Interaction of the koinobiont parasitoid *Microplitis rufiventris* of the cotton leafworm, *Spodoptera littoralis*, with two entomopathogenic rhabditids, *Heterorhabditis bacteriophora* and *Steinernema carpocapsae*

Atwa A. Atwa1,2a, Esmat M. Hegazi3b, Wedad E. Khafagi4c, and Gehan M. Abd El-Aziz4d

1Deanship of Scientific Research, King Abdulaziz University, Jeddah, Kingdom of Saudi Arabia
2Plant Protection Research Institute, Dokki, Giza, Egypt
3Department of Economic Entomology, Faculty of Agriculture, Alexandria University, Alexandria, Egypt
4Plant Protection Research Institute, Alexandria, Egypt

**Abstract**

Entomopathogenic nematodes are generally considered beneficial organisms. However, they can affect beneficial insects such as parasitoids. The interaction between the entomopathogenic nematodes *Heterorhabditis bacteriophora* Poinar (Rhabditida: Heterorhabditidae) and *Steinernema carpocapsae* Weiser, and the parasitoid *Microplitis rufiventris* Kokujev (Hymenoptera: Braconidae) was investigated in the laboratory. In non-parasitized hosts, *Spodoptera littoralis* Boisduval (Lepidoptera: Noctuidae) larvae exposed to *H. bacteriophora* showed a higher percent mortality than those exposed to *S. carpocapsae*. Both nematodes were able to invade and propagate in non-parasitized *S. littoralis* larvae and those parasitized by *M. rufiventris*. Both nematode species reproduced in *Microplitis*-parasitized hosts, but there was a higher number of nematodes in non-parasitized larvae. *S. carpocapsae* yielded higher numbers of infective juveniles than *H. bacteriophora*. Generally, the number of nematodes harvested increased as their host’s size increased. The interaction between the nematodes and parasitoid favored the nematodes when the nematodes were inoculated during the parasitoid egg stage or the young parasitoid larvae, thus giving the nematodes a better chance to grow and reproduce, resulting in the death of the parasitoid larvae. Conversely, when the nematodes were inoculated during the late larval instar of the parasitoid, the competition partially favored the wasp, thus giving approximately 50% of the wasps a better chance to develop, emerge, and reproduce, providing evidence that both nematodes and wasps could reproduce in the same host. Egg maturation of female wasps derived from nematode-infected hosts was not significantly different than those from control hosts. The combined application of nematodes and parasitoids may be beneficial if the detrimental effects of the nematodes on the parasitoid could be avoided by precisely timing the application strategies. It is clear that *Microplitis* larvae and the nematodes share the host larva and engage in a trophic interaction with each other. Intraguild predation is briefly discussed.
Introduction

The cotton leafworm, *Spodoptera littoralis* Boisduval (Lepidoptera: Noctuidae), is a major plant pest that causes substantial economic losses worldwide. Most control strategies involve chemical insecticides, but this approach is becoming less attractive (Wang et al. 1995) due to resistance, cost, and the lack of availability of pesticides (Mosallanejad and Smagghe 2009). Therefore, biological control has the potential to be a useful strategy. *Microplitis rufiventris* Kokujev (Hymenoptera: Braconidae) is a dominant parasitoid of *S. littoralis*, *S. exigue*, and *Helicoverpa zea* in Egypt (Hammad et al. 1965). This wasp is a specialist endoparasitoid of earlier instars of *S. littoralis* (late 1st to 3rd instar larvae), when they still live in clusters near the place of egg deposition. However, 3rd instars are preferred. Later instars (4th through 6th) disperse, hide under the soil surface in the daytime, and are active at night. As a result, 4th instars are less suitable and less easily accessible than earlier instars for *M. rufiventris* larval development (Hegazi et al. 1977). There is no literature to suggest that *M. rufiventris* females normally attack 5th or 6th instar hosts in the field. The non-preference of later instars has been explained by physiological and host defense traits (Hegazi and Khafagi 2005). The parasitoid oviposits a single egg per host and has three instars that feed on the host hemolymph (Hegazi and Führer 1985).

Among the alternative measures to chemical control of insect pests, in recent years attention has focused on biological control using entomopathogenic nematodes of the families Steinernematidae and Heterorhabditidae (Gaugler 1981; Kaya 1985; Poinar 1986 Georgis et al. 2006). These nematodes have a mutualistic symbiosis with a bacteria (*Xenorhabdus* spp. and *Photorhabdus* spp. for *steinernematids* and *heterorhabditids*, respectively) (Poinar 1990). The third stage, infective juveniles, of the nematodes carries the symbiotic bacteria (*Xenorhabdus in Steinernema*) in a special intestinal vesicle (Poinar 1979; Akhurst 1983; Bird and Akhurst 1983), whereas *Photorhabdus* are primarily located in the anterior part of the guts of the *Heterorhabditis indica* infective juvenile (Boemare et al. 1996). The infective juvenile nematodes are attracted to the insects (Gaugler et al. 1980) and enter via the mouth, anus, or spiracles (Mrácek et al. 1988). *Heterorhabditis* infective juveniles are also able to enter through the insect’s cuticle (Bedding and Molyneux 1982). They penetrate the hemocoel and release the symbiotic bacteria into the insect’s hemolymph. The bacteria then multiply and kill the insect host within 24 hr. Products based on *Steinernema (=Neoaplectana) carpocapsae* Weiser (Rhab-
ditida: Steinernematidae), *S. feltiae* (=bibionis) Bovien (Rhabditida: Steinernematidae) and *Heterorhabditis bacteriophora* (=helioidis) Poinar (Heterorhabditidae) are the most widely commercialized and have almost entirely been marketed as inundative applications in high value niche and specialty markets (Ehlers 1996; Georgis et al. 2006). The pests commonly controlled include soil or root-dwelling pests, and the use of nematodes against above ground pests remains negligible, despite a demand for effective microbial sprays against foliar pests (Cross et al. 1999; Copping and Menn 2000). *S. littoralis* has been shown to be susceptible to nematode infection (Sikora et al. 1979), so we therefore selected *S. littoralis* and its braconid, *M. rufiventris*, as a model system to study the interaction between entomopathogenic nematodes and this koinobiont parasitoid.

**Materials and Methods**

**Insects**

**Rearing of *S. littoralis* and *M. rufiventris***. Cultures of *S. littoralis* and the parasitoid *M. rufiventris* were obtained from a laboratory colony established in 2009 at the Department of Entomology, Faculty of Agriculture, Alexandria University. The colony of *S. littoralis* and *M. rufiventris* originated from field-collected individuals from crops that included cotton in Alexandria, Egypt. However, feral individuals were added to the colonies twice a year to maintain genetic diversity. Larvae of *S. littoralis* were reared on an artificial diet (Hegazi et al. 1977) at 27 ± 1 °C, 60–65% RH, and a 14:10 L:D photoperiod. The *M. rufiventris* colony was maintained using 3rd instar *S. littoralis* larvae as hosts, according to methods described by Hegazi and El-Minshawy (1979). Development of *M. rufiventris* from egg to larval maturity is 8–9 days at 27° C and 65 ± 5% RH (Hegazi and Führer 1985). Under these conditions, the parasitoid egg hatches within a day to a mandibulate 1st instar, which roams inside the host for approximately 4 days, eliminating competitors before developing to a 2nd instar. The 2nd instar molts within 12–16 hr into a final 3rd instar, which lasts for 3 days. The last instar exits the host larva and pupates in a silken cocoon near the host. The host larva does not feed or develop further and dies within 3–12 days (Hegazi and Fuhrer 1985). Mating in *M. rufiventris* wasps occurs as soon as both sexes are present (Hegazi et al. 1977), thus male and female wasps grouped together in glass vials (25 x 100 mm) for 24 hr were presumed to have mated. Groups of presumed mated females (hereafter referred to as mated females) were maintained together with the accompanying males throughout the test period. The wasps were provided with fine droplets of honey diluted (1:1) with distilled water daily to ensure maximum reproductive success.

**Nematode cultures**

The greater wax moth, *Galleria mellonella* (L) (Lepidoptera: Pyralidae), used as a host for nematodes, was obtained from infested hives and reared on an artificial diet at a constant temperature of 27 ± 2° C and 65 ± 5 % RH, as described by Singh (1994). The final instar larvae (25 days old) were utilized for mass rearing of entomopathogenic nematodes. The entomopathogenic nematodes *H. bacteriophora* (Isolate EBN 10K) and *S. carpocapsae* (Isolate EGB5) were isolated from soil samples in El-Nubaria, Behera, and El Badrashin, Giza, Egypt, respectively (Atwa 1999). The nematodes were cultured in the last instar larvae of *G. mellonella*, according to the methods reported by Kaya and Stock (1997). The infective juveniles of both nematodes were harvested in nematode White traps as described by White (1927) at 25 ± 1° C.
A stock suspension of infective juveniles in distilled water was stored at 10°C for 2 weeks before use. Plastic Petri dishes (9 cm x 1.5 cm) lined with filter paper were inoculated with 1200 infective juveniles in 1 mL of water per dish and given 30 minutes to distribute on the filter paper. Each dish was provided with fresh diet (1 x 1 x 2 cm). Five newly molted 3rd instar S. littoralis larvae were added to one dish inoculated with H. bacteriophora nematodes, and a second set of five S. littoralis larvae was added to the second dish with S. carpocapsae nematodes. As a control, the mortality of S. littoralis larvae was followed on filter paper inoculated with 1 mL of distilled water without nematodes in a Petri dish. Ten replicates were used for each treatment.

The dishes were maintained in a climate control chamber at 25°C and allowed to incubate for 24 hr, after which the larvae were transferred to rearing cups with a fresh diet. Dead S. littoralis larvae, 2–3 days post-treatment, from each replicate were transferred to White traps, and the number of infective juveniles produced was counted (Woodring and Kaya 1988). The infective juveniles were collected daily for one week (Shamseldean et al. 1999). The total number of infective juveniles per White trap was divided by the number of nematode-infected S. littoralis larvae to obtain the yield per larva.

**Bioassay methodology**

A series of experiments was conducted to determine how M. rufiventris might interact with the H. bacteriophora or S. carpocapsae nematodes when two species that are competing for the same prey attack the same S. littoralis larva. Groups of 30 3rd instar S. littoralis larvae (determined by the presence of a molted head capsule) were prepared. These larvae were presented individually to female wasps. Oviposition by females (1–2 days old) was observed for individual female in 15 x 60 mm Petri dishes (5–7 females/dish), and only one oviposition was allowed per host larva. Nematode treatments started on days 0, 3, 5, or 7 after parasitism (i.e., times to coincide with the occurrence of egg stage, mid 1st, 3rd, and late 3rd instar of the parasitoid in the host, respectively). The weights for each parasitized S. littoralis host in each test category and those of age-matched non-parasitized hosts were performed to test the hypothesis that infective juvenile production is related to the initial weight of the hosts upon nematode infection.

After each treatment, 10% of each treated group was dissected to ascertain the parasitoid’s developmental stage. If 70% of dissected insects showed the same immature stage, nematode treatment of the corresponding test group would be designated for that stage of development. The larvae were allowed to feed on diet ad libitum for 24 hr, after which time the treated parasitized larvae were reared on a nematode-free diet under the environmental conditions mentioned above until the host died, pupated, or the parasitoid emerged. Mortality counts were recorded daily for five to seven days from the initiation of the experiment. The nematode yield per larva was recorded as mentioned above. As a control, we also determined the mortality of non-parasitized larvae on filter paper inoculated with 1 mL distilled water without nematodes in a Petri dish. Newly formed parasitoid cocoons from nematode-treated host larvae were collected and checked daily for adult emergence. The externally unaffected (normal) parasitoid females obtained from the treated hosts were collected, grouped in pairs, and placed in glass vials (10.3 by 2.3 cm). This was achieved by pairing a female, which resulted from a treated and surviving host larva,
with two males grown using normal laboratory cultures. Honey droplets were smeared on the inner surface of the lid of the rearing vials. Ten females (1 day old) from each nematode-treated group were removed and submersed in 70% ethanol for 10 minutes. Their reproductive tracts were dissected in saline solution. The ovaries were dissected under a binocular dissecting microscope at 40x into the egg tube, reservoir, and calyx. Similarly sized developing eggs were gently teased separately from the egg reservoir and calyx. To standardize the egg counting, only eggs that possessed a distinct opaque area (380–390 μm in length) were counted.

**Statistics**
The experimental design was completely randomized and balanced (equal numbers of subjects were assigned randomly to each treatment group). The data presented as percentages were normalized using a logarithmic transformation. Data were subjected to analysis of variance (one-way ANOVA) for determination of differences between means. Where significant differences occurred, a least significant differences test was applied for mean separation. The level for significance testing was set at \( p < 0.05 \) (Winer et al. 1991). Duncan’s multiple range test or Student’s t-test were applied to significant differences for mean separation. Parameter estimates are given as mean ± 1 SEM unless otherwise stated. (Steel and Torrie 1986).

### Results

**Effect of nematodes on the mortality of *S. littoralis* larvae**
Mortality of non-parasitized *S. littoralis* larvae after exposure to nematodes occurred only between 24 and 72 hr. *S. littoralis* larvae infected with *S. carpocapsae* retained their color, whereas those infected with *H. bacteriophora* developed a red-brown color, which is characteristic of *Heterorhabditis*-infected *G. mellonella* (Woodring and Kaya 1988). The rate of larval mortality (Figure 1) in the *H. bacteriophora* treatments was significantly higher (97.6%) than in the *S. carpocapsae* treatments (81.6%) (\( F = 23.68, \text{df} = 2.27, p < 0.01 \)). The mortality and production of parasitoid progeny for the parasitized *S. littoralis*

![Figure 1. Mean (± SE) percent mortality of 3rd instar Spodoptera littoralis larvae exposed to Steinernema carpocapsae and Heterorhabditis bacteriophora nematodes. Bars bearing the same letter are not significantly different by ANOVA (\( p < 0.01 \)). High quality figures are available online.](image)

**Table 1.** Effects of the timing of the nematode application against *Spodoptera littoralis* parasitized larvae on the percent (± SE) parasitized host mortality and hosts producing parasitoids.

| Trial               | Measurement                  | Non-parasitized | Parasitoid stage at treatment |
|---------------------|------------------------------|-----------------|-------------------------------|
|                     | Weight of hosts (mg)         | Egg             | Mid-1st | Early 3rd | Late 3rd |
| Control             |                              | 2.94 ± 0.08     | 6.62 ± 0.3 | 19.4 ± 1.0 | 27.97 ± 1.0 |
|                     | (2.94 ± 0.08)*               | (2.94 ± 0.08)   | (15.55 ± 0.8) | (60.52 ± 1.3) | (199.23 ± 3.2) |
| *Steinernema carpocapsae* | Hosts produced parasitoids (%) | 99.92 ± 0.05   | 11.76 ± 3.7 | 10.9 ± 1.1 | 16.19 ± 3.6 | 53.2 ± 6.5 |
|                     | Mortality (%)                | 0.8 ± 0.5       | 88.34 ± 3.7 | 89.1 ± 1.1 | 83.81 ± 3.6 | 46.8 ± 6.5 |
| *Heterorhabditis bacteriophora* | Hosts produced parasitoids (%) | 99.9 ± 0.05    | 0.0 ± 0.0   | 0.0 ± 0.0  | 0.0 ± 0.0  | 52.19 ± 8.9 |
|                     | Mortality (%)                | 0.8 ± 0.5       | 100 ± 0.0   | 100 ± 0.0  | 100 ± 0.0  | 47.4 ± 8.6 |

*Numbers in parenthesis represent weights of age-matched non-parasitized hosts.
larvae infected by nematodes at 0, 3, 5, or 7 days post-parasitism are shown in Table 1. Host mortality occurred between 24 and 120 hr. Analysis of variance showed that the percent mortality resulting from the treatments was significantly different (for *Steinernema*, *F* = 111.202, df = 4.45, *p* > 0.01; for *H. bacteriophora*, *F* = 186.89, df = 4.45, *p* < 0.01).

When the nematode *S. carpocapsae* was applied on parasitized hosts either on day 0 or 3 post-parasitism, the percent mortality (Table 1) was slightly increased compared to non-parasitized hosts (Figure 1). However, a significant reduction in the larval mortalities was detected when the nematode treatments were applied on day 7 post-parasitism (Table 1). *H. bacteriophora* nematode treatments on *S. littoralis* larvae on day 0, 3, or 5 post-parasitism resulted in 100% mortality. In contrast, when nematodes were applied on hosts containing late 3rd instars of parasitoid larvae (day 7 post-parasitism), host mortality was significantly reduced by more than 53%. The number of nematode-treated hosts that produced wasps was used to determine the sensitivity of *M. rufiventris* eggs or wasp larvae to nematode infection (Table 1). The mean number of wasps that completed their development under various post-parasitism treatments was significantly different (for *S. carpocapsae*, *F* = 124.729, df = 4.45, *p* < 0.01; for *H. bacteriophora*, *F* = 203.793, df = 4.45, *p* < 0.01). Exposure of developing wasps in the egg stage, mid 1st, and early 3rd larval stages to *S. carpocapsae* infective juveniles via their hosts resulted in a low percentage of hosts that produced wasps (11.7, 10.9, and 16.19%, respectively). However, full-grown parasitoid larvae were partially protected from nematode infection, as 6.5 ± 2.53% completed their development and successfully emerged from treated hosts. When the *H. bacteriophora* infective juveniles were applied on *S. littoralis* hosts either on day 1, 3, or 5 post-parasitism,
none of the hosts produced parasitoids. In contrast, when *H. bacteriophora* infective juveniles were applied to hosts containing older parasitoid larvae (late 3rd instars), 52.19% of parasitized hosts produced parasitoids. Some of the wasp larvae that emerged from nematode-parasitized *Spodoptera* larvae (9.6 ± 2.6%) did not form cocoons and died within 24–48 hr of emergence.

**Nematode yields in parasitized *S. littoralis* larvae**

The yield of nematode infective juveniles in parasitized and non-parasitized *S. littoralis* larvae is shown in Figures 2 and 3. Significant differences were detected in *S. carpocapsae* nematode among parasitized hosts (*F* = 284.409, *df* = 3.36, *p* < 0.01). Application of infective juveniles against host larvae on days 0, 3, 5 or 7 post-parasitism produced (± SE) 4094 ± 163, 1169 ± 135, 8790 ± 231, and 1825 ± 261 infective juveniles per larva, respectively. Corresponding age-matched non-parasitized hosts produced significantly higher numbers of nematodes, with 5045 ± 249, 13,328 ± 821, 48,388 ± 1928, and 87,747 ± 3456 infective juveniles per larva, respectively. In all the *S. carpocapsae* treatments, the number of nematodes harvested was significantly higher in non-parasitized larvae compared to parasitized larvae (t0.05 = 3.3, *df* = 18, t0.05 = 14.607, *df* = 18, t0.05 = 60.708, *df* = 18 and t0.05 = 4.787, *df* = 18, for hosts treated at days 0, 3, 5, or 7 post-parasitism, respectively). *H. bacteriophora* reproduced in the *S. littoralis* host larvae but at a lower multiplicative rate than *S. carpocapsae*. Significant differences were recorded in nematode yields between the different post-parasitism ages of parasitized host larvae (*F* = 193.985, *df* = 3.36, *p* < 0.01). When *H. bacteriophora* infective juveniles were applied on day 0, 3, 5, or 7 post-parasitism, the larvae produced 951 ± 55, 3033 ± 205, 1471 ± 74 and 7289 ± 345 infective juveniles, respectively, versus 1033 ± 66.9, 3913 ± 483, 10,814 ± 673, and 37,184 ± 4152 infective juveniles per age-matched non-parasitized host, respectively. When comparing the nematode yields between parasitized and non-parasitized *S. littoralis* larvae, there were no significant differences in nematode propagation when the nematodes were applied on hosts either 0 or 3 days post-parasitism. However, significant differences were detected when the nematodes were applied on day 5 (t0.05 = 17.79, *df* = 18) or day 7 post-parasitism (t0.05 = 7.18, *df* = 18).

**Egg maturation of female wasps derived from nematode-infected hosts**

Adult *M. rufiventris* females developing from nematode treated *S. littoralis* larvae on days 5 or 7 post-parasitism that appeared morphologically normal were able to find and attack their hosts. The dissected ovaries of mated, but host-deprived, parasitoid females (1 day old) that resulted from nematode treated hosts...
did not show a significantly different number of mature oocytes (Figure 4) compared to control females. The dissection of some mature, full-grown parasitoid larvae derived from nematode-treated hosts containing early 3rd or late 3rd instar parasitoid larvae upon treatment showed that they were nematode free. The oviducts of the adult wasps derived from the control, *Heterorhabditis*- and *Steinernema*-treated hosts, contained a non-significant number of mature eggs (96.7 ± 4.09, 102.8 ± 3.5, and 98.3 ± 1.8 eggs/female, respectively).

**Discussion**

When *S. carpocapsae* and *H. bacteriophora* were screened for their efficacy against 3rd instar larvae of *S. littoralis* using a filter paper bioassay, both nematode species were able to invade and propagate in the larvae tested. The obtained results indicate that *H. bacteriophora* induced significantly higher percent mortality than *S. carpocapsae*. The results are consistent with previous results reported by Abdel-Kawy et al. (1992), who stated that the 3rd and 4th instar of *S. littoralis* larvae were susceptible to infection even at the lowest inoculum levels. Sikora et al. (1979) showed that most developing stages of *S. littoralis* were susceptible to *S. carpocapsae* infection, with the exception of the pre-pupal stage. However, neonate larvae of *S. exigua* were also significantly susceptible to nematode infection of *S. feltiae* (Kaya 1985).

The number of nematode generations inside the host can change according to the different hosts, the size of the host, food availability, the number of infective juveniles that penetrated the host, and the environmental conditions (Griffin et al. 2005; Bazman et al. 2008). The multiplication of nematodes against *S. littoralis* larvae was examined by comparing the number of nematodes produced between parasitized and non-parasitized larvae. The obtained results suggest that the number of nematodes harvested was directly proportional to the weight of the larvae.

The nematode yields were higher in non-parasitized *S. littoralis* larvae than in parasitized larvae. The parasitoid *M. rufiventris* is a polyDNA virus carrying braconid wasp. *S. littoralis* larvae parasitized by this wasp exhibit reduced growth (Hegazi et al. 2005). Therefore, the higher yields of infective juveniles in control hosts may be attributed to larger body weights of non-parasitized larvae compared to parasitized larvae. In all cases, *S. carpocapsae*-infected larvae yielded a higher number of nematodes than *H. bacteriophora*-infected larvae.

The results suggest that in *S. littoralis* larvae, *S. carpocapsae* reproduced more than *H. bacteriophora*. When the eggs and young larvae of *M. rufiventris* were exposed to *S. carpocapsae* or *H. bacteriophora* via *S. littoralis* larvae, the combined application resulted in a higher percent mortality of host larvae compared to the use of either the parasitoid or nematode alone. A higher host-insect mortality was previously observed when entomopathogenic nematodes were combined with parasitoids. For example, Mbata and Shapiro-Ilan (2010) reported that a combination of the nematode *H. indica* and the parasitoid *Habrobracon hebetor* increased the mortality of *Polidia interpunctella*. Additionally, Dillon et al. (2008) observed that the interaction between the nematodes *H. downesi* or *S. carpocapsae* and the parasitoid *B. hylobii* enhanced the mortality of the host insect, *Hylobius abietis*.

Entomopathogenic nematodes are known to have an adverse effect on the development of some parasitoids (e.g., Head et al. 2003;
Lacey et al. 2003). When the nematodes *Heterohabditis downesi* were applied to the gregarious ectoparasitoid *B. hylobii*, which feeds on larvae of the weevil *Hylobius abietis*, the nematodes parasitized the parasitoid larvae, and there was a reduction in parasitoid cocoon formation and fewer cocoons that were enclosed (Everard et al. 2009). It is clear from the experiments reported here that *S. carpocapsae* and *H. bacteriophora* have an impact on the internal developmental stages of *M. rufiventris*. Parasitoid eggs were adversely affected by the nematode treatments, which were initiated concomitant with parasitization. Additionally, if the wasps were in the 1\textsuperscript{st} to early 3\textsuperscript{rd} larval instar stage when *S. littoralis* larvae were parasitized by nematodes, all the wasp larvae in larvae inoculated with *H. bacteriophora* and most inoculated with *S. carpocapsae* died from starvation. The results suggest that nematodes and their associated bacteria rapidly occupy the host larva, dramatically altering its quality for the other organism (*M. rufiventris*). This alternation in resources would effectively starve the young wasp larvae that are not directly killed by nematodes (Everard et al. 2009).

The secondary metabolites produced by entomopathogenic nematodes might also act as antagonistic factors that hinder the development of the young wasps. Therefore, the nematode exclusively developed in the host and induced high mortalities. Parasitoid death due to premature death of the host is the most common consequence of a host-parasitoid-pathogen interaction and has been reported in several laboratory studies using nematodes, primarily *S. carpocapsae*. This is particularly clear in cases where the parasitoid itself is not infected by the nematodes, which was the case for the wasp in our study and the endoparasitoid braconids *Glyptapanteles miliaris* (Kaya 1978), *Apanteles ultor* (Triggiani 1985), and *Myxexoristops* sp. (Mrácek and Spitzer 1983). In our study, the premature death of the host is the most likely cause of parasitoid failure in the experiments where the nematodes were applied on day 0, 3, or 5 post-parasitism. The effect of nematodes on the young stages of *M. rufiventris* appears to be directly antagonistic, the costs of which may be measured in terms of the loss of progeny that fail to complete development, reduced adult size, and increased development time (data not shown). However, when the nematode treatments were performed on the *S. littoralis* larvae on day 7 post-parasitism, the nematodes were not effective in preventing all the parasitoid larvae from emerging, but the emergence time was delayed by two days and costs were less. The full-grown parasitoid larvae were almost completely protected from nematode penetration within their hosts. The reduced sensitivity of late stage parasitoid larvae to nematode infection may make the two compatible in an integrated control program for *S. littoralis* because the nematodes do not kill all the parasitoids.

These findings suggest that the various developmental stages of the parasitoids have varying susceptibilities to entomopathogenic nematodes, thus confirming an earlier observation that later parasitoid stages are less affected by entomopathogenic nematodes than earlier parasitoid stages (Kaya 1987). The latter parasitoid stages may have already developed an effective immunity strategy against nematode infection. The effects of nematode infection were more evident in parasitoid adults that were exposed to nematodes while early 3\textsuperscript{rd} instar. The survival of these adults was significantly reduced ($p < 0.05$) (data not shown). 8–10\% percent of the apparently normal resultant adults died within a few days after emergence. When host larvae were introduced to these adults, no parasitiza-
tion occurred. However, nematode-treatments against late 3rd instar wasp larvae were not effective in preventing significant numbers of wasp larvae from completing their development and emerging.

The *S. littoralis* larvae develop through six larval instars. The first three larval instars feed in groups, leaving the opposite epidermis of the leaf intact. *M. rufiventris* can attack these hosts. The 4th to 6th instar larvae disperse and spend the day in the ground under the host plant, where entomopathogenic nematodes may exist, feeding on plant leaves at night and early in the morning. There is little information on the response of parasitoids to nematode-infected hosts. In the present work, both parasitoid and nematodes targeted the *Spodoptera* larvae, and so there is potential for competition or intraguild predation when two species competing for the same prey attack and consume the food of the one another. Intraguild predation occurs when two species that share a host also engage in a trophic interaction (predation or parasitism) with each other (Rosenheim et al. 1995). Specifically, the entomopathogenic nematodes *H. bacteriophora* or *S. carpocapsae* infect the parasitized host larvae did not infect the mature parasitic larvae, and the presence of nematodes along with the younger parasitic larvae decreased the chance of wasp survival to adulthood. Nonetheless, using both the parasitoid and entomopathogenic nematodes together results in greater overall mortality on *S. littoralis* larvae than either agent inflicts alone. In entomopathogenic nematode-parasitized hosts there was a significant reduction in wasp’s cocoon formation, or no cocoons eclosed at all. Entomopathogenic nematodes are known to interact antagonistically with other competitors, such as entomopathogenic fungi (Barbercheck and Kaya 1991) and parasitoids (Sher et al. 2000; Stuart et al. 2006). The ichneumonids *Mastrus ridibundus* and *Liotryphon caudatus* avoided codling moth hosts previously exposed to *S. carpocapsae* nematodes (Lacey et al. 2003).

No significant differences were observed between the number of mature eggs in the oviducts of 1-day-old females derived from treated hosts and those from non-treated hosts. This study provides evidence that both nematodes and wasps can reproduce in the same host. Therefore, the costs for the parasitoid associated with nematodes attacking the parasitized hosts are dependent on the timing of the application. The ability of these nematodes to avoid the full-grown wasp larvae and survive nematode treatments in parasitized hosts enhances the complementarity of entomopathogenic nematodes and *M. rufiventris*. The interactions between nematodes and wasps in a single host are relevant to application strategies (Barbercheck and Kaya 1991). We conclude from this study that when nematodes and parasitoids are applied concurrently, both compete for the same host, the costs of which are possibly more severe for the parasitoid.

The dual application of parasitoids and nematodes may result in a more efficient control of insects when they are applied sequentially and with the proper timing. Additional studies are needed to further define the interactions in the parasitoid-nematode community in an agro-ecosystem.

**Acknowledgments**

The authors would like to thank the Alexander Von Humboldt Foundation for the research grant supporting this work.
References

Abdel-Kawy AGM, El-Bishry MH, El-Kifl TAH. 1992. Controlling the leopard moth borer, *Zeuzera pyrina* by three entomopathogenic nematode species in the field. *Bulletin of the Faculty Agriculture, Cairo University* 43: 769–780.

Akhurst RJ. 1983. *Neoaplectana* species: Specificity of association with bacteria of the genus *Xenorhabdus*. *Experimental Parasitology* 55: 258–263.

Atwa AA. 1999. Interaction of certain insecticides and entomopathogenic nematodes in controlling some insect pests on fruit and vegetable crops. M.Sc. Thesis, Faculty of Agriculture, University of Ain-Shams, Cairo, Egypt.

Barbercheck ME, Kaya HK. 1991. Competitive interactions between entomopathogenic nematodes and *Beauveria bassiana* (Deuteromycotina: Hyphomycetes) in soilborne larvae of *Spodoptera exigua* (Lepidoptera: Noctuidae). *Environmental Entomology* 20: 707–712.

Barbercheck ME, Kaya HK. 1991. Competitive interactions between entomopathogenic nematodes and *Beauveria bassiana* (Deuteromycotina: Hyphomycetes) in soilborne larvae of *Spodoptera exigua* (Lepidoptera: Noctuidae). *Environmental Entomology* 20: 707–712.

Bazman I, Ozer N, Hazir S. 2008. Bionomics of the entomopathogenic nematode, *Steinernema weiseri* (Rhabditida: Steinernematidae). *Nematology* 10: 735–742.

Bedding RA, Molyneux AS. 1982. Penetration of insect cuticle by infective juvenile of *Heterorhabditis* spp. (Heterorhabditidae; Nematoda). *Nematologica* 28: 354–359.

Bird AF, Akhurst RJ. 1983. The natural of the intestinal vesicle in nematodes of the family Steinernematidae. *International Journal of Parasitology* 13: 599–606.

Boemare NE, Laumound C, Mauléon H. 1996. The nematode-bacterium complexes: biology, life cycle and vertebrate safety. *Biocontrol Science and Technology* 6: 333–345.

Copping LG, Menn JJ. 2000. Biopesticides: a review of their action, applications and efficacy. *Pest Management Science* 56: 651–676.

Cross JV, Soloman MG, Chandler D, Jarrett P, Richardson PN, Winstanley D, Bathon H, Hüb J, Keller B, Langenbruch GA, Zimmerman G. 1999. Biocontrol of pests of apple and pears in northern and central Europe: 1. Microbial agents and nematodes. *Biocontrol Science and Technology* 9: 125–149.

Dillon AB, Moore CP, Downes MJ, Griffin CT. 2008. Evict or infect? Managing population of the large pine weevil, *Hyllobius abietis*. Using bottom-up and top-down approach. *Forest Ecology Management* 255: 2634–2642.

Ehlers RU. 1996. Current and future use of nematodes in biocontrol: practice and commercial aspects with regard to regulatory policy issues. *Biocontrol Science and Technology* 6: 303–316.

Everard A, Griffin CT, Dillon AB. 2009. Competition and intraguild predation between
the braconid parasitoid *Bracon hylobii* and the entomopathogenic nematode *Heterorhabditis downesi*, natural enemies of the large pine weevil, *Hylobius abietis*. *Bulletin of Entomological Research* 99: 151–161.

Gaugler R. 1981. The biological control potential of neoapectanid nematodes. *Journal of Nematology* 13: 241–249.

Gaugler R, Lebeck L, Nakagaki B, Bousch GM. 1980. Orientation of the entomogenous nematodes *Neoaplectana carpocapsae* to carbondioxide. *Environmental Entomology* 9: 649–652.

Georgis R, Koppenhofer AM, Lacey LA, Belair G, Duncan LW, Grewal PS, Samish M, Tan L, Torr P, van Tol RWHM. 2006. Successes and failures in the use of parasitic nematodes for pest control. *Biological Control* 38: 103–123

Griffin CT, Boemare NE, Lewis EE. 2005. Biology and behaviour. In: Grewal PS, Shapiro-Ilan D. Editors. *Nematodes as Biocontrol agents*. CABI.

Hammad SM, El-Minshawy AM, Salama A. 1965 Studies on *Microplitis rufiventris* Kok. (Hymenoptera: Braconidae). *Bulletin of the Entomological Society of Egypt* 49: 215–219.

Hegazi EM, El-Minshawy AM, Hammad SM. 1977. Mass rearing of the Egyptian cotton leafworm, *Spodoptera littoralis* (Boisd.) on semi-artificial diet. In: *Proceedings of the Second Arab Pesticide Conference*. pp. 61–70. Egypt Press.

Hegazi EM, El-Minshawy AM. 1979. Laboratory technique for mass rearing of *Microplitis rufiventris* (Kok) (Braconidae; Hymenoptera) and internal parasites of the cotton leafworm, *Spodoptera littoralis* (Boisd.) (Noctuidae: Lepidoptera). *Bollettino del Laboratoria di Entomologia Agraria. "Filippo Selvestri" di Potici* 36: 205–210.

Hegazi EM, Führer E. 1985. Instars of *Microplitis rufiventris* (Hymoptera; Braconidae) and their relative developmental speed under different photoperiods. *Entomophaga* 30: 231–243.

Hegazi EM, Khafagi WE. 2005. Developmental interaction between suboptimal instars of *Spodoptera littoralis* (Lepidoptera: Noctuidae) and its parasitoid *Microplitis rufiventris* (Hymenoptera: Braconidae). *Archives of Insect Biochemistry and Physiology* 60: 172–184.

Kaya HK, Stock SP. 1997. Techniques in insect nematology. In: Lacey LA, Editor, *Biological Techniques Series: Manual of techniques in insect pathology*. pp. 281–324. Academic Press.

Kaya HK. 1978. Interaction between *Neoaplectana carpocapsae* (Nematoda: Steinernematidae) and *Apanteles militaris* (Hymenoptera: Braconidae). *Journal of Invertebrate Pathology* 13: 358–364.
Kaya HK. 1985. Entomopathogenic nematodes for insect control in IPM system. In: Hass MA, Herzog DC, Editors. *Biological Control in Agricultural IPM Systems*. pp. 283–302. Academic Press.

Kaya HK. 1987. Diseases caused by nematodes. In: Fuxa JR, Tanada Y, Editors. *Epizootiology of Insect Diseases*. pp. 453–470. Wiley.

Lacey LA, Unruh TR, Headrick HL. 2003. Interactions of two parasitoids (Hymenoptera: Ichneumonidae) of codling moth (Lepidoptera: Tortricidae) with the entomopathogenic nematode *Steinernema carpocapsae* (Rhabditida: Steinernematidae). *Journal of Invertebrate Pathology* 83: 230–239.

Mabata GN, Shapiro-Ilan D. 2010. Compatibility of *Heterorhabditis indica* (Rhabditida: Heterorhabditidae) and *Habrobracon hebetor* (Hymenoptera: Braconidae) for biological control of *Plodia interpunctella* (Lepidoptera: Pyralidae). *Biological control* 54: 75–82.

Mosallanejad H, Smagghe G. 2009. Biochemical mechanisms of methoxyfenozide resistance in the cotton leafworm *Spodoptera littoralis*. *Pest Management Science* 65: 732–736.

Mrácek Z, Hanzad R, Kodrik D. 1988. Sites of penetration of juvenile steinernematids and heterorhabditids (Nematoda) into the larvae of *Galleria mellonela* (Lepidoptera). *Journal of Invertebrate Pathology* 52: 477–478.

Mrácek Z, Spitzer K. 1983. Interaction of predators and parasitoids of the sawfly, *Cephalica abietis* (Pamphilidae: Hymenoptera) with its nematode *Steinernema kraussei*. *Journal of Invertebrate Pathology* 42: 397–399.

Poinar GO Jr. 1986. Entomopathogenic nematodes. In: Franz BD, Editor. *Biological Plant and Health Protection*. pp. 95–121. Fisher-Verlag.

Poinar GO Jr. 1979. *Nematodes for Biological Control of Insects*. CRC Press.

Poinar GO Jr. 1990. Taxonomy and biology of Steiner nematid and Heterorhabditididae. In: Gaugler R, Kaya HK. Editors. *Entomopathogenic nematodes in biological control*. pp. 23–61. CRC Press.

Rosenheim JA, Kaya HK, Ehler LE, Marois JJ, Jaffee BA. 1995. Intraguild predation among biological control agents – theory and evidence. *Biological Control* 5: 303–335.

Shamseldean MM, Abd-Elgawad MM, Atwa AA. 1999. Factors affecting pathogenicity of an Egyptian strain of *Heterorhabditis indica* (Nematoda: Heterorhabditidae) infecting the cotton leafworm *Spodoptera littoralis* (Lepidoptera: Noctuidae). *International Journal of Nematology* 9(1): 90–94.

Sher RB, Parrella MP, Kaya HK. 2000. Biological control of the leafminer, *Liriomyza trifolii* (Burgess): implications for intraguild predation between *Diglyphus begini* Ashmead and *Steinernema carpocapsae* (Weiser). *Biological Control* 17: 155–163.

Sikora RA, Salem IEM. Klingauf F. 1979. Susceptibility of *Spodoptera littoralis* to the entomogenous nematode *Neoaplectana carpocapsae* and the importance of environmental factors in an insect control program. *Rijksuniversiteit Faculteit Landbouwwetenschappen* Ghent, 44: 309–322.
Singh SP. 1994. Technology for production of natural enemies. Project Directorate of Biological Control, Bangalore, India. *Technical Bulletin No. 4: 221*.

Steel GD, Torrie JH. 1984. *Principles and Procedures of Statistics*. McGraw-Hill

Stuart RJ, Barbercheck ME, Grewal PS, Taylor RAJ, Hoy CW. 2006. Population biology of entomopathogenic nematodes: Concepts, issues, and models. *Biological Control* 38: 80–102.

Triggiani O. 1985. Influenza dei nematode della famiglia Steinernematidae e Heterorhabditidae sul parassitoide *Apanteles ultiOr* Rhd (Hymenoptera: Braconidae). *La Difesa delle piante* 2: 293–300.

Wang Y, Campbell JF, Gaugler R. 1995. Infection of entomopathogenic nematodes *Steinernema glaseri* and *Heterorhabditis bacteriophora* against *Popillia japonica* (Coleoptera: Scarabaeidae) larvae. *Journal of Invertebrate Pathology* 66: 178–184.

White GF. 1927. A method for obtaining infective nematode larvae from culture. *Science* 66: 302–303.

Winer BJ, Brown DR, Michel SKM. 1991. *Statistical principles in experimental design*, third edition. MacGraw-Hill.

Woodring JL, Kaya HK. 1988. Steinernematid and Heterorhabditid nematodes: a handbook of biology and techniques. *Arkansas Agricultural Experiment Station Southern Cooperative Series Bulletin* volume 331.