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Abstract
Entomopathogenic ascomycete fungi are ubiquitous in soil and on phylloplanes, and are important natural enemies of many soil-borne arthropods including larval western corn rootworm, Diabrotica virgifera virgifera, which is a major pest of corn. We measured the prevalence of Beauveria bassiana and Metarhizium anisopliae sensu lato in ten cornfields in Iowa, USA by baiting with larval insects. B. bassiana and M. anisopliae s.l. were present in 60% ± 6.3% and 55% ± 6.4% of soil samples, respectively. Subsequent laboratory bioassays found that some M. anisopliae s.l. strains collected from cornfields killed a greater proportion of D.v. virgifera larvae than a standard commercial strain.

Keywords
Bioassays, Cornfield, Beauveria bassiana, Galleria mellonella, Metarhizium, Tenebrio molitor

Disciplines
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Comments
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Entomopathogenic fungi in cornfields and their potential to manage larval western corn rootworm *Diabrotica virgifera virgifera*

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**Abstract**

Entomopathogenic ascomycete fungi are ubiquitous in soil and on phylloplanes, and are important natural enemies of many soil-borne arthropods including larval western corn rootworm, *Diabrotica virgifera virgifera*, which is a major pest of corn. We measured the prevalence of *Beauveria bassiana* and *Metarhizium anisopliae* sensu lato in ten cornfields in Iowa, USA by baiting with larval insects. *B. bassiana* and *M. anisopliae* s.l. were present in 60% ± 6.3% and 55% ± 6.4% of soil samples, respectively. Subsequent laboratory bioassays found that some *M. anisopliae* s.l. strains collected from cornfields killed a greater proportion of *D.v. virgifera* larvae than a standard commercial strain.

1. Introduction

Western corn rootworm, *Diabrotica virgifera virgifera* (Coleoptera: Chrysomelidae), is a major pest of corn in the United States and Europe, with larvae that live in the soil and feed on corn roots (Gray et al., 2009). Current management relies in part on corn that produces toxins of *Bacillus thuringiensis*; however, development of Bt resistance and predominant use of conventional corn hybrids in Europe requires the exploration of other management tactics (Gray et al., 2009; Gassmann et al., 2011). Entomopathogenic fungi, especially hypocrealean Ascomycetes, occur widely in the soil (Scheepmaker and Butt, 2010) and kill a variety of insect species (Hajek and St. Leger, 1994), including larval and pupal *D.v. virgifera* (Kuhlmann and van der Burgt, 1998; Pilz et al., 2008). In a two-year field study, we sampled cornfields in Iowa for the natural occurrence of entomopathogenic fungi *Metarhizium anisopliae* sensu lato (Hypocreales: Clavicipitaceae) and *Beauveria bassiana* (Hypocreales: Cordycipitaceae). We also measured mortality of *D.v. virgifera* larvae when challenged with naturally occurring *M. anisopliae* s.l. and *B. bassiana* strains isolated from cornfields, and with two commercial strains of these pathogens.

2. Materials and methods

2.1. Survey

Between 9 and 18 October, 2008 and between 9 and 18 October, 2009, soil samples were obtained from five cornfields in Iowa that were planted to corn for at least two years; different fields on the same five farms were sampled in both years, for a total of ten fields. Six samples (each separated by >15 m) were collected from each field, with a sample consisting of the root mass and surrounding soil from one corn plant (ca. 5 L). Soil samples were sifted through a 1.5 cm sieve and stored at 4 °C. Three insect species were used for baiting entomopathogens: *Diabrotica virgifera virgifera*, *Galleria mellonella* and *Tenebrio molitor*. Baiting was conducted between 24 October and 7 November 2008, and between 21 October 2009 and 16 January 2010 (with more time used in 2009 because of initial contamination of *G. mellonella* with *B. bassiana*). *T. molitor* larvae were obtained commercially (Jax Outdoor Gear and Earl May Nursery & Garden Center, Ames, Iowa). Fifth instar *G. mellonella* and early third instar *D.v. virgifera* larvae were reared in the laboratory following King and Hartley (1985) and Jackson (1985), respectively. The *D.v. virgifera* larvae were from a non-diapausing laboratory strain obtained from United States Department of Agriculture’s North Central Agricultural Research Laboratory.

Each soil sample was divided among three 500 ml containers (Reynolds Del-Pak 16 oz Container, Johnson Paper & Supply Co, Minneapolis, Minnesota). Containers with *D.v. virgifera* received
50 ml of soil and containers with *G. mellonella* and *T. molitor* received 150 ml. Ten pieces of corn root, 1 cm long, were added from each sample, soil was moistened with deionized water following *Goettel and Inglis* (1997), and six insects of a single species were placed in each container. For each species, 36 insects were exposed to soil from each of the five sites per year, for a total of 360 insects per species and 1080 insects per year. Controls were run simultaneously with soil samples, and each 500 ml control container consisted of six insects exposed to either autoclaved field soil or moistened paper towels. In 2008, for each insect species, two replicates were established for each type of control, for a total of 210 control insects. In 2009, 10 replicates were run for each insect species by type of control, for a total of 357 control insects.

Containers were held in an incubator (27 °C, 0/24 L/D) and checked every 2 to 3 d for a total of 11 d. Dead larvae were placed on modified White traps (*Kaya and Stock*, 1997), which allowed fungal infections to become expressed. Fungi were identified by placing the cadavers to selective media for further culturing (*Goettel and Inglis*, 1997), with *M. anisopliae* s.l. cultured on the medium of *Veen and Ferron* (1966) and *B. bassiana* on the medium of *Doberski and Tribe* (1980).

### 2.2. Bioassays

Bioassays used three strains each of *M. anisopliae* s.l. and *B. bassiana*. *B. bassiana* strains D2008 and D2009 were isolated from *D. v. virgifera* larval cadavers at site D in 2008 and 2009, respectively (*Fig. 1A and 1B*). Strain GHA was derived from the commercial product *BotaniGard ES* (*Laverlam International, Butte, Montana*). *M. anisopliae* s.l. strains C20091 and C20092 were each isolated from individual *D. v. virgifera* larval cadavers at site C in 2009 (*Fig. 1D*, and *M. brunneum* (F52 strain) (=*M. anisopliae* sensu lato) was derived from the commercial product *Met* EC (*Novozymes Biologicals, Salem, Virginia*). Strains obtained from *D. v. virgifera* were isolated by transferring external spores from the cadavers to selective media for further culturing (*Goettel and Inglis*, 1997).

Bioassays used sieved (0.6 mm mesh), air-dried field soil. The GHA strain (*B. bassiana*) was produced using solid substrate fermentation (*Jaronski and Jackson*, 2012) and represented the sixth passage since culturing in insects. Previously, this strain had no change in efficacy after 15 *in vitro* cycles (*Brownbridge et al.*, 2001). The remaining five strains were cultured on *G. mellonella* larvae. Live larvae were infected by dipping into a concentrated spore solution (>1 × 10⁶ spores per ml) for 5 s, incubating in sealed Petri dishes in the dark at 27 °C until the insects died (3–5 d), and placing the cadaver in a modified White trap. Conidia were harvested by vortexing three to seven conidia-covered cadavers with 8 ml of 0.1% *Tween* for 2 min, sonicating 10 min, and vortexing. Conidial viability was determined and soil was prepared by mixing each fungal solution with soil to achieve a uniform distribution of inoculum at the desired concentration and 25% water holding capacity following *MacDonald and Ellis* (1990). Bioassays were conducted in 45 ml containers (*Souffle Cup*, *Solo Cup Company, Highland Park, Illinois*) containing three germinated corn seedlings, 30 g of moistened soil and six second instar *D. v. virgifera* larvae. Containers were covered with fine mesh and a vented lid, and placed between two trays lined with moist paper towels. *M. anisopliae* s.l. was applied at 6.1 × 10⁵ and 6.1 × 10⁶ conidia per g of soil and *B. bassiana* was applied at 1.33 × 10⁶ and 1.33 × 10⁷ conidia per g of soil, with the high concentrations selected to achieve 50% mortality based on preliminary experiments with F52 and GHA. Ten days after larvae were placed in bioassay containers, soil and corn roots were carefully inspected, and the number of live larvae counted. Six replicates were run (except the high concentration of F52, which had four replicates), with each replicate consisting of three containers per strain per concentration, for a total of 210 containers and 1260 larvae. Six control containers, which were identical to experimental containers but did not receive fungal spores,

Fig. 1. Mortality from entomopathogenic fungi. Data are present for *B. bassiana* in (A) 2008 and (B) 2009, and for *M. anisopliae* sensu lato in (C) 2008 and (D) 2009. Bar heights are sample means and error bars are the standard error of the mean. Letters on the x-axis indicate site (i.e., farm) sampled. The y-axis describes the average proportion of insects killed by each pathogen.
were run with each replicate, for a total of 72 control containers and 432 control larvae.

2.3. Data analysis

For the survey, larval mortality was analysed separately for each year with a test of independence (PROC CATMOD; SAS, 2009). Some control G. mellonella were killed by naturally occurring B. bassiana strains, thus mortality of G. mellonella from B. bassiana in cornfield soil was corrected by subtracting the average number of insects killed per control container (0.5 insects in 2008 and 0.4 insects in 2009). Factors in the analysis were site, insect (G. mellonella, T. molitor and D. v. virgifera), and pathogen (B. bassiana, M. anisopliae s.l.). Contrasts among factors were made with a G test (Sokal and Rohlf, 1995). For the percentage of cups with entomopathogenic fungi, accompanying standard deviations are calculated following Sokal and Rohlf (1995).

Mortality in bioassays was calculated as the number of larvae initially placed in bioassay containers minus live larvae recovered from containers after 10 d. Mortality in bioassays was adjusted for control mortality using the correction of Abbott (1925). Data were analysed separately for B. bassiana and M. anisopliae s.l. using two-way analyses of variance (PROC GLM), and data were transformed by the arcsine of the square root to ensure normality of the residuals. Pairwise comparisons were conducted within strains at a concentration (PDHIF option in PROC GLM), with \( z = 0.017 \) based on the Dunn–Šidák correction for three pairwise comparisons (Sokal and Rohlf, 1995).

3. Results and discussion

3.1. Survey

For larval mortality from fungal infections, there was a significant interaction between pathogen and site in 2008 \((df = 4; \chi^2 = 16.17; P = 0.0028)\) and a significant effect of site in 2009 \((df = 4; \chi^2 = 15.71; P = 0.0034)\) (Fig. 1). Significant variation in occurrence of pathogens across sites is consistent with previous work, which found spatial heterogeneity among natural populations of soil entomopathogens (Chandler et al., 1997; Meyling and Eilenberg, 2007). Both pathogens were present in all fields in 2008 and four of five fields in 2009. Among all soil samples, B. bassiana was present in 60% ± 6.3% and M. anisopliae s.l. was present in 55% ± 6.4%.

The insect species used for baiting influenced mortality imposed by pathogens in both years \((df = 2; \chi^2 \geq 13.72; P \leq 0.0034)\) (Fig. 1), with greater mortality for G. mellonella than either T. molitor and D. v. virgifera \((df = 1; G \geq 11.47; P \leq 0.001)\) and greater mortality for T. molitor than D. v. virgifera \((df = 1; G \geq 23.92; P \leq 0.0001)\) (Fig. 1). This may be due in part to better adaptation of D. v. virgifera to soil pathogens because larvae live in the soil (Gray et al., 2009), and in part to the high sensitivity of the other bait species to infections (Goettel and Inglis, 1997). This work highlights the importance of the insect species used to recover pathogens, and previous work has shown that the insect species used can influence the fungal strains recovered (Klingen et al., 2002). Additional molecular work is needed to understand the genetic diversity of strains found here.

3.2. Bioassays

There was a significant interaction between concentration and fungal strain \((df = 2.96; F = 4.68; P = 0.01)\) for M. anisopliae s.l. (Fig. 2B). At the high concentration, the two field-isolated strains of M. anisopliae s.l. caused significantly greater mortality of D. v. virgifera larvae than FS2. This suggests potential host adaptation because fungal strains were collected from D. v. virgifera larvae. No other differences among strains were found (Fig. 2A and 2B).

3.3. Conclusions

Either M. anisopliae s.l. or B. bassiana were found in 76.7% ± 3.3% of soil samples from cornfields. These entomopathogenic fungi have broad host ranges, but strains can vary in their effectiveness against different insect species (Hajek and St. Leger, 1994; Jaronski, 2007). Because of their occurrence in cornfields and their ability to kill D. v. virgifera larvae, these pathogens may be an important natural source of mortality for this pest and may be developed for biological control.

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