Influence of temperature and soil type on the histopathology of *Meloidogyne incognita* on snap beans

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**ABSTRACT**

Three groups of seedlings of snap bean cultivars Contender and Conquest were planted in Coto clay, Fraternidad silty clay, and San Antón silt loam; inoculated with 10,000 eggs and second stage juveniles of *Meloidogyne incognita* and placed in growth chambers at 15, 20 and 25°C for 45 days. Multinucleate giant cells with thick cell walls and dense cytoplasm surrounded by proliferation of small cells, breakdown of cell walls, and malformation of vascular bundles were observed. No giant cells were observed in bacterial nodules, even though several of them were invaded by nematodes. Cytoplasmic content of the giant cells decreased as temperature increased. Nematode numbers within roots increased as the temperature increased. There was a tendency for the formation of oval or rectangular giant cells around the vascular bundles and round giant cells in the cortex. Giant cells with the largest number of nuclei were observed at 20°C, and fewer at 25°C. Giant cell was largest at 20 and 25°C, especially in cv Contender. Necrosis or mechanic injury due to migration of juveniles was not observed. The nematodes appeared to develop best in Fraternidad soil.

**INTRODUCTION**

Snap beans, *Phaseolus vulgaris* L., are among the most popular fresh vegetables in Puerto Rico (16). Production during 1983-84 was 86,364 kg, whereas importations of canned string beans amounted to 2,387,273 kg (3). In the United States, phytoparasitic nematodes cause losses of 5% of the total snap bean production (9). *Meloidogyne incognita* (Kofoid and White) Chitwood is one of the most important nematodes affecting this crop (13, 18).

It has been reported that the behavior of *Meloidogyne* species is affected by environmental factors such as temperature and soil texture (14, 17). Temperature is of particular importance and often affects the nematode action within the plant.

The histopathology of root-knot nematodes has been studied in several crops (1, 2, 4, 10), but research on nematode effects on snap bean cells has been limited (6). The objective of this study was to determine the influence of temperature and soil type on the histopathology of root-knot nematodes in beans.

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FIG. 1.—Histopathology of roots of snap bean, cv Conquest at 15° C, 45 days after inoculation with *M. incognita*. A) Longitudinal section showing giant cells (gc) close to vascular bundles (vb), proliferation of nuclei (n), thick-cell walls (cw) and cell wall dissolution (d). B) Longitudinal section exhibiting hyperplasia (h) around giant cells (gc) and the aggregation of cytoplasm particles.
FIG. 2.—Histopathology of roots of snap bean, cvs Contender and Conquest at 15° C, 45 days after inoculation with *M. incognita*. A, B) Longitudinal sections of cv Conquest (A) and Contender (B) revealing adult females (af) and immature females (if), dense cytoplasmic content, giant cells (gc), thick-cell walls (cw), mechanic injury (mi) due to female growth and malformation of vascular bundles (vb). C, D) Transverse sections exhibiting very large giant cells (gc) in cv Contender (C) and small ones in cv Conquest (D).
MATERIALS AND METHODS

Experiments were conducted from January to August 1982, to compare the histological effects on snap bean cultivars Contender and Conquest attributable to *M. incognita* when the plants were grown in three different soil types: Coto clay (pH 7.72, organic matter, 3.5%) from Isabela; Fraternidad silty clay (pH 7.31, organic matter, 3.12%) from Lajas; and San Antón silt loam (pH 7.89, organic matter, 2.87%) from Santa Isabel at three constant temperatures: 15, 20 and 25° C.

Pre-germinated bean seeds were planted in 10 cm plastic pots filled with soil previously sterilized with methyl bromide. Pots were maintained in the greenhouse for 7 days, then inoculated with 10,000 eggs and 2nd stage juveniles of *M. incognita* per pot, extracted from infected tomato roots, *Lycopersicon esculentum* cv Rutgers. Inoculum was obtained by the Hussey and Barker method (7). Plants that served as control received tap water only. The plants were then transferred to growth chambers under constant temperature for 45 days. Five plants served as replication for each treatment and were distributed completely randomized in the growth chamber.

Forty-five days after inoculation, histological analyses of root sections were made to determine the effect of the nematode on the root tissue. Samples of infected root pieces were preserved in FAA (formalin aceto-alcohol) according to Johansen’s method (8). Roots were cut into 1-cm pieces, dehydrated in tert-butyl alcohol and infiltrated in paraplast (melting point 56-57° C). Root pieces were sectioned transversally and longitudinally to a thickness of 12 μ with a rotary microtome. Sections were mounted on glass slides, stained with safranine 0 and fast green, and examined microscopically.

RESULTS AND DISCUSSION

Snap bean plants grown at 15° C developed normal roots and foliage both in nematode inoculated and noninoculated plants. Inoculated plants exhibited small, almost undetectable nematode galls. Bacterial nodules were large on all plants. At 20° C, plants grown in Coto clay exhibited small root systems and thin roots, especially in inoculated plants. Root development in noninoculated plants appeared normal. Nematode galls and bacterial nodules were clearly observed in inoculated plants.

At 25° C, all roots from inoculated plants showed numerous nematode galls and bacterial nodules. The latter were smaller at 25° C than at 15 and 20° C. In general, root systems were smaller at 25 than at 20° C. Plants grown in Coto clay showed relatively small root systems. On many occasions nematodes invaded the base of the stems.

Histological sections revealed the presence of many giant cells in all the inoculated plants at all temperature and all soil types. These cells
were larger than normal cells, multinucleate and with dense cytoplasm and thick cell walls (figs. 1A and B). We observed hyperplastic zones and excessive proliferation of completely disorganized small cells surrounding the giant cells (fig. 1B). The presence of this type of giant cells confirms the susceptibility of both cultivars to M. incognita. Martínez's findings on dry bean (10) are similar to ours. According to Fassuliotis (6), giant cells in resistant cultivars are relatively small and are confined to the nematode head area. Plant tissues most frequently invaded by nematodes were the root cortex, pericycle and vascular bundles. These results are consistent with those of Martínez (10), Negrón (11) and Toro (15).

Nematodes were observed in bacterial nodules, but giant cell formation, or structural changes were not associated with them (fig. 3A). Similar results were obtained in soybean by Hussey and Barker (7).

Temperature was the determining factor in nematode development. At 15° C, we observed few juveniles, sausage stage and immature females. No mature females (with egg masses) nor males were observed. At 20° C, almost all developmental stages were observed in all soil types, but juveniles were found only in cv Contender grown in Coto and Fraternidad soils. Immature and mature females were present in all soil types. At 25° C, second stage juveniles were found in all soil types but were more numerous in Coto (fig. 3B) and Fraternidad soils. Sausage stage nematodes were found occasionally in roots of cv Contender grown in Coto and Fraternidad soils. A few immature females were found in all soil types. Mature females were found more frequently in Coto and Fraternidad soils.

No mature females were found at 15° C in any of the soil types, but at 20° C, egg masses containing many eggs were found in all soil types. In the Coto and San Antón soils, eggs had completed only the first stages of embryonic development whereas in Fraternidad soil juveniles had developed. At 25° C, there were many females with large egg masses containing numerous eggs. Many of them had well developed juveniles, and recently hatched 2nd stage juveniles toward vegetative tissues (fig. 3B). Fassuliotis et al. (6), studying susceptible snap bean cultivars, found that changes in soil temperature from 16 to 21° C or 28° C affected female development and increased egg production.

The number of individuals per gall varied from 1 to 9. More nematodes per gall were observed in Fraternidad soil at 15° C in both cultivars, and in cv Contender grown in Coto and Fraternidad soils at 20 and 25° C.

The number of giant cells was high in all treatments irrespective of the stage of nematode development. Giant cells varied in shape from round to ovoid or rectangular, depending on the invaded tissue. A tendency to the formation of long cells was observed when invaded tissues were associated to or near the vascular bundle (fig. 1A), whereas the giant cells were round in the cortex (fig. 2B). Román (12) working with
Fig. 3.—Histopathology of roots of snap bean cvs Conquest and Contender at 25°C, 45 days after inoculation, with *M. incognita*. A) Transverse section of a portion of a bacterial nodule (bn), cv Conquest, exhibiting 2nd stage juvenile (j). B) Longitudinal section, cv Contender, exhibiting 2nd stage juveniles (j) migrating along cortical tissue.
sugarcane roots, observed that giant cells were long in vascular system
tissues. Toro (15) found similar results in several other crops. At 25° C,
giant cells appeared empty with little or no cytoplasm. Cell disorganiza­
tion was evident in root tips. Long cell proliferation was also observed
around giant cells in cv Conquest (fig. 1B). Apparently the stage of de­
velopment of the nematodes and the number of specimens within the
galls affected the appearance and cytoplasmic content of giant cells. The
greater the number of nematodes within a gall and the more advanced
the nematode stage of development, the lower the cytoplasmic content
in giant cells.

The number of nuclei per giant cell varied from 3 to 40. Giant cells
with excessive nuclei proliferation were abundant at 20° but less numer­
ous at 25° C. Multinucleate giant cells were most frequent in cv Conquest.
These differences may be due to cultivars genotype. Giant cells with
large numbers of nuclei were generally long or rectangular, suggesting
a possible association with vascular bundles.

In transverse sections, the space occupied by the nematodes and giant
cells varied between 10-20% at 15° C and between 10-50% at 20° C and
25° C in relation to the total space of the section. No distinctive differ­
ences were observed at 15° C, but giant cells in cv Contender were larger
at 20 and 25° C (fig. 2C, D).

Mechanical injury due to migration of juveniles was not observed in
the root sections. It was observed that cells surrounding female bodies
had mechanical injury due to nematode growth (fig. 2B).

The absence of necrosis in root sections confirmed the susceptibility
of snap bean cultivars Contender and Conquest. Fassuliotis (5) noted
hypersensitivity reactions in root tips of resistant cultivars and observed
that migration of juveniles caused meristem and cortical parenchyma
necrosis.

At the histological level, the effect of soil type on the nematode his­
topathology could not be ascertained since studied parameters resulted
variable and somewhat erratic. However, a tendency to a better
nematode development in Fraternidad soil was observed.

**RESUMEN**

La influencia de la temperatura y el tipo de suelo sobre la histopatología
del Meloidogyne incognita en la habichuela tierna

El efecto a nivel histológico del nematodo nodulador, Meloidogyne in
cognita (Kofoid and White) Chitwood se estudió en dos cultivares de
habichuelas tiernas (Phaseolus vulgaris L.) sembradas en 3 tipos de suelos
y mantenidas a 3 temperaturas. Las plantas de las cultivares Contender y
Conquest se sembraron en tres grupos de tiestos. En cada grupo se usó uno
de los siguientes tipos de suelo; Coto franco-arcilloso, Fraternidad arcillo­
limoso y San Antón franco-limoso. Las plántulas se inocularon con 10,000
huesos y segundos estadios juveniles de M. incognita y se colocaron en cámaras de crecimiento a 15, 20 y 25° C. Estudios histológicos de las raíces de las plantas, 45 días después de la inoculación, revelaron la formación de células gigantes con citoplasma multinucleado, de paredes gruesas, y rodeadas generalmente, por una proliferación de células muy pequeñas. Se observó ruptura de paredes celulares, deformación de los haces vasculares y daño mecánico a las célulasadyacentes al cuerpo de la hembra del nemátodo. Se observaron nematodos invadiendo nódulos bacterianos, pero no así la formación de células gigantes dentro de éstos. Aumentos en la temperatura generalmente ocasionaron disminución en el contenido citoplásmico de las células gigantes. La cantidad de nematodos dentro de las raíces y la etapa de su desarrollo aumentó proporcionalmente con los aumentos en temperatura. Se observó una tendencia a la formación de células gigantes de forma ovalada o rectangular en las zonas cercanas a los haces vasculares y redondeadas en la corteza. Las células gigantes con mayor cantidad de núcleos predominaron a 20°C, mientras que a 25°C disminuyeron. El tamaño de las células gigantes aumentó a 20° y 25° C, especialmente en la cv Contender. No se observó necrosis ni daño mecánico que se pueda atribuir a la migración de etapas juveniles. Se observó una tendencia a un mejor desarrollo del nemátodo en el suelo Fraternidad.

LITERATURE CITED
