Abstract

Sclerotinia homoeocarpa isolates were collected from golf courses in Japan and the United States (2016–2017). Japanese isolates were collected during a monitoring study and the U.S. isolates were collected due to field failure. Five succinate dehydrogenase inhibitor (SDHI) active ingredients (boscalid, fluopyram, fluxapyroxad, isofetamid, and penthiopyrad) were examined using in vitro sensitivity assays to determine cross-resistance. Sequence analysis revealed a point mutation leading to an amino acid substitution (H267Y) and a silent mutation (CTT to CTC) at codon 181 in the SdhB subunit gene. Isolates with the B–H267Y (n = 10) mutation were resistant to boscalid and penthiopyrad and had increased sensitivity to fluopyram. SdhB silent mutation 181C>T isolates (n = 2) were resistant to boscalid, isofetamid, and penthiopyrad. Sequence analysis revealed 3 mutations leading to an amino acid substitution (G91R, n = 5; G150R, n = 1; G159W, n = 1) in the SdhC subunit gene. Isolates harboring the SdhC (G91R or G150R) mutations were resistant to boscalid, fluxapyroxad, isofetamid, and penthiopyrad.

Resistance of Sclerotinia homoeocarpa Field Isolates to Succinate Dehydrogenase Inhibitor Fungicides

James T. Popko, Jr., Stockbridge School of Agriculture, University of Massachusetts, Amherst, MA 01003; Hyunkyu Sang, Department of Plant, Soil, and Microbial Sciences, Michigan State University, East Lansing, MI 48824; Jaemin Lee, Stockbridge School of Agriculture, University of Massachusetts, Amherst, MA 01003; Toshikiko Yamada and Yoichiro Hoshino, Field Science Center for Northern Biosphere, Hokkaido University, Sapporo, Hokkaido 0606-0808, Japan; Geunhwa Jung,† Stockbridge School of Agriculture, University of Massachusetts, Amherst, MA 01003; and Field Science Center for Northern Biosphere, Hokkaido University, Sapporo, Hokkaido 0606-0808, Japan

Dollar spot, caused by Sclerotinia homoeocarpa (F.T. Bennett), is one of the most common and economically important diseases of cool-season turfgrass species worldwide (Smiley et al. 2005). Cultural practices to manage dollar spot are often ineffective. Therefore, multiple fungicide applications are often needed throughout the growing season to manage dollar spot and maintain high turf quality. However, repeated use of fungicides has led to the development of resistance in S. homoeocarpa populations (Smiley et al. 2005). Resistance to the benimidazole (FRAC #1) and dicycarboximide (FRAC #2) fungicide classes and reduced sensitivity to the sterol demethylation inhibitor (DMI, FRAC #3) class have been reported in S. homoeocarpa field populations across North America (Cole et al. 1968; Detweiler et al. 1983; FRAC 2017b; Golembiewski et al. 1995). The succinate dehydrogenase inhibitor (SDHI, FRAC #7) fungicides have become an important penetrant class for dollar spot control (FRAC 2017b; Vincelli et al. 2017). Currently, five active ingredients (boscalid, fluopyram, fluxapyroxad, isofetamid, and penthiopyrad) are registered for use on turfgrass, and more are in the registration process. Reports of SDHI resistance in other frequently treated plant-pathogenic fungal species have led to heightened alertness for SDHI resistance in S. homoeocarpa due to the heavy dependence on SDHI fungicides for dollar spot control in turfgrass (Sierotzki and Scalliet 2013).

The SDHI fungicide class has expanded rapidly with the discovery of new molecules with broad spectrum fungal control that are very useful to the crop protection industry (Sierotzki and Scalliet 2013). Currently, the SDHI class contains 20 different molecules in 10 different chemical groups (FRAC 2017a). SDHI fungicides have a common site-specific mode of action that blocks the succinate dehydrogenase complex composed of four subunits (SdhA, SdhB, SdhC, and SdhD) and inhibits mitochondrial electron transport (Matsson and Hederstedt 2001). Succinate dehydrogenase catalyzes the oxidation of succinate to fumarate and the reduction of ubiquinone to quinone, and inhibition leads to decreased energy production and arrested fungal growth (Matsson and Hederstedt 2001). The SDHI class carries a moderate to high resistance risk, which has been reported in 16 different fungal pathogens (FRAC 2017b). The genetic mechanisms responsible for SDHI resistance have been well-studied in both lab and field mutants. Mutations causing resistance in plant pathogenic fungi have been reported in 10 different positions in the SdhB, SdhC, and SdhD genes (FRAC 2017b; Sierotzki and Scalliet 2013). In addition, differential effects of gene mutations on resistance have been reported in Aspergillus oryzae, Botrytis cinerea, Corynespora cassicola, and Podosphaera xanthii (Ishii et al. 2011; Shima et al. 2011; Veloukas et al. 2013). In S. homoeocarpa, ancestral resistance to SDHIs has been reported since 2004; however, resistance has not been corroborated with in vitro fungicide sensitivity assays or molecular confirmation (Anthony and Kerns 2017; Gilstrap and Vargas 2005).

In this study, S. homoeocarpa isolates were initially collected from 12 golf courses in a fungicide resistance monitoring study conducted in Japan. One golf course contained two isolates with reduced boscalid sensitivity collected from a putting green in 2016, and nine additional S. homoeocarpa isolates were obtained from a different green at the same Japanese golf course in 2017. While these experiments were being conducted, a golf course in Rhode Island (RI), USA reported (boscalid) field failure, and nine S. homoeocarpa isolates were collected from a fairway in 2017. In vitro sensitivity and molecular experiments were initiated to compare isolates from the Japanese monitoring study and the RI golf course along with a diverse group of SDHI-sensitive reference isolates. Specific objectives were: 1) to quantify in vitro sensitivities of S. homoeocarpa field isolates to boscalid, fluopyram, fluxapyroxad, isofetamid, and penthiopyrad, 2) to sequence the SdhB, SdhC, and SdhD genes of field isolates for potential point mutations associated with SDHI resistance, 3) to understand cross-resistance among five active ingredients mentioned above according to mutation patterns, and 4) to validate the function of point mutations.
mutations in $SdhB$ and $SdhC$ genes using a genetic transformation system.

Materials and Methods

Origin of fungal isolates. Twenty-five $S. homoeocarpa$ field isolates were used in this study. Eleven isolates (5.1, 5.2, 5.5, J-5, J-10, J-12, J-13, J-15, J-16, J-17, and J-19) were collected from two putting greens at the Takehara Golf Course (Hiroshima, Japan), which were suspected to be SDHI resistant based on the results of the 2016 in vitro sensitivity monitoring study. Eight isolates (M-1, M-2, M-5, M-7, M-8, M-9, M-10, and M-11) were collected from a golf course fairway in Rhode Island that reported SDHI field efficacy failure and were suspected to be SDHI resistant. Three isolates, 3.1, 6.1, and 6.3, were collected from 2 golf courses in Japan and included because they were sensitive to SDHI fungicides in the 2016 Japanese monitoring study. Isolate JTS30 was sampled from the UMass Turf Research Center (South Deerfield, MA) in an area that had not been exposed to fungicides, and isolate HRS10 was sampled from a golf course in Amherst, MA that had been treated with fungicides, but showed a similar level of fungicide sensitivity to JTS30 (Sang et al. 2015). Isolates JTS30 and HRS10 are sensitive to the DMI, dicarboximide, and SDHI fungicide classes. Isolate CT45 collected from Wethersfield, CT is a dicarboximide-resistant isolate harboring a mutation in the histidine kinase $Shos1$ gene and is also resistant to the DMI class (Sang et al. 2017). CT45 represented an isolate that may be found in a golf course setting regularly exposed to fungicides.

SDHI in vitro fungicide sensitivity assays. In vitro sensitivity of 25 $S. homoeocarpa$ isolates to 5 SDHI active ingredients was assayed. 1,000 μg mL$^{-1}$ discriminatory concentrations of the commercial grade fungicides boscalid (Emerald 70WG, BASF), fluopyram (Fluopyram 50SC, Bayer Crop Science), fluxapyroxad (Xemexlar 2.51SC, BASF), isofetamid (Kabuto 3.38C, PBI Gordon), and penthiophen (Velista 50WDG, Syngenta) were used for measuring percentage (%) of relative mycelium growth (RMG) (Jo et al. 2006). Discriminatory concentrations were selected based on preliminary screening results that showed consistent growth differences among isolates. Discriminatory concentrations were previously used to effectively determine resistance profiles across boscalid, fluopyram, fluxapyroxad, and penthiophen in Botrytis cinerea (Hu et al. 2016). All isolates were transferred from slant tubes in −4°C and grown on PDA for 3 days at 22°C. After 3 days of incubation, two perpendicular colony diameters were measured using 16EX digital calipers (Mahr). Each SDHI amended PDA colony diameter was divided by a nonamended PDA colony diameter and multiplied by 100 to calculate RMG%. Two colony diameters were measured on each petri plate, and two petri plates were used in two repeated experiments (8 colony diameter measurements total). The repeated experiments were not significantly different, and all colony measurements were combined for statistical analysis.

For statistical analysis, $S. homoeocarpa$ isolates were grouped into three categories based on target gene mutation patterns ($SdhB$ H267Y, $SdhC$ G91R, and no mutation on either of the genes) with a minimum of 5 isolates per group. Analysis of variance was conducted on target gene mutation groups, and Fisher’s protected least significant difference (LSD) was conducted to compare RMG% values among mutation groups for each fungicide using the JMP software package, version 10.0 (SAS Institute Inc.).

DNA extraction, polymerase chain reaction (PCR) amplification, and sequencing. DNA of 25 isolates was extracted using a CTAB method (Huvely et al. 2012). Three PCR primer sets, F$_{SdhB}$/R$_{SdhB}$, F$_{SdhC}$/R$_{SdhC}$, and F$_{SdhD}$/R$_{SdhD}$, were used to amplify the $SdhB$, $SdhC$, and $SdhD$ genes identified from a draft genome of $S. homoeocarpa$ (Green et al. 2016). The PCR products of $SdhB$, $SdhC$, and $SdhD$ were amplified using Phusion $taq$ polymerase (New England Biolabs) by touch-down PCR programs with different annealing temperatures at 72, 68, and 67°C, respectively, decreasing at increments of 1.0°C per cycle for the first eight cycles and 64, 60, and 59°C of the remaining 45 cycles. A total 20 μL of volumes includes 1x Phusion HF Buffer, 0.2 dNTPs, 0.4 μM primers, 1 U Phusion $taq$ polymerase, 3% DMSO, and <250 ng of gDNA. Amplicons were purified using PureLink PCR Micro kit (Invitrogen). The purified amplicons were sequenced by Macrogen Corporation (Boston, MA).

Plasmid construction and generation of $SdhB$ or $SdhC$ mutants from a SDHI-sensitive isolate. The 1,500-bp upstream region and full length of $SdhB$ (941 bp) was amplified from gDNA of isolate 5.1 harboring the B-H267Y mutation using primer F$_{Apal}$/up1500SdhB and R$_{SacI}$/SdhB. The 1,500-bp upstream region and full length of $SdhC$ (759 bp) was also amplified from gRNA of isolate M1 harboring C-G91R mutation using primer F$_{Apal}$/up1500SdhC and R$_{NotI}$/SdhC. Each amplicon purified by Zymo Gel DNA recovery kit (Zymo research, Irvine, CA, USA) and plasmid pYHN3-MCS (Sang et al. 2017) was digested with Apal/SacI for $SdhB$ gene or Apal/NotI for $SdhC$ gene. Two digested products were purified and ligated to generate the plasmid pYHN3-SdhB(H267Y) or pYHN3-SdhC(G91R). The plasmid DNA (5 μg) was used for a polyethylene glycol (PEG)-mediated transformation in the protoplasts from isolate HRS10. The protoplast generation and PEG-mediated transformation were conducted according to Sang et al. (2017). After 7 days, hygromycin resistant transformants were grown on 100 μg mL$^{-1}$ of hygromycin amended regeneration agar medium (239.6 g of sucrose, 0.5 g of yeast extract, 15 g of agar per liter). Two $S. homoeocarpa$ mutants HRS10(SdhB(H267Y)) and HRS10(SdhC(G91R)) and one control isolate HRS10 were used for in vitro sensitivity assays to five SDHI fungicides (boscalid, fluxapyroxad, isofetamid, penthiophen, and fluopyram). The diameter of fungal colonies from three individual plates was measured after the isolate, and mutants grew for 144 h on PDA amended with 1,000 μg mL$^{-1}$ of boscalid, fluxapyroxad,

---

### Table 1. Primers used in this study

<table>
<thead>
<tr>
<th>Primers name</th>
<th>Primer sequence (5’-3’)</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>F$_{SdhB}$</td>
<td>CGGCCCTCTGGTTGCGCAATTGA</td>
<td>Amplification and sequencing primers for SdhB gene (941 bp)</td>
</tr>
<tr>
<td>R$_{SdhB}$</td>
<td>TACACAAGGCGCATCCGCA</td>
<td></td>
</tr>
<tr>
<td>F$_{SdhC}$</td>
<td>TCTCCGATCAAGACGATA</td>
<td>Amplification and sequencing primers for SdhC gene (759 bp)</td>
</tr>
<tr>
<td>R$_{SdhC}$</td>
<td>TCTGTCTGGAAGCCTCAT</td>
<td></td>
</tr>
<tr>
<td>F$_{SdhD}$</td>
<td>TGAGGATCAGGGAGCTAC</td>
<td>Amplification and sequencing primers for SdhD gene (683 bp)</td>
</tr>
<tr>
<td>R$_{SdhD}$</td>
<td>CTGCTCACATAATCTCGTTTC</td>
<td></td>
</tr>
<tr>
<td>F$_{Apal}$/up1500SdhB</td>
<td>ATGGGCAGCCCAAAACACTTATTCGCTTC</td>
<td>Amplification of upstream region and SdhB gene (2,441 bp)</td>
</tr>
<tr>
<td>R$_{SacI}$/SdhB</td>
<td>ATGGGACGGCTTAAAAGGCACTCTCTCTCTGATCC</td>
<td></td>
</tr>
<tr>
<td>F$_{Apal}$/up1500SdhC</td>
<td>ATGGATGAGGGCCTCCGAAAAGAGTGAC</td>
<td>Amplification of upstream region and SdhC gene (2,259 bp)</td>
</tr>
<tr>
<td>R$_{NotI}$/SdhC</td>
<td>ATTTGGGGCCCGCTAGAAGACCGCAACCAAG</td>
<td></td>
</tr>
</tbody>
</table>
or isofetamid, and for 72 h on PDA amended with 1,000 μg ml⁻¹ of penthiopyrad or fluopyram. The diameter of fungal colonies on PDA (control) plates was measured after 48 h. Fisher’s protected least significant difference (LSD) was conducted to compare diameter (mm) among three strains to each fungicide and control. The 1,500 bp of upstream region and full length of SdhB, SdhC, and SdhD from HRS10 were deposited in GenBank with accession numbers MG708132, MG708133, and MG708134, respectively. All primers used in this study are listed in Table 1.

Prediction of protein structure and ligand binding sites of SdhB and SdhC. I-TASSER (Iterative Threading ASSEMBly Refinement) (https://zhanglab.ccmb.med.umich.edu) was used to predict protein structure and ligand binding sites SdhB and SdhC in S. homoeocarpa (Roy et al. 2012; Yang and Zhang 2015; Zhang 2009). The full length of amino acid sequences of SdhB and SdhC from draft genome sequences of isolate HRS10 (Green et al. 2016) were used for the prediction. PyMOL was used to generate the protein structure.

Results

Sequence analysis of SdhB, SdhC, and SdhD genes. To find the genetic determinant conferring SDHI resistance in S. homoeocarpa, the SDHI target genes (SdhB and SdhC) from all 25 isolates were sequenced. Since no sequence differences in the SdhD gene were detected in 8 representative isolates, the rest of the isolates were not sequenced. Four single nucleotide changes that resulted in amino acid substitutions were detected (Figs. 1 and 2), and one nucleotide that caused a silent mutation was also detected. Ten Japanese isolates contained a mutation in SdhB (CAC to TAC) at codon 267 that resulted in substitution of histidine by tyrosine (H267Y) (Fig. 1). Five Rhode Island isolates contained a mutation in SdhC (GGA to CGA) at codon 91 that resulted in substitution of glycine by arginine (G91R) (Fig. 2). One Japanese isolate (J-19) contained a change in SdhC (GGG to TGG) at codon 159 that resulted in substitution of histidine by tyrosine (G159W) (Fig. 2). Two Japanese isolates (M-2 and M-8) more closely resembled the C-G91R group for all active ingredients except for isolates J-19, M-2, M-8, and M-10. These isolates were excluded due to the small group size.

The C-G91R mutation group expressed significantly higher mean RMG% values on all SDHI active ingredients than the B-H267Y mutation group and sensitive reference isolate/no mutation group (Table 2). The B-H267Y mutation group also expressed higher RMG% values than the sensitive reference isolate/no mutation group on all SDHI active ingredients except fluopyram (Table 2). Increased sensitivity to fluopyram was observed from all isolates in the B-H267Y mutation group only (Table 2). Isolate J-19 displayed a similar cross-resistance profile to isolates in the C-G91R mutation group, and isolate M-10 with both a silent mutation 181C>T and SdhC (G91R) (Fig. 2). One Japanese isolate (J-19) contained a change in SdhC (G91R) (Fig. 2). One Japanese isolate (J-19) contained a change in SdhC (G91R) (Fig. 2). One Japanese isolate (J-19) contained a change in SdhC (G91R) (Fig. 2).

Generation of SdhB_H267Y and SdhC_G91R mutants and fungicide sensitivity assay. A plasmid with the 1,500-bp of upstream and full length of SdhB gene containing H267Y or SdhC gene containing G91R was transformed into the SDHI sensitive HRS10 to test if the H267Y mutation in SdhB or G91R mutation in SdhC confers SDHI resistance in S. homoeocarpa. The mutants HRS10(SdhB_H267Y) exhibited resistance to boscalid and penthiopyrad but not to fluopyram, isofetamid, and fluopyram. The mutant HRS10(SdhC_G91R) displayed resistance to boscalid, fluxapyroxad, penthiopyrad, and isofetamid but not to fluopyram. Furthermore, the

Table 2. The percentage (%) of relative mycelium growth of 25 field Sclerotinia homoeocarpa isolates from Japan and the United States to five SDHI fungicides

<table>
<thead>
<tr>
<th>Isolate/isolate group</th>
<th>Target gene mutation</th>
<th>Origin</th>
<th>Mean RMG (%)w ± standard error</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Boscalid</td>
</tr>
<tr>
<td>3.1, 6.1, 6.3, CT45, HRS10, JTS30</td>
<td>No mutation</td>
<td>Japan, CT, USA, MA, USA</td>
<td>33.4± ± 1.5c</td>
</tr>
<tr>
<td>5.1, 5.2, J-5, J-7, J-10, J-12, J-13, J-15, J-16, J-17</td>
<td>SdhB H267Y</td>
<td>Japan</td>
<td>66.0± ± 1.7b</td>
</tr>
<tr>
<td>M-1, M-5, M-7, M-9, M-11</td>
<td>SdhC G91R</td>
<td>RI, USA</td>
<td>77.9± ± 2.2a</td>
</tr>
</tbody>
</table>

P value

<table>
<thead>
<tr>
<th>Additional isolates</th>
<th>Mean RMG (%)w ± standard error</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Boscalid</td>
</tr>
<tr>
<td>M-2</td>
<td>SdhB 181 silent</td>
</tr>
<tr>
<td>M-8</td>
<td>SdhB 181 silent</td>
</tr>
<tr>
<td>M-10</td>
<td>SdhB 181 silent + SdhC G159W</td>
</tr>
<tr>
<td>J-19</td>
<td>SdhC G159R</td>
</tr>
</tbody>
</table>

* Relative mycelial growth (RMG) (%) was calculated by dividing the mean colony diameter of each isolate on PDA amended with each SDHI fungicide by the mean colony diameter of each isolate grown on nonamended PDA.

† Mean RMG% is composed of the isolates in the group.

‡ Means followed by the same letter within each column are not significantly different according to Fisher’s protected least significant difference test (α = 0.05).

§ *** = significance at P < 0.001.
mutant HRS10(SdhCG91R) showed higher levels of resistance to boscalid and penthiopyrad than the mutant HRS10(SdhBH267Y) (Fig. 3).

Prediction of ligand binding sites in SdhB and SdhC. The protein structure and ligand binding sites of SdhB and SdhC in S. homoeocarpa were predicted. In the SdhB subunit, predicted ligand binding sites were R220, S221, W224, H267, and I269. The mutated binding site (H267) was found in isolates 5.1, 5.2, J-5, J-7, J-10, J-12, J-13, J-15, J-16, and J-17. This protein was described as 3vraF in PDB (Protein Data Bank). In SdhC subunit, predicted sites were

---

![Fig. 3. Sensitivity of Sclerotinia homoeocarpa isolates HRS10 and mutants HRS10(SdhBH267Y) and HRS10(SdhCG91R) to five SDHI active ingredients (boscalid, penthiopyrad, fluxapyroxad, isofetamid, and fluopyram). (A) The pictures were taken after the isolate and mutants grew for 144 h on PDA amended with 1,000 µg ml⁻¹ of boscalid, fluxapyroxad, and isofetamid, and for 72 h on PDA amended with 1,000 µg ml⁻¹ of penthiopyrad and fluopyram. The picture of control plate without treatments was taken after 48 h. (B) Diameter (mm) of fungal colonies on PDA without (control) and amended with SDHI fungicides. Means followed by the same letter are not significantly different according to Fisher’s protected least significant difference test (α = 0.05).](image_url)
N87, R88, G91, L94, S95, H146, G147, G150, and H153. The binding site G91 mutated was found in isolates M-1, M-5, M-7, M-9, and M-11, and the binding site G150 was mutated in isolate J-19. This protein was described as 3ae3C in PDB (Fig. 4).

**Discussion**

The current study reports the occurrence of *S. homoeocarpa* field isolates exhibiting differential resistance to SDHI fungicides for the first time and elucidates the molecular mechanisms of SDHI resistance in *S. homoeocarpa*. We observed five different mutations in the *SdhB* (H267Y and a silent mutation 181C>T) and *SdhC* (G91R, G150R, and G159W) subunit genes in *S. homoeocarpa* isolates from two golf courses. The B-H267Y mutation was determined to be a direct factor in resistance to boscalid and penthiopyrad, and the C-G91R was confirmed to be a direct factor conferring resistance to boscalid, fluxapyroxad, isofetamid, and penthiopyrad through the genetic transformation system. In addition, isolate J-19 displayed resistance to boscalid, fluxapyroxad, isofetamid, and penthiopyrad, which contained the C-G150R mutation. Isolate M-10 contained double mutations (C-G159W and a silent mutation 181C>T in *SdhB*), and displayed moderate resistance to boscalid, and weak resistance to fluxapyroxad and isofetamid. Lastly, two isolates (M-2, M-8) with the silent mutation *SdhB* 181C>T displayed resistance to boscalid, fluxapyroxad, isofetamid, and penthiopyrad. The presence of multiple target gene mutations causing differential cross-resistance in *S. homoeocarpa* field isolates from two separate continents is cause for concern.

In total, 16 different plant-pathogenic fungal species have reported to be resistant to SDHI fungicides due to mutations in three of four SDHI subunit genes (FRAC 2017b). Reports of SDHI resistance in other frequently treated plant pathogens have led to heightened alertness for SDHI resistance in *S. homoeocarpa* due to the over-reliance on fungicides for dollar spot control in turfgrass (Sierotzki and Scalliet 2013). Location and climatic conditions greatly influences disease occurrence, and 3 to 10 applications per year are commonly made to control dollar spot. Fungicide rotations/mixtures are an essential component of successful resistance management strategies because resistance to the benzimidazole and DMI classes are commonly found along with occasional resistance to the dicarboximide class (Putman et al. 2010; Sang et al. 2017). The current status of fungicides that effectively control dollar spot is fairly limited, and therefore delaying the onset of resistance to the SDHI class is critical. Differences in the SDHI cross-resistance profile of the isolates examined in this study suggest that fluopyram may be effective on populations that harbor the target gene mutations presented in this work; however, many other target gene mutations exist in other plant pathogens and may develop in *S. homoeocarpa* populations later (FRAC 2015; Sierotzki and Scalliet 2013). Several studies have reported high percentages of boscalid resistance in the absence of fluopyram resistance (Avenot et al. 2012; De Miccolis Angelini et al. 2014; Hu et al. 2016; Ishii et al. 2011; Veloukas et al. 2013).

SDHI field resistant isolates harboring the B-H267Y mutation and *SdhB*H267Y mutants showed resistance to boscalid and penthiopyrad, increased sensitivity to fluopyram, and no resistance to fluxapyroxad and isofetamid (Table 2; Fig. 3). The histidine at codon 267 is a highly conserved residue, which is associated with the (3Fe-4 S) high-potential nonheme iron sulfur-redox (S3) center (Skinner et al. 1998). Horsefield et al. (2006) suggested the conserved histidine residue of *SdhB* might strongly interact with the histidine and serine residue at codon 27 of *SdhC*, blocking ubiquinone from accessing the quinone-binding site (Q-site) based on the structural and computational analysis of Q-site from *Escherichia coli* using carboxin. Therefore, the histidine residue was suggested to be essential for carboximide binding (Avenot et al. 2008), and the mutation in this residue might affect direct fungicide binding. The mutation from the conserved histidine residue to tyrosine in *SdhB* gene was also reported in SDHI field resistant isolates of other plant-pathogenic fungi, such as *Alternaria alternata*, *Botrytis cinerea*, *Corynespora cassicola*, and *Pyrenophora teres* (Avenot et al. 2008; Miyamoto et al. 2010; Stammel et al. 2014; Veloukas et al. 2013; Yin et al. 2011).

The increased sensitivity response to fluopyram in isolates containing the *SdhB* H267Y mutation may be due to fluopyram possessing a benzamide ring that is better equipped to bind to the Q-pocket of resistant isolates containing the mutation (histidine to tyrosine) (Veloukas et al. 2013). Other fungal species with the same substitution also displayed increased sensitivity to benzamide fungicides (Avenot et al. 2012; Ishii et al. 2011; Scalliet et al. 2012; Shima et al. 2011; Veloukas et al. 2013). However, increased sensitivity to fluopyram was not confirmed by the *SdhB*H267Y mutants of *S. homoeocarpa* due to possession of both wild and mutated *SdhB* genes in the mutant cells. In the presence of fluopyram, the wild *SdhB* gene might be dominant in the fungal cells, and this may explain why mutants had the same growth rate as the wild-type isolate. Results from the genetic transformation assay suggest that despite statistical differences from the in vitro fungicide assay, mean RMG% differences of ~10% on fluxapyroxad and isofetamid amended media did not correlate to a true resistance response (Table 2; Fig. 3).

SDHI field resistant isolates containing the C-G91R mutation and *SdhCG91R* mutants modified from the sensitive isolate showed high resistance to boscalid, fluxapyroxad, and penthiopyrad, moderate resistance to isofetamid, and no resistance to fluopyram (Table 2). According to the FRAC SDHI working group meeting in 2016, the C-G79R mutation in *P. teres* confers moderate resistance in the field, and this mutation was first detected in 2013 in Europe. This position is homologous to the mutation present in the five resistant isolates from Rhode Island (M-1, M-5, M-7, M-9, and M-11) and represents the first report of a mutation in this position in North America. In *S. homoeocarpa*, the C-G91R mutation exhibited the highest resistance compared with the SDHI-sensitive isolate group and the widest
spectrum of resistance across active ingredients (boscalid, fluxapyroxad, isofetamid, and pentaipyrazophenyl) in the class (Table 2). Furthermore, the active ingredients most-severely affected are pyrazole-4-carboximide (fluxapyroxad and pentaipyrazophenyl) or pyridine-carboximide (boscalid) fungicides. Lastly, HRS10(SdhCG91R) mutants grew more on isofetamid-amended media than the HRS10(SdhH267Y) mutants and HRS10 and suggest the 15% increase in RMG% observed from the in vitro fungicide assay correlates to a true resistance response.

Isolate J-19 was the only isolate determined to have the C-G150R mutation and resulted in a very similar cross-resistance profile to the C-G91R mutation (Table 2). There are no previous reports of a mutation in this position of the ShdhC gene in any other fungal species, and this appears to be the first report. The predicted binding site simulation confirms that a mutation at this position is likely to alter SDHI fungicide binding affinity. Isolate M-10 was the only isolate determined to have the C-G159W mutation and showed high resistance to boscalid and increased fluopyram sensitivity as the B-H267Y mutation isolates (Table 2). Moreover, isolate M-10 exhibited weak or no resistance to fluxapyroxad, isofetamid, or pentaipyrazophenyl (Table 2).

In addition, isolates M-2, M-8, and M-10 were discovered to have a silent mutation in ShdhB (CTT to CTC) at codon 181 that did not result in an amino acid substitution. Isolates M-2 and M-8 both showed resistance to boscalid, isofetamid, and pentaipyrazophenyl, but no resistance to fluopyram. Silent mutation has not been reported to cause SDHI resistance in any other fungal pathogens, and this is the first report. Kimchi-Sarfaty et al. (2007) suggested that a silent mutation in nucleotide triplets can result in a different three-dimensional structure of the protein due to speed-dependent folding of an amino acid chain, influenced by slow processing of ribosomes. Also, Yamashita and Fraaije (2018) observed Zymoseptoria tritici isolates collected from the field that were resistant to isofetamid and fluopyram, but did not contain target gene mutations within the ShdhB, C, or D subunits. The function of the silent mutation in S. homoeocarpa should be validated using a genetic transformation system.

Sang et al. (2015) described that multidrug resistant isolates of S. homoeocarpa exhibited reduced sensitivity to propiconazole (DMI), iprodione (dicarboximide), and boscalid. These isolates harbor overexpression of an ABC efflux transporter ShPDR1 which is involved in decreased sensitivity to these different fungicide classes and was confirmed by a heterologous expression system in yeast (Sang et al. 2015). The field study of Allen-Perkins et al. (2017) also indicated that applications of boscalid selected DMI-resistant S. homoeocarpa isolates that overexpressed ShPDR1 and supported existing evidence that ABC-transporters are involved in resistance to chemically unrelated fungicides. Future research in field conditions is needed to determine the impact of qualitative (target gene mutations) and quantitative (gene overexpression) resistance on S. homoeocarpa population sensitivity in relation to SDHI fungicides.

Detection of differential resistance to SDHIs signals that turfgrass industry must implement sound fungicide resistance management strategies to prolong the longevity of this important fungicide class in S. homoeocarpa. S. homoeocarpa field isolates have already developed resistance to four different classes of single-site fungicides, multiple fungicide resistance, and multidrug resistance conferred by xenobiotic detoxification (Jo et al. 2006; Putman et al. 2010; Sang et al. 2017). Future research should focus on developing an integrated pest management approach by implementing effective cultural practices, weather-based disease forecasting, and initiating a long-term monitoring of resistant populations using molecular diagnostic tools. Furthermore, it is essential for academic and industry scientists to develop a better understanding of the mechanisms of differential resistance to SDHIs and management of field populations with multiple fungicide resistance. Lastly, recommendations for dollar spot control must emphasize rotation of different fungicide classes with single-site modes of action. Furthermore, multisite fungicides (chlorothalonil [FRAC 8MS] and fluazinam [FRAC #29]) should be used as tank-mix options with single-site mode of action fungicides under periods of intense disease pressure. Hopefully, the sum of these practices can delay fungicide resistance to the SDHI fungicide class.

Acknowledgments

We thank golf course superintendents in Japan for sending dollar spot samples. The contents are solely the responsibility of the authors and do not necessarily represent the official views of the USDA or NIFA.

Literature Cited


