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Tratamento de sementes afeta a penetração, colonização e reprodução de *Meloidogyne incognita* em algodão

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-NOTE-

The root-knot nematode [*Meloidogyne incognita* (Kofoid and White)] is the most threatening species in cotton production, mainly in areas with coarse-texture soils (MONFORT et al., 2007). Nowadays, it is widespread in Brazilian cotton plantations in Bahia, Goiás, Paraná and São Paulo states, causing yield losses that range from 10 to 40% (ASMUS & INOMOTO, 2007). Crop rotation is the most effective measure for the management of *M. incognita* in cotton although its application, in Brazil and the United States, is sometimes restrict by the low number of non-hosts available for use, such as peanut (*Arachis hypogaea*) and brachiaria grasses (*Brachiaria spp*.) (ASMUS & INOMOTO, 2007; STARR et al., 2007). Also, genetic resistance is of limited value due to the low availability of high yielding nematode-resistant cultivars (STARR et al., 2007). As a consequence of the constrained utilization of crop rotation and cotton resistant cultivars, the application of granular nematicides, as aldicarb, carbofuran and terbuphos, in the planting furrow is yet widely used, in spite of their elevated costs in comparison to the modest results in terms of yield increase (120 to 180kg of cotton lint) (ASMUS & INOMOTO, 2007).

An abamectin formulation (Avicta) was recently available as a seed treatment for cotton and vegetable crops. Such method is a low-cost treatment, in which abamectin is placed at a close proximity to plant-parasitic nematodes, minimizing some undesirable characteristics of this substance, as low water solubility...
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In addition, comparing to the traditional granular nematicides, the use of seed treatment represents significantly lower environmental risk and toxicity to humans. Taking into account the increasing importance of seed treatment for nematode management, two greenhouse experiments were carried out in order to evaluate the effect of cotton seed treatment with abamectin on *M. incognita* penetration, colonization, and reproduction.

Experiment 1. *Meloidogyne incognita* race 4 was collected from a cotton field in Campo Verde, Mato Grosso state, and maintained alternately on tomato and cotton plants in a greenhouse. Nematodes were extracted from tomato roots using blender and flotation centrifugal technique (COOLEN & D’HERDE, 1972), using 0.5% NaOCl and centrifuged at 580g. The aqueous suspension obtained, containing eggs and juveniles (J2), was used as inoculum. Plastic pots (7.6cm high x 5.5cm diam., 180cm$^3$) were filled with steam-treated sandy clay soil (57% sand, 4% silt and 39% clay, pH 6.3, 15g organic matter dm$^{-3}$), and 1.25mL of the inoculum, containing 1,000 freshly extracted specimens (918 eggs and 82 J2), were transferred to a central 2-cm hole in the substrate. The surface was covered with the same substrate and the pots were kept for three days (average temperature: 25°C) to allow the hatching of J2. After this period, one seed of a susceptible cotton cultivar, Fibermax 966, not treated with abamectin, was used as inoculum. Plastic pots (7.6cm high x 5.5cm diam., 180cm$^3$) were filled with steam-treated sandy clay soil (57% sand, 4% silt and 39% clay, pH 6.3, 15g organic matter dm$^{-3}$), and 1.25mL of the inoculum, containing 1,000 freshly extracted specimens (918 eggs and 82 J2), were transferred to a central 2-cm hole in the substrate. The surface was covered with the same substrate and the pots were kept for three days (average temperature: 25°C) to allow the hatching of J2. After this period, one seed of a susceptible cotton cultivar, Fibermax 966, not treated or treated with abamectin (150μg of abamectin per seed), both from Syngenta Proteção de Cultivos Ltda. (São Paulo SP, Brazil), was placed in the same central hole as the nematodes. The germination occurred three days after sowing. The infected plantlets were then assigned to evaluate penetration, colonization and reproduction, using 8 replications per date of sampling.

The effect on the J2 penetration was evaluated 3, 9 and 15 days after germination (dag), when treated and non-treated seedlings were removed from the substrate and their root systems gently washed under tap water. The J2 in the whole root system were then stained using acid fuchsin (DAYKIN & HUSSEY, 1985) and counted under stereomicroscope. The effect on nematode colonization was evaluated at 27dag, by counting the total number of galls and egg masses per root system. For this evaluation, Phloxine B (aqueous solution containing 15mg L$^{-1}$) was used to stain the egg masses. From the remaining plants, half were transferred to 500-cm$^3$ plastic pots (11.6cm high x 7.4cm diam) containing steam-treated substrate, in order to assess the nematode reproduction at 50dag. The other plants were transferred to 1,500-cm$^3$ clay pots for assessing the reproduction at 100dag. Eggs and J2 of *M. incognita* were extracted from the whole root system (50dag) and 10g of roots (100dag), using COOLEN & D’HERDE (1972), and the root population was estimated. The experiment was arranged in a complete randomized design, with two treatments and eight replicates for each sampling date. The data were analyzed by ANOVA, using SAS statistical software (SAS Institute, 2003).

Experiment 2. The aim was to confirm the results obtained in experiment 1 concerning the effect of abamectin on J2 penetration. Freshly extracted inoculum of *M. incognita* was obtained from tomato roots and inoculated (682 eggs and 318 J2 per pot) into steam-treated substrates as described previously. The germination occurred four days after sowing and the plants were evaluated 1, 3, and 9dag. The juveniles in the roots were stained using acid fuchsin and counted under stereomicroscope. The experiment was arranged in a complete randomized design, with two treatments and eight replicates for each sampling date.

The seed treatment caused a decrease in the *M. incognita* J2 penetration (87.1 and 81.9%, 9dag in the Experiments 1 and 2) as evaluated in experiment 1 (Table 1), result that was confirmed in the experiment 2 (average number of J2 counted in the roots 1, 3 and 9dag: 1.9, 1.9 and 3.5 for treated plants and 5.4; 12.9 and 19.3 for non-treated ones). When abamectin is applied to cotton seed, it concentrates on the seed coat (FASKE & STARR, 2007) and is highly effective on the soil nematodes in the vicinity. Thereafter, only few J2 are able to penetrate the roots from treated seeds.

Abamectin is a mixture of two closely related compounds, avermectin B1a and avermectin B1b, produced by the soil fungus *Streptomyces avermitilis*. Commercial formulations of abamectin are available for foliar spray applications to control insects and mites. The use as a nematicide is limited by its low solubility in water, tightly binding to soil particles and vulnerability to microbial degradation (WISLOCKI et al., 1989). However, it proved to be effective when applied in close proximity to nematodes. Using an *in vitro* assay of nematode mobility, LD50 value of 1.56μg mL$^{-1}$ was calculated for *M. incognita* based on 2h exposure (FASKE & STARR, 2006).

In both experiments, only infective juveniles were found at 3 and 9dag. The post infection development of juveniles was not affected by abamectin, as 94% of juveniles found within the roots from treated seeds at 15dag were swollen forms, a value very close to the 90% obtained from roots of non-treated seeds (data not shown). Such result corroborates that abamectin is not taken up from soil into plants (WISLOCKI et al., 1989). Lower colonization and reproduction of *M. incognita* were attained in cotton plants from treated seeds in experiment 1 (Table 1). Roots from treated seeds were clearly less galled than from non-treated seeds. These results were likely indirect effects, considering that the population of *M.*...
incognita was diminished by abamectin and consequently fewer J2 were able to penetrate and initiate colonization in the roots. After infection, population increased sharply even more in cotton plants from treated seeds: between 50 and 100 dag. M. incognita population built up 22-fold in plants from treated and 8-fold in plants from non-treated seeds. Probably healthier cotton roots from treated seeds provided more suitable food source for the nematodes.

The results obtained were congruent with the data provided by MONFORT et al. (2006), who demonstrated that root galling was less severe and nematode reproduction was lower in cotton plants from seed treated with 10, 50, 75 and 100 μg of abamectin per seed, at 45 days after cotton sowing. However, in another greenhouse trial (FASKE & STARR, 2007), seed treatment (150 μg per seed) did not reduced galling or nematode reproduction, at 21, 28 and 35 days after cotton sowing. The majority of abamectin applied remains on the seed coat (FASKE & STARR, 2007), and it seems that the seed deposition next to the inoculation site, as used in the experiments here described, improved the efficiency of the treatment. As the roots grow out beyond the soil volume effectively protected by abamectin, the living nematodes are able to infect and colonize them. The relative success in reducing nematode penetration, as demonstrated in the experiments, confirmed that seed treatment with abamectin is a valuable tool in nematode management in cotton culture. Abamectin seed treatment has allowed variable results in suppressing M. incognita in cotton fields (MONFORT et al., 2006). This limitation may be overcome by the adoption of a measure to compensate the immobility of abamectin in soil, eg the concomitant application of soluble nematicides, which reduces nematode population not affected by abamectin. Therefore, it is expected that rather than a substitute to soluble nematicides, seed treatment with abamectin might actually be viewed as a complementary method designed to reduce the total amount of nematicide applied on the ground.

### Table 1 - Effect of the seed treatment with abamectin (150 μg seed⁻¹) on the penetration (3, 9 and 15 days after germination, dag), colonization (27 dag) and reproduction (50 and 100 dag) of Meloidogyne incognita. Experiment 1.

<table>
<thead>
<tr>
<th></th>
<th>3dag</th>
<th>9dag</th>
<th>15dag</th>
<th>27dag</th>
<th>50dag</th>
<th>100dag</th>
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<td></td>
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<td>galls</td>
<td>Egg masses</td>
<td>ng²</td>
<td>RF³</td>
<td>ng</td>
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<td>11.1a</td>
<td>9.6a</td>
<td>41a</td>
<td>34a</td>
<td>1,353a</td>
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<tr>
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<td>86.1b</td>
<td>88.2b</td>
<td>140b</td>
<td>136b</td>
<td>8,763b</td>
</tr>
</tbody>
</table>

1 J2: second stage juveniles; 2 ng: Number of J2 and eggs per gram fresh root; 3 RF: reproduction factor.

### REFERENCES


