Foraging behavior and virulence of some entomopathogenic nematodes

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Abstract

At present the biological control as a pest control technology is becoming more desirable. Biological formulations on basis of entomopathogenic nematodes is one of the effective means for the protection of agricultural and forest plants from harmful insects. Nowadays, the use of entomopathogenic nematodes as biological control agents is a key component in IPM system. The foraging strategies of entomopathogenic nematodes (EPNs) vary between species. This variation is consistent with use of different foraging strategies between ambush, cruise and intermediate to find their host insects. In order to ambush prey, some species of EPNs nictate, or raise their bodies of the soil surface so they are better poised to attach passing insects, other species adopt a cruising strategy and rarely nictate. Some species adopt an intermediate strategy between ambush and cruise. We compared in laboratory the foraging strategies of the entomopathogenic nematode species: Steinernema carpocapsae, Heterorhabditis bacteriophora and the recently described species Steinernema tbilisiensis and assessed their virulence against mealworm beetle, Tenebrio molitor L. (Coleoptera: Tenebrionidae). The tests showed that S. tbilisiensis adopts both foraging strategies.

Keywords: Biological control, Entomopathogenic nematode, Infective juveniles, Foraging strategies, IPM system

1. Introduction

Insect-pathogenic nematodes of the family Heterorhabditidae and Steinernematidae have been known for decades as effective biological agents against insect pests. These nematodes can actively locate, infect and kill a wide range of insect species. Only the third-stage infective juvenile (IJs) can survive outside the insect host and move from one insect to another. Insect mortality, due to nematode infection, is caused by a symbiotic bacterium [1]. Heterorhabditid nematodes have a symbiotic association with Photorhabdus bacteria whereas Steinernematids are associated with Xenorhabdus [2]. After gaining access to the host haemocoel, the bacteria multiply, killing the host within 24-48 h, and convert
the insect into a suitable environment for development and reproduction of the nematodes’ parasitic stages [3].

The foraging strategies of entomopathogenic nematodes vary between species, influencing their soil depth distributions and host preferences. Infective juveniles use strategies to find hosts that vary from ambush to cruise foraging [4].

Ambushers – IJs mostly remain in the same spot for a long period of time waiting for the prey to cross the boundary of their strike area. Chemical cues are not important for them. Nematodes belonging to the category of ambushers are also capable to nictate, i.e. to stand on their tails with more than 75% of the body held straight. Nictation is a relatively stationary tactic which is applied by infective juveniles [5]. Many Steinernema are able to jump by forming a loop with their bodies [6]. Other species adopt a cruising strategy and rarely nictate. Instead, they roam through the soil searching for potential hosts. Crusiers move continuously in the environment in search of hosts hence they may become preys themselves. They largely use long-range chemical cues (carbon dioxide, vibration and other chemical cues) to discover the location of resources [7]. Ambush predators such as Steinernema carpocapsae infect more insects on the soil surface, while cruising predators like Heterorhabditis bacteriophora infect insects that live deep in the soil.

Based on these data we tested and compared the ability of infective Juveniles of three species of EPNs – Steinernema carpocapsae, Heterorhabditis bacteriophora and the autoctone species Steinernema tbilisiensis [8] - to find the host insect Tenebrio molitor L. on 2-dimensional substrates and in sand columns and determined their virulence against it.

2. Material and methods

The recently described species of entomopathogenic nematode, S. tbilisiensis, was isolated from soil samples of the deciduous forest located in the Tbilisi area. Morphological and morphometric data as well as phylogenetic analysis show that S. tbilisiensis belongs to the group Steinernema affine/intermedium. S. tbilisiensis has been attributed to the group S. affine/intermedium on the basis of spicule and gubernaculum structure. The new species differs from other species of the S. affine/intermedium group in the following diagnostic characters: the spicule of S. tbilisiensis is the smallest; and the gubernaculum of S. tbilisiensis is shorter than in other species of the S. affine/intermedium group. Infective juveniles of S. tbilisiensis are distinguished by having a relatively long body (L=866 µm), the position of excretory pore (EP=72 µm), the length of the esophagus (ES=140 µm), the length of the ABW (25 µm). Infective juveniles of S. tbilisiensis have 4 lateral lines like S.beddingi, but the number of lines is 6 in S. affine, S. sichuanense and S. intermedium. Also analysis of rDNA (28S and ITS) gene sequences depict this Steinernema species as a distinct and unique entity.

The mealworm beetle Tenebrio molitor like all holometabolic insects has four life stages: egg, larva, pupa, and adult. The larva of this species has 9 to 20 instars. After the final one it becomes a pupa. Larvae of T. molitor were maintained in laboratory condition at the Ilia State University at room temperature (20–22°C). This insect is typically fed on cereal bran or flour (wheat, oats, maize) supplemented with fresh fruits and vegetables (carrots, potatoes, lettuce) for moisture together with protein sources such as soybean flour, skimmed milk powder or yeast. Also larvae of T. molitor are able to utilize the small amounts of water contained in dry feeds but the productivity of water-deprived is low.
(one generation per year). It is preferable to provide them with a source of water for better productivity (up to 6 generations per year) and to prevent cannibalism. Relative humidity is linked positively with fertility and adult activity. It is necessary to monitor fresh feeds as they may turn mouldy [9, 10].

For cultivation of T. molitor at all stages of their development, larvae were placed in the vessels (60 x 40 cm) with a wide bottom. Dishes were filled with bran and wheat flour mix and with fresh carrot/apple pieces offered at least three times a week. All dishes with larvae were maintained in an environmental chamber at 27 C, 75% RH. Pupae and larvae with sizes from 1.5 to 3.2 cm were used for the experiments.

Nematodes were reared at 25°C in last instar larvae of the wax moth, Galleria mellonella, according to procedures described by Woodring and Kaya [11]. The IJs that emerged from cadavers were recovered using modified White traps and stored at 7°C for 7-14 days before use [12].

Host location and parasitism. Ambushing nematodes would be more effective in seeking hosts on 2-dimensional substrates allowing nictation, whereas cruising species are more effective in a sand column [13].

We compared the proportion of infective juveniles that located and established in a host on a filter paper, sand surface and at the bottom of sand columns. For the filter paper assay dewyfilter paper discs were placed on 10 cm diameter Petri dish whereas for sand substrate 8g sand with 15% moisture was equally distributed on the same size Petri dish. On each substrate 10 host insects’ pupae and larvae were individually placed.

For cruiser nematodes sand columns were prepared by placing three T. molitor pupae and three larvae individually at the bottom of a 2-cm diameter and 18-cm high sand columns which were then filled up to 15 cm with sand.

In all situations insects in the stage of pupa and larva were individually exposed to only one nematode species. The dose of nematode suspension for the 10 insects placed on 2-dimensional nictation substrate made 1000 IJ/ml water, while for the three insects placed in sand columns -300 IJ/ml water (i.e. 100 IJ per insect).

Polyethylene was placed on Petri dishes and glass vials in order to protect them from drying out and two-winged insects. Then they were placed in the incubator at 25°C for 24 hours.

Insects invaded by nematodes were collected after 24 h. Before identifying the number of nematodes invading insects, they were incubated for 48 h at 25°C. The average number of nematodes was determined on the basis of the host insect. The behavior of all three species of nematodes was also observed on two-dimensional substrate for 20, 30 and 50 minutes after the start of the test. Insect’s mortality was recorded, and the number of nematodes established in each insect was determined by dissection. Presence of nematodes inside the insects was checked as indicator of nematode infection.

Control variants were identical to the treatments except that no IJs were added. Each treatment was replicated five times and included untreated control dishes. All experiments were carried out under laboratory conditions at temperature 23°C and 80% RH. Much of this work has been focused on nematode species behavioral interactions with hosts.

3. Results and analysis

H. bacteriophora and S. tsbiliensis caused 100% insects mortality on both surface and at the bottom of sand column after 72 h exposure. However, infective juveniles of S. carpocapsae were more effective on filter paper and sand surface (100% mortality) than in sand columns where insects mortality was 55.5%.
Through the observation under a microscope for 20, 30 and 50 minutes after the start of the test, it was stated that the behavioral interaction of the three species of nematodes on 2-dimensional substrate differed during their contact with the host insect.

About 35-40% of *S. carpocapsae* underwent nictation on sand surface, while others were crawling. Nematode nictation on the filter paper decreased relatively and was observed in about 15-20%. Besides, nematodes found a host insect on a filter paper more easily than on sand surface.

After 48 h of incubation, through dissection of each insect it was stated that infective juveniles of *S. carpocapsae* located and established in host insects’ pupae and larvae more effectively (on average 40-43%) than on sand surface (34-36%). Besides, infective juveniles had a weak reaction on a chemical cues of a host insect, had difficulty contacting with insects placed on the bottom of sand columns and establishing in their bodies. The number of nematodes invading insects did not exceed 3-4% (Figure 1).

The ability to nictate on a sand surface and a filter paper, ineffectiveness in sand columns and a weak reaction to a host’s chemical cues indicate that *S. carpocapsae* in the process of foraging show the properties of ambusher nematodes.

![Bar graph showing average percentage of *S. carpocapsae* established in host insects](image1)

**Fig. 1.** Average percentage of *S. carpocapsae* established in host insects

Only 4-5% of *H. bacteriophora* invaded a host’s pupae and larvae on the filter paper, and an average of 6-8% on a sand substrate.

The performance of nematodes were almost the same on the two types of substrates. Infective juveniles did not nictate on a filter paper. However, this feature was weakly expressed on a sand substrate while approaching and contacting with a host insect. At the same time, nematodes had tracking ability to host’s chemical cues in sand columns, infective juveniles could easily find a remote host and were more effective at establishing in insects in sand columns rather than on filter paper or sand surface. 23-25% of infective juveniles established in insects placed on the bottoms of the sand columns (Figure 2).

Consequently, we believe that *H. bacteriophora* has revealed cruiser nematode’s characteristic behavior that mainly uses chemical cues of a prey to find its location.

The obtained results match the data of Alatorre-Rosas & Kaya [14] which state from the experiments that *H. bacteriophora* has
ability to infect host insects located vertically at 35 cm and horizontally at 30 cm distances and constantly search host insects.

Fig. 2. Average percentage of H. bacteriophora established in host insects

About 20-25% of the new local entomopathogenic nematode S. tbilisiensis experienced nictation on a filter paper and sand surface. They almost equally (14-15%) invaded and established in both hosts on a filter paper and infected them. However, the number of infective juveniles on sand surface invading insects (10-9%) was relatively low. Nematodes positively reacted to the host cues placed on the bottom of sand columns, where the number of infective juveniles reached 14-13% (Figure 3).

Fig. 3. Average percentage of S. tbilisiensis established in host insects

Cruiser nematodes' characteristic response to the host’s signals, ability to nictate and at the same time their effective performance on the sand surface suggests
that *S. tbilisiensis*’ reaction is intermediate between two different strategies and in the process of foraging reveals the qualities of both categories of nematodes. In the three cases the number of nematodes invading pupae was 7-10% more than those invading insect larvae. In the control experiments insect mortality rate was insignificant.

**Fig. 4. Insects infected by entomopathogenic nematodes**

### 4. Conclusion

The action of IJs that located and established in insect on both surface and in sand columns differed among nematodes.

*H. bacteriophora* and *S. tbilisiensis* caused 100% insects mortality on both surface and at the bottom of sand column after 72 h exposure. However, infective juveniles of *S. carpocapsae* were more effective on filter paper and on sand surface than in sand columns where they caused only 55% insects mortality.

The filter paper bioassay is a rapid and simple method to screen for nematode virulence, but removes any environmental barriers to infection, while the sand column bioassays are closer to field conditions. Several previous studies have indicated that the sand column bioassay is a better standard tool for predicting EPN efficacy in a field trial, especially when soil-dwelling insect pests are considered [15, 16, 17].

Ambusher category nematodes search for a host mainly on the soil surface, while cruiser category nematodes are associated with insects inhabiting the lower layers of soil. Therefore, the use of a filter paper and sand cups for ambusher nematodes and sand columns for cruiser nematodes is considered to be the most favorable. In the two types of substrates and sand columns target insects and entomopathogenic nematodes are free in performance.

The foraging strategy of entomopathogenic nematode species can be predicted on the basis of its response to host volatile cues and dispersal behaviour on a 2-dimensional nictation substrate.

The effectiveness of EPNs typically depends on a combination of nematode foraging strategy, insect host species, host location, soil conditions (soil type, pH, soil moisture, etc.), climate and application methods [18].

Laboratory experiments have stated that a cruiser category nematode species does not nictate and completely rely on chemical cues. In contrast, ambusher nematodes nictate and in the process of foraging are less dependent on a host’s chemical cue.

In laboratory an ambusher and a cruiser entomopathogenic nematodes species can coexist. In the field, many other factors will influence the population dynamics, and this higher diversity should promote the coexistence of nematodes species with different foraging behavior [19].

The species of entomopathogenic nematode *S. tbilisiensis*, found in Georgia in the environs of Tbilisi, has shown an intermediate foraging strategy, sharing some
characteristics of both ambush and cruise foragers.

Previous laboratory experiments have demonstrated that *S. tbilisiensis* can be successfully applied as biological control agent. Therefore, the possibility of using this species in biological control programs is of considerable interest.

**References**


