten replications. Actual data are presented for nematode reproduction based on log_{10} (x+1) transformed values for analysis. Percentage of galling based on arcsin transformed values for analysis. Means in the same columns followed by the same letter do not differ significantly according Fisher's LSD test ($P \le 0.05$).

* Galling based on percentage of total root system galled: 0 = no galls to 100% = totally galled (Barker, 1985).

Cultivar/ accession	Root lesion index*	Final nematode population in roots	Nematodes per gram of root
Saba	17 a	402 020 bc	8 070 bc
Yangambi Km 5	24 ab	106 440 <i>a</i>	1 730 a
FHIA-01 (Goldfinger)	24 ab	347 170 <i>b</i>	8 300 <i>b</i>
PV 03.44 (EMB 402)	27 abc	411 270 bcdef	9 250 bcd
Bluggoe	33 bcd	515 090 cdef	17 250 def
GCTCV 119	36 bcd	326 780 bc	8 870 bcd
PA 03.22 (EMB 404)	43 cd	429 050 bcde	9 380 bcd
Pisang Jari Buaya	40 de	557 870 cdefg	10 970 cde
Pisang Lilin	47 de	299 260 bcd	9 320 bcd
GCTCV 215	52 <i>def</i>	451 380 bcdef	10 130 bcd
Gruesa	55 <i>def</i>	903 500 fg	32 140 f
Brier Pequeña Enana	57 <i>def</i>	769 500 efg	23 880 ef
Williams	86 <i>ef</i>	883 950 defg	26 080 ef
Johnson Negrita	76 <i>f</i>	916 150 fg	27 220 f
Grande Naine	79 f	1 162 200 g	29 420 f

Table 4. Root lesion index and reproduction of Pratylenchus goodeyi on Musa cultivars and accessions 180 days after inoculation with 2000 nematodes per plant.

Data are means of eight replications. Actual data are presented for nematode reproduction based on log_{10} (x+1) transformed values for analysis. Percentage of lesion index of the root based on arcsin transformed values for analysis. Means in the same columns followed by the same letter do not differ significantly according Fisher's LSD test ($P \le 0.05$).

* Lesion index of the root, based on the length of roots with lesions expressed in percentage of root system (Pinochet, 1988).

and warmer conditions. Parasitism (nematodes per g root) is considered to be high. Cv. Saba consistently showed the lowest values of parasitism to both root-knot species (slightly over 4000 nematodes per g root).

Limited information is available on the evaluation of *Musa* germplasm for root-knot nematode resistance. In Indonesia, Hadisoeganda (1994) tested 30 local banana cultivars and found them all to be susceptible, meanwhile, in the Philippines, nine local diploid (AA/BB) or triploid (AAA/AAB/BBB) cultivars were reported to be resistant to *M. incognita* (Davide & Marisagan, 1985). Of these, Paa Dalaga (*Musa* BB) is closely related to Saba (*Musa* BBB), which was found to be susceptible in our study.

Our efforts have so far been unsuccessful in finding resistance for use in banana breeding. Future research should consider testing some of the clones found to be resistant in the Philippines, as well as screening untested banana accessions from the INIBAP collection (the most complete and well characterized *Musa* collection), preferably those accessions with *M. balbisiana* parentage that appear to show better potential as sources of root-knot nematode resistance.

Nearly all plant material tested were good hosts for *P. goodeyi* (Table 4). The most susceptible bananas were the five local commercial cultivars (Gruesa, Brier Pequeña Enana, Williams, Johnson Negrita, and Grande Naine) cultivated in the Canary Islands.

A recently released dessert banana, FHIA 01 (Goldfinger), was the best performing commercial cultivar. Yangambi Km 5 was the most interesting accession evaluated in this study as it showed a lower root lesion index than many cultivars/ accessions, and significantly lower population build-up and number of nematodes per g root than the rest of the tested material. Although it is generally regarded as a host, its relative host suitability was considerably lower than that of the other tested banana material. This accession was reported to have moderate resistance to R. similis and P. goodeyi in West Africa (Price, 1994; Fogain & Gowen, 1998). In Costa Rica, Fallas and Marban (1994) found that susceptibility to R. similis was lower in an accession of Yangambi than in other Musa material. However, it is not certain that the accession evaluated by these authors was indeed Yangambi Km 5 (several accessions of Yangambi are available). Yangambi Km 5 is the only known triploid source of resistance to both species of migratory endoparasitic nematodes (R. similis and P. goodeyi). Most sources of resistance and immunity to R. similis are found in wild and commercial diploids of Musa AA (Wehunt et al., 1978; Pinochet & Rowe, 1979; Stover & Buddenhagen, 1986; Pinochet, 1992). The better host suitability of Yangambi Km 5 to P. goodeyi found in our study as compared to that reported by Fogain and Gowen (1998) could be due to differences in screening procedures or in pathogenicity between African and Canary Island populations of *P. goodeyi*, or, more likely, to a combination of both.

In the future, much screening will be needed to detect resistance against *P. goodeyi* at a level superior to that currently available. Furthermore, resistance will need to be sufficiently broad to apply to most existing forms of the nematode pathogen. Mass screening is one choice. A good starting point would be to test several Yangambi accessions and related clones. Another approach would be to evaluate improved diploids from the Honduran program (Stover & Buddenhagen, 1986; Rowe & Rosales, 1994), which from the breeding standpoint are pollen fertile and already incorporate several desired disease resistant traits including *R. similis* resistance.

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