Preventive application of an entomopathogenic fungus in cover crops for wireworm control

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Abstract Efficacy of the Metarhizium brunneum Petch (Hypocreales: Clavicipitaceae) strain ART2825 for control of wireworms (Agriotes obscurus (L.), Coleoptera: Elateridae) was examined in a semi-field pot experiment. Pots were treated in late summer during sowing of spring oat as a cover crop. Survival of wireworms was assessed four weeks after their release in October 2013, and 30 weeks after release in April 2014. Viability and persistence of the fungus was determined by counting colony forming units from substrate samples and microsatellite analyses of recovered Metarhizium isolates. The number of colonies detected in the substrate in October 2013 increased with increasing concentrations of applied conidia, and no significant reduction was observed at the second evaluation date in April 2014. Increasing conidia application rates significantly increased mycosis and reduced wireworm survival, to a level comparable to that of treatment using insecticide-coated oat seeds. The preventive application of M. brunneum conidia to reduce wireworm populations in cover crops, preceding a damage-sensitive crop like potatoes, may be a promising biocontrol strategy.

Keywords Coleoptera · Elateridae · Hypocreales · Clavicipitaceae · Arable crops · Potato · Biological control

Introduction

Wireworms are the soil dwelling larvae of click beetles (Coleoptera: Elateridae) and several species, including those in the genus Agriotes Eschscholtz, are major pests in the northern hemisphere. They feed on the roots, shoots, tubers and bulbs of many arable and vegetable crops, including maize and other cereals, potatoes and carrots (reviewed in Parker and Howard 2001; Ritter and Richter 2013; Traugott et al. 2015). In many of these crops, wireworms are not necessarily the most destructive insects, but are among the most difficult to control, particularly because of their long life cycle and overlapping generations, their polyphagous nature and their adaption to a wide range of
agricultural ecosystems, including grassland and arable crops (Furlan 1998; Ritter and Richter 2013; Sonnemann et al. 2014; Sufyan et al. 2014; Traugott et al. 2013, 2015). Wireworms are also able to react quickly to changes in abiotic conditions with vertical migration to depths of 50 cm or more (Fisher et al. 1975; Furlan 1998, 2004; Schaerffenberg 1942). This behaviour often reduces the efficacy of short-term control attempts, which are typically applied to upper soil layers.

Chemical control of wireworms has been most successful with persistent insecticides, like organochlorines, organophosphates, carbamates and phenylpyrazols (reviewed in Barsics et al. 2013; Traugott et al. 2015; Vernon and van Herk 2013), some of which provide sufficient control for several seasons after a single application (Vernon et al. 2009, 2013). As a result, the preventive application of cereal seeds coated with phenylpyrazol became widely adapted in some European countries, the goal being to reduce wireworm populations and the risk of damage in subsequent sensitive crops like potatoes (Jossi 2001). Pre-planting of organochlorine-coated wheat seeds as a catch crop before planting of potatoes has also been shown to reduce wireworm damage to potatoes in experiments in Canada (Vernon et al. 2016). However, this control strategy was later abandoned when the insecticide was de-registered.

The ban or withdrawal of many of these widely used soil insecticides and the trend to more conservative soil cultivation techniques led to both an increase of wireworm populations and the risk of damage in subsequent sensitive crops like potatoes (Jossi 2001). Pre-planting of organochlorine-coated wheat seeds as a catch crop before planting of potatoes has also been shown to reduce wireworm damage to potatoes in experiments in Canada (Vernon et al. 2016). However, this control strategy was later abandoned when the insecticide was de-registered.

Materials and Methods

Fungal inoculum and host insects

*M. brunneum* strain ART2825 was isolated in 2007 from an infected *Agriotes obscurus* larva in the rearing facility at Agroscope in Zurich, Switzerland. The strain has shown high virulence towards larvae of *A. obscurus* and *A. lineatus* (Eckard et al. 2014; Kölliker et al. 2011). After a host passage, the fungus was cultivated on a selective medium containing antibiotics and dodine (Strasser et al. 1996). Conidia harvested from Petri dishes were used for mass-production of fungus colonised barley kernels (FCBKs) (Aregger 1992). Number of conidia per gram of FCBKs was determined by shaking samples in 0.1% (v/v) aqueous Tween®80 and counting the resulting conidia densities with a haemocytometer. Germination rate was examined by plating approximately 300 conidia on selective media and counting colony forming units (CFUs) after incubation for two weeks at 23 °C in the dark.

*Agriotes obscurus* larvae used in the experiment originated from a laboratory livestock established according to Kölliker et al. (2009). Larvae used in the experiment were about one year old and had reached at least the 7th larval stage (minimum width of head capsule 1.16 mm according to Klausnitzer 1994).

Experimental set-up

A semi-field pot experiment was set up in an experimental bed at Agroscope in Zurich, Switzerland (47°25′40.13″N, 8°30′58.10″E). Pots (28 cm diameter and 24 cm height, surface area 615 cm²) were left above ground and watered as needed. They were filled with 14 kg of a mixture of field soil (32.5% clay,
28.8% silt, 43.7% sand, pH 7.78) and sand (3:1 v:v). Pots were arranged in a randomized complete block design, with six blocks, six treatments, and two pots per treatment in each block, to enable sub-sampling of all treatments at two different dates (see below).

FCBKs were applied in mid-August 2013 in four different concentrations, 0.05, 0.5, 5 and 50 g per pot, corresponding to a field application rate of 8–8000 kg of FCBKs or an estimated concentration of $10^{12}$–$10^{15}$ conidia per hectare. FCBKs were thoroughly mixed into the upper 10 cm of the pot substrates with a flower rake. Subsequently, 30 kernels of spring oat (variety “Expander”, ASS, Moudon, Switzerland) were sown per pot ($=150$ kg ha$^{-1}$). We included two controls without FCBKs: in the first, oat seeds of the same variety coated with Regent® TS (BASF, 500 g l$^{-1}$ Fipronil at a rate of 100 ml per 100 kg of seeds) were planted as an insecticide treatment, and in the second, pots with untreated spring oat served as control. Two weeks after set up, when oat seedlings reached the three-leaf stage, seven lab-reared *Agriotes obscurus* were released per pot.

Data collection and analysis

The first sub-sample of 36 pots was evaluated at the beginning of October 2013, six weeks after application of FCBKs and four weeks after wireworm release. This presumably coincided with the end of the wireworms’ seasonal activity period, when decreasing temperatures drove them to deeper substrate layers for hibernation. The second sub-sample of pots was analysed in April 2014, 32 weeks after application and 30 weeks after wireworm release, which corresponded to the time point suitable for potato planting in central Europe.

The substrate of each pot was thoroughly examined to recapture released wireworms or cadavers. Alive and dead individuals were transferred singly into separate cups filled with peat, provided with a carrot slice as food source, and incubated at 23 °C and 75% RH in the dark for six weeks. Wireworms with a healthy appearance and normal escaping behaviour after six weeks of incubation (negative phototaxis, quickly burrowing into the substrate upon replacement into cups) were counted as survivors. Only cadavers showing infection with *Metarhizium* spp., i.e. out-growth of white mycelium from intersegmental integuments and formation of green conidia layers, were counted as mycosed (=killed by the FCBK treatment). Where mycosed cadavers were found, one *Metarhizium* spp. isolate per pot was obtained by dilution plating. Isolates were maintained on selective medium plates at 23 °C in the dark.

Abundance of the fungus in the substrate was assessed by counting the number of CFUs per gram of substrate as described in previous studies (Kessler et al. 2003; Pilz et al. 2010). Following recapture of wireworms, five soil portions of about 10 g were taken from the substrate of each pot using a spoon, combined in a plastic bag, and mixed thoroughly. A sub-sample of 20 g per pot was suspended in 100 ml of a sodium pyrophosphate solution (1.8 g l$^{-1}$ Na$_4$P$_2$O$_7$·10H$_2$O) and shaken in an Erlenmeyer flask for four hours at 120 rpm. After sedimentation for five seconds, aliquots of 100 μl of the supernatant were plated on each of three Petri dishes with selective medium. Petri dishes were incubated at 23 °C and 75% RH in the dark. CFUs were counted after ten days and numbers of CFUs per gram soil substrate were calculated. For analysis, we used the mean of the three Petri dishes from each pot. Five and four single colonies were randomly selected from Petri dishes of control and insecticide treatments (soil samples) of the second evaluation date for molecular genetic analyses. They were transferred to selective medium plates and maintained at 23 °C in the dark.

Wireworm recapture, survival and mycosis rates were analysed with generalised linear mixed-effect models with rates (as binary variable: positives vs. negatives) as response variable and the number of FCBKs per pot (log-transformed), the harvest date (autumn vs. spring), their interaction, and the presence of insecticide (only in one of the controls) as explanatory variables. To allow interpretation of the interaction term, the numerical variable “number of FCBKs per pot” was scaled (i.e. centred on the mean and divided by its SD; Schielzeth 2010). We assumed underlying binomial distributions and used a logit link function. Numbers of CFUs per gram soil substrate were analysed with a linear mixed model with the amount of FCBKs per pot (scaled), the harvest date (autumn vs. spring), their interaction, and the presence of insecticide (only in one of the controls) as explanatory variables. To allow interpretation of the interaction term, the numerical variable “number of FCBKs per pot” was scaled (i.e. centred on the mean and divided by its SD; Schielzeth 2010). We assumed underlying binomial distributions and used a logit link function. Numbers of CFUs per gram soil substrate were analysed with a linear mixed model with the amount of FCBKs per pot (scaled), the harvest date (autumn vs. spring), and their interaction as explanatory variables. Here, insecticide treated pots were excluded from the analysis. In all models, we included the treatment pair nested within blocks as random effects. All statistical analyses were performed with RStudio (Version 1.0.136, RStudio
Identification of re-isolated fungal strains

One single colony per recovered *Metarhizium* isolate was transferred into liquid complete medium (Riba et al. 1986) and incubated in an Erlenmeyer flask at 150 rpm for three days. The mycelium was harvested by filtration (Enkerli et al. 2001) and frozen at −70 °C. Frozen mycelium was lyophilized and homogenized with a FastPrep FP120 homogenizer (MP Biomedicals, Santa Ana, CA, USA) using 0.15 g of glass beads (1 mm diameter) at a speed of 6 m s⁻¹ for 10 s. Genomic DNA was extracted using the NucleoSpin Plant II DNA extraction kit according to the manufacturer’s protocol (Macherey–Nagel, Germany).

Microsatellite marker analysis

Twelve microsatellite markers were used to compare genotypes of the collected isolates (Enkerli et al. 2005; Oulevey et al. 2009). Multiplex PCRs were performed with four sets of three markers each (set 1: Ma2049, Ma2060, Ma2069, annealing temperature 58 °C; set 2: Ma2065, Ma2098, Ma2269, annealing temperature 58 °C; set 3: Ma142, Ma417, Ma2224, annealing temperature 56 °C; set 4: Ma097, Ma195, Ma2077, annealing temperature 50 °C) in reaction volumes of 20 μl. Reactions contained 10 ng genomic DNA, 5× GoTaq® Flexi Reaction Buffer and 0.25 units of GoTaq® Flexi DNA Polymerase (Promega, WI, USA), 4 (sets 1 and 2) or 3 (sets 3 and 4) mM MgCl₂ (Oulevey et al. 2009), 0.2 μM of each dNTP and 0.2 mM forward and labelled reverse primer (FAM and HEX, Microsynth, Switzerland; NED, Applied Biosystems, CA, USA). Touch-down PCR consisted of 2 min initial denaturation at 94 °C followed by 12 cycles of touch-down amplification consisting of 30 s at 94 °C, 60 s at multiplex specific annealing temperature (see above) plus 12 °C with a decrease of 1 °C per cycle, and 60 s at 72 °C. Subsequently, 22 cycles of 30 s at 94 °C, 60 s at multiplex specific annealing temperature and 60 s at 72 °C were performed. The amplification was terminated with a final elongation at 72 °C for 15 min. Amplified DNA-fragment sizes (allele sizes) were determined on an ABI 3500 x L Genetic Analyzer (Applied Biosystems, CA, USA) using 50 cm capillaries, POP-7 polymer and GeneScan™ 400HD ROX™ size standard. A Swiss strain of *M. brunneum* (ARSEF 7524) was used as reference for allele sizes. Analysis of data was performed using GeneMarker software version 2.4.0 (SoftGenetics, PA, USA).

Results

Abundance of *Metarhizium* CFUs in the substrate

Evaluation of pot substrates revealed the presence of *Metarhizium* colonies in every treatment, including the untreated control and the insecticide treatment (Table 1). The number of CFUs detected increased significantly with the concentration of conidia applied, but there was no difference between the collection dates (Table 2).

Wireworm recapture, survival and mycosis

Recapture rates of wireworms (alive and dead) were not influenced by FCBK treatments (Table 3a), but significantly fewer wireworms were recaptured from insecticide-treated pots (Fig. 1 shows the first evaluation date). Recapture rates were also significantly different between collection dates. An average wireworm recapture rate of 81% (5.6 ± 1.2 individuals out of seven released) was achieved with the first sub-sample, evaluated in October 2013, four weeks after releasing wireworms. This rate dropped to an average of only 42% (2.9 ± 1.9 individuals out of seven released) in the second sub-sample, evaluated in April 2014, 30 weeks after wireworm release.

More than 33% of the wireworms recaptured at the first collection date died during laboratory incubation following recapture, mainly those from the insecticide treatment and the higher concentration fungal treatments (Fig. 2). Consequently, overall survival of wireworms after incubation showed a different pattern to that of wireworm recapture. Increasing concentrations of FCBK treatments clearly exhibited a negative effect on wireworm survival (Table 3b). This effect was similar at both collection dates, although less pronounced at the second collection in April 2014, as the significant interaction term indicates. The impact of the insecticide treatment was still significant and comparable to that of the higher-concentration FCBK treatments.
The main cause of mortality for all but the insecticide treatment, where no mycosed larvae were found, was *Metarhizium* mycosis (Fig. 3). Treatments with higher FCBK concentrations produced a significantly greater number of mycosed cadavers (Table 3c). The number of mycosed cadavers retrieved from the second sub-sample was very low (eight mycosed cadavers from a total of 105 recaptured individuals) and, correspondingly, the model showed a significant effect of evaluation date.

Identification of re-isolated *Metarhizium* strains

At the first evaluation date, 24 out of 36 pots yielded at least one mycosed wireworm cadaver. *Metarhizium* was isolated from one cadaver per pot, resulting in 24 isolates. At the second evaluation date, seven mycosed cadavers were detected in the pot substrate of two treatments ($10^{13}$ and $10^{14}$ conidia ha$^{-1}$) only, and *Metarhizium* was isolated from each cadaver. Additionally, *FCBK* fungus colonised barley kernels; the number of applied conidia per gram of substrate was estimated based on a mean yield of $1.2 \times 10^9$ conidia per gram of FCBKs applied to 14 kg of pot substrate.

### Table 1

Mean number ± SE of CFUs (colony forming units) per gram pot substrate six weeks (first evaluation date, early autumn) and 32 weeks (second evaluation date, spring) after application

| Treatment     | Field concentration | Number of applied conidia per gram substrate | Number of CFUs recovered per gram substrate |
|---------------|---------------------|---------------------------------------------|-------------------------------------------|
| Control       | Ambient             | 0                                           | $25 \pm 17$                                |
| Insecticide   | 50 g Fipronil/100 kg seeds | 0                                           | $70 \pm 40$                                |
| FCBK 1        | $10^{12}$ conidia ha$^{-1}$ | 400                                         | $153 \pm 55$                               |
| FCBK 2        | $10^{13}$ conidia ha$^{-1}$ | 4000                                        | $2467 \pm 475$                             |
| FCBK 3        | $10^{14}$ conidia ha$^{-1}$ | 40,000                                      | $20,414 \pm 6554$                          |
| FCBK 4        | $10^{15}$ conidia ha$^{-1}$ | 400,000                                     | $182,945 \pm 81,717$                       |

### Table 2

Coefficients, SE, z-values and p-values from a linear mixed effects model for the number of fungal colonies detected in pot substrates six weeks (first evaluation date) and 32 weeks (second evaluation date) after application. Conidia = conidia concentration applied per gram of pot substrate

| Value | SE  | z-value | p-value |
|-------|-----|---------|---------|
| (Intercept) | 52,104 | 28,945 | 1.80    | 0.083  |
| Conidia  | 89,604 | 28,780 | 3.11    | 0.005  |
| Date     | −10,904 | 17,536 | −0.62   | 0.539  |
| Conidia × date | −17,728 | 17,684 | −1.00   | 0.325  |

### Table 3

a–c Coefficients, SE, z- and p-values from generalised linear mixed effects models for the number of (a) re-captured, (b) surviving, and (c) mycosed wireworms four weeks (first evaluation date) and 30 weeks (second evaluation date) after release; FCBK = concentration of fungus colonized barley kernels applied; the insecticide treatment was excluded from the analysis of wireworm mycosis (3c)

| a (Intercept) | 3.76 | 0.43 | 8.67 | <0.001 |
| FCBK         | −0.39 | 0.35 | −1.14 | 0.253  |
| Date         | −1.97 | 0.23 | −8.67 | <0.001 |
| Insecticide  | −1.30 | 0.31 | −4.16 | <0.001 |
| FCBK × date  | −0.08 | 0.21 | −0.40 | 0.69   |
| b (Intercept) | −0.31 | 0.35 | −0.90 | 0.369  |
| FCBK         | −1.61 | 0.44 | −3.68 | <0.001 |
| Date         | −0.05 | 0.21 | −0.23 | 0.815  |
| Insecticide  | −1.96 | 0.37 | −5.34 | <0.001 |
| FCBK × date  | 0.51  | 0.26 | 2.00  | 0.046  |
| c (Intercept) | 1.40  | 0.57 | 2.45  | 0.014  |
| FCBK         | 1.59  | 0.60 | 2.66  | 0.008  |
| Date         | −2.05 | 0.45 | −4.54 | <0.001 |
| FCBK × date  | −0.41 | 0.46 | −0.91 | 0.366  |

at the second evaluation date, five *Metarhizium* isolates were randomly chosen from substrate samples of the untreated control and four isolates from the insecticide treatment, to check for the natural presence of the *M. brunneum* strain applied during the experiment.
Microsatellite marker analyses allowed discrimination of nine different genotypes among the 40 isolates (31 isolates from cadavers and nine from soil samples, Table 4, supplementary Table S1). All isolates obtained from wireworm cadavers of fungal treatments displayed a genotype identical to the genotype of the applied strain (ART2825), irrespective of conidia concentration and evaluation date (Table 4). In contrast, genotypes of isolates obtained from four cadavers collected from the untreated control were different from the genotype of the applied strain. Nevertheless, analysis of the isolates from substrate samples revealed that the strain used for FCBK production was also present naturally in control and insecticide treatments.

**Discussion**

We applied *M. brunneum*-FCBKs to pot substrates in late summer, together with planting of spring oat as a cover crop, and assessed the abundance of *Metarhizium* colonies and the recapture, survival and mycosis of wireworms released into pots. FCBK treatments significantly increased the abundance of *Metarhizium* colonies six weeks after treatment. Native *Metarhizium* colonies were also detected in all substrates, including those of the untreated control and the insecticide treatment. The presence of eight native *Metarhizium* genotypes in a rather limited number of samples showed that *Metarhizium* fungi were not only...
abundant, but also genetically diverse in the field soil used for the experiment (see Table 4, supplementary Table S1). Similar abundance and diversity of *Metarhizium* spp. has been reported for soils of different habitats and geographical regions around the world (Steinwender et al. 2014; Enkerli et al. 2016; Kepler et al. 2015; Castro et al. 2016).

*Metarhizium* strains were re-isolated from about one third of the wireworm cadavers recovered from FCBK treatments. All were genetically identical to the applied strain, ART2825 (Table 4). Four *Metarhizium* strains re-isolated from cadavers of control pots, however, were genetically different from ART2825, showing that several naturally occurring *Metarhizium* genotypes may have competed for the host resource. Nonetheless, the applied strain ART2825 was the preferential source of infection, which may indicate its competitiveness, or simply be a consequence of the

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**Fig. 3** Number of *A. obscurus* larvae with *Metarhizium* infection in November 2013, re-captured from pots four weeks after release and incubated individually in peat-filled cups for another six weeks. Boxes show the 1st (bottom line), 2nd (median, bold line), and 3rd (top line) quartiles of the data distribution, whisker ends represent the lowest and the highest data point within the 1.5 interquartile range of the lower and the upper quartile, respectively. Seven larvae were released per pot

**Table 4** Origin and genotype of the 40 *Metarhizium* isolates obtained from mycosed cadavers and substrate samples collected six weeks (first evaluation date) and 32 weeks (second evaluation date) after application, respectively

| Treatment | Evaluation date | Number of colonies from cadavers | Number of colonies from substrate | Genotypes detected |
|-----------|----------------|---------------------------------|----------------------------------|--------------------|
| Control   | 1              | 4                               | –                                | C, D, E (×2)       |
|           | 2              | –                               | 5                                | A, B, F, G, H      |
| Insecticide| 1              | –                               | –                                | –                  |
|           | 2              | –                               | 4                                | A, B (×2), I       |
| FCBK 1    | 1              | 2                               | –                                | A                  |
|           | 10^12 conidia ha^{-1} | 2 | – | – |
| FCBK 2    | 1              | 5                               | –                                | A                  |
|           | 10^13 conidia ha^{-1} | 2 | 4 | – |
| FCBK 3    | 1              | 8                               | –                                | A                  |
|           | 10^14 conidia ha^{-1} | 2 | 3 | – |
| FCBK 4    | 1              | 5                               | –                                | A                  |
|           | 10^15 conidia ha^{-1} | 2 | – | – |

*FCBK 1–4* treatments with fungus colonized barley kernels; “–”= not evaluated; the FCBK production strain, *Metarhizium brunneum* ART2825, is designated as genotype A
large number of conidia applied. The incidence of the strain ART2825 in the substrate samples from untreated controls and the insecticide treatment may be of natural origin, since the strain was originally isolated from the region of the study.

Several studies indicate that a sufficient amount of EPF inoculum for insect pest control ranges from \(10^5\) to \(10^9\) CFUs per gram of soil (Bruck 2005; Ekesi et al. 2002; Ferron 1981; McDowell et al. 1990). Based on these data, Jaronski (2010) estimated \(10^{14}\)–\(10^{15}\) conidia per hectare as the required dose for broadcast application and incorporation to a depth of 10 cm. We reached this level of CFUs per gram of substrate with the highest dose applied \((10^{15} \text{conidia ha}^{-1})\), which reduced wireworm survival in our experiment. A similar effect on wireworm survival was also achieved with a ten times lower application rate, which suggests that \(10^4\) *Metarhizium* CFU per gram of soil may already provide a sufficient level of wireworm control (Figs. 2, 3). Furthermore, improved application methods may also help to reduce application rates. Jaronski (2010) proposed up to a 100-fold reduction of the field application rates by the use of banded instead of broadcast application of *Metarhizium* conidia against sugar beet root maggot. A similar strategy may be tested by applying FCBKs and sowing of the cover crop in strips. Furthermore, a switch from *Metarhizium* formulated as corn grit granules to recently developed microsclerotia of *Metarhizium* enables a reduction of field application rates by a factor of eight (Jaronski 2010; Jaronski and Jackson 2008).

Our results confirmed the persistence of *M. brunneum* conidia for up to eight months, without being affected by winter conditions. Therefore, an application of FCBKs into the cover crop several months before planting of potatoes in the main season should provide for an extended exposure period of the pest insects to the fungus. Wireworms typically show seasonal migration between upper and lower soil layers, depending on their moulting and feeding cycles, but also on environmental conditions in upper soil layers (reviewed in Traugott et al. 2015). The combination of these factors results in two activity “peaks” of many semivoltine wireworm species in the northern hemisphere, during which they are more frequently found in the topsoil near the soil surface. One occurs in March or April, after hibernation, and a second in in August or September, after the dry season. Consequently, application of the EPF into a cover crop in late summer, and persistence of conidia over winter, should allow the pest population to be targeted twice. However, we were not able to detect a further decrease in wireworm survival rates at the second evaluation date, 30 weeks after release. One explanation may be that the second evaluation was performed too early. Wireworms may still have been burrowed in lower layers of the pot substrates for hibernation, without having come into sufficient contact with conidia for a second time.

It has been shown that lower temperatures slow down fungal infections of wireworms (Kabaluk and Ericsson 2007). Accordingly, an EPF application during summertime in the northern hemisphere, as performed in our experiment, should promote fungal growth and efficacy. During this period, however, upper soil layers may suffer desiccation, which may affect viability of conidia beneath the soil surface. A green plant cover, as provided by spring oat in our experiment, might mitigate such drought stress and foster persistence of the fungus. There is evidence in the literature that the abundance of *Metarhizium* is higher in soils with a dense plant cover, like grasslands or field margins (Keller et al. 2003; Schneider et al. 2012). There are, however, many factors that would contribute to making such habitats a favourable environment, including microclimatic and edaphic factors, as well as a better supply of potential host insects, less input of fungicides and herbicides and less disturbance by soil cultivation (Jaronski 2007; Keller et al. 2003). Their complex interactions are still not understood and need to be further investigated (Jaronski 2007; Schneider et al. 2012).

The combined application of the entomopathogenic fungus together with the cover crop may also constitute a simple, yet effective “attract and kill” strategy. Carbon dioxide is emitted from germinating seeds and other plant tissues, and CO₂ gradients are therefore a basic means of orientation of insect larvae towards potential food sources (Doane et al. 1975; Klingler 1957). These gradients have been shown to influence the vertical migrations of wireworms in soil (Jung et al. 2014; Sonnemann et al. 2014). In our experimental set-up, the germinating oat seedlings also produced CO₂, and potentially attracted wireworms into conidia-contaminated soil layers. Current research has investigated this means of orientation as part of an “attract and kill” strategy in several ways. Schumann et al. (2013, 2014) lured western corn rootworm (*Diabrotica virgifera virgifera* LeConte,
Coleoptera: Chrysomelidae) larvae to an insecticide or to a microbial control agent with the help of artificial CO₂ sources. Vernon et al. (2016) sowed insecticide-treated wheat as a lethal trap crop for wireworms into furrows at potato planting. The CO₂ released from germinating seeds and/or wheat roots attracted wireworms and facilitated contact with the insecticide.

In conclusion, high concentrations of *M. brunneum* conidia caused significant wireworm mortality in our pot experiment, after an incubation period of about one to two months. *M. brunneum* conidia persisted in the substrate for up to eight months. Spring oat as a cover crop on treated substrates may have provided favourable conditions for fungal persistence. Contrary to expectations, however, this did not lead to a further decrease in wireworm survival rates between the first and the second evaluation date. It remains to be shown if an extended period of exposure of wireworms to *Metarhizium* conidia has the potential to increase efficacy of the biocontrol treatment, and thus facilitate the reduction of field application rates.

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