Phytoprotection

Canadian entomopathogenic nematode isolates: virulence against black cutworm (Lepidoptera: Noctuidae)
Isolats canadiens de nématodes entomopathogènes : virulence contre les vers-gris noirs (Lepidoptera: Noctuidae)
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Volume 93, Number 1, 2013

URI: https://id.erudit.org/iderudit/1018982ar
DOI: https://doi.org/10.7202/1018982ar
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Publisher(s)
Société de protection des plantes du Québec (SPPQ)

ISSN
0031-9511 (print)
1710-1603 (digital)

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Cite this article

Article abstract
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Canadian entomopathogenic nematode isolates: virulence against black cutworm (Lepidoptera: Noctuidae)

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Received 2012-06-28; accepted 2013-04-09

PHYTOPROTECTION 93 : 43-46

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Keywords: Biological control, cutworm, entomopathogenic nematodes, golf courses, Steinernema carpocapsae, Steinernema feltiae, turfgrass.

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Black cutworms used in this study originated from a population infesting a golf course green (Montebello, QC, Canada, 45°39’N, 74°57’W). The BCW colony was maintained in an environmental chamber following the method described in Reese et al. (1972). All experiments were conducted on fifth-instar larvae. Canadian EPN isolates were collected in the provinces of Quebec and Ontario and included four *S. feltiae* (Filipjev) [1Sf (27), 2Sf (37), 3Sf (13), 4Sf (18)], five *S. carpocapsae* [5Sc (34), 6Sc (34), 7Sc (28), 8Sc (27), 9Sc (8)], and one *S. kraussei* (Steiner) [10Sk (3)]; numbers in parentheses correspond to golf courses described in Simard et al. (2007). Prior to performing each assay, indigenous EPN isolates and commercial formulations of *S. feltiae* (CSf; Plant Products Co. Ltd., Laval, QC) and *S. carpocapsae* (CSc; Natural Insect Control Inc., Stevensville, ON) were reproduced once on *Galleria mellonella* larvae (Dutky et al. 1964). Emerging infective juveniles (IJJs) were collected and stored in tap water at 6°C for 2 wk before use. For all experiments, the virulence of each EPN isolate was confirmed on 15 *G. mellonella* larvae before use. In all cases, mortality rate ranged between 90 and 100% at a concentration of 25 IJs/larva.

BCW larvae were placed at the centre of 9-cm-diam petri dishes whose bottom and top contained two 10-cm-diam Whatman No. 2 filter papers. Petri dishes were prepared as follows: 0.5 mL of distilled water on each filter paper, 1 mL of EPNs + distilled water on each filter paper, and 2 g of creeping bentgrass clippings (*Agrostis palustris* Huds. ‘Alpha’) (Gloco, Anjou, QC). Controls consisted of 1.5 mL of distilled water on each filter paper. Petri dishes were placed randomly in an environmental chamber at 24 ± 1°C, 16L:8D, for 72 h. Mortality was recorded 48 and 72 h after treatment application.

The commercial formulations of *S. feltiae* and *S. carpocapsae* were tested against BCW at the following concentrations: 0, 50, 250, 500 and 1000 IJs/larva/petri dish. For each species and concentration, the experimental unit was 10 larvae per dish with three replications (N = 3 x 10), and the experiment was repeated once for a total of 60 larvae per species per concentration. A factorial analysis of variance (ANOVA; 2 nematode species x 5 concentrations x 2 tests) followed by a protected LSD test were used to compare insect mortality between treatments ($\alpha = 0.05$) (SAS Institute Inc. 1999). The effects of nematode species (48 h: $F = 103.40$, df $= 1$, $P < 0.0001$; 72 h: $F = 70.44$, df $= 1$, $P < 0.0001$) and concentrations (48 h: $F = 53.20$, df $= 4$, $P < 0.0001$; 72 h: $F = 89.37$, df $= 4$, $P < 0.0001$) were significant, but the data were pooled since no significant difference was found between tests (48 h: $F = 1.86$, df $= 1$, $P = 0.1806$; 72 h: $F = 0$, df $= 1$, $P = 1.0$). The data were normally distributed and not transformed.

In another test, the virulence of ten indigenous EPN isolates against BCW was compared with the commercial formulations of *S. feltiae* and *S. carpocapsae* at the following concentrations: 0.50 and 250 IJs/larva/petri dish. For each isolate and concentration, the experimental unit was ten larvae per dish with five replications (N = 5 x 10). A factorial analysis of variance (ANOVA; 2 concentrations x 13 isolates) followed by a protected LSD test were used to compare insect mortality between treatments ($\alpha = 0.05$) (SAS Institute Inc. 1999). Both effects, i.e. concentration (48 h: $F = 127.38$, df $= 1$, $P < 0.0001$; 72 h: $F = 158.53$, df $= 1$, $P < 0.0001$) and isolates (48 h: $F = 34.27$, df $= 12$, $P < 0.0001$; 72 h: $F = 35.54$, df $= 12$, $P < 0.0001$), were significant. The data were normally distributed and not transformed.

For the greenhouse tests, 1 mo before nematode inoculation, creeping bentgrass was seeded in plastic pots (11 cm diam, 9.5 cm high) filled with a 80% sand:20% peat (v:v) soil mixture. Turfgrass was trimmed to a 25-mm height 3 d prior to nematode inoculation, and one BCW larva was introduced per pot. The commercial formulation of *S. carpocapsae* was suspended in water and applied using spray bottles calibrated to deliver 50, 250, 500 and 1000 IJs/pot in a standard 3-mL volume per pot. An additional 3 mL of water was applied to drench the nematodes into the soil. For each concentration, the experimental unit was ten larvae with four replications (N = 4 x 10). The experiment was repeated once with three replications (N = 3 x 10). Insect mortality was recorded 5 d post-application. Data were analyzed together since no significant difference was observed between experiments ($F = 0.30$, df $= 1$, $P = 0.5890$).

In the Petri dish tests, both commercial formulations of *S. carpocapsae* and *S. feltiae* caused significant mortality in fifth-instar larvae of BCW (Fig. 1). *Steinernema carpocapsae* showed higher virulence than *S. feltiae*, the two formulations causing 98% and 70% mortality, respectively, at the 1000 IJs concentration after 72 h. No significant increase in mortality was found between concentrations of *S. carpocapsae* at 500 and 1000 IJs/larva/petri dish.

When the indigenous isolates were included, significant differences were observed among EPN isolates after 48 h (50: $F = 9.19$, df $= 12$, $P < 0.0001$; 250: $F = 56.37$, df $= 12$, $P < 0.0001$) and 72 h (50: $F = 8.51$, df $= 12$, $P < 0.0001$; 250: $F = 32.87$, df $= 12$, $P < 0.0001$) for both concentrations tested (Fig. 2). After 48 h, the highest mortality (80%) was obtained with isolate 6Sc at 250 IJs/larva/petri dish. At the low concentration, isolate 6Sc was also the most virulent with 32% mortality. After 72 h, isolates 6Sc (94%) and CSc (85%) caused the highest mortality. Virulence against BCW was similar for Canadian *S. feltiae* isolates and the commercial formulation. The virulence of the *S. carpocapsae* isolate 6Sc (higher than the commercial formulation at the low concentration) is highly desirable for a biological control agent and its potential should be investigated further under natural soil conditions.

*Steinernema carpocapsae* (commercial formulation) was efficient against BCW larvae in greenhouse trials with mortality rates higher than 50% recorded for all concentrations 5 d post-exposure ($F = 14.75$; df $= 4$, $P < 0.0001$). Average BCW mortality (%) for each *S. carpocapsae* concentration was 11%, 53%, 66%, 76% and 88% at 0, 50, 250, 500 and 1000 IJs/larva/pot, respectively. BCW mortality at the highest concentration of 1000 IJs/larva/pot was not significantly different from that recorded at 500 IJs/larva/pot.

Bioassays performed in the laboratory and greenhouse demonstrated the susceptibility of fifth-
Figure 1. Mortality (±SD) of *Agrotis ipsilon* larvae exposed to commercial formulations of *Steinernema carpocapsae* (CSc) and *S. feltiae* (CSf). Within the same species and evaluation time, nematode concentrations followed by the same letter are not significantly different (protected LSD test; *p* < 0.05).

Figure 2. Mortality (±SE) of *Agrotis ipsilon* larvae caused by indigenous EPN isolates (*Steinernema feltiae*: 1Sf, 2Sf, 3Sf, 4Sf; *S. carpocapsae*: 5Sc, 6Sc, 7Sc, 8Sc, 9Sc, and *S. kraussei*: 10Sk) compared to the *S. carpocapsae* (CSc) and *S. feltiae* (CSf) commercial formulations at 50 and 250 IJs/larva. Within the same concentration and evaluation time, nematode isolates followed by the same letter are not significantly different (protected LSD test; *p* < 0.05).
instar BCW larvae to EPNs, particularly to the species *S. carpocapsae*. The virulence of *S. feltiae* was consistently lower than that of *S. carpocapsae* isolates. These results are directly in line with results published previously on the susceptibility of BCW larvae to EPNs (Baur et al. 1997; Ebssa and Koppenhöfer 2011). None of the Canadian EPN isolates outperformed the two commercial formulations tested in controlling BCW. One indigenous *S. carpocapsae* isolate (6Sc) performed as well as the commercial formulation. The *S. kraussei* isolate (originally described on a sawfly host by Mráček 1977) did not exhibit any significant virulence to BCW larvae.

In the turf and lawn industry, EPNs are usually applied at a rate of 250,000 IJs m⁻² (Grewal et al. 2005). Based on field experiments for BCW control in turfgrass in New Jersey, USA (Ebssa and Koppenhöfer 2011), *S. carpocapsae* applied at this rate provided a high and consistent control of BCW larvae on turfgrass. Under Quebec’s weather conditions, we believe that *S. carpocapsae* could be a useful biological tool to control BCW on golf course greens where damage is recurrent. These habitats are quite suitable for the use of EPN applications for the following reasons: 1) low turfgrass mowing height facilitates contact with BCW larvae; 2) irrigation is readily available on golf greens to prevent EPN desiccation; 3) EPN application could be made at the end of the day to minimize EPN exposure to ultraviolet light and improve contact with BCW larvae, which move on the turfgrass surface during the night; 4) adequate equipment is available and employees are well trained. However, the high standards required on golf course greens could limit nematode application to lower maintenance sites and/or those where pesticide restrictions are implemented.

**ACKNOWLEDGEMENTS**

The authors thank Nathalie Dauphinais, Yvon Fournier, Marie-Eve Gosselin, and Nicolas Turgeon for their dedicated technical assistance. This research was conducted through a collaborative research agreement between the Canadian Golf Course Superintendents Association, the Coalition for Responsible Golf, the Canadian Turfgrass Research Foundation, and Agriculture and Agri-Food Canada’s Matching Investment Initiative program.

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