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June 2004

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Mazzola, M.; Funnell, Deanna L.; and Raaijmakers, J.M., "Wheat Cultivar-Specific Selection of 2,4-Diacetylphloroglucinol-Producing Fluorescent *Pseudomonas* Species from Resident Soil Populations" (2004). *Papers in Plant Pathology*. 37.

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Wheat Cultivar-Specific Selection of 2,4-Diacetylphloroglucinol-Producing Fluorescent *Pseudomonas* Species from Resident Soil Populations

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Received: 6 October 2003 / Accepted: 22 December 2003 / Online publication: 24 August 2004

Abstract

An emerging body of evidence indicates a role for plant genotype as a determinant of the species and genetic composition of the saprophytic microbial community resident to the rhizosphere. In this study, experiments were conducted to determine the capacity of five different wheat cultivars to enhance resident populations and support introduced strains of 2,4-diacetylphloroglucinol (2,4-DAPG)-producing fluorescent pseudomonads, a group of bacteria known to provide biological control of several soilborne diseases. When soils were cropped with three successive 28-day growth cycles of wheat, the 2,4-DAPG-producing strains were consistently recovered from the rhizosphere of the cultivar Lewjain, and commonly were present at populations higher than those recovered from other wheat cultivars. Based on restriction fragment length polymorphism and sequence analyses of *phlD*, a key gene involved in 2,4-DAPG production, two previously undefined *phlD*+ genotypes, referred to as genotypes PfZ and PfY, were discovered. Wheat cultivar Lewjain was the primary source of genotype PfY while cultivar Penawawa yielded the majority of genotype PfZ. Based on 16S rDNA sequence analysis, both new *phlD* genotypes were classified as *P. fluorescens*. Comparison of the rhizosphere competence of 2,4-DAPG-producing *P. fluorescens* Q2-87 (genotype B) and *P. fluorescens* LR3-A28 (genotype PfY) showed that both strains persisted at similar populations in the rhizosphere of all cultivars tested over a 30 day period

when introduced as a seed inoculant. However, when strain LR3-A28 was applied as a soil inoculant, this strain was recovered at higher populations from the rhizosphere of wheat cultivar Lewjain than from the rhizospheres of two other cultivars. No cultivar effects were shown for strain Q2-87. Collectively, these results add further to evidence indicating a degree of specificity in interactions between plant cultivars and specific members of the saprophytic microbial community. Furthermore, as 2,4-DAPG-producing fluorescent *Pseudomonas* spp. have a central role in the spontaneous reduction in severity of take-all disease of wheat in response to continuous wheat monoculture, we postulate that the use of specific cultivars, such as Lewjain, which possess a superior capacity to enhance resident soil populations of these bacteria may have potential to reduce the length of the monoculture period required to induce natural suppressiveness of soils toward this disease.

Introduction

The relative contribution of soil type and plant in determining composition of soil microbial communities has been debated. However, considerable evidence suggests that in the specific environment of the rhizosphere the plant plays a significant role in influencing attributes of the bacterial community [5, 16, 17, 24, 31]. Mazzola and Gu [28] demonstrated that composition of the fluorescent pseudomonad population in soil and the rhizosphere of apple was modified through prior cultivation of apple orchard soils with wheat. Cropping of soils with any of five different wheat cultivars enhanced populations of fluorescent pseudomonads recovered from these soils in a similar fashion. However, the genetic composition of this bacterial community recovered from soil and the rhizosphere of apple, and its capacity to

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inhibit *in vitro* growth of the pathogen *Rhizoctonia solani* AG 5, was dependent upon the wheat cultivar cropped prior to planting apple [8, 28].

Numerous studies have demonstrated the capacity of specific nonsymbiotic rhizobacteria to protect roots from infection by plant pathogenic fungi [51, 53]. Among these, fluorescent *Pseudomonas* spp. with the capacity to produce the antibiotic 2,4-diacetylphloroglucinol (2,4-DAPG) have been shown to provide biological control of several important soilborne fungal plant pathogens when introduced as an inoculant [10, 21], and also have a functional role in the natural plant-induced development of soil suppressiveness to certain soilborne diseases (reviewed in Weller et al. [52]). One system that has been explored extensively is the spontaneous decline in the severity of take-all of wheat, incited by *Gaeumannomyces graminis* var. *tritici*, which often is observed after a severe outbreak of the disease for one or two crops early in a wheat monoculture cropping sequence. This phenomenon has been termed take-all decline. Although numerous biological and chemical elements have been put forth as the operative factors responsible for take-all suppressive soils [4, 52], a broad series of studies has provided convincing evidence for the predominant role of 2,4-DAPG-producing fluorescent pseudomonads in take-all decline in certain geographic regions [29, 38, 40, 41, 48].

The existence of disease suppressive soils was first recognized over 100 years ago, and the mechanisms by which pathogen suppression occurs have been the focus of study for nearly four decades. Although much progress has been made toward understanding the biology of constitutive and induced suppressive soils [2, 24, 41, 49], significant difficulty has been encountered in moving this knowledge into field level disease management. As the induction of soil suppressiveness is mediated through transformations in soil microbial communities [19, 28, 40, 52], it is plausible that opportunities exist to manage the phenomenon, and perhaps hasten the onset of the disease-suppressive state.

It was previously shown that, in response to cropping of apple orchard soils with wheat, the species and genetic composition of the fluorescent pseudomonad population will vary in a cultivar-dependent manner [27, 28]. Achieving a threshold population density of 2,4-DAPG-producing fluorescent pseudomonads in wheat-field soils is essential to the development of take-all decline [41]. Thus, identification of wheat genotypes with a superior capacity to select for resident populations of 2,4-DAPG-producing fluorescent pseudomonads could enhance the viability of a management scheme to utilize induced soil suppressiveness as a means to control take-all of wheat.

The objective of this research was to determine whether wheat cultivars differed in their capacity to select, both quantitatively and qualitatively, for 2,4-DAPG-producing fluorescent pseudomonads from resident soil

microbial communities. Population densities of 2,4-DAPG-producing pseudomonads were monitored by colony hybridization followed by PCR. The genotypic composition of the 2,4-DAPG-producing pseudomonad population recovered from different wheat cultivars was characterized by restriction fragment length polymorphism analysis of the *phlD* gene followed by sequence and phylogenetic analyses of newly selected *phlD*⁺ genotypes. Finally, the capacity of two genotypically distinct 2,4-DAPG-producing strains to colonize different wheat cultivars was assessed.

Methods

Soils. Studies were conducted in soils obtained from the Columbia View Research and Demonstration (CV) orchard located 19.2 km north of East Wenatchee, WA; the Wenatchee Valley College Auvil Research and Demonstration (WVC-A) orchard located 6.4 km east of East Wenatchee, WA; and the Wenatchee Valley College Sunny Slope (WVC-SS) orchard located in Wenatchee, WA. Microbial attributes of these soils, including composition of the fluorescent *Pseudomonas* spp. population, have been extensively characterized [22, 23, 27, 28]. Soils were collected in October of 2000, 2001, and 2002 at a depth of 10 to 30 cm. By October 2002, apple had been cropped for 23, 7, and 46 years at the CV, WVC-A, and WVC-SS, respectively. The WVC-A site had been cropped to continuous wheat monoculture prior to orchard establishment in 1995.

Wheat Cultivars and Cropping of Soils. Wheat used in these studies included the cultivars Eltan, Hill-81, Lewjain, Madsen, and Penawawa. All cultivars are soft white winter wheat, with the exception of Penawawa, which is a soft white spring wheat. All seeds were obtained from the Washington State Crop Improvement Association, Foundation Seed Service, Washington State University, Pullman, WA. Seeds were not treated prior to planting.

Soils were dispensed into plastic pots (3.8 L volume, 16 cm diameter) and seeds (10 g) of an individual wheat cultivar were sown in each pot with a total of four pots per cultivar. Seeds were covered with soil to a depth of 1 cm. Pots were arranged in a completely randomized design on a greenhouse bench and plants were grown for 28 days at 18 to 22°C with supplemental lighting to maintain a minimum photoperiod of 12 h. Plants were watered at 3-day intervals and were not fertilized during the course of plant growth. At the conclusion of the 28-day growth cycle, plant shoots were excised at the soil surface and discarded. The contents of an individual pot were decanted and cultivated by hand to disrupt the wheat root system. Soils were dispensed back into the

original pot and were reseeded with the same wheat cultivar. A total of three wheat growth cycles each 28 days in duration were used in all experiments.

Isolation of Fluorescent *Pseudomonas* spp. Fluorescent pseudomonads were recovered from the root systems of wheat harvested after the completion of the third growth cycle. Pots were inverted and wheat plants were separated from the soil. A composite sample consisting of five plants was selected for each pot, and soil loosely adhering to the root systems was detached by shaking. A 0.5-g composite root sample was excised from the five plants and placed in 10 mL sterile distilled water. Roots and associated rhizosphere soil were vortexed for 60 s. Serial dilutions of the resulting root wash were plated on King's medium B (KB) agar amended with ampicillin ($100 \mu\text{g mL}^{-1}$), chloramphenicol ($13 \mu\text{g mL}^{-1}$) and cycloheximide ($75 \mu\text{g mL}^{-1}$) (KB⁺) [45]. Plates were incubated at 28°C for 72 h, at which time fluorescent colonies were counted. For each of the four root wash suspensions, 15 fluorescent pseudomonads were randomly selected and transferred to fresh KB⁺ agar plates. Direct detection of 2,4-DAPG-producing fluorescent pseudomonads in the population recovered from wheat root systems was conducted through extraction and amplification of DNA from individual isolates as described below.

Detection and Quantification of 2,4-DAPG-Producing Fluorescent *Pseudomonas* spp. 2,4-DAPG-producing fluorescent pseudomonads were identified via detection of *phlD*, which is a key gene in the biosynthesis of 2,4-DAPG. For each wheat cultivar, total DNA was extracted from the 60 individual fluorescent pseudomonad isolates using a modification of the method developed by Raaijmakers et al. [40] as previously described [28]. The oligonucleotide primers Phl2a and Phl2b [40] developed from conserved sequences within *phlD* were used in amplification reactions. DNA was amplified in 50- μL volumes containing 1 μL of the heat-lysed cell suspension, 1 \times PCR buffer, 200 μM , of each deoxynucleoside triphosphate, 1.5 mM MgCl_2 , each primer at 0.1 μM , and 2.5 U AmpliTaq Gold (Applied Biosystems). Samples were amplified in a DNA GENE Amp 9700 thermal cycler (PerkinElmer, Norwalk, CT) using conditions previously described [40]. Each PCR included a positive (*Pseudomonas fluorescens* strain Q2-87 [37]) and negative (*phl* *P. fluorescens*) control. Amplified DNA products were verified by electrophoresis of 10- μL aliquots of PCR mixtures through 1.2% agarose gels in 1 \times TBE buffer and visualization following ethidium bromide treatment.

Alternatively, in separate experiments, 2,4-DAPG-producing fluorescent pseudomonads recovered from wheat roots were quantified by colony hybridization

according to the methods developed by Raaijmakers et al. [40]. The *phlD* probe was generated by randomly primed labeling of PCR fragments with the nonradioactive digoxigenin system (Roche Diagnostics, Mannheim, Germany) using DNA from the 2,4-DAPG-producing strain *P. fluorescens* Q2-87 as the template. The probe was detected with anti-digoxigenin-AP-FAB fragments, and *phlD*⁺ colonies were visualized using nitroblue tetrazolium/5-bromo-4chloro-3-indoyl phosphate substrate according to protocols provided by the supplier. Colonies that reacted positively were purified by plating on KB⁺ agar, DNA was isolated using the cell lysis method, and the presence of the *phlD* gene was confirmed by means of the PCR protocol described above.

Genotypic Diversity. Genetic composition of the 2,4-DAPG-producing fluorescent pseudomonad population recovered from wheat cultivars grown in WVC-A soil was characterized by restriction fragment length polymorphism analysis of the *phlD* gene. This gene is highly polymorphic [43]. The primers B2BF and BPR4 [30] were used to amplify DNA from *phlD*⁺ isolates. These primers are highly conserved across isolates possessing the *phlD* gene and possess a lower level of identity than the Phl2A and Phl2B with other known prokaryotic sequences [30]. Amplification reactions were conducted using the conditions previously described [30]. Genotypes were defined by digesting the resulting *phlD* DNA fragments individually with *Hae*III, *Taq*I, or *Hpa*II (Invitrogen Carlsbad, CA) in reactions containing 10 μL of the amplified product. Digestion products were separated on 1.5% agarose gels in 1 \times TBE and banding patterns were visualized by ethidium bromide staining. Isolates were classified based on the genotypes erected by McSpadden Gardener et al. [30].

Sequence Analysis of the *phlD* Gene and 16S rDNA. The *phlD* genes from isolates representing two apparent new genotypes were subjected to sequence analysis. Individual DNA fragments, resulting from amplification reactions which employed the Phl2a and Phl2b primers, were cloned into the vector pCR2.1-TOPO (Invitrogen, Carlsbad, CA) and the resulting plasmid transformed into *E. coli* TOP10. Plasmid DNA was purified from *E. coli* using an Invitrogen S.N.A.P. miniprep kit. Sequencing was conducted on an ABI 377 automated sequencer using the dye terminator cycle sequencing ready reaction kit (PE Biosystems, Warrington, England). Each 20- μL reaction contained 2 μg of template DNA and 10 pmol of the T7 or M13 reverse primer. The *phlD* sequences from genotypes designated PfY and PfZ were deposited in GenBank under accession numbers AY391780 and AY391779, respectively. The primers 8f and 1406r [1] were used to amplify the 16S rRNA gene from these same strains as previously de-

scribed [8], and partial 16S rDNA sequences were obtained using the methods described above.

Phylogenetic Analysis. Alignment of the newly sequenced *phlD* genes with *phlD* homologs present in the database (GenBank) was performed with GeneDoc 2.6 [32] and ClustalW 1.8 [50]. Sequences were trimmed to identical length. Subsequent phylogenetic analysis of the *phlD* genes was performed with MEGA [11] using the neighbor-joining method with the Kimura-2-parameter model. For comparison purposes, similar analyses were performed with the maximum parsimony method. Confidence of the topology of the phylogenetic trees for the *phlD* gene and protein sequences was performed by bootstrap analysis on 1000 resamplings [6].

Colonization of Wheat Roots by Introduced 2,4-DAPG-Producing Fluorescent *Pseudomonas* Strains. The capacity of the 2,4-DAPG-producing strains Q2-87 and LR3-A28 to colonize different wheat cultivars when introduced as a soil or seed inoculant was assessed. *P. fluorescens* strain Q2-87 was originally isolated from wheat grown in a take-all suppressive soil from a field near Quincy, WA, and is a representative of BOX genotype B. Strain LR3-A28 was isolated in this study from roots of the wheat cultivar Lewjain grown in WVC-A orchard soil and, based on comparison with *phlD* genotypes previously described, represents a new *phlD* genotype. Spontaneous rifampicin-resistant mutants were selected by plating wild-type strains on KB⁺ agar amended with rifampicin (100 µg mL⁻¹). Resulting colonies were cultured two additional times on rifampicin-amended KB⁻ agar. Isolates that possessed a stable phenotype and exhibited rapid growth on this medium were selected, and an isolate exhibiting growth rate in 1/10-strength and full-strength KB broth equivalent to the wild-type strain was selected for use in these studies. Strains were cultivated overnight at 28°C in KB broth; cells were pelleted by centrifugation and washed twice in sterile distilled water. Cells were resuspended in sterile distilled water and 0.5% (w/v) methylcellulose for use in soil and seed treatments, respectively.

Seeds of the wheat cultivars Eltan, Lewjain, and Penawawa were treated individually by mixing 25 g of seed with a 10-mL cell suspension of Q2-87 or LR3-A28. Treated seeds were air dried in a laminar-flow hood and seed populations of the respective strains were determined as previously described [26]. WVC-Auvil orchard soil was dispensed into tapered plastic tubes (21 cm depth, 3.8 cm top diameter; Stuewe & Sons, Inc., Corvallis, OR); three wheat seeds were placed on the soil surface and covered with vermiculite to a depth of 1 cm. At emergence, each tube was thinned to a single wheat plant. Plants were grown at 24°C with a 16-h photoperiod. Fourteen seedlings were harvested at 10, 20, and 30

days after planting, and populations of the introduced strain were determined as previously described [25].

For soil applications, bacterial strains were introduced to individual samples of WVC-Auvil orchard soil as an atomized mist resulting in an initial population density of approximately 10⁴ and 10⁵ cfu g⁻¹ soil for strain LR3-A28 and Q2-87, respectively. Soils were dispensed into tapered plastic tubes and each cell was planted with five seeds of the wheat cultivar Eltan, Lewjain, or Penawawa, with 14 replicates for each wheat cultivar–bacterial strain treatment. Plants were grown for 28 days at 24°C with a 16-h photoperiod. At harvest, plants and soil were removed from an individual cell and a 0.5 g bulk root sample with firmly adhering soil was collected. Shoots were excised and soil with the remaining wheat root system was decanted back into the cell. Each cell was replanted with five seeds of the same wheat cultivar and the growth/harvest cycle was repeated five times. Root samples with associated soil were placed in 10 mL sterile water and vortexed for 60 s. Serial dilutions of the resulting root wash were plated onto KB agar amended with rifampicin (100 µg mL⁻¹) and cycloheximide (75 µg mL⁻¹). Colonies were enumerated after incubation for 72 h at 28°C.

Statistical Analysis. Population data were log-transformed prior to analysis. Percent data were transformed to arcsine-square root values prior to analysis. Data were subjected to analysis of variance and means separation using the Student–Newman–Keuls method. Statistical analyses were conducted using Sigma-Stat, version 2.0 (SPSS, San Rafael, CA).

Results

Size of the fluorescent *Pseudomonas* spp. population recovered from the rhizosphere of wheat was dependent upon the orchard soil in which wheat was cropped. Populations recovered from the rhizosphere of wheat cropped in WVC-Auvil and CV soils, averaging 7.82 and 6.81 × 10⁷ cfu g⁻¹ root, respectively, did not differ significantly. However, rhizosphere populations recovered from wheat cropped in WVC-SS soils were significantly ($P < 0.001$) lower and averaged 9.14 × 10⁵ cfu g⁻¹ root. For a given orchard soil, fluorescent pseudomonad rhizosphere populations did not vary significantly among the wheat cultivars examined (data not shown). *PhlD*⁺ isolates were not detected (limit of detection = 10⁴ cfu g⁻¹ soil) among fluorescent pseudomonad populations recovered from any of the orchard soils prior to cropping with wheat.

As determined by direct amplification of isolates randomly selected after the third wheat cycle from serial dilutions ($n = 60$ per wheat cultivar; $n = 300$ per soil),

the fluorescent pseudomonad population recovered from the rhizosphere of wheat (all cultivars combined) cropped in WVC-Auvil soil possessed a significantly ($P < 0.05$) greater proportion of 2,4-DAPG-producing strains relative to the population recovered from wheat cropped in CV or WVC-SS orchard soil. *phlD*⁺ strains comprised 5.0%, 2.3%, and 10.7% of the isolates evaluated from the rhizosphere of wheat cropped in CV, WVC-SS, and WVC-A soils, respectively.

From these populations, the relative proportion of *phlD*⁺ isolates recovered from the wheat rhizosphere varied in a cultivar-dependent manner. The fluorescent *Pseudomonas* spp. population isolated from the rhizosphere of Lewjain commonly possessed a greater proportion of *phlD*⁺ isolates relative to other wheat cultivars, regardless of the soil in which wheat was cropped (Fig. 1). In one experiment conducted in WVC-Auvil soil, *phlD*⁺ isolates represented 28% (17 of 60 isolates) of the fluorescent pseudomonad isolates examined from the Lewjain rhizosphere as detected by direct PCR. In comparison, using the same method, 2,4-DAPG-producing strains were only recovered from the rhizosphere of Madsen in a single experiment conducted in WVC-Auvil soil. *PhlD*⁺ isolates never exceeded 5% of the population recovered from the rhizosphere of Eltan and were not detected in the population examined by direct PCR amplification of individual isolates recovered from this cultivar when grown in WVC-SS orchard soil.

Similar results were obtained in studies where a *phlD* probe was used in colony hybridizations to examine the fluorescent pseudomonad community. Populations of *phlD*⁺ fluorescent pseudomonads recovered from the wheat rhizosphere differed among the orchard soils examined. *PhlD*⁺ isolates were consistently detected in populations recovered from the rhizosphere of Lewjain regardless of the orchard soil in which wheat was cultivated (Table 1). With the exception of Lewjain, rhizosphere populations of *phlD*⁺ fluorescent pseudomonads recovered from wheat cropped in WVC-SS orchard soil were below the limits of detection (Table 1). As observed in direct detection by amplification of DNA from randomly selected isolates, colony hybridization experiments demonstrated that wheat cropped in WVC-Auvil and CV orchard soils sustained greater densities of 2,4-DAPG-producing fluorescent pseudomonads than did wheat cultivated in WVC-SS soil.

Genotype Diversity of *phlD*⁺ Isolates from Different Wheat Cultivars. The genotypic diversity of the isolated *phlD*⁺ fluorescent pseudomonad population was determined by RFLP analysis of the *phlD* gene and classified based on previously erected genotypes [30]. The results showed that the genetic composition of the *phlD*⁺ population varied among soils and, in certain instances, was dependent on the wheat genotype. All *phlD*⁺ isolates

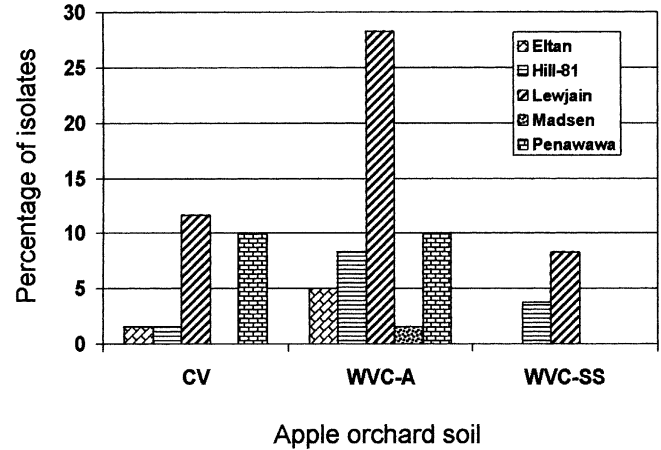


Figure 1. Percentage of *phlD*⁺ isolates determined by direct PCR of individual fluorescent *Pseudomonas* isolates ($n = 60$) recovered from the rhizosphere of different wheat cultivars at the end of three successive 28-day cycles of wheat cropping in Columbia View (CV), Wenatchee Valley College–Auvil (WVC-A), or Wenatchee Valley College–Sunnyslope (WVC-SS) apple orchard soils.

recovered from the rhizosphere of wheat cropped in CV orchard soil were defined as genotype A regardless of the wheat cultivar. The few isolates recovered from the rhizosphere of wheat cropped in WVC-SS all were identified as genotypes A or B based upon RFLP analysis of the *phlD* gene. In contrast, the population recovered from the rhizosphere of wheat cropped in WVC-Auvil orchard soil consisted of four different genotypes including genotypes A and B, and two previously undefined genotypes referred to as PfY and PfZ (Figs. 2 and 3). Isolates defined as genotype A were recovered from the rhizosphere of all wheat cultivars grown in WVC Auvil soil. Wheat cultivar Lewjain was the primary source of isolates defined as genotype PfY while Penawawa yielded the majority of

Table 1. Population densities (cfu g⁻¹ root) of *phlD*⁺ fluorescent pseudomonads recovered from the rhizosphere of five different wheat cultivars grown in apple orchard soils

Wheat cultivar	Orchard ^x		
	CV	WVC-A	WVC-SS
Eltan	ND	ND	ND
Hill-81	2.09 × 10 ⁴ a ^y	ND	ND
Lewjain	2.15 × 10 ⁵ b	2.75 × 10 ⁵ a	12,500
Madsen	3.63 × 10 ⁴ a	ND	ND
Penawawa	4.80 × 10 ⁴ ab	3.20 × 10 ⁵ a	ND

^xCV: Columbia View Research and Demonstration Orchard, Orondo; WVC-A: Wenatchee Valley College Auvil Research and Demonstration Orchard, E. Wenatchee; WVC-SS: Wenatchee Valley College Sunnyslope Orchard, Wenatchee, WA.

^yPopulations were determined after completion of three successive 28-day wheat cropping cycles by colony blot hybridization using the *phlD* probe. Each value is a mean consisting of six replicates. Statistical analyses were performed on log-transformed population densities. Means in the same column followed by the same letter are not significantly ($P > 0.05$) different. ND: Not detected; limit of detection = 10⁴ cfu g⁻¹ root.

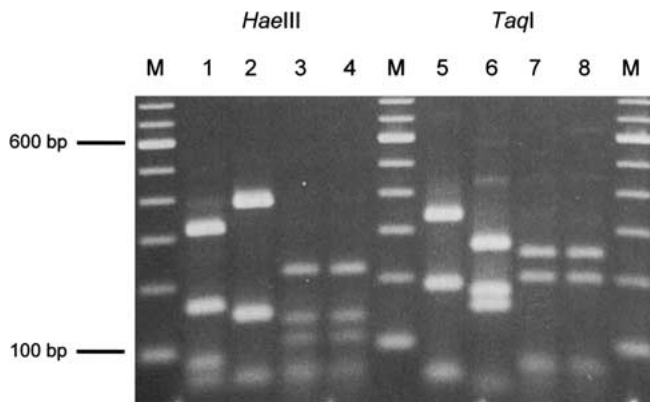


Figure 2. Restriction fragment length polymorphism analysis of the 629-bp *phlD* fragment amplified in reactions with the B2BF and BPR4 [29] primer pair. DNA was digested with *Hae*III or *Taq*I and products were separated on a 2.0% agarose gel. The sizes of individual fragments were compared with 100-bp ladder (M). PCR was performed using template from lysed cells of the following isolates, lanes 1 and 5: Q2-87 (genotype B); lanes 2 and 6: HR3-A13 (PfY); lanes 3 and 7: PR3-A3 (PfZ); lanes 4 and 8: PR3-A52 (PfZ); lane 4: Q2-87.

isolates defined as genotype PfZ (Fig. 2). Sequence analysis of the 16S rDNA placed representative isolates from both genotypes within *P. fluorescens*.

Sequence and Phylogenetic Analyses of phlD+ Genotypes PfY and PfZ. For two isolates representing *phlD* genotypes PfY and PfZ, a 745-bp fragment of the

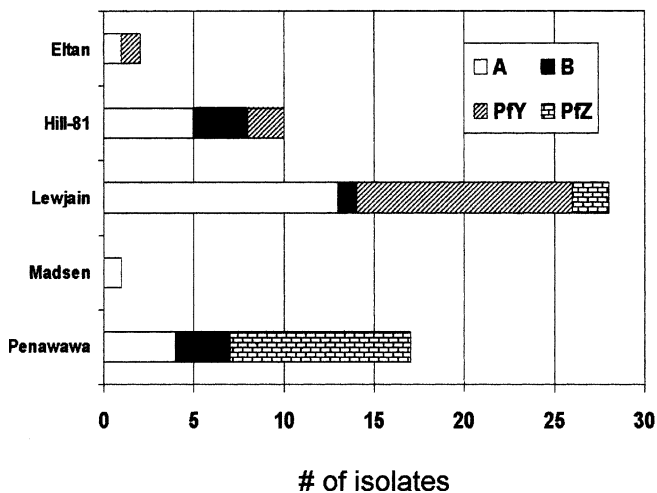


Figure 3. Distribution of *phlD*+ genotypes recovered from the rhizosphere of five wheat cultivars grown in Wenatchee Valley College–Auvil orchard soil. The population includes those fluorescent *Pseudomonas* obtained by plating serial dilutions of root washes on KB⁺ and randomly selecting isolates for subsequent amplification with the Phl2a and Phl2b primers to determine the presence of the *phlD* gene, as well as isolates detected by probing colony blots with the *phlD* probe. Genotypes A and B indicate BOX genotypes as defined by McSpadden Gardner et al. [30], and genotypes PfY and PfZ indicate new *phlD* genotypes.

Table 2. Bacterial species, strains, and *phlD* GenBank accession numbers used in the phylogenetic analyses

Species	Strain/isolate	Code	Accession number
<i>P. fluorescens</i>	42-36	Pf4236	AF396857
	42-27	Pf4227	AF396856
	39-8	Pf398	AF396855
	37-27	Pf3727	AF396854
	22-27	Pf2227	AF396853
	19-41	Pf1941	AF396852
	19-30	Pf1930	AF396851
	19-7	Pf197	AF396850
	18-33	Pf1833	AF396849
	11-18	Pf1118	AF396848
	7-37	Pf737	AF396847
	6-28	Pf628	AF396846
	3-1	Pf31	AF396845
	F113	Pff113	AJ278811
	PILH1	PfPILH1	AJ278810
	PITR2	PfPITR2	AJ278809
	CM1'A2	PfCM1A2	AJ278808
	Q65C-80	PfQ65c80	AJ278807
	CHA0	PfCHA0	AJ278806
	Pf-5	PfPf5	AF214457
	UP61	PfUP61	AF214108
	Q8r1-96	PfQ8r196	AF207693
	M1-96	PfM196	AF207692
	Q2-87	PfQ287	U41818
HR3-A13	PfY	This study	
PR3-A52	PfZ	This study	
<i>P. putida</i>	R62	PpR62	AJ515696
	872	Pc872	AJ515694
<i>P. corrugata</i>	711	Pc711	AJ515693
	709	Pc709	AJ515692
	R117	AfR117	AJ515695
<i>Acidovorax facilis</i>	R81	CaR81	AJ515697
<i>Commamonas acidovorans</i>			AJ515662 to
Uncultured bacteria (n = 30)			AJ515691

phlD gene was amplified, cloned, sequenced, and aligned to known *phlD* genes. Currently, 60 bacterial *phlD* sequences are present in the database of which 30 (GenBank accession numbers AJ515662 to AJ515691) were derived from uncultured bacteria (Table 2). Our phylogenetic analysis focused initially on comparison of the *phlD* gene sequences of PfY and PfZ isolates with 24 *phlD* gene sequences of other *P. fluorescens* isolates. These 24 sequences were selected as their sizes are approximately 650 to 700 bp, whereas the other *phlD* sequences present in the database are only approximately 230 bp. After alignment and trimming, *P. fluorescens* strain UP61 (accession number AF214108) was excluded from the phylogenetic analyses since its *phlD* sequence is substantially different from the other 23 *phlD* sequences, including those obtained from PfY and PfZ, thereby creating multiple gaps scattered across the entire sequence. Subsequent neighbor-joining and maximum-parsimony analyses showed that the *phlD* genes found in PfY and PfZ isolates were significantly different from any of the reported *phlD* sequences (Fig. 4A). Interestingly,

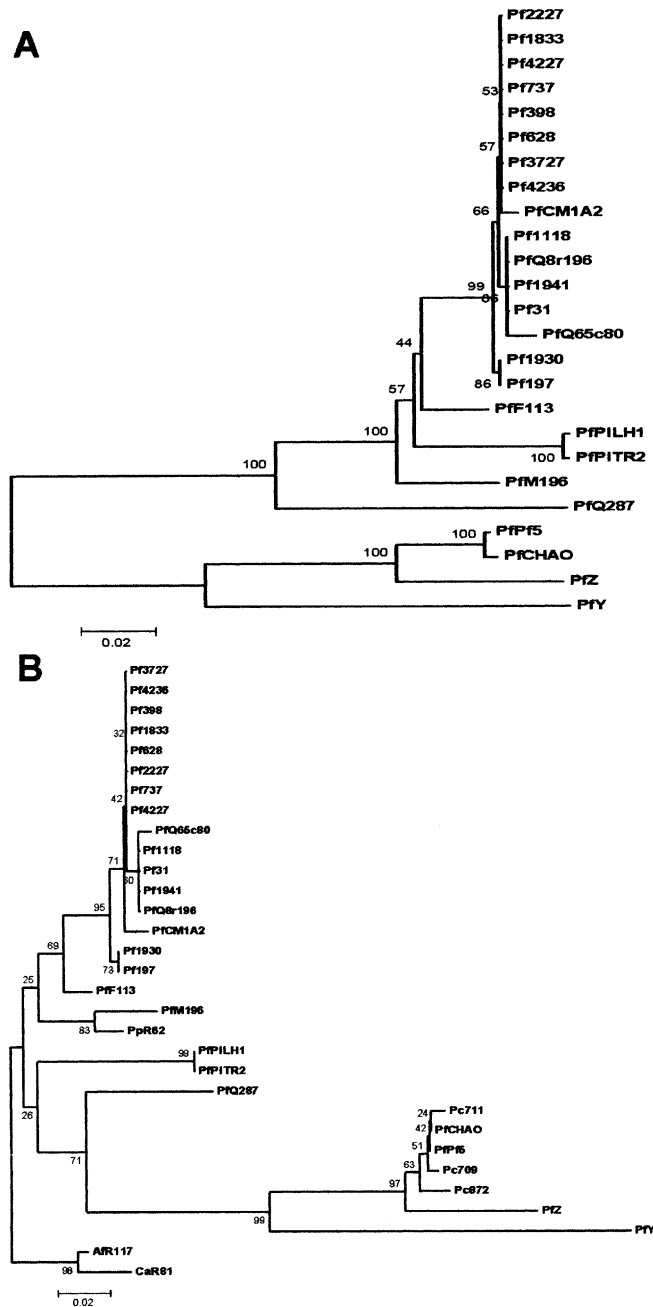


Figure 4. Phylogenetic trees generated with neighbor-joining using 670-bp *phlD* sequences from 25 strains of *Pseudomonas fluorescens* (A) and 210-bp *phlD* sequences from a total of 31 strains representing *P. fluorescens*, *P. putida*, *P. corrugata*, *C. acidovorans*, and *A. facilis* (B). The phylogenetic tree shown in panel A is unrooted, whereas the tree shown in panel B is rooted with the *phlD* sequence of *C. acidovorans* strain R81 (CaR81). Codes used for the bacterial species and strains/isolates are given in Table 2. Bootstrap values are indicated for each of the branches.

PfY and PfZ clustered close to each other and to *phlD* genes found in BOX genotype A strains CHA0 [10] and Pf-5 [33]. Also, analysis of the deduced protein sequences

revealed that *phlD* from PfY and PfZ isolates are different from known *PhlD* sequences and closely related to those from strains CHA0 and Pf-5 (data not shown). After this initial analysis covering a relatively large part of the *phlD* gene, phylogenetic analysis was also performed with an additional six *phlD* sequences obtained from strains other than *P. fluorescens*, including three *Pseudomonas corrugata* isolates, one *Pseudomonas putida* isolate, one *Acidovorax facilis* isolate, and one *Comamonas acidovorans* isolate (Table 2). After alignment, *phlD* sequences were trimmed to 210 bp. In the phylogenetic analysis, the *phlD* sequence from *C. acidovorans* isolate R81 was used to root the tree. The results showed again that PfY and PfZ represent new *phlD* genotypes (Fig. 4B). Interestingly, the *phlD* genes obtained from the three *P. corrugata* isolates were highly similar to the *phlD* genes of strains CHA0 and Pf-5. Neighbor-joining and maximum-parsimony analyses yielded similar results. Also analysis of the deduced protein sequences revealed that PfY and PfZ are different from known *PhlD* sequences and closely related to those from strains CHA0 and Pf-5 (data not shown). Final analysis with all 60 *phlD* sequences, including those obtained from uncultured bacteria, confirmed that PfY and PfZ represent two new *phlD* genotypes (data not shown).

Colonization of Wheat Cultivars by Two Genotypically Different 2,4-DAPG Producers. When introduced as a seed inoculant, 2,4-DAPG-producing fluorescent *Pseudomonas* strains Q2-87 (BOX genotype B) and LR3-A28 (genotype PfY) characteristically persisted in the rhizosphere at similar populations among the three wheat cultivars examined. Bacterial strains were established at initial populations of $\sim 4 \times 10^7$ cells seed⁻¹. Rhizosphere populations of strain Q2-87 or LR3-A28 exhibited a continuous decline over a 30-day period, and this trend transpired in a similar fashion for all wheat cultivars (data not shown). For strain Q2-87, populations declined from ~ 4 log cfu cm⁻¹ root at 10 days after planting to 2 log cfu cm⁻¹ root at 30 days. Strain LR3-A28 followed a similar pattern declining from ~ 5 log to 2.5 log cfu cm⁻¹ root over the same time period. No significant differences in populations of these two strains were detected among wheat cultivars at any sampling point.

Similar population dynamics of strain Q2-87 applied directly to soil were detected in soils cropped with five successive cycles of the same wheat cultivar. Populations of strain Q2-87 declined from a maximum of 5 log cfu cm⁻¹ root after cycle 1 to a population below the level of detection for all wheat cultivars at the end of cycle 5 (Fig. 5A). No significant differences among three wheat cultivars in populations of Q2-87 were noted at any sampling. In contrast, rhizosphere populations of strain LR3-A28 recovered from Eltan and Penawawa remained rather static over the course of successive wheat cycling

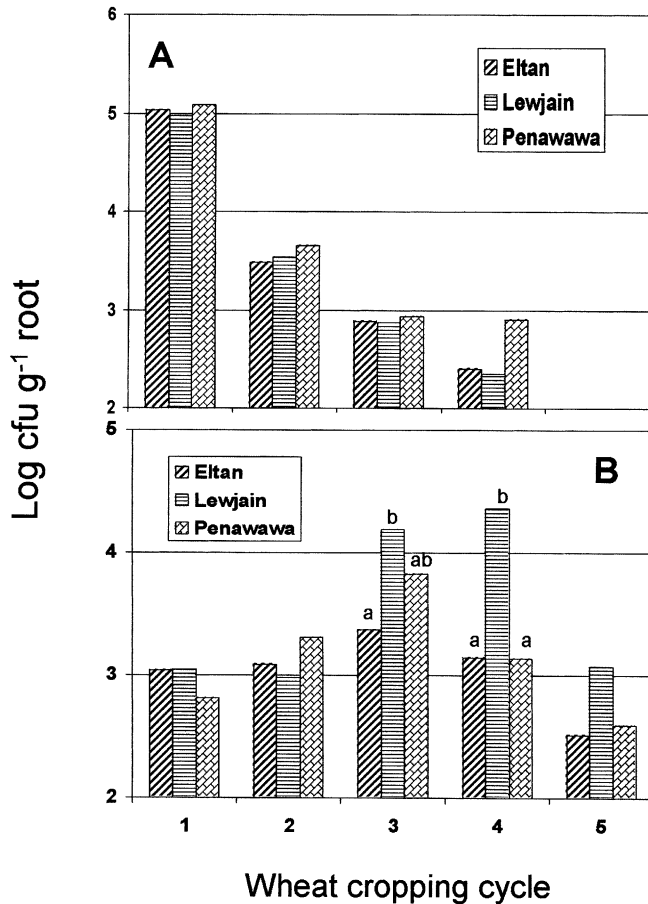


Figure 5. Mean population density of *Pseudomonas fluorescens* strain Q2-87 (A) and strain LR3-28A (B) recovered from the roots of wheat grown in Wenatchee Valley College–Auvil orchard soil for five successive cycles of 28 days each (A). Each mean consists of 14 replicates. Strain Q2-87 was not recovered from any wheat cultivar at the completion of cycle 5. For strain Q2-87, no significant ($P = 0.05$) differences were detected at any sampling period. For strain LR3-28A, means for the same cycle designated with the same letter are not significantly ($P = 0.05$) different.

(Fig. 5B). Moreover, populations of strain LR3-A28 recovered from Lewjain increased over the initial four wheat cycles to a density of $4.36 \log \text{cfu g}^{-1} \text{root}$ in cycle 4 (Fig. 5B). Significant differences among wheat cultivars were observed at cycles 3 and 4 when the population of strain LR3-A28 recovered from Lewjain was significantly larger than that recovered from either Eltan (cycle 3 and 4) or Penawawa (cycle 4). After three and four successive growth cycles, strain LR3-A28 established more than 10-fold higher rhizosphere population densities in the rhizosphere of cultivar Lewjain than strain Q2-87 (Fig. 5A, B).

Discussion

Plants have an influential role in determining the species and genetic composition of soil microbial com-

munities [7, 12]. Previous studies have demonstrated that composition of the fluorescent pseudomonad population will vary significantly within the rhizosphere of different plant hosts cultivated in the same soil [17] and will change dramatically overtime in response to changes in the crop species [23, 27]. The impact of plant species on composition of microbial communities is accepted, and there is emerging evidence that host genotype will have a significant influence on the structure and function of important elements of the saprophytic soil microbial community. As observed in this and previous studies [8], fluorescent pseudomonad populations are enhanced in a similar fashion by cropping soils with wheat regardless of the cultivar employed. However, the genetic and species composition of the fluorescent pseudomonad community recovered from the rhizosphere of wheat was found to vary among cultivars when grown in the same soil [8, 28]. In addition, expression of the 2,4-DAPG biosynthesis gene *phlA* in *P. fluorescens* CHA0 was found to differ among cultivars of maize [34].

As take-all decline functions in part through the enrichment of 2,4-DAPG-producing fluorescent *Pseudomonas* spp. and their activity in the wheat rhizosphere (reviewed in Weller et al. [52]), it is plausible that a wheat cultivar possessing the capacity to selectively enhance resident populations of these bacteria would encourage disease decline more readily than cultivars that do not select for these strains. In the current study, it was observed that wheat cultivars possessed a differential capacity to select for resident populations of 2,4-DAPG-producing fluorescent pseudomonads. When using either of two assessment methods, *phlD*⁺ isolates invariably were detected among the population of fluorescent *Pseudomonas* spp. recovered from the rhizosphere of Lewjain. In several instances the population of 2,4-DAPG-producing fluorescent pseudomonads recovered from this cultivar was significantly larger than that present in the rhizosphere of four other wheat cultivars examined. This trend was observed irrespective of soil in which wheat cycling was conducted. For all other wheat cultivars, 2,4-DAPG-producing fluorescent pseudomonads were not detected in one or more of the three soils examined, and recovery of *phlD*⁺ isolates from the rhizosphere of cultivars Eltan or Madsen was atypical. A similar finding was obtained by Gu and Mazzola [8], where 2,4-DAPG-producing fluorescent pseudomonads, defined as genotype 9 by the methods employed in that study, represented 17–33% of the population recovered from the rhizosphere of apple seedlings planted in soils previously cropped with Lewjain, Penawawa, or Symphony, but represented only 0–7% of the isolates from apple grown in Eltan, Hill-81, or Madsen cropped soils.

These findings suggest that a continuous wheat monoculture cropping system which employed Lewjain

could conceivably achieve a state of take-all decline more rapidly than with any of the other four wheat cultivars examined. However, one confounding factor to employing such a strategy may involve the genetic composition of the 2,4-DAPG-producing fluorescent *Pseudomonas* spp. population resident to a soil. In all three soils examined in this study, isolates representing BOX genotype A were recovered from roots of all five wheat cultivars tested. Interestingly, two other *phlD* genotypes, referred to as PfY and PfZ, were isolated. Based on RFLP and sequence analyses of *phlD*, PfY and PfZ represent two new *phlD*⁺ genotypes which are closely related to the *phlD* genotypes represented by *P. fluorescens* strains CHA0 and Pf-5. 16S rDNA sequence analysis confirmed that PfY and PfZ isolates were closely related to strains CHA0 and Pf-5. Wheat cultivar Lewjain was the primary source of genotype PfY while cultivar Penawawa yielded the majority of genotype PfZ. In contrast to the results obtained in this study, BOX genotype D isolates were previously found to predominate on roots of wheat grown in field soils exhibiting take-all decline in Washington [29, 42] and have been shown to displace genotype A isolates from soil and the rhizosphere when co-inoculated at equivalent populations [14]. Because of their superior colonization abilities, genotype D isolates provide a level of take-all control superior to genotype A isolates. Thus, even should the cultivar Lewjain enhance populations of 2,4-DAPG-producing isolates quantitatively, initial selection of an inferior genotype could result in a delay in the development of take-all decline. In this context, the ability of PfY and PfZ genotypes to control soil-borne disease of wheat, and in particular take-all, needs to be established to further support this hypothesis.

As has been noted previously, the development of take-all decline may be influenced by multiple attributes of the agro-ecosystem [44, 52]. In this study, it was rare to recover *phlD*⁺ isolates from the rhizosphere of wheat grown in WVC-SS orchard soil, and populations of 2,4-DAPG-producing fluorescent pseudomonads did not approach the threshold densities ($>10^5$ cfu g⁻¹ root) necessary to achieve take-all decline [41]. In contrast, populations of *phlD*⁺ fluorescent pseudomonads in the rhizosphere of one or more cultivars were greater than this threshold population when wheat was cropped for three successive cycles in CV or WVC-A orchard soils. This finding suggests that take-all decline could be expressed earlier in WVC-A and CV soils relative to WVC-SS soil.

Root colonization by introduced biocontrol rhizobacteria has been the subject of intense study as in many systems it has been deemed an essential element in determining the ultimate success of biological disease control [20]. The maintenance of a threshold population by an introduced biocontrol agent is known to be a factor in the ability to achieve effective disease suppression in

certain systems [9, 39], including the application of phenazine or 2,4-DAPG-producing fluorescent pseudomonads for the control of take-all of wheat [3, 41]. In the current study, no differences were observed among wheat cultivars in colonization of the rhizosphere by strain Q2-87 when introduced as either a seed or soil inoculant. Colonization of the wheat rhizosphere by strain LR3-A28 when applied to seed was similar for all cultivars. However, when applied to soil which was subsequently cropped to successive wheat cycles, strain LR3-A28 preferentially increased in the rhizosphere of Lewjain relative to Eltan and Penawawa for a period of time, prior to declining to similar populations during the final cropping cycle. Similarly, Okubara et al. [35] suggested a wheat cultivar variable response in colonization by *phlD*⁺ fluorescent pseudomonads based on the observation that the 2,4-DAPG producing strain Q8r1-96 was maintained in the rhizosphere of certain, but not all, wheat cultivars at significantly higher populations than strain Q2-87. Specific genotypes of 2,4-DAPG-producing fluorescent pseudomonads appear to preferentially colonize certain crop species [13, 36]. These and other studies [46, 47] indicate that linking a specific plant genotype with an appropriate genotype of a biocontrol rhizobacterium may enhance the efficacy of the disease control system.

Several recent studies have demonstrated the impact of vegetation on composition of the resident soil microbiota [7, 12, 17] and have indicated an important role for host genotype in determining the efficacy of resident and introduced microorganisms for the control of soilborne diseases [18, 28, 34, 46]. Management of the resident soil microbial community holds promise as a means to alleviate the activity of soilborne plant pathogens [28]. Such a system could employ specific cropping sequences to induce soil suppressiveness. However, a major limitation to the use of this management strategy is the time frame required to achieve an economically viable level of disease control. Again, the evidence garnered to date has demonstrated a plant-genotype-dependent capacity to select for specific functional microbial elements that contribute to soil suppressiveness [15, 28]. We postulate that selection of the appropriate plant genotype could substantially reduce the time frame required to induce suppressiveness to a specific soilborne disease. This may allow for the effective exploitation of the phenomenon as a component of an alternative disease management protocol in agricultural ecosystems.

Acknowledgments

This material is based upon work supported by the USDA National Research Initiative Competitive Grants Program under agreement number 2001-35319-09922.

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