Population dynamics and biocontrol efficacy of the nematophagous fungus *Hirsutella rhossiliensis* as affected by stage of the soybean cyst nematode

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**Abstract**

Monitoring the population dynamics of a biocontrol agent in soil is important for understanding, predicting, and increasing its efficacy. In this study, the population dynamics and the efficacy of a promising biocontrol agent against nematode, the fungus *Hirsutella rhossiliensis*, were investigated in greenhouse experiments with quantitative real-time polymerase chain reaction (PCR) and bioassay. To explore the effects of the fungus on nematode inoculum, soil infested with eggs or second-stage juveniles (J2) of *Heterodera glycines* was used. The results showed that the quantity of *H. rhossiliensis* DNA based on real-time PCR decreased over time, regardless of eggs or J2 as inoculum. The quantity of *H. rhossiliensis* DNA (femtagram (fg)/g soil) was highest (3.83 ± 1.96 × 10^7 fg/g soil) when the fungus was first added to soil and then decreased rapidly to 3.24 ± 1.22 × 10^7 fg/g in egg-infested soil and 7.34 ± 2.94 × 10^7 fg/g in J2-infested soil 17 days after planting, and then declined gradually between 17 and 59 days after planting. The data based on the bioassay showed that the percentage of J2 parasitized by *H. rhossiliensis* decreased throughout the experiment in both egg- and J2-infested soils. *H. rhossiliensis* controlled H. glycines more effectively in J2-infested soil than in egg-infested soil. At the end of the experiments (59 days after planting), nematode suppression and plant growth promotion in J2-infested soil were 79% and 55%, respectively, which were higher than the 34% and 2% in egg-infested soil. In soil that was not inoculated with the fungus, the number of H. glycines was 10 times higher in J2-infested soil than in egg-infested soil, indicating the greater nematode inoculum potential in soil infested with J2. DNA yield of *H. rhossiliensis* was statistically higher in J2-infested soil than in egg-infested soil in earlier period (prior to day 31) (p < 0.05), which is consistent with the hypothesis that the numbers of J2 present during and soon after fungal inoculation are critical for maintaining the population density and biocontrol efficiency of the fungus.

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**1. Introduction**

The soybean cyst nematode (SCN), *Heterodera glycines* Ichinohe, is the most important pathogen of soybean (Wrather and Koenning, 2006). A number of widely used and effective chemical nematicides for controlling it have been banned in recent years due to environmental concerns (Rosskopf et al., 2005). This has fuelled increased interest in finding alternative methods for nematode control, one of which is biocontrol by natural enemies. One such natural enemy is the nematophagous fungus, *Hirsutella rhossiliensis* Minter and Brady, which occurs naturally in agricultural fields, and parasitizes vermiciform stages of many species of soil-borne, plant-parasitic nematodes (Ciancio et al., 1986; Jaffee and Muldoon, 1989; Jaffee et al., 1991; Hay, 1995; Velvis and Kamp, 1995; Liu and Chen, 2000). Because natural and artificial infestations cause high levels of nematode mortality in the field and in the laboratory, *H. rhossiliensis* is an attractive candidate as a potential biocontrol agent (Jaffee and Muldoon, 1989; Liu and Chen, 2001; Chen and Liu, 2005).

*Hirsutella rhossiliensis*, an endoparasitic fungus of nematodes, initiates infection by producing adhesive conidia that adhere to the cuticles of passing nematodes. Only the conidia that are attached to the phialides are infectious (Mclnnis and Jaffee, 1989), and one conidium is generally sufficient to infect a nematode. Once the fungus penetrates the nematode, it proliferates in the nematode body and produces new conidia within a few days (Lackey et al., 1992); these conidia then initiate a new round of infection. Although *H. rhossiliensis* significantly suppressed plant-parasitic nematodes in greenhouse and laboratory assays (Velvis and Kamp, 1996; Amin, 2000; Liu and Chen, 2001; Chen and Liu, 2005), the fungus did not reduce population densities of cyst and root-knot nematodes.
nematodes in some field trials (Tedford et al., 1993; Jaffee et al., 1996; Chen et al., unpublished data). Because the fungus provided inconsistent nematode suppression in different studies, we suspect that a better understanding of its ecology after being introduced into soil is critical for its successful use as an inundative commercial biocontrol agent. Understanding how the fungus does or does not suppress nematode numbers requires detailed knowledge of the population dynamics of both the fungus and its host. However, due to the limitations in methods for quantitatively monitoring the fungus in soil, most studies have inferred changes in _H. rhossiliensis_ numbers by measuring changes in numbers of nematodes parasitized by the fungus, changes in numbers of resident nematodes or changes in plant growth. As a result, we lack direct measures of _H. rhossiliensis_ population dynamics in soil.

When inoculum of a biocontrol agent is added to soil, many factors determine whether the agent will establish and control pest nematodes. Generally, soil is considered resistant to the introduction of new microorganisms (Cook and Baker, 1983; Stirling, 1991; Pereira et al., 1993; Jaffee, 1999). Soil organisms, soil abiotic factors (including pH, soil moisture, porosity) and formulation of the agent affected the biocontrol potential of _H. rhossiliensis_ (Tedford et al., 1992; Jaffee, 1999; Jaffee and Zasoski, 2001). In addition, as an endoparasitic fungus, _H. rhossiliensis_ is considered obligately dependent on nematodes for nutrition in nature, and its parasitism on nematodes was positively correlated with host density (Jaffee et al., 1993; Jaffee and Muldoon, 1995). Lackey and co-workers (1994) reported that the fungus was more effective in soil infested with J2 than in soil infested with egg masses or cysts, but substantial variability occurred among trials. In previous studies, _H. rhossiliensis_ isolate OWVT-1, obtained from a field continuously planted with soybean for more than 27 years, reduced egg and second-stage juvenile (J2) densities of SCN by more than 90% in the greenhouse (Liu and Chen, 2001). However, in the field, the fungus showed inconsistent suppression of SCN (Chen et al., unpublished data). In our preliminary experiment, the fungus also failed to suppress SCN in soil infested with eggs.

The objectives of the present study were to monitor the fate of _H. rhossiliensis_ inoculum after it was added to soil and then to explore the possible effect of the predominant stage of SCN in soil on the efficacy of _H. rhossiliensis_. We used real-time polymerase chain reaction (PCR) to quantify _H. rhossiliensis_ DNA in soil (Zhang et al., 2006) and a bioassay to quantify _H. rhossiliensis_ parasitism of nematodes.

### 2. Materials and methods

#### 2.1. Fungal and nematode inoculum

_Hirsutella rhossiliensis_, isolate OWVT-1, was stored on potato dextrose agar (PDA; Oxoid Ltd., Basingstoke, Hampshire, England) at 4 °C. Fungal inoculum (a mycelial slurry) was prepared by growing the fungus in a liquid medium (Zhang et al., 2006) on an orbital shaker at 150 rpm for 7 days at 25 °C. The fungal colonies were collected and blended following the method of Liu and Chen (2005). The mycelial slurry was collected, and colony forming units (CFUs) per gram of fresh mycelium was determined by plating a series of slurry dilutions on plates with the same nutrition components as the liquid medium. The number of colonies formed on agar after 2 weeks at 25 °C was recorded (CFUs per gram of fresh mycelium was 1.6 × 10⁶ in experiment 1 and 2.5 × 10⁶ in experiment 2).

Soybean cyst nematode race 4 (originally collected from a soybean field in the suburb of Beijing, China) was cultured on soybean cultivar Zhonghuang 13 in autoclaved soil in the greenhouse. The eggs and J2 were prepared from the newly formed cysts (Liu and Chen, 2001).

#### 2.2. Soil

Soil (52% sand, 33% silt, 15% clay and 1.03% organic matter at pH 6.7) was collected from a corn field in Gaomi County, Shandong Province, China. _H. glycines_ was not detected in the soil after wet sieving and sucrose centrifugation (Jenkins, 1964), and _H. rhossiliensis_ was not observed on resident nematodes or on _H. glycines_ J2 added to and extracted from the soil. The field soil was passed through a 2-mm-aperture sieve and stored at room temperature (22–25 °C) for 2 months before being used. For the greenhouse experiment, soil moisture was adjusted to 9% (9 g water/100 g dry soil) and heated in a microwave (1-kg lot of soil in a plastic bag at 800 W for 1.5 min) to eliminate nematodes and other microfauna but to allow most fungi and bacteria to survive (Chen et al., 1995).

#### 2.3. Evaluation of the efficacy of _H. rhossiliensis_ in soil infested with nematode eggs (experiment 1)

A fresh mycelial slurry was added to the field soil (1000 mg wet weight of mycelia/100 g of dry soil), and nematode eggs (5000 eggs/100 cm³ soil) were added at the same time. The soil was mixed thoroughly and placed in 12-cm-diameter pots (240 cm³). Soil without fungal inoculation was used as the control. Soybean seeds of Sturdy variety were surface-disinfested with 0.1% NaClO for 3 min. Four seeds were sown per pot, and pots were covered with a plastic film to reduce evaporation. Pots were completely randomized and maintained in a growth chamber at 20–25 °C at a photoperiod of 12 h of light and 12 h of dark. After 7 days, the plastic film was removed and the seedlings were thinned to 2 plants/pot. Thereafter, pots were watered daily. There were four replications for each treatment (plus or minus added mycelia). At 14, 28, 42, 56 and 70 days after planting, four pots from each treatment were randomly selected and destructively sampled (see Section 2.5).

#### 2.4. Evaluation of the efficacy of _H. rhossiliensis_ in soil infested with nematode eggs or J2 (experiment 2)

The procedures for fungal inoculation and soybean plantation in experiment 2 were identical to those in experiment 1, but the eggs and J2 of _H. glycines_ were used as nematode inoculum, respectively. Four treatments included in experiment 2 were: (1) egg-inoculation—nematode eggs (5000 eggs/100 cm³ soil) were added to soil at the same time as the fungus inoculating; (2) J2-inoculation—J2 (5000 J2/100 cm³ soil) in 6 ml sterilized water were added to each pot after thinning (7 days after planting); (3) egg-control (egg-CK)—soils with eggs and no fungal inoculum and (4) J2-control (J2-CK)—soils with J2 and no fungal inoculum. For all treatments, _H. glycines_ J2 (3000 J2 in 4.5 mM KCl/100 cm³ soil) were added to each pot 3 days before that pot was destructively sampled. Nematodes were then extracted and examined along with resident nematodes (see Section 2.5). The other procedures were identical to those in experiment 2.

#### 2.5. Sampling

Four replicate pots per treatment were destructively sampled every 2 weeks after seeding. The soil and plants in each pot were gently removed and placed on a plastic sheet. Mature females and cysts were dislodged from the roots, and roots were carefully removed from soil. Rhizosphere soil (defined as the soil attached to roots after gentle shaking by hand) was collected by crumbing the soil attached to roots lightly and dried at room temperature for 1 h. The roots were rinsed with tap water and root and shoot dry mass were determined. The bulk soil of each pot was thoroughly mixed.
A subsample of 100 g of soil was used for extraction of eggs and J2 using the sucrose floatation-centrifugation technique (Jenkins, 1964; Liu and Chen, 2001). All extracted nematodes were observed at 100× magnification with an inverted microscope. Nematode eggs and J2 were counted, and the percentage of J2 parasitized by fungi was recorded. Nematodes with attached fungal conidia or those that were colonized by fungal mycelium were considered to be parasitized.

From each pot, 0.5 g of rhizosphere soil (in experiment 1) and bulk soil (in experiment 2) were used for DNA extraction with the MoBio UltraClean soil DNA isolation kit (MoBio Laboratories, Solana Beach, CA, USA) according to the manufacturer’s protocol with minor revision: that is, beating at 4200 rpm for 90 s in a mini-bead beater (Biospec Products, Bartlesville, OK, USA) instead of vortexing to break up cells. The DNA was stored at −20°C before use. Soil DNA was diluted 10-fold and subjected to real-time PCR as described in the next section.

2.6. Real-time PCR quantification of H. rhossiliensis in soil

Hirsutella rhossiliensis DNA in soil was quantified by real-time PCR with specific primers (F419 and R480) and probe (P442) as described previously (Zhang et al., 2006). The TaqMan™ fluorescent PCR was performed in 25-μl reaction mixtures that contained 1× buffer; 2.5 mM MgCl2; 100 μM each dNTP and 1 U Taq DNA polymerase (Promega, Madison, WI, USA) and 2 μl of DNA template. The optimized thermal cycling consisted of a denaturation step of 95°C for 10 min followed by 45 cycles of 95°C for 15 s and 60°C for 1 min. Reactions were carried out in an ABI Prism 7700 Sequence Detector (PE Applied Biosystems, Foster, CA, USA). The amount of initial template DNA was estimated by determining the threshold cycle, the cycle number at which statistically significant increases in the reporter fluorescence were detected. Standard curves were constructed based on real-time PCR amplification of five known H. rhossiliensis genomic DNA concentrations ranging from 1 × 107 to 1 × 103 femtograms (fg).

2.7. Statistical analysis

To determine the biocontrol effectiveness of the fungus, the percent suppression of H. glycines and the percent plant growth promotion were calculated at the end of the experiments. Percent suppression was equal to (1 − x) × 100, where x was the number of nematode eggs per unit of soil in the fungal inoculation treatment divided by the number of nematode eggs per unit of soil in the control. Percent growth promotion was equal to (y − 1) × 100, where y was the plant dry weight in the fungal inoculation treatment divided by the plant dry weight in the control (Lackey et al., 1994). Before being subjected to statistical analysis, nematode population density and fungal DNA yield data were log-transformed and the percentages of J2 parasitized by the fungus were arcsine-transformed to improve variance homogeneity. Analyses of variance were performed using the GLM procedure of SAS for Windows version 8.0. Differences were considered significant at P < 0.05.

3. Results

3.1. Effectiveness of fungal inoculation in soil infested with nematode eggs (experiment 1)

In experiment 1, fungal treatment did not increase plant shoot or root weight (Table 1). Numbers of nematode eggs per cm³ soil did not differ between treatments before day 56 but were less in the fungal treatment than in the control on day 56 and 70 (Table 1).

3.2. The activity and dynamics of H. rhossiliensis in soil infested with nematode eggs (experiment 1)

At each sampling time, J2 were extracted from the soil and examined for fungal parasitism. The percentage of J2 parasitized by H. rhossiliensis was low for all sampling times, ranging from 0.5% to 2.3% (Table 1). No J2 were parasitized by the fungus in soils that were not inoculated with the fungus.

The quantity of H. rhossiliensis DNA as determined by real-time PCR was highest on day 1 and gradually declined over time in soil with fungal inoculation (Fig. 1). H. rhossiliensis was not detected by real-time PCR in soils that were not inoculated with the fungus.

3.3. Biocontrol effectiveness of H. rhossiliensis on H. glycines in soil infested with eggs or hatched J2 (experiment 2)

In experiment 2, soil was infested with eggs and J2 of H. glycines at the same concentration level. Shoot weight did not differ between egg-CK and egg-inoculation treatments until day 59 after planting and were higher in egg-inoculation treatment than in egg-CK treatment on day 59 (F = 7.66; df = 3, 12; P = 0.004; Table 2). J2-inoculation treatment had significantly higher shoot weight than egg-CK treatment after day 17 (F = 7.36; df = 3, 12; P = 0.0044; Table 2). The lowest shoot weight among all treatments was observed in the J2-CK treatment. Root weight in egg-CK, egg-inoculation and J2-inoculation treatments showed no significant difference but were significantly higher than that in J2-CK on day

Table 1

<table>
<thead>
<tr>
<th>Days after planting</th>
<th>Treatmenta</th>
<th>Dry shoot weight (mg)b</th>
<th>Dry root weight (mg)b</th>
<th>1000 eggs gram soilb</th>
<th>Eggs per cm³ soilb</th>
<th>J2 per cm³ soilb</th>
<th>% Parasitized by the fungus</th>
</tr>
</thead>
<tbody>
<tr>
<td>14</td>
<td>CK</td>
<td>233 ± 5 a</td>
<td>33 ± 12 a</td>
<td>17.7 ± 8.0</td>
<td>20.0 ± 9.0</td>
<td>4.0 ± 0.8</td>
<td>0.0 ± 0.0</td>
</tr>
<tr>
<td></td>
<td>Inoculation</td>
<td>223 ± 5 a</td>
<td>43 ± 11 a</td>
<td>15.4 ± 6.9</td>
<td>16.6 ± 7.0</td>
<td>3.8 ± 1.3</td>
<td>2.3 ± 1.2</td>
</tr>
<tr>
<td>28</td>
<td>CK</td>
<td>216 ± 6 a</td>
<td>43 ± 10 a</td>
<td>14.3 ± 5.0</td>
<td>13.7 ± 7.2</td>
<td>2.7 ± 1.9</td>
<td>0.0 ± 0.0</td>
</tr>
<tr>
<td></td>
<td>Inoculation</td>
<td>197 ± 7 a</td>
<td>37 ± 8 a</td>
<td>14.4 ± 2.9</td>
<td>13.8 ± 2.6</td>
<td>4.7 ± 0.5</td>
<td>2.1 ± 0.8</td>
</tr>
<tr>
<td>42</td>
<td>CK</td>
<td>231 ± 31 a</td>
<td>41 ± 6 a</td>
<td>14.0 ± 7.1</td>
<td>16.4 ± 4.6</td>
<td>4.7 ± 1.3</td>
<td>0.0 ± 0.0</td>
</tr>
<tr>
<td></td>
<td>Inoculation</td>
<td>216 ± 44 a</td>
<td>40 ± 7 a</td>
<td>13.2 ± 6.2</td>
<td>13.1 ± 4.0</td>
<td>4.3 ± 0.9</td>
<td>0.5 ± 0.4</td>
</tr>
<tr>
<td>56</td>
<td>CK</td>
<td>250 ± 27 a</td>
<td>44 ± 5 a</td>
<td>13.2 ± 10.0</td>
<td>16.4 ± 0.5</td>
<td>7.2 ± 1.5</td>
<td>0.0 ± 0.0</td>
</tr>
<tr>
<td></td>
<td>Inoculation</td>
<td>214 ± 13 a</td>
<td>48 ± 11 a</td>
<td>9.3 ± 2.8</td>
<td>10.0 ± 3.0</td>
<td>5.4 ± 1.2</td>
<td>1.3 ± 0.6</td>
</tr>
<tr>
<td>70</td>
<td>CK</td>
<td>229 ± 33 a</td>
<td>56 ± 16 a</td>
<td>8.8 ± 1.8</td>
<td>10.8 ± 3.2</td>
<td>4.0 ± 1.6</td>
<td>0.0 ± 0.0</td>
</tr>
<tr>
<td></td>
<td>Inoculation</td>
<td>220 ± 59 a</td>
<td>42 ± 13 a</td>
<td>5.2 ± 2.4</td>
<td>5.4 ± 2.1</td>
<td>3.0 ± 1.7</td>
<td>1.1 ± 0.8</td>
</tr>
</tbody>
</table>

a Treatment: without fungal inoculation (CK), with fungal inoculation (Inoculation).

b Means ± SD (n = 4). Values followed by the same letter in the column within the same days after planting are not significantly different at P > 0.05 according to LSD test. Egg and J2 densities were transformed to log10(x) values before being subjected to statistical analysis.
No J2 was observed to be parasitized by the fungus in egg-CK and J2-CK treatments. The quantitative data from real-time PCR showed that the quantity of *H. rhossiliensis* DNA was highest (3.83 ± 1.96 × 10^8 fg/g soil) on day 1, and decreased significantly to 7.34 ± 2.94 × 10^7 fg/g soil in J2-inoculation treatment ($F = 4.001; df = 4, 15; P < 0.0001$) and 3.24 ± 1.22 × 10^7 fg/g soil in egg-inoculation treatment ($F = 78.51; df = 4, 15; P < 0.0001$) on day 17 (Fig. 2B). The quantity of *H. rhossiliensis* DNA showed no significantly decreasing from day 17 to day 31 in J2-inoculation treatment, and from day 17 to day 45 in egg-inoculation treatment. The lowest DNA yield was observed on day 45 and day 59 in J2-inoculation treatment with 11.1 times lower than day 1, and on day 59 in egg-inoculation treatment with 19.5 times lower than day 1 (Fig. 2B). The DNA yield was significantly higher in J2-infested soil than in egg-infested soil on day 17 ($F = 8.94; df = 1, 6; P = 0.0243$) and day 31 ($F = 19.09; df = 1, 6; P = 0.0047$) (Fig. 2B). *H. rhossiliensis* was not detected by real-time PCR in egg-CK and J2-CK treatments.

### 4. Discussion

Maintaining an introduced fungus in the soil is the key to maximizing its effect on the nematode population. In our study, results based on real-time PCR showed that the quantity of *H. rhossiliensis* DNA declined over time in eggs-infested or J2-infested soil. In experiment 2, the quantity of *H. rhossiliensis* DNA decreased by 5.2 and 11.8 times from day 1 to day 17, and then by 2.1 and 1.7 times from day 17 to day 59 in J2-infested soil and egg-infested soil, respectively, which indicated that the establishment and survival of the fungus were greatly affected by some biotic antagonism or abiotic environmental factors after the fungus was introduced to soil, especially in the earlier period.

The results in experiment 2 showed that nematode population density in J2-CK treatment was significantly higher than that in egg-CK treatment 45 days after planting in experiment 2, and that suppression of soybean cyst nematodes by *H. rhossiliensis* was significantly less in egg-infested soil than in J2-infested soil. These data shared some features with Lackey et al. (1994), who reported that the fungus was more effective in controlling sugar beet cyst nematodes in soil infested with J2 than in soil infested with egg masses or cysts. Lackey et al. (1994) inferred that the J2 in eggs re-

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**Table 2**

<table>
<thead>
<tr>
<th>Days after planting</th>
<th>Treatmenta</th>
<th>Dry shoot weight (mg)b</th>
<th>Dry root weight (mg)b</th>
<th>1000 eggs per gram plantb</th>
<th>Eggs per cm³ soilb</th>
</tr>
</thead>
<tbody>
<tr>
<td>17</td>
<td>Egg-CK</td>
<td>234 ± 13 a</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Egg-inoculation</td>
<td>236 ± 22 a</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>J2-CK</td>
<td>233 ± 26 a</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>J2-inoculation</td>
<td>227 ± 39 a</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>31</td>
<td>Egg-CK</td>
<td>251 ± 27 a</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Egg-inoculation</td>
<td>276 ± 11 a</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>J2-CK</td>
<td>189 ± 23 c</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>J2-inoculation</td>
<td>231 ± 40 b</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>45</td>
<td>Egg-CK</td>
<td>274 ± 20 a</td>
<td>70 ± 18 a</td>
<td>22 ± 8 c</td>
<td>33 ± 10 c</td>
</tr>
<tr>
<td></td>
<td>Egg-inoculation</td>
<td>273 ± 45 a</td>
<td>64 ± 8 a</td>
<td>15 ± 5 c</td>
<td>22 ± 4 d</td>
</tr>
<tr>
<td></td>
<td>J2-CK</td>
<td>165 ± 29 b</td>
<td>40 ± 5 b</td>
<td>590 ± 82 a</td>
<td>512 ± 129 a</td>
</tr>
<tr>
<td></td>
<td>J2-inoculation</td>
<td>280 ± 17 a</td>
<td>80 ± 8 a</td>
<td>40 ± 8 b</td>
<td>60 ± 12 b</td>
</tr>
<tr>
<td>59</td>
<td>Egg-CK</td>
<td>286 ± 84 b</td>
<td>87 ± 7 a</td>
<td>22 ± 8 c</td>
<td>37 ± 16 c</td>
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<td></td>
<td>Egg-inoculation</td>
<td>399 ± 67 a</td>
<td>82 ± 8 a</td>
<td>11 ± 4 d</td>
<td>25 ± 10 c</td>
</tr>
<tr>
<td></td>
<td>J2-CK</td>
<td>181 ± 52 c</td>
<td>47 ± 6 b</td>
<td>649 ± 297 a</td>
<td>588 ± 12 a</td>
</tr>
<tr>
<td></td>
<td>J2-inoculation</td>
<td>272 ± 48 b</td>
<td>81 ± 8 a</td>
<td>88 ± 31 b</td>
<td>126 ± 31 b</td>
</tr>
</tbody>
</table>

a Treatment: egg-infested and without/with fungal inoculation (eggs-CK/eggs-inoculation); J2-infested and without/with fungal inoculation (J2-CK/J2-inoculation).

b Means ± SD (n = 4). Values followed by the same letter in the column within the same days after planting are not significantly different at P > 0.05 according to LSD test. Egg and J2 densities were transformed to log10(x) values before being subjected to statistical analysis.
Infested with eggs
The percentage of and quantity of rhossiliensis the conidia of through the soil, and thus reduced their chance of encountering remained in eggs until roots were near, traveled short distances through the soil, and thus reduced their chance of encountering the conidia of H. rhossiliensis. The small number of J2 and limited movement through the soil would minimize not only the chance of invading soybean roots but also the chance of encountering the conidia of H. rhossiliensis, which resulted in a lower nematode inoculum potential and poor suppression efficiency of the fungus on the nematode. The small number of J2 and their limited movement might also account for the persistently low percentage of J2 parasitized by H. rhossiliensis (0.5–2.3%) in experiment 1, where fungal parasitism was determined by extracting nematodes directly from soil without addition of bait J2. The quantity of H. rhossiliensis DNA, which ranged from $3.10 \pm 1.60 \times 10^8$ to $3.52 \pm 0.63 \times 10^9$ fg/g soil in experiment 1, was comparable to $3.83 \pm 1.96 \times 10^8$ to $1.96 \pm 6.39 \times 10^8$ fg/g soil in experiment 2, indicating that the failure to parasitize in experiment 1 did not result from the absence of the fungus. Fungal parasitism of nematodes in experiment 2, estimated by adding fresh J2 to the soil 3 days before sampling and then extracting and examining the J2, was higher (12–48%) than that in experiment 1 (0.5–2.3%). One interpretation is that the determination of fungal parasitism based on basing overestimated the chance of encountering the conidia due to the addition of fresh J2 (Lackey et al., 1994). Much of the uncertainty associated with the bioassay is eliminated by real-time PCR because the latter does not depend on nematode movement through soil and detects all fungal propagules directly (Zhang et al., 2006).

In experiment 2, real-time PCR assay showed that the amount of H. rhossiliensis DNA was statistically higher in J2-inoculation treatment than in egg-inoculation treatment in the earlier period (prior to day 31) after being introduced to soil. The result was partially consistent with the investigation over a period of 2 year in an agricultural soil by Persmark et al. (1996) showing that there is a positive relationship between the number of nematodes and the endoparasitic fungi in agricultural soil. The higher DNA yield of H. rhossiliensis in J2-infested soil during this period indicated a larger biomass than in egg-infested soil. Under natural conditions, H. rhossiliensis is an endoparasite of nematodes and depends on nematodes for nutrition (Jaffee and Muldoon, 1989; Jaffee et al., 1993). Previous laboratory studies have confirmed that the probability of parasitism of nematodes by H. rhossiliensis increases with increased host density (Jaffee et al., 1993; Jaffee and Muldoon, 1995). The J2 of nematode is the entity which H. rhossiliensis parasitizes. The presence of J2 at high concentrations in the earlier period provided a plentiful nutrient source for the fungus, and consequently resulted in higher biomass in J2-infested soil. The quantity of H. rhossiliensis DNA in J2-infested soil decreased over time and showed no significant difference with egg-infested soil 45 days after planting, which could be attributed to the low nematode propagation as a result of effective nematode suppression by the fungus. It seems possible that the numbers of J2 present during and soon after fungal inoculation affected the establishment of the fungus in soil.

To achieve successful and reproducible biological control, we must understand the ecological interactions affecting the control agent and the target (Whipps, 1997, 2001). Real-time PCR provides an effective way to quantify biocontrol agents after they are added to soil. In our study, the results based on real-time PCR assay showed that the DNA yield of H. rhossiliensis decreased over time regardless of eggs or J2 as inoculum, and that the DNA yield of H. rhossiliensis was statistically higher in J2-infested soil than in egg-infested soil in the earlier period. Thus, the fungus showed greater efficiency in suppressing nematodes in J2-infested soil than in egg-infested soil. We also note that nematode densities were initially higher in J2-infested soil, suggesting that the nematode density could affect the ability of H. rhossiliensis to establish in soil. Further studies are necessary to determine how various biotic and abiotic factors affect H. rhossiliensis DNA quantity in soil. The relationship between DNA quantity and nematode control by H. rhossiliensis also requires additional research. With the resulting knowledge, researchers should be better able to predict conditions that favor biocontrol by the fungus.

Acknowledgments

This work was jointly supported by National Outstanding Youth Foundation (30625001), National 863 Plan of China (2006AA10A211) and Beijing Municipal Science and Technology Commission (D0706005040331 and D0705002040191). The authors also thank Prof. Bruce A. Jaffee (the University of California at Davis) for serving as pre-submission reviewers and for his valuable comments and suggestions.

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