Colonization of Potato by Colletotrichum coccodes: Effect of Soil Infestation and Seed Tuber and Foliar Inoculation

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ABSTRACT

Colonization of potato (Solanum tuberosum) tissue, including roots, stolons, and above and below ground stems, by Colletotrichum coccodes, the causal agent of black dot, was evaluated following soil infestation, inoculation of seed tubers and foliage, and every combination thereof, in field trials over two growing seasons in North Dakota and Minnesota. A total of 107,520 isolations for C. coccodes performed across four site-years allowed for an extensive comparison of fungal colonization of the host plant and disease severity. The black dot pathogen was detected in potato stems at the first sampling date in all four site-years, as early as 14 days prior to emergence. Colonization of above and below ground stems occurred at a higher frequency than in roots and stolons in all four site-years, resulting in significantly higher relative area under the colonization progress curves (RAU CPCs) (α = 0.05). Although fungal colonization and disease incidence were higher in inoculated and/or infested treatments, sufficient natural inoculum was present to result in substantial levels of disease in noninoculated and noninfested plots. However, noninoculated and noninfested plots displayed the lowest RAU CPC values across three of four site-years and those treatments with multiple inoculation events tended to have higher RAU CPC values. Isolates belonging to vegetative compatibility group (VCG) 2 and -5 were recovered from plants sampled in 2004 more frequently than isolates belonging to VCG 1 and -3. A significant difference in disease incidence on stems was observed only in North Dakota in 2004 and Minnesota in 2003 (α = 0.05). Noninoculated and noninfested plots displayed the lowest disease incidence, whereas those treatments with more than one inoculation and/or infestation event tended to have higher disease incidence. Results of this study, including the disease severity and yield data, provide a better understanding of colonization of potato plants by C. coccodes and its impact.

Black dot, caused by Colletotrichum coccodes (Wallr.) S. Hughes, is a disease that occurs wherever potato crops (Solanum tuberosum L.) are grown. Although the most economically important hosts of C. coccodes are potato, tomato, and pepper, this pathogen is able to infect a wide range of plant species, a majority of them members of the Solanaceae family. Several weed species, including some solanaceous species, also have been identified as hosts (33,37). On potato, black dot generally is considered to be primarily a tuber blemish disease resulting in symptoms similar to silver scurf caused by Helminthosporium solani Durieu & Mont.; however, the pathogen also can infect roots, stolons, stems, and foliage (1,3,13,21,27,36,50). Infected areas of tubers are silver to brown in color with microsclerotia present. Deep sunken lesions (18) and tuber shriveling (20) also have been noted with severe infections. Dark lesions form on infected leaf and stem tissue and contribute to wilting and defoliation (9,23). In severe cases, the cortical tissue of infected below ground stems and stolons may slough off, resulting in a frayed or stringy appearance (9). Upon vine desiccation and disintegration of the cortex, an area of amethyst coloration may be observed in association with the remnants of the vascular bundles (13).

Black dot has been recognized as a disease of potato since the early part of the 20th century (13). It generally had been considered to be of minor importanceт (24,35,40,46,48), having little impact on commercial potato production in most growing areas. Several factors have contributed to this view, many of which may be related to misdiagnosis. As with other typical anthracnose pathogens, C. coccodes often infects the host early and symptoms are not expressed until much later in the growing season (1), at which point they are often mistaken for normal plant senescence and saprophytic colonization. C. coccodes also can cause early dying in potato similar to other diseases such as Verticillium wilt, caused by Verticillium dahliae Kleb, and V. albo-atrum Reinke & Berthold, as well as early blight, caused by Alternaria solani Sorauer, and often occurs in conjunction with these diseases. Co-infection with V. dahliae has been shown to result in greater reductions than observed with either pathogen alone (51). Although yield losses of up to 30% due to black dot have been documented, it has proven difficult to reproduce these losses across growing seasons under field conditions, even when differences in black dot symptoms were noted among treatments (27,28,50). Yield losses due to black dot also have been documented in the absence of symptom expression (3).

Because of the aforementioned yield and quality losses, as well as losses reported by commercial potato growers, black dot research has garnered renewed interest in recent years as a developing threat to potato production and crop quality (25,52). The apparent emergence of black dot as an important disease of potato simply may be due to increased awareness of the factors outlined above. Changes in tillage and other cropping practices during the later part of the past century also may have promoted the accumulation of soilborne sclerotial inoculum (12,41,47), because the pathogen’s longevity in the soil has been demonstrated to extend 5 to 13 years in the absence of a potato crop (4,14). Although crop rotation may be successful in reducing soil inoculum, the wide host range of C. coccodes, including both weed and rotational crop hosts, as well as the longevity of the microsclerotia render crop rotation a fairly impractical control measure (9,15,33,37). Additionally, the large, heavily melanized microsclerotia of C. coccodes are not killed effectively by currently registered soil fumigants (11,46,52). Although seed treatment and in-furrow fungicide applications have not been successful at reducing black dot incidence or increasing yield in infested soil, foliar fungicide applications of the QoI fungicide azoxystrobin have proven effective in reducing black dot severity on stems and progeny tubers and increasing yield in the Columbia Basin of the United States (5,28). Long-term survival of the pathogen in the soil, limited number of useful fungicides, a long latent period, a wide host range, varying and unpredictable effects of the disease, and confusion regarding disease etiology illustrate why effective control of black dot can be difficult.

A more complete understanding of disease epidemiology and etiology is needed.
to develop successful strategies for effective management of black dot of potato. The pathogen typically is introduced into noninfested soils via contaminated seed tubers, becomes established on the current-season crop, and subsequently builds up in the soil on infected plant debris (3,24,38). Soilborne inoculum may infect tubers, stolons, roots, and below ground stems (38,50,51). The airborne phase of C. coccodes also can cause above ground stem and foliar infections via windblown inoculum originating from the soil, debris of previously infected plants, or current-season foliar infections, often exacerbated by wounds caused by windblown soil (3,21,23,27). Although considerable research has been performed comparing the effects of some inoculum sources, this research has not taken into account all potential infection sites and the importance, frequency, and timing of both above and below ground host tissue colonization (1,6,11,21,27,28,38,39). Additionally, much of the early research was performed before differences in aggressiveness were characterized among vegetative compatibility groups (VCGs) (2,19,31,32,34). As a result, there are gaps in knowledge concerning the influence, significance, and relative importance of individual or combinations of infection courts upon pathogen colonization and disease development in various host tissues.

Yield losses are known to occur as a result of C. coccodes infections (3,21,27,50) but the relationship among disease severity, particularly with infection of specific host tissues, inoculation and/or infestation sites, the extent of these losses, and the interactions among these factors has not been investigated. The objectives of this research were to determine infection frequency of C. coccodes in specific plant tissues as affected by the site of inoculation and/or infestation and to determine the effects of such infections on black dot disease severity as well as yield and market value of the potato crop.

MATERIALS AND METHODS

Field trials. Field trials were conducted in 2003 and 2004 at the Northern Plains Potato Growers Association Irrigated Research Site in central North Dakota. The plot area in 2003 had been pasture land with no previous potato crop but, in 2004, the experiment was the second crop of potato in 3 years. The same trial also was conducted both years in commercial potato fields in west-central Minnesota which previously had been cropped to potato and were presumed to have indigenous levels of the black dot fungus. Trials were planted on 24 April and 29 May 2003 and 15 and 29 April 2004 in Minnesota and North Dakota, respectively. Certified seed tubers of cv. Russet Burbank were used in all four site-years, with the same lot used at both sites in a given year. The trials were managed using standard agronomic practices employed in each region. Fungicides, including chlorothalonil, ethylenebisdithiocarbamates, and fluazinam, were applied to the entire trial as a foliar spray to prevent late blight (Phytophthora infestans, (Mont.) de Bary) and to minimize development of early blight (A. solani). All trials were conducted using overhead irrigation and water was applied at intervals necessary to meet the evapotranspiration demands of the crop. Treatments consisting of four-row blocks were arranged in a randomized complete block design with four replications. The distance between rows was 0.91 m, in-row seed tuber spacing was 0.3 m, and row length was 12.2 and 13.7 m in 2003 and 2004, respectively.

Quantification of C. coccodes in soil. The indigenous level of C. coccodes in the soil each year at the central Minnesota site was quantified using dilution plating techniques as previously described (10), with the following modifications. Soil cores were removed at 0- to 20-cm depths in a grid pattern (equidistance within the trial border), air dried and ground before being combined, and mixed thoroughly. In total, five 5-µg subsamples each of nondiluted and diluted (1:10 with sterile soil) cores were evenly dispersed onto Sorenson’s NP-10 semiselective medium (16). Plates were incubated at 25 ± 2°C for 14 days in the dark, soil particles were washed from the plates under running tap water, and colonies were counted using a stereomicroscope at x65 magnification.

C. coccodes inoculations. Eight isolates of C. coccodes collected from tubers, stems, roots, and stolons of commercial potato plants from across the United States were collected and inoculated using a suberized tuber and foliage for each of four site-years. In 2003, VCG designation was not known prior to performing the trial and, therefore, each C. coccodes VCG is not equally represented. Subsequent testing revealed that, among these eight isolates, two belonged to VCG1, two to VCG2, one to VCG5, and three to VCG6 (19). In 2004, isolates were specifically chosen representing VCG1 to -5: two isolates each of VCG1, 2, and 5 and one each of VCG3 and 4. In either case, each VCG was equally represented in the inoculum mixture; that is, twice the volume was added for each of the single isolates of VCG3 and -4 compared with the two isolates which were used for each of the other VCGs.

The importance of inoculation and/or infestation site was examined in a similar manner at all four site-years. Soil infestations, seed tuber, and foliar inoculations were performed individually and in every combination thereof, resulting in eight treatments, including a noninoculated and noninfested control. In the Minnesota 2003 trial, soil was infested with C. coccodes-colonized rye seed (34). C. coccodes was grown on solid 10% clarified V8 juice (CV8) medium (26) for 7 to 9 days in the dark at 25 ± 2°C. Conidia and microsclerotia were scraped from cultures in sterile water and used to inoculate sterile rye seed. The rye was incubated in the dark at 25 ± 2°C for 4 weeks, air dried for 6 days, and subsequently placed in furrow at planting (IFAP) at a rate of 1.9 g/m. In the remaining three site-years, a C. coccodes-infested agar slurry was utilized to inoculate the soil. Isolates of C. coccodes were grown on CV8 for 2 to 3 weeks in the dark at 25 ± 2°C. Agar cultures were homogenized in a blender and the microsclerotial concentration of each isolate was standardized to 102 CFU/ml. In 2003 in North Dakota, 4 liters of microsclerotia–agar suspension was mixed with 22 liters of vermiculite and applied IFAP at a rate of 164 ml/m of row. At both sites in 2004, the ensuing fungal slurry was applied directly to the field at a rate of 80 ml/m² and tilled into the soil at a depth of 7.5 to 10.0 cm prior to planting.

Inoculum applied to seed tubers was prepared by growing isolates in 10% CV8 liquid medium for 2 to 3 weeks. Fungal cultures were centrifuged at 5,000 rpm for 5 to 7 min and resuspended to an adjusted concentration of 10² spores/ml in a 0.25% gelatin solution to aid in spore adhesion and prevent desiccation. This suspension was sprayed onto suberized seed tubers until each tuber was coated (approximately 60 ml per 450 g). Noninoculated tubers were sprayed with a sterile solution of 0.25% gelatin. The tubers were air dried for 5 to 10 min, placed in paper bags, and stored at 12 ± 2°C and 80 to 85% relative humidity (RH) for no longer than 24 h prior to planting. Foliar inoculations were performed using a microsclerotial suspension prepared utilizing the same procedure as was described for soil inoculations in 2004. At 6 to 8 weeks after planting, the basal portion of plants in each row of the four-row plot, for applicable treatments, were sandblasted with silica sand at 245 kPa of pressure to create wounds for infection (23). A 10⁻² microsclerotia/ml suspension at 15 ml/m was applied to the resulting wounded portion of the canopy using a hand sprayer at 137 kPa of pressure.

Tissue colonization. In all four site-years, the frequency of colonization of C. coccodes was determined throughout the growing season by destructively sampling five plants from each treatment–replication combination at approximately 7-day intervals. The process was initiated 7 days post emergence and continued for 12 weeks in 2003 and 14 days preemergence and continued for 16 weeks in 2004. Three stems per hill (stems originating from a single seed tuber) were assayed on each sampling date by excising a 2- to 3-mm stem segment approximately 10 cm above and below the soil line. A single stolon and root segment, 5 to 7 mm in length, also was collected from each stem. In total, 46,080
and 61,440 isolations for *C. coccodes* were made in 2003 and 2004, respectively. All tissue samples were placed onto culture plates containing solid Sorenson’s NP-10 medium. Cultures were examined for the presence of *C. coccodes* after 3 to 4 weeks of incubation at 25 ± 2°C in the dark. The number of infections per tissue segment was recorded and infection frequency was expressed as percentage per stem. The area under the colonization progress curve (AUCPC) was calculated using weekly colonization data (44):

\[
\text{AUCPC} = \frac{1}{n} \sum_{i=1}^{n} \left[ (W_i + W_{i+1}) / 2 \right] (t_{i+1} - t_i)
\]

where \( W_i \) = percentage of *C. coccodes* colonization at the \( i \)th observation, \( t_i \) = time in days at the \( i \)th observation, and \( n \) = total number of observations. AUCPC values were standardized to enable comparisons among site-years. Standardization was achieved by dividing the AUCPC values for each treatment of the replicated trials from each site-year by the total area of the graph, resulting in relative area under the colonization progress curve (RAUCPC).

**Presumptive VCG analysis.** Monoclonal isolates collected from all tissues and treatments at both sites in 2004 were selected for presumptive VCG analysis using amplified fragment length polymorphism (AFLP) markers (19). Sections of *C. coccodes* grown from tissues sampled at week 1, 2, 3, 7, and 8 were transferred to solid media containing 1.5% agar for hyphal tip or monoconidial isolation by micromanipulation. Permanent cultures were established on silica gel crystals stored at –80°C in a 7.5% skim milk solution using microsclerotia scraped from homogeneous cultures of *C. coccodes* grown on CV8 medium amended with ampicillin at 50 mg/ml for 5 to 7 days (45).

**Disease incidence.** Black dot incidence on stems was assessed visually throughout the growing season. The number of stems in the center two rows of each four-row plot was recorded approximately 3 weeks after emergence. Incidence of black dot infection was assessed by determining the number of infected, wilted, or dead stems with obvious microsclerotial formation characteristic of *C. coccodes* commencing 62 to 115 days after planting (DAP) and continuing for 1 to 3 and 5 to 11 weeks in 2003 and 2004, respectively. Incidence was expressed as the percentage of stems exhibiting black dot disease symptoms.

**Assessment of tuber yield and quality.** The center two rows of each replicated treatment, 9.1 m in length at all four site-years after destructive sampling was completed, were harvested between 125 and 160 DAP. Total yield and United States Department of Agriculture grade data were collected at the end of each growing season for each treatment. In 2004, French fry color and quality ratings also were performed on 25 randomly selected tubers per replication.

**Statistical analysis.** Two-factor analyses of variance (ANOVA) were performed on RAUCPC generated from in vitro tissue assays within each site-year using Proc GLM of SAS (version 9.1; SAS Institute, Cary, NC) with tissue type assayed and inoculation and/or infestation site as main effects. One-way ANOVAs were performed on black dot stem incidence as well as yield grade and processing data, when applicable, across each site year. In all instances, means were differentiated using Fisher’s protected least significant difference (LSD) test (\( \alpha = 0.05 \)). Pearson’s correlation was utilized to compare all combinations of *C. coccodes* colonization at the point in the growing season when frequency was approximately 40 to 50% at each site-year, black dot stem incidence at the final data collection date at each of the four site-years, and total yield. Then, \( \chi^2 \) tests of homogeneity were performed to evaluate the frequency of presumptive VCG recovery across sites (Minnesota and North Dakota), tissues (above and below ground stems, roots, and stolons) and weeks (1, 2, 3, 7, and 8) during which *C. coccodes* isolates were obtained, as well as across all eight treatments (\( \alpha = 0.05 \)). Fisher’s exact tests were performed when underlining assumptions of the \( \chi^2 \) test were not met (\( \alpha = 0.05 \)).

**RESULTS**

**Quantification of *C. coccodes* in soil.** In Minnesota in 2003, the indigenous *C. coccodes* population was 69 propagules per gram (ppg) dry weight of soil and, in 2004, the population was less than 1 ppg dry weight. Levels of indigenous *C. coccodes* in the soil were not determined for each year at the North Dakota site because it was a newly developed potato research site.

**C. coccodes tissue colonization.** *C. coccodes* was detected in stems of potato plants at the first sampling date in all four site-years, including 14 days prior to emergence at both sites in 2004. The progression of *C. coccodes* colonization was variable, in some cases substantially, in noninoculated and noninfested plants among the four site-years when this study was performed (Fig. 1). At the North Dakota site in 2003, the frequency of colonization remained relatively low and unchanged until the last three collection dates of the season. Similar trends were observed at this site in 2004 but *C. coccodes* colonization frequencies began to increase earlier and were higher at the end of the growing season compared with 2003. *C. coccodes* colonization frequency was highest and progressed more rapidly in Minnesota in 2003 when compared with the other three site-years. At this site, *C. coccodes* colonization of noninoculated and noninfested plants was nearly 40% at 28 days after emergence (DAE) and exceeded 80% at 49 DAE compared with between nearly 0 and 50% during that same time period in the other three site-years. Colonization by *C. coccodes* at the Minnesota site in 2004 was similar to that of the North Dakota site that same year.

A significant interaction was observed between the main effects of inoculation and/or infestation site and tissue sampled in colonization rate as expressed by the RAUCPC in the 2004 North Dakota trial (\( P < 0.0001 \)) but not in the 2003 North Dakota trial (\( P = 0.065 \)) or the Minnesota trial in 2003 (\( P = 0.748 \)) and 2004 (\( P = 0.998 \)). The interaction at the 2004 North Dakota trial in 2003 was significant (\( P < 0.05 \)).

[Fig. 1. Percentage of *Colletotrichum coccodes* colonization assayed in vivo from tissue of noninoculated potato plants grown in field trials performed in North Dakota (ND) and Minnesota (MN) in 2003 and 2004. Colonization frequency represents the mean of above and below ground stem, root, and stolon tissue.]
Dakota trial was due, in part, to those treatments with inoculated seed tubers having higher *C. coccodes* colonization frequencies of roots than stolons while those treatments without seed tuber inoculation had higher colonization frequencies of stolons compared with roots (data not shown). There also were significant differences among the main effects of inoculation and/or infestation site as well as tissues sampled in all four site-years (Tables 1 and 2). *C. coccodes* colonization frequencies measured by RAUCPC were significantly different in all site-years among inoculation and/or infestation treatments (Table 1). Similar patterns of tissue colonization were observed in both years in North Dakota. At this site, RAUCPC values of noninoculated and noninfested controls were significantly lower than nearly all inoculated and/or infested treatments. Treatments with multiple inoculation and/or infestation sites also tended to have significantly higher RAUCPC values than those treatments with single inoculation or infestation events. Plants from soil infested + seed tuber + foliar inoculated-treatments displayed the highest level of colonization, although not always significantly so. When comparing multiple inoculation and infestation events, plants from treatments in which seed tubers were inoculated, in combination with either soil infestation or foliar inoculation, tended to have higher colonization levels compared with the combination of soil infestation and foliar inoculation. Again, these differences were not always significant. In Minnesota in 2003, although differences among RAUCPC values were significant, the range of these values was small (Table 1). The noninoculated and noninfested control did not display the lowest RAUCPC values, and additional inoculation and/or infestation events did not consistently increase colonization as was observed in North Dakota, presumably due to high indigenous soil populations present that year (69 ppg of soil). However, at this site in 2004, with relatively low indigenous soil populations (<1 ppg of soil), trends were similar to those observed in North Dakota.

Across site-years, colonization was detected at the first sampling date but progressed more quickly in above and below ground stem tissue than in roots and stolons, resulting in significantly higher RAUCPC values for these tissues (Fig. 2A–D; Table 2). In North Dakota in 2003, below ground stem tissue was infected at significantly higher frequencies than above ground stem tissue while colonization frequencies of above and below ground stems were the same in 2004 at this site (Fig. 2A and B; Table 2). In both years in North Dakota, colonization of stolon tissue was significantly greater than that of root tissue (Fig. 2A and B; Table 2). In Minnesota in 2003, differences in colonization were significantly different among all tissues, with above ground tissue colonization greatest, followed by below ground stems, stolons, and roots (Fig. 2C; Table 2). In 2004 at this same site, there was no significant difference between above and below ground stem colonization or between roots and stolons; however, stems were colonized at significantly higher frequencies than roots and stolons (Fig. 2D; Table 2).

**Presumptive VCG analysis.** In total, 91 *C. coccodes* isolates collected from both sites in 2004 were evaluated via AFLP analysis to determine a presumptive VCG (19). AFLP analysis is not able to differentiate the uncommon VCG4 from other VCgs; therefore, only VCG1, -2, -3, and -5 were detected among isolates collected from sample plants. Across all isolates, the frequency of recovery was not evenly distributed among these four VCgs. Overall, the frequency of recovery of isolates from VCG2 and -5 (28 and 40%, respectively) was substantially greater than that of VCG1 and -3 (10 and 13%, respectively). However, χ² and Fisher’s exact analyses revealed that there was no significant difference in the frequency of recovery of each VCG across the two sites (P = 0.1371; P = 0.1106), across the 5 weeks isolates were characterized (P = 0.1970; P = 0.3369), or among above and below ground stems, stolons, or roots (P = 0.8988; P = 0.9719). A significant difference was observed in the frequency of isolates recovered from each of the eight treatments using both analyses (P = 0.0008; P = 0.0001) (Fig. 3A–D). Interestingly, although the number of isolates recovered per treatment–VCG combination was low, the noninoculated and noninfested control was the only treatment from which no isolates belonging to VCG5 were recovered (Fig. 3D). The mean of isolates belonging to VCG5 recovered from treatments containing inoculated seed tubers, alone or in any combination, ranged from 46 to 73%, whereas those with infested soil and inoculated foliage alone had a mean of 17 and 9%, respectively. Similarly, the treatment containing the combination of soil infestation and foliar inoculation yielded 29% of isolates belonging to VCG5. The inverse was true for isolates belonging to VCG2 (Fig. 3B). The frequency of *C. coccodes* isolates belonging to VCG2 recovered from treatments with inoculated seed tubers was much lower (mean of 13%) than the frequency of this VCG recovered from infested soil or foliar inoculated treatments at 67 and 82%, respectively. The combination of soil infestation and foliar inoculation yielded 71% of isolates belonging to VCG2. Isolates belonging to VCG1 and -3 represented less than 25% of the total for any individual treatment (Fig. 3A and C).

**Black dot disease incidence.** Black dot disease incidence on stems tended to be significantly different only among inoculation and/or infestation treatments when disease incidence was high (Fig. 4A–D). In central North Dakota in 2003, black dot disease incidence ranged from 9.6 to

### Table 1. Relative area under the Colletotrichum coccodes colonization progress curve among inoculation and/or infestation sites across all potato tissues sampled

<table>
<thead>
<tr>
<th>Site of inoculation and/or infestation</th>
<th>North Dakota</th>
<th>Minnesota</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2003</td>
<td>2004</td>
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<tr>
<td>No inoculation and infestation</td>
<td>0.10 (f)</td>
<td>0.20 (f)</td>
</tr>
<tr>
<td>Seed tuber inoculation</td>
<td>0.11 (ef)</td>
<td>0.25 (bc)</td>
</tr>
<tr>
<td>Soil infestation</td>
<td>0.12 (de)</td>
<td>0.21 (ef)</td>
</tr>
<tr>
<td>Foliar inoculation</td>
<td>0.15 (bc)</td>
<td>0.22 (de)</td>
</tr>
<tr>
<td>Soil infestation + seed tuber inoculation</td>
<td>0.14 (bc)</td>
<td>0.27 (ab)</td>
</tr>
<tr>
<td>Seed tuber + foliar inoculation</td>
<td>0.16 (bc)</td>
<td>0.28 (a)</td>
</tr>
<tr>
<td>Soil infestation + foliar inoculation</td>
<td>0.13 (cd)</td>
<td>0.24 (cd)</td>
</tr>
<tr>
<td>Soil infestation + seed tuber + foliar inoculation</td>
<td>0.19 (a)</td>
<td>0.28 (a)</td>
</tr>
<tr>
<td><em>P</em> value</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
</tr>
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</table>

* Values in a column followed by the same letter are not statistically different based on Fisher’s protected least significant difference (α = 0.05). *P* value represents the probability of observing a greater value in the *F* test.

### Table 2. Relative area under the Colletotrichum coccodes colonization progress curve among potato tissues sampled across all noninoculated and noninfested and inoculated and/or infected treatments

<table>
<thead>
<tr>
<th>Tissue</th>
<th>North Dakota</th>
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<th>2004</th>
<th>Minnesota</th>
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<tr>
<td></td>
<td></td>
<td>2003</td>
<td>2004</td>
<td>2003</td>
</tr>
<tr>
<td>Above ground stem</td>
<td></td>
<td>0.16 (b)</td>
<td>0.34 (a)</td>
<td>0.75 (a)</td>
</tr>
<tr>
<td>Below ground stem</td>
<td></td>
<td>0.20 (a)</td>
<td>0.34 (a)</td>
<td>0.63 (b)</td>
</tr>
<tr>
<td>Roots</td>
<td></td>
<td>0.07 (d)</td>
<td>0.13 (c)</td>
<td>0.42 (d)</td>
</tr>
<tr>
<td>Stolons</td>
<td></td>
<td>0.12 (c)</td>
<td>0.15 (b)</td>
<td>0.52 (b)</td>
</tr>
<tr>
<td><em>P</em> value</td>
<td></td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

* Values in a column followed by the same letter are not statistically different based on Fisher’s protected least significant difference (α = 0.05). *P* value represents the probability of observing a greater value in the *F* test.
18.5% at the final data collection date 113 DAP (Fig. 4A). Although nearly twice as much black dot was observed in soil infested + seed tuber + foliar-inoculated plots compared with noninoculated and noninfested plots, there was no significant difference among treatments at any of the four data collection dates. In 2004 at this site, a significant difference was observed among treatments at the last data collection date 138 DAP, with disease incidence ranging from 37.9 to 55.4% (Fig. 4B), substantially higher than that observed in 2003. The noninoculated and noninfested control displayed the least amount of black dot stem incidence, although not significantly different from all inoculated and/or infested treatments.

In the 2003 Minnesota trial, a significant difference was observed among treatments in black dot stem incidence at both data collection dates. Disease incidence ranged from 24.9 to 47.2% at the last data collection date 119 DAP, and the noninoculated and noninfested treatment had significantly lower disease incidence than inoculated and/or infested treatments with the exception of seed tuber inoculation (Fig. 4C). There were no significant differences observed among any of the inoculated and/or infested treatments (Fig. 4C). At this site in 2004, no significant differences were observed among treatments at any data collection date, and black dot disease incidence was low, ranging from 7.7 to 14.7% at the last data collection date at 139 DAP, similar to what was observed in North Dakota in 2003 (Fig. 4D).

Yield and tuber quality assessments. No significant differences in total yield were observed among sites of inoculation and/or infestation in either year at the North Dakota site (Table 3). A significant difference in total yield was observed only among site of inoculation and/or infestation at the 2003 Minnesota site. Although the noninoculated and noninfested treatment did result in the highest total yield, these differences were significant only when compared with treatments with more than one site of inoculation and/or infestation, with the exception of the soil infested + seed tuber + foliar-inoculated treatment. However, differences in total yield were not significantly different in Minnesota in 2004, and no significant differences were observed in marketable yield, including tuber size and quality, or French fry quality among the four site-years (*data not shown*).

**Fig. 2.** Percentage of *Colletotrichum coccodes* colonization assayed in vivo from above and below ground stem, root, and stolon tissue of potato plants grown in field trials performed in A and B, North Dakota and C and D, Minnesota in A and C, 2003 and B and D, 2004. Colonization frequency represents the mean of all plants in noninoculated and noninfested and inoculated and/or infested treatments.
The relationship among black dot stem incidence at the last data collection date, C. coccodes colonization, and total yield was variable among site-years, according to Pearson’s correlation analyses (Table 4). A consistent and significant correlation was observed among all three comparisons only at the 2003 Minnesota site. At this site, black dot stem incidence at 122 DAP and colonization frequency at 60 DAP ($r = 0.83; P = 0.012$), black dot stem incidence and total yield ($r = -0.88; P = 0.004$), and colonization frequency and yield ($r = -0.76; P = 0.030$) all had a highly significant relationship. A significant negative relationship was also observed between C. coccodes colonization frequency 96 DAP and total yield ($r = -0.64; P = 0.087$) at the North Dakota 2003 site as well as black dot stem incidence at 138 DAP and C. coccodes colonization frequency 89 DAP ($r = 0.84; P = 0.010$) at the North Dakota 2004 site. Although some trends were observed among these variables at other site-years, none were significant (Table 4).

DISCUSSION

Although several previous research studies have examined C. coccodes colonization and the development of black dot symptoms in potato (1,6,7,22,31,39,50,51), the results reported here provide a comprehensive comparison of colonization, disease development, and yield. Some of the previous studies concentrated on the development of disease without evaluating the frequency of C. coccodes colonization (1,39). Research conducted with soil infestations of C. coccodes on two cultivars commonly grown in the United Kingdom demonstrated that black dot symptoms appeared at a high rate in root tissue (60 to 90%) at the first assessment date 5 weeks after planting regardless of inoculum level (low versus high) but little or no disease was visible on below ground stems (39). Similar research focused on tuber-borne inoculum determined that symptoms on roots and stolons could be detected within 1 week after inoculating seed tubers, around the time of emergence, whereas symptoms on stems did not appear until approximately 7 to 10 weeks after inoculation (1).

Among studies that have examined C. coccodes colonization of host tissue, in plants assayed from 37 commercial potato fields in Idaho, colonization of both basal and apical stem sections by C. coccodes was correlated with the amount of patho-

![Fig. 3. Frequency of recovery of Colletotrichum coccodes isolates belonging to A, vegetative compatibility group (VCG)1; B, VCG2; C, VCG3; and D, VCG5 among sources of inoculation and/or infestation. Isolates were recovered from potato plants produced in North Dakota and Minnesota trials in 2004 from above and below ground stem, root, and stolon samples taken at five sampling dates across the season.](image)
gen recovered from the soil (7). However, subsequent research determined that, under growth-chamber conditions, colonization by *C. coccodes* at the base of the stem was not affected by soil inoculum density (51). Research performed under commercial growing conditions in the Columbia Basin of central Washington reported that *C. coccodes* was isolated at the first sampling date, as early as 15 days after emergence in above ground stems, and later, 22 days after emergence, in below ground stems; however, a larger number of CFU typically were isolated from below ground stems on subsequent sampling dates (22). More recent research performed under greenhouse conditions by the same group determined that the pathogen moved more quickly downward from a single inoculation point on the above ground stem than toward the apex of the plant (31). Root and stolon tissue was not assayed in any of these studies. Under field trial conditions in Scotland, *C. coccodes* colonization of root tissue produced from disease-free micropropagated plants was similar to that in roots produced from both visually blemish-free and blemished seed tubers when evaluated early in the growing season but was substantially lower at later sampling dates (6). One study has evaluated *C. coccodes* colonization in roots as well as above and below ground stem tissue in inoculated plants under field conditions but did so only once during the growing season at 90 DAP (50). At that point, no differences in colonization frequency among these plant tissues were apparent across the five cultivars evaluated in five trials. To our knowledge, the studies reported here represent the first attempt to evaluate colonization of potato tissue by *C. coccodes* using multiple inoculation and/or infestation sites and all affected tissues, including roots and stolons as well as above and below ground stems, across the entire growing season.

The results reported here illustrate a different picture of tissue colonization than previously has been described. Colonization of stem tissue by *C. coccodes* above and below ground was higher than the colonization frequency of stolons and roots at all four site-years of the study. This trend was true regardless of whether the infection originated from soil infestation, seed tubers, or foliar inoculation. This is in contrast to previous studies which have demonstrated that black dot disease symptoms can be detected first in root tissue.

![Graph A](image1)

**Fig. 4.** Incidence of black dot symptoms in potato stems on the final data collection date for each trial: A, 113 days after planting (DAP) at the North Dakota 2003 trial; B, 138 DAP at the North Dakota 2004 trial; C, 119 DAP at the Minnesota 2003 trial; and D, 139 DAP at the Minnesota 2004 trial. Bars with the same letter are not statistically different based on Fisher’s protected least significant difference ($\alpha = 0.05$).
compared with other plant tissues evaluated (139). Both of these studies evaluated symptom expression, and not tissue colonization of the fungus, the most likely reason for the discrepancies. It is commonly accepted that infections by *C. coccodes* remain latent for an extended time period, and this may be more evident in thicker stem tissue than in finer root and stolon tissue. Among the previous research which evaluated colonization, comparisons in timing of colonization were made only between above and below ground stems, and indicated that above ground stems were colonized approximately a week earlier than below ground stem tissue (22). In the first year of this study, colonization was detected in all tissues sampled at the Minnesota site and in above ground stems and stolons at the North Dakota site at the first sampling date 14 days after emergence. Because the point at which initial infection had taken place presumably was missed, sampling was initiated earlier in the second year of the study. Again, *C. coccodes* stem colonization was detected at both sites in both below ground stems and roots approximately 14 days prior to emergence, which never has been reported previously. Also, colonization was recorded at the first sampling date in all tissues sampled in three of the four site-years. The contrast in these results from previous studies might be attributed to differences in the levels of seed tuber inocula and soil infestation (3,11,14,17), cultivar susceptibility (1,27,48,50,51), or environmental factors (38,46,50).

These studies also corroborate previous work indicating that soil infestations and foliar and seed tuber inoculations are all capable of initiating *C. coccodes* infections. Previous research conducted by inoculating foliage under greenhouse conditions found a correlation between leaf lesions and wilt, as well as wilt and yield for plants, but seed tuber inoculations or soil infestations were not evaluated (3). The effect of foliar inoculations and soil infestations have been investigated individually in the field and greenhouse (27,50). Although some significant differences in stem death and wilting were observed between foliar and noninoculated plants under field conditions as well as between soil and noninfested plants in the greenhouse (27), greater stem colonization occurred with foliar inoculations compared with soil infestations under field conditions (50); however, these sources were not evaluated in the same trial and direct comparisons are not possible.

Previous studies which compared the effect of soil and seedborne inoculum indicated that soil inoculum may cause more black dot than seedborne inoculum (11,28,29,39). Under field conditions in the United Kingdom, varying levels of inoculum applied to seed tubers resulted in increases in black dot infection on stem bases and roots but not consistently across seed tuber disease levels and cultivars, whereas soil infestation more consistently increased black dot infection (39). Seed tuber and soilborne inoculum was investigated in individual field trials which did not allow for direct comparisons to be made, and no combination of the inoculation and/or infestation sources was evaluated (39). Also, under field conditions, soil infestation was reported to result in decreased yield and increased black dot incidence on progeny tubers compared with either light or severe seedborne pathogen levels (11) but no wilt severity or *C. coccodes* colonization levels were examined. More recently, where similar levels of natural soil infestations were present, progeny tubers from seed tubers with low levels of black dot displayed more black dot symptoms than tubers produced from seed tubers with higher levels of black dot (28). This indicates that soil inoculum may cause more infections in progeny tubers than seedborne inoculum. Finally, soil infestations performed in the greenhouse were determined to result in increased sclerotial development on roots and stems when compared with seedborne inoculum (29). However, differences in research methods as well as the type of data collected in the aforementioned research make comparisons between studies difficult. Therefore, gaps remain in our understanding of *C. coccodes* colonization of potato plants.

In all four site-years, sufficient indigenous inoculum was available to establish substantial disease levels in noninoculated and noninfested treatments. Despite this, at three of the four site-years, lower colonization frequency was observed in noninoculated and noninfested treatments compared with inoculated and/or infested treatments, indicating that the inoculations and/or infestations were effective in increasing black dot colonization to varying degrees. Levels of *C. coccodes* in the soil were evaluated prior to planting the trial in both years at the central Minnesota site. Although colonization frequencies were affected by inoculation and/or infestation in both years, no consistent trends were observed in 2003, likely due to the relatively high level of naturally occurring inoculum (approximately 69 ppg of soil). However, in 2004, when the indigenous soil inoculum was <1 ppg, plants from noninoculated and noninfested treatments displayed the lowest levels of colonization, followed by plants from single-inoculation or -infestation and multiple-inoculation and/or -infestation treatments, respectively. Interestingly, it is apparent from these data that inoculum potential <1 ppg of soil was sufficient to establish stem infections as high as 50% midway through the season, while higher levels of inocula present the previous year were effective in

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### Table 3. Total yield (mT/ha) among *Colletotrichum coccodes* inoculation and/or infestation sources across all potato tissues sampled

<table>
<thead>
<tr>
<th>Site of inoculation and/or infestation</th>
<th>North Dakota 2003</th>
<th>North Dakota 2004</th>
<th>Minnesota 2003</th>
<th>Minnesota 2004</th>
</tr>
</thead>
<tbody>
<tr>
<td>No inoculation and infestation</td>
<td>7.94</td>
<td>7.26</td>
<td>9.05 a</td>
<td>8.84</td>
</tr>
<tr>
<td>Seed tuber inoculation</td>
<td>8.08</td>
<td>7.45</td>
<td>8.40 ab</td>
<td>9.33</td>
</tr>
<tr>
<td>Soil infestation</td>
<td>7.98</td>
<td>7.78</td>
<td>8.65 ab</td>
<td>9.38</td>
</tr>
<tr>
<td>Foliar inoculation</td>
<td>7.60</td>
<td>6.95</td>
<td>7.94 bc</td>
<td>9.16</td>
</tr>
<tr>
<td>Soil infestation + seed tuber inoculation</td>
<td>8.16</td>
<td>7.09</td>
<td>7.99 bc</td>
<td>9.00</td>
</tr>
<tr>
<td>Seed tuber + foliar inoculation</td>
<td>8.17</td>
<td>6.51</td>
<td>7.47 c</td>
<td>9.42</td>
</tr>
<tr>
<td>Soil infestation + foliar inoculation</td>
<td>7.98</td>
<td>7.66</td>
<td>7.98 bc</td>
<td>9.21</td>
</tr>
<tr>
<td>Soil infestation + seed tuber + foliar inoculation</td>
<td>7.86</td>
<td>7.46</td>
<td>8.35 ab</td>
<td>9.37</td>
</tr>
<tr>
<td>P value</td>
<td>0.895</td>
<td>0.064</td>
<td>0.012</td>
<td>0.847</td>
</tr>
</tbody>
</table>

* Values in a column followed by the same letter are not statistically different based on Fisher’s protected least significant difference (α = 0.05). P value represents the probability of observing a greater value in the *F* test.

### Table 4. Relationship between black dot stem incidence (Incidence), *Colletotrichum coccodes* colonization frequency (Frequency), and total yield of potato (Yield) as determined by Pearson’s correlation coefficient

<table>
<thead>
<tr>
<th></th>
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</tr>
</thead>
<tbody>
<tr>
<td>Incidence vs. Frequency</td>
<td>0.04</td>
<td>0.84*</td>
<td>0.83**</td>
<td>0.38</td>
</tr>
<tr>
<td>Incidence vs. Yield</td>
<td>−0.01</td>
<td>−0.22</td>
<td>−0.88**</td>
<td>−0.28</td>
</tr>
<tr>
<td>Frequency vs. Yield</td>
<td>−0.64*</td>
<td>−0.42</td>
<td>−0.76**</td>
<td>0.37</td>
</tr>
</tbody>
</table>

* Asterisks indicate Pearson correlation coefficients; ** and * = significant at the α = 0.05 and 0.10 levels, respectively; n = 8.

7 Black dot stem incidence was evaluated at 113, 138, 122, and 139 days after planting for the North Dakota 2003 and 2004 and Minnesota 2003 and 2004 trials, respectively. *C. coccodes* colonization frequency was evaluated at 96, 89, 60, and 101 days after planting for the North Dakota 2003 and 2004 and Minnesota 2003 and 2004 trials, respectively.
raising infection frequencies to nearly 95% at the same time in the growing season. Although levels of soilborne inoculum in naturally occurring infestations of C. coccodes and their relationship to disease development have not been examined in detail, increasing soilborne inoculum was reported to increase black dot disease severity, including foliar necrosis and chlorosis as well as sclerotial development on roots and stems, under greenhouse conditions (29). Also, among 37 potato fields in Idaho, the levels of C. coccodes in the soil were highly correlated with both basal and apical stem colonization (7). A later survey of Idaho potato fields confirmed these reports. C. coccodes levels ranged from 0.2 to 211 ppg of soil and tuber tissue infection was highly correlated with the field soil inoculum levels (3). Soil infestation levels at the central Minnesota trial site in the present study fall within the range documented above.

Inconsistencies in the ability of C. coccodes to affect yield or cause disease are not unexpected and have been reported on numerous occasions with black dot greenhouse and field research (3). Variable results in yield reduction were reported between field experiments performed over 2 years in Idaho when comparing cultivar reaction to foliar inoculations (27). A later study successfully demonstrated that C. coccodes infections significantly reduced yield under both greenhouse and field conditions and that these yield losses could be correlated to wilt, although asymptomatic C. coccodes infections also led to yield reductions (3). In field experiments examining soil infestations and seedborne inoculum on black dot disease development in two cultivars, significant reductions in total tuber yield were observed in only 1 year of this study, when the crop was planted with seed tubers severely infected with C. coccodes, even though plants were noticeably colonized by C. coccodes (39). Differences in tuber weight reductions were reported under greenhouse conditions when comparing foliar to root inoculations (2). In that study, root inoculations reduced tuber weights more than foliar inoculations but disease progression and colonization were not evaluated. In the current study, root colonization by C. coccodes generally lagged behind above ground infections; therefore, it is clear why yield reductions were not detected in most site years. In contrast, foliar inoculations decreased yield more than both seed tuber inoculation and soil infestation, although these differences were significant at only one site-year. This disparity is most likely due to the presence of natural inoculum, differences in inoculation methods, and environmental factors (which were controlled under greenhouse conditions), in addition to difficulties measuring yield reductions caused by this pathogen.

Differences in aggressiveness among VCGs of C. coccodes (2,19,32,34) may play a role in these inconsistencies because most studies involving the impact of black dot on yield of potato were performed before vegetative compatibility was reported in this fungus. One recent research study has taken into account C. coccodes VCGs (29). Two isolates of VCG2 led to higher disease incidences when originating from soil than from seed tubers, whereas the opposite was true for a third isolate belonging to VCG1. These results support the findings reported here, in which higher frequencies of isolates of VCG2 were recovered from plants that had grown in infested soil when compared with those from inoculated seed tubers. Some of the past contradictory yield results also may be attributed to differences in cultivar susceptibility. Data generated in both field and greenhouse trials demonstrated that later-maturing cultivars are more likely to suffer yield reductions than earlier-maturing cultivars (2,27). Also, recently reported results of colonization of control cultivars and breeding selections by C. coccodes grown in naturally infested soil indicated that differences among cultivars or selections exist and that these differences were significantly affected by environmental conditions (30). Although there are other examples of contradictory reports concerning the effect of C. coccodes on yield of potato, the aforementioned research results provide an ample illustration of the difficulties that lie in quantifying direct affects of this pathogen. The effects of black dot on yield reported here are consistent with the observations made in the several earlier studies. The Minnesota 2003 site was the only one to have significant yield loss compared with the nonoinoculated and non-infested control. At this site, the significant negative correlation between black dot incidence at the time of haulm desiccation on 22 August and yield in the late-maturing cv. Russet Burbank may provide some indication of yield loss. A similar comparison was made with soil inoculum, C. coccodes colonization levels, and wilt on 23 August in previous research performed in Idaho (7). An association of this type potentially may act as a predictor of season-end black dot incidence and, ultimately, effect on tuber yield. These relationships clearly should be investigated in further studies.

C. coccodes still often is considered to be a weak pathogen, attacking plants following periods of stress or causing blemsishes on tubers; however, results reported here and from similar work demonstrate that the picture is much broader. The effects of C. coccodes on yield and tuber quality ultimately will be tied to a variety of factors, such as inoculum potential, environmental conditions, cultural practices, cultivar, and pathogen VCG. Although black dot may not cause reductions in yield and tuber quality in all instances, C. coccodes remains a serious threat to commercial potato production and the seed potato industry, particularly in areas where Verticillium wilt is a concern and interactions between the two pathogens occur (8,35,42,43,51). The data reported here may be useful in establishing the proper timing of fungicides such as azoxystrobin, which is highly efficacious (28). Because the infections that are most likely to become symptomatic occur early in the growing season, fungicide applied immediately following symptom development may be the most effective.

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LITERATURE CITED

Effects of wounding and wetting duration on Verticillium wilt, and powdery scab on Russet Burbank potato. Plant Dis. 78:1075-1078.


Miller, P. M. 1955. V8 juice agar as a general purpose medium for fungi and bacteria. Phytopathology 45:461-462.


