Diversity, Virulence, and 2,4-Diacetylphloroglucinol Sensitivity of *Gaeumannomyces graminis* var. *tritici* Isolates from Washington State

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**ABSTRACT**


We determined whether isolates of the take-all pathogen *Gaeumannomyces graminis* var. *tritici* become less sensitive to 2,4-diacetylphloroglucinol (2,4-DAPG) during wheat monoculture as a result of exposure to the antibiotic over multiple growing seasons. Isolates of *G. graminis* var. *tritici* were baited from roots of native grasses collected from noncropped fields and from roots of wheat from fields with different cropping histories near Lind, Ritzville, Pullman, and Almota, WA. Isolates were characterized by using morphological traits, *G. graminis* variety-specific polymerase chain reaction and pathogenicity tests. The sensitivity of *G. graminis* var. *tritici* isolates to 2,4-DAPG was determined by measuring radial growth of each isolate. The 90% effective dose value was 3.1 to 4.4 µg ml⁻¹ for 2,4-DAPG-sensitive isolates, 4.5 to 6.1 µg ml⁻¹ for moderately sensitive isolates, and 6.2 to 11.1 µg ml⁻¹ for less sensitive isolates. Sensitivity of *G. graminis* var. *tritici* isolates to 2,4-DAPG was normally distributed in all fields and was not correlated with geographic origin or cropping history of the field. There was no correlation between virulence on wheat and geographical origin, or virulence and sensitivity to 2,4-DAPG. These results indicate that *G. graminis* var. *tritici* does not become less sensitive to 2,4-DAPG during extended wheat monoculture.

Additional keywords: biological control, *Pseudomonas fluorescens*, take-all.
order to maximize the effectiveness of TAD for growers who crop wheat continuously or in rotation with barley. The specific objectives of this study were to determine the range of sensitivities to 2,4-DAPG among isolates of *G. graminis var. tritici* from TAD and non-TAD soils and to determine whether *G. graminis var. tritici* isolates with reduced sensitivity to 2,4-DAPG become enriched in wheat monoculture soil as a result of exposure to the antibiotic over multiple growing seasons.

**MATERIALS AND METHODS**

**Isolates, media, and storage.** Six strains of *G. graminis var. tritici* were used as controls throughout this study: R3-111a-1, ARS-A1, MV-116, MV-119, L-109, and L-116 (14). R3-111a-1 originally was isolated in 1980 from wheat grown in a TAD field near Moses Lake, WA. ARS-A1 was isolated in 1990 from wheat grown on the USDA-ARS Palouse Conservation Field Station near Pullman. MV-116 and MV-119 were isolated in 1986 from wheat grown on the Washington State University (WSU) Mount Vernon Northwestern Washington Research & Extension Center, Mt. Vernon, WA. L-109 and L-116 were isolated in 1986 from the TAD field at the WSU Lind Dryland Research Station, Lind, WA.

*G. graminis var. tritici* isolates were routinely cultured on one-fifth strength potato dextrose agar (1/5× PDA): potato dextrose broth (PDB, 4.5 g), (BD, Sparks, MD), agar (20 g) (Sigma Chemical Co., St. Louis, MO), and water (1 liter). The *G. graminis var. tritici* semiselective medium (R-PDA) (14) was used to isolate *G. graminis var. tritici* from roots. For R-PDA, peeled potato slices (40 g) were boiled for 10 min and the filtered extract was increased to a volume of 1 liter with water. Dextrose (4 g liter⁻¹) and agar (18 g liter⁻¹) were added and the pH of the medium was adjusted to 6.5 before autoclaving. Rifampicin (100 µg ml⁻¹) (Sigma Chemical Co.) and Rizolex (Tolclofos-Methyl; purity 98%) (1 µg ml⁻¹) (Chem Service, West Chester, PA) were added after autoclaving when the medium cooled to below 60°C.

Isolates were stored by a variety of methods. R3-111a-1 and ARS-A1 were stored on 1/5× PDA amended with rifampicin (100 µg ml⁻¹) at 4°C and once a year isolates were taken out of storage, cultured on 1/5× PDA, and tested for virulence on wheat as described below. Each isolate was reisolated from diseased roots, grown on 1/5× PDA with rifampicin, and again placed at 4°C. Isolates MV-116, MV-119, L-109, and L-116 were stored in glycerol as hyphal fragments at –80°C. *G. graminis var. tritici* isolates collected during this study were stored at room temperature on slants of 1/5× PDA, with and without mineral oil, and as colonized 1/5× PDA agar plugs in autoclaved distilled water (15).

**Isolation and culture conditions of *G. graminis var. tritici*.** *G. graminis var. tritici* was isolated from wheat grown in fields in eastern and central Washington State that had undergone at least 3 years of wheat or barley monoculture or crop rotation. *G. graminis var. tritici* was also isolated from native grasses growing in noncropped (virgin) fields. Table 1 shows a list of the fields from which *G. graminis var. tritici* isolates were collected. The Lind TAD (LD) field is located on the WSU Lind Dryland Research Station (33). LD isolates were from healthy looking wheat randomly selected throughout the field. LDP isolates were also from the Lind TAD field but were isolated from wheat with symptoms of take-all within distinct take-all patches. The Lind virgin (LV) site has never been cropped and is located about 100 m from the Lind TAD field (33). The Pullman conducive field (PC) (32) is located on the USDA-ARS Palouse Conservation Field Station near Pullman. The Almota TAD (ADB) field is located on a commercial farm near the Port of Almota and is direct seeded. The Ritzville TAD (RD) field is located on a commercial field near Ritzville, WA, and is irrigated.

Most isolates were collected by using a baiting method and a tube assay essentially as described by Cook and Naiki (11). Roots were washed under running tap water to remove soil and then blotted dry with a paper towel. Roots from a single plant were excised, rolled into a ball, and placed on the top of a 6-cm column of autoclaved vermiculite inside a plastic tube (15 cm long, 2.5 cm diameter, Stuewe and Sons Inc., Corvallis, OR); 100 tubes were hung in a plastic rack. The roots were covered with a 1-cm layer of autoclaved vermiculite; three wheat seeds (spring wheat cv. Penawawa) were sown and then covered with a 1-cm layer of vermiculite. Each tube received 12 ml of water and the rack with tubes was covered with a sheet of plastic until the shoots emerged. The tubes were watered with 12 ml of water twice a week and with dilute (1/3 strength) Hoagland's nutrient solution (macro-element only) once a week. The racks were placed in a growth room at 16°C, with a 16 h photoperiod. If *G. graminis var. tritici* is present in the ball of roots, the fungus will infect the wheat seedlings. After 3 to 8 weeks, plants were removed from the plastic tubes, and roots were washed to remove the vermiculite and inspected for take-all lesions. Roots with lesions were washed for 2 h in running tap water and then surface-disinfested by immersion in a solution of 0.5% silver nitrate for 1 min, followed by rinsing twice in distilled water. Roots were placed on autoclaved Whatman no. 1 filter paper (Whatman Inc., Florham Park, NJ). Isolates were again cultured on PDA, then washed under running tap water to remove the filter paper.

**TABLE 1. Cropping history of fields and geographic origin of *Gaeumannomyces graminis var. tritici***

<table>
<thead>
<tr>
<th>Location and isolate</th>
<th>Isolate</th>
<th>Number of isolates</th>
<th>Cropping history</th>
<th>Host</th>
<th>Presence of phlD⁺ isolates</th>
<th>Soil suppressive to take-all</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pullman conducive</td>
<td>PC1–13</td>
<td>13</td>
<td>Crop rotation</td>
<td>Wheat</td>
<td>–</td>
<td>–</td>
<td>This study</td>
</tr>
<tr>
<td>Almota TAD</td>
<td>ADB 1–8; 10–15; 20–23</td>
<td>18</td>
<td>7 years com., 4 years continuous wheat</td>
<td>Wheat</td>
<td>+</td>
<td>+</td>
<td>This study</td>
</tr>
<tr>
<td>Ritzville TAD</td>
<td>RD 1–6; 8–47</td>
<td>46</td>
<td>3 years continuous wheat</td>
<td>Wheat</td>
<td>+</td>
<td>+</td>
<td>This study</td>
</tr>
<tr>
<td>Lind virgin</td>
<td>LV 1–30</td>
<td>30</td>
<td>Non-cropped</td>
<td>Native grasses</td>
<td>–</td>
<td>–</td>
<td>This study</td>
</tr>
<tr>
<td>Lind TAD⁺</td>
<td>LD 1–40</td>
<td>40</td>
<td>39 years continuous wheat</td>
<td>Wheat</td>
<td>+</td>
<td>+</td>
<td>This study</td>
</tr>
<tr>
<td>Lind TAD patch</td>
<td>LDP 1–30</td>
<td>30</td>
<td>39 years continuous wheat (inside patch)</td>
<td>Wheat</td>
<td>+</td>
<td>+</td>
<td>This study</td>
</tr>
<tr>
<td>MV116, MV119</td>
<td>2</td>
<td></td>
<td></td>
<td>Wheat</td>
<td></td>
<td>(24)</td>
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<tr>
<td>L109, L116</td>
<td>2</td>
<td></td>
<td></td>
<td>Wheat</td>
<td></td>
<td>(24)</td>
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<tr>
<td>R3-111a-1</td>
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<td></td>
<td></td>
<td>Wheat</td>
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<td>(24)</td>
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<tr>
<td>ARS-A1</td>
<td>1</td>
<td></td>
<td></td>
<td>Wheat</td>
<td></td>
<td>D. M. Weller, unpublished</td>
<td></td>
</tr>
</tbody>
</table>

- *The Pullman conducive field was direct-seeded and rain-fed and located on the USDA-ARS Palouse Conservation Field Station. The Almota TAD field was direct-seeded and rain-fed and located on a commercial farm. The Ritzville TAD field was conventionally tilled and irrigated and is located on a commercial farm. The Lind virgin and TAD fields are located on the WSU Lind Dryland Research Station. The Lind TAD field is irrigated and has been both conventionally cultivated and direct-seeded during its history.
- + indicates wheat grown in the soil supports threshold population densities of 2,4-DAPG-producing *Pseudomonas fluorescens* (>10⁸ colony forming units [CFU] root⁻¹) (phlD⁺ isolates). – indicates wheat grown in the soil supports population densities of phlD⁺ isolates below the detection limit of 5 × 10⁹ CFU root⁻¹.
- LD isolates were from “healthy-looking” wheat plants randomly selected throughout the field. LDP isolates were from wheat plants within distinct patches.
Park, NJ) to remove excess water. Roots with lesions were cut into pieces (<1 cm) and placed on R-PDA. Plates were incubated at room temperature in the dark for up to 2 weeks until hyphae grew out from the pieces of root. Hyphae of putative *G. graminis* var. *tritici* isolates were selected based on morphological characteristics of hyphae and color change in the medium around the hyphae, and then hyphal tips were transferred to R-PDA. *G. graminis* var. *tritici* alters the color of rifampicin in R-PDA from orange to purple, and this reaction occurs in as little as 24 h (14). Over 250 putative isolates of *G. graminis* var. *tritici* were isolated and stored as described above.

**DNA extraction and Gaeumannomyces graminis variety-specific polymerase chain reaction.** *G. graminis* var. *tritici* mycelium was incubated at room temperature for 1 week in 1/5x PDB (BD, Sparks, MD) either in 1.5-ml tubes on a shaker at 150 rpm or in slowly shaken petri dishes. The mycelium was harvested by centrifugation at 5,000 rpm for 5 min and then washed twice in sterile distilled water. DNA was extracted by using a FastDNA Kit (Bio101, Carlsbad, CA) (28) and eluted in 100 µl of DNA elution solution. DNA was amplified with *G. graminis* variety-specific primers, NS5 and GGTP-RP (16). The polymerase chain reaction (PCR) volume was 50 µl: it contained 200 µM dNTP mixture, 2 mM MgCl2, 10 µl of 5x Green GoTaq Flexi buffer (Promega Corporation, Madison, WI), 0.4 µM of each primer; 50 ng of DNA, and 1 unit of *Taq* polymerase (Promega Corporation, Madison, WI). The PCR was performed in an MJ Research PTC-200 thermal cycler (Bio-Rad, Hercules, CA). An initial denaturation step at 95°C for 5 min was followed by 35 cycles consisting of denaturation at 95°C for 45 s, annealing at 52°C for 1 min, and extension at 72°C for 1 min. After 35 cycles, samples were incubated at 72°C for 10 min for complete extension of PCR products. PCR amplicons were visualized by gel electrophoresis in 1.5% agarose. A single 410-bp PCR fragment indicated *G. graminis* var. *tritici*, a single 310-bp fragment indicated *G. graminis* var. *avenae*, and no PCR product indicated *G. graminis* var. *graminis* or a genus other than *Gaeumannomyces* (16).

To amplify avenacinase-like genes with variety-specific avenacinase-like gene primers (34), the PCR volume was 50 µl and contained 25 pmol of each of the three variety-specific 5' primers and 75 pmol of 3' AV-3 primer. An initial denaturation step at 95°C for 5 min was followed by 35 cycles consisting of denaturation at 95°C for 45 s, annealing at 64°C for 45 s, and extension at 72°C for 2 min. To obtain the sequence of partial avenacinase-like genes, PCR products were cleaned using DTR gel filtration cartridges (EdgeBio, Gaithersburg, MD). Sequencing reactions were performed by using a dye-terminator cycle sequencing reaction (ABI-Prism, Foster City, CA) according to the manufacturer’s protocol. Reactions were cleaned again using DTR gel filtration cartridges and then completely dried. The reactions were run on an ABI 377 sequencer at the Center for Integrated Biotechnology Bioinformatics Core Facility, Washington State University, Pullman.

**In vitro 2,4-DAPG sensitivity assay.** *G. graminis* var. *tritici* isolates were tested for sensitivity to 2,4-DAPG (Toronto Research Chemicals Inc., Toronto, Canada) in agar plate bioassays. A 4-mm-diameter agar plug was cut with a cork borer from the margin of a 1-week-old culture of an isolate of *G. graminis* var. *tritici* grown on 1/5x PDA. The agar plug was then transferred with the mycelium facing downward to the center of a 36-mm-diameter well containing 5 ml of 1/5x PDA in a 6-well cell culture cluster (Corning Inc., Corning, NY). Each well contained medium amended with either 0, 1, 2, 3, 4, or 5 µg ml⁻¹ of 2,4-DAPG dissolved in methanol. This range of concentrations was used because of the previous findings of Mazzola et al. (24), who demonstrated that *G. graminis* var. *tritici* isolates that were not inhibited by 3 µg ml⁻¹ of 2,4-DAPG were not suppressed on wheat by a *phiD* strain of *P. fluorescens*. Hyphal growth was measured daily from 3 to 7 days after inoculation. Radial growth was measured from the center of the plug to the edge of the mycelium. Each measurement was performed in four different directions per well. Radial growth measurements were converted to percent growth inhibition by comparing growth on medium amended with the antibiotic to that on medium with no 2,4-DAPG (control, methanol only). Hyphal growth measurement data at day 6 provided the greatest difference between the control and 2,4-DAPG treatments, and thus was used for the final analysis of growth inhibition. Growth inhibition data were transformed to probits and linear regression was performed on transformed inhibition data versus 2,4-DAPG concentration using SigmaPlot 8.0 (SPSS Inc., Chicago, IL). Points from all four replicates were pooled together (24 points: four replicates × six dosages) to run the regression. The 90% effective dose (ED₉₀) values were calculated from the predicted regression equation. Three 2,4-DAPG sensitivity categories (sensitive, moderately sensitive, and less sensitive) were generated based on quartiles from the frequency distribution of ED₉₀ values (1st, 2nd and 3rd combined, and 4th).

**Virulence assay.** *G. graminis* var. *tritici* isolates were tested for virulence on wheat by the tube assay described above except that instead of adding a ball of roots to the assay, two 9-mm agar plugs, cut with a cork borer, from a 1-week-old culture of an isolate grown on 1/5x PDA were added to the column of vermiculite. Virulence tests of each isolate were replicated five times with a replicate consisting of five individual tubes sown with three seeds each. The experiment was arranged in a randomized complete block design. Tubes were incubated and watered as described above. After 4 weeks, plants were removed, washed, and rated for disease on a 0 to 8 scale: 0 = healthy, 8 = dead or nearly so (27).

**Production of perithecia and measurements of ascospores.** To stimulate perithecia formation, plants infected with *G. graminis* var. *tritici* were wrapped in wet paper towels, placed in a plastic bag, and incubated in a plant growth room (16°C, 16 h photoperiod) for 8 to 12 weeks. Perithecia were observed under a dissecting microscope and plants with mature perithecia were washed briefly in sterile water before selecting perithecia. Three perithecia of each isolate were transferred to a microscope glass slide, crushed, and then examined at ×40 and ×100 magnifications with an Olympus BX41TF microscope (Olympus, Center Valley, PA). For each isolate of *G. graminis* var. *tritici*, pictures of at least 30 ascospores were taken and images were saved as JPEG files. The length of each ascospore was measured using the ImageJ program (available online) (8).

**Statistical analysis.** The ED₉₀ values for isolates from different locations were analyzed by analysis of variance (ANOVA) and means were separated by Tukey’s honestly significant difference test (Statistix 9.0, Analytical Software, St. Paul, MN) to determine if there was any difference in sensitivity to 2,4-DAPG among isolates from different locations. Disease severity of isolates from the various locations was compared by ANOVA, and means were separated by a protected least significant difference test (*P* = 0.05) by SAS 9.1.3 (SAS Institute, Inc., Cary, NC). Linear regression analyses (SigmaPlot 8.0) were performed to determine the relationship between the ED₉₀ of 2,4-DAPG and the virulence of *G. graminis* var. *tritici* isolates.

**RESULTS**

Isolation of *G. graminis* var. *tritici* from fields with different cropping histories. *G. graminis* var. *tritici* was isolated from roots of grasses from noncropped virgin sites and from wheat grown in fields that had undergone crop rotation or three or more years of continuous wheat. Roots of wheat grown in soil from the Pullman conducive and Lind virgin fields do not contain threshold population densities of 2,4-DAPG-producing *P. fluorescens* (>10⁶ CFU g⁻¹ of root) (density needed to suppress take-all; 32), whereas soils from the other fields contained threshold densities (Table 1). Over 250 putative *G. graminis* var. *tritici* isolates were
isolated from roots on R-PDA and tentatively identified as *G. graminis* var. *tritici* on the basis of a color change on R-PDA and morphological characteristics such as colony color and curling back of the hyphae on the edge of the colony (14). On R-PDA, the color of the medium changes from orange to purple around the *G. graminis* var. *tritici* mycelium. We focused on 177 new *G. graminis* var. *tritici* isolates: 134 isolates from fields with three or more years of continuous wheat or barley, 13 isolates from the Pullman take-all conducive field, and 30 isolates from the Washington noncropped (virgin) soil (Table 1).

**Identification of *G. graminis* var. *tritici* by morphological characteristics and PCR.** Of the 183 isolates used in this study (177 new and 6 control isolates), 135 were tested for ability to produce perithecia. Perithecia characteristic of *G. graminis* var. *tritici*, were produced by 119 isolates on infected wheat roots (cv. Penawawa), but were not formed by the other isolates after 12 weeks of incubation. Ascospore length ranged from 61 to 90 µm (average 78.3 µm) (data not shown), which is typical of *G. graminis* var. *tritici*. Among the isolates, the color of the hyphae and the growth habit on 1/5x PDA (data not shown) varied, which is also typical of *G. graminis* var. *tritici* (22). We also observed simple hypophodia on the outer layers of wheat seedling coleoptiles (data not shown) produced by control (ARS-A1, R3-111a-1) and new (i.e., ADB-15) isolates of *G. graminis* var. *tritici*.

We then performed *G. graminis* variety-specific PCR with NSS and GGT-RP primers, which amplify 18S ribosomal DNA of *G. graminis*. All 183 *G. graminis* var. *tritici* isolates used in this study generated a single 410-bp PCR product indicative of var. *tritici* (Fig. 1).

It was previously reported (34) that nucleotide differences in avenacinase-like genes can distinguish among *G. graminis* vars. *avenae*, *graminis*, and *tritici* based upon the size of PCR amplicons. All of the 183 *G. graminis* var. *tritici* isolates from Washington State responded positively with *G. graminis* variety-specific mixed primers (each of the three variety-specific primers and the 3′ common primer). The size of the PCR amplicons was approximately 1,100-bp (data not shown). Because Rachawong et al. (34) reported that amplification of the var. *tritici* and var. *graminis* avenacinase-like genes generated 870- and 1,086-bp PCR products, respectively, we sequenced the primer’s binding site of the avenacinase-like genes from six of our *G. graminis* var. *tritici* isolates (Fig. 2A and B) in order to determine the basis of the discrepancy. The var. *tritici*-specific primer sequence is 5′-TCCTCGGGCCCGTGTAGGGC-3′. Compared with the isolates of Rachawong et al. (34), Washington State isolates had two different nucleotides in the primer binding site (C to T). However, the primer binding site sequences were identical in the Washington isolates and the American Type Culture Collection (ATCC, Manassas, VA) type strain ATCC28230 (Fig. 2B). Unexpectedly, the sequence of the var. *graminis*-specific primer (5′-ACCC-CCGTCCTCCGTA-3′) was complementary to the binding site of the avenacinase-like gene in *G. graminis* var. *tritici* isolates from Washington and strain ATCC28230 (Fig. 2A). As a result, PCR with the *G. graminis* variety-specific primers generated amplicons of approximately 1,100 bp with Washington isolates.

**Sensitivity of *G. graminis* var. *tritici* to 2,4-DAPG.** Of the 183 *G. graminis* var. *tritici* isolates used in this study, 155 were tested for sensitivity to 2,4-DAPG (Fig. 3). Regardless of source, isolates varied in sensitivity to 2,4-DAPG. In all soils, sensitivity of isolates to 2,4-DAPG was normally distributed. For isolates from all soils, the range of ED$_{50}$ of 2,4-DAPG was 3.1 to 11.1 µg ml$^{-1}$. Based on ED$_{50}$ frequency distributions, *G. graminis* var. *tritici* isolates were divided into three sensitivity categories: sensitive (ED$_{50}$; 3.1 to 4.4 µg ml$^{-1}$) (37 isolates), moderately sensitive (ED$_{50}$: 4.5 to 6.1 µg ml$^{-1}$) (75 isolates), and less sensitive (ED$_{50}$: 6.2 to 11.1 µg ml$^{-1}$) (37 isolates). The ratio of sensitive, moderately sensitive, and less sensitive isolates was approximately 1:2:1. The average ED$_{50}$ was 4.6 µg ml$^{-1}$ for ADB isolates; 5.8 µg ml$^{-1}$ for LD isolates; 5.8 µg ml$^{-1}$ for LDP isolates; 5.1 µg ml$^{-1}$ for LV isolates; 5.5 µg ml$^{-1}$ for PC isolates; and 5.3 µg ml$^{-1}$ for RD isolates. Sensitivity of isolates to 2,4-DAPG was not correlated with cropping history (Fig. 4). Six isolates were exceptions to the above pattern: ADB-11 was insensitive to 2,4-DAPG (ED$_{50}$ >11.1 µg ml$^{-1}$) and isolates ADB-14, LD-5, LDP-14, LDP-21, and LV-2 were extremely sensitive to 2,4-DAPG (ED$_{50}$ <1 µg ml$^{-1}$). Exact ED$_{50}$ values for these isolates were not calculated because they were outside of the range of linear regression of growth inhibition. These isolates are the focus of another ongoing study.

**Virulence of *G. graminis* var. *tritici* isolates.** Of the 183 *G. graminis* var. *tritici* isolates used in this study, 135 were tested for virulence on wheat (cv. Penawawa). Most of these isolates caused significant disease. Exceptions included the eight isolates LD-27, LDP-14, LD-28, ADB-14, ADB-11, LD-24, LD-33, and PC-2, which had disease ratings of less than 1. Disease ratings ranged from 0 to 5.78. Virulence tests divided *G. graminis* var. *tritici* isolates into three groups based on the frequency distribution, weakly virulent (rating 0 to 2.83) (34 isolates), moderately virulent (rating 2.84 to 4.19) (67 isolates), and strongly virulent (rating 4.2 to 5.78) (34 isolates); the ratio was approximately 1:2:1, respectively. Populations of isolates from different locations did not differ significantly in virulence (Fig. 5). In addition, linear regression analyses showed that among *G. graminis* var. *tritici* isolates from Washington State fields there was no correlation between virulence (as determined in the tube assay) and sensitivity to 2,4-DAPG in vitro (Fig. 6).

**DISCUSSION**

In the Pacific Northwest and The Netherlands, TAD develops because of the buildup of populations of 2,4-DAPG-producing *P. fluorescens* and production of the antibiotic in the wheat or barley rhizosphere (38). We frequently have been asked whether isolates of *G. graminis* var. *tritici* with reduced sensitivity to 2,4-DAPG become enriched in monoculture wheat and barley fields as a...
result of exposure to the antibiotic. To address this question, we compared the ED$_{90}$ for 2,4-DAPG against isolates of *G. graminis* var. *tritici* from well-studied TAD and non-TAD fields. ED$_{90}$ values for all isolates tested ranged from 3.1 to 11.1 µg ml$^{-1}$, and like Mazzola et al. (24), we found that isolates showed differential sensitivity to 2,4-DAPG. However, sensitivity to the antibiotic was normally distributed among isolates regardless of the source (monoculture versus nonmonoculture fields), and the average ED$_{90}$ values for isolates from the fields tested did not differ statistically with the exception of ADB isolates, which had a lower average ED$_{90}$ value than those of the LD and LDP isolates (Fig. 4).

Of particular interest were the results from Lind because the Lind TAD and virgin fields are located only about 100 m apart. The average ED$_{90}$ for isolates from the non-cropped Lind virgin site (no detectable 2,4-DAPG producers) was 5.1 µg ml$^{-1}$, whereas the average ED$_{90}$ for isolates from the Lind TAD field (threshold *phlD* population) was 5.8 µg ml$^{-1}$. Thus, after 39 years of continuous wheat monoculture, significant changes in sensitivity to 2,4-DAPG of isolates in the Lind TAD field had not occurred. In addition, within the Lind TAD field, the average ED$_{90}$ for isolates inside and outside of take-all patches was identical, suggesting that the patches do not result from pathogen isolates that are less sensitive to 2,4-DAPG. TAD effectively suppresses the development of severe take-all in the field, but the robustness of its expression varies among fields and years. Fluctuations in take-all incidence and severity are a normal but poorly understood part of this dynamic process (22). In Washington TAD fields, small take-all patches typically occur even in strongly suppressive fields. In years when TAD is weakly expressed, patches are much larger and more numerous.

Collectively, our results indicate that (i) differential sensitivity to 2,4-DAPG is a natural characteristic of *G. graminis* var. *tritici* populations, (ii) production of 2,4-DAPG in the rhizosphere does not exert a selection pressure on *G. graminis* var. *tritici* in Washington State TAD fields, and (iii) isolates with reduced sensitivity to the antibiotic do not appear to emerge in a field even after decades of wheat monoculture and exposure to the antibiotic. To our knowledge, this is the first study addressing the question of emergence of resistance in a target pathogen to an introduced or indigenous biocontrol agent.

**Fig. 2.** Alignment of variety-specific polymerase chain reaction primers with avenacinase-like gene sequence in *Gaeumannomyces graminis*. The highlighted sequence indicates a different template DNA sequence from the variety-specific primer. **A,** *G. graminis* var. *graminis*-specific primer binding sequence; and **B,** *G. graminis* var. *tritici*-specific primer binding sequence. Sequences from ATCC strain 28230; *Ggt* CB1, M1, and CH1; reference strains ARS-A1 and R3-111a-1; and isolates from eastern Washington fields, ADB-3, ADB-11, PC-1, and LDP-21.

**Fig. 3.** Six-well plate assay used to test the sensitivity of isolates of *Gaeumannomyces graminis* var. *tritici* to 2,4-diacyethylphloroglucinol (2,4-DAPG). 1/5× potato dextrose agar was amended with either 0, 1, 2, 3, 4, or 5 µg of 2,4-DAPG ml$^{-1}$. A plug of the fungus was placed in the center of the well. Radial growth of each isolate well was measured over a period of 3 to 7 days after inoculation. The picture was taken at 6 days after inoculation.
Several factors may explain why isolates of *G. graminis* var. *tritici* with reduced sensitivity to 2,4-DAPG have not emerged or become enriched in Washington State TAD soils. For example, the pathogen may be exposed to inhibitory concentrations of the antibiotic only during its parasitic phase in sites, such as infection courts and lesions, where *phlD* + bacteria proliferate. During the rest of the pathogen’s life cycle, while growing saprophytically or surviving in crown and root tissues, 2,4-DAPG may not be present or is present at extremely low levels. This scenario would reduce the selection pressure on *G. graminis* var. *tritici* for the emergence of isolates with less sensitivity to the antibiotic, as compared to if the pathogen was exposed to the antibiotic throughout its life cycle.

Regression analysis revealed no correlation between antibiotic sensitivity and virulence (Fig. 6), suggesting that reduced sensitivity to 2,4-DAPG does not affect the pathogen’s virulence. However, the impact of reduced antibiotic sensitivity on the saprophytic growth and survival of *G. graminis* var. *tritici* has not been assessed. Elucidating the fine details about the spatial and temporal accumulation and subsequent degradation of 2,4-DAPG in the rhizosphere environment and in plant debris will help to clarify the extent to which the pathogen is exposed to the antibiotic. However, quantifying antibiotics in the rhizosphere of field-grown plants currently requires extractions from multiple pooled root systems, and detecting 2,4-DAPG in individual microsites is currently beyond the capacity of commonly used biochemical techniques.

A second factor may relate to the wheat or barley varieties grown a TAD field. Washington State growers often change wheat varieties for economic reasons or pest control. It is well documented that both the crop species and varieties (including of wheat) differentially enrich and support *phlD* + populations (4,13, 25) and genotypes (23,25,30), and modulate 2,4-DAPG accumulation in the rhizosphere (4,26). For example, Okubara and Bonsall (26) reported significant differences in the accumulation of 2,4-DAPG on the roots of wheat cultivars Tara, Buchanan, and Finley by *P. fluorescens* Q8r1-96, a strain typical of those primarily responsible for TAD in Washington State (38). It is likely, therefore, that the concentration of the antibiotic to which isolates of *G. graminis* var. *tritici* are exposed in the field can vary significantly from year to year due to changes in varieties. In addition, we think that the supportiveness of a variety to 2,4-DAPG producers and antibiotic accumulation is an important contributor to the robustness of the suppressiveness observed in a TAD field.

Finally, the mechanism of action of 2,4-DAPG may hamper emergence of less sensitive isolates in the field. Although the exact mechanism of action of 2,4-DAPG is not known, the antibiotic has antiviral, antibacterial, antifungal, antihelminthic and phytotoxic properties (38). Thus, given the breadth of its activity...
against such a wide range of organisms, it is not likely that 2,4-DAPG has a single site of action. This would reduce the probability of mutants with less sensitivity emerging in the G. graminis var. tritici population.

Identification of G. graminis varieties on the basis of pathogenicity tests and morphological characteristics can be time consuming and sometimes inconclusive because of overlap in characteristics with G. graminis var. avenae (attacks oat and turf grasses), var. graminis (attacks Bermuda grass, rice and other grasses) and Phialophora (19,22). G. graminis var. graminis and var. tritici have similar ascospore length, ranging from 70 to 105 µm, whereas G. graminis var. avenae has slightly longer ascospores (100 to 130 µm). G. graminis var. avenae and tritici produce simple hynopodia on infected plant tissues, whereas var. graminis produces lobed hypopodia (19,22). For example, Yeates et al. (40) identified G. graminis var. tritici isolates based on ascospore length, yet they attacked oat.

Several molecular approaches have been developed to facilitate identification of varieties of G. graminis: i.e., restriction fragment length polymorphism using a mitochondrial DNA probe (3,20, 21); restriction patterns of PCR products using internal transcribed spacer (ITS) region and 18S rDNA (17,37); random amplified polymorphic DNA (6,18); and specific PCR assays based on nuclear rDNA (5,16) or avenacin and avenacin-like gene sequences (34). However, some of these molecular methods may not be reliable across a broad collection of isolates (19). We conducted PCR on DNA of our isolates with NS5 and GGT-RP primers, which are reported to amplify 185 ribosomal DNA of G. graminis var. tritici as a single 410-bp fragment (16). Our isolates, which caused typical take-all symptoms and had hyphal characteristics, growth on R-PDA, and ascospores typical of G. graminis var. tritici, generated a single 410-bp PCR product. Thus, we are confident that the isolates selected for our studies were typical G. graminis var. tritici.

Rachdawong et al. (34) reported sequence variation in the avenacinase-like gene in G. graminis varieties that could distinguish vars. avenae, graminis, and tritici. They developed three variety-specific forward primers and one common reverse primer (AV3). Varieties were identified based on different PCR ampli- cons sizes: 617 bp for G. graminis var. avenae, 870 bp for var. tritici, and 1,086 bp for var. graminis. We tested these primers early in our study but unexpectedly found that DNA from our new G. graminis var. tritici isolates from Washington and six control isolates generated PCR amplicons of approximately 1,100 bp in size with a mix of the three variety-specific primers. Alignment of the sequences of the avenacinase-like genes of different isolates showed variation, especially in the region of the binding site of the G. graminis variety-specific primers (Fig. 2). Washington isolates have nucleotide variation in the avenacinase-like gene as compared to isolates tested by Rachdawong et al. (34). These sequence variations resulted in the 1,100-bp PCR amplicons from the Washington G. graminis var. tritici isolates. However, Washington isolates had the same sequence as G. graminis var. tritici type strain ATCC82820 (Fig. 2). In general, our results highlight the need to test a broad range of G. graminis var. tritici isolates from multiple locations when developing specific primers for G. graminis. We suggest that at least two different molecular methods be employed to identify a G. graminis variety.

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LITERATURE CITED


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