Biocontrol and plant growth-promoting activity of rhizobacteria from Chinese fields with contaminated soils

Xuefei Wang,1,2 Dmitri V. Mavrodi,3 Linfeng Ke,1 Olga V. Mavrodi,2 Mingming Yang,2 Linda S. Thomashow,4 Na Zheng,1 David M. Weller4** and Jibin Zhang1*

1State Key Laboratory of Agricultural Microbiology, College of Life Science and Technology, Huazhong Agricultural University, Wuhan, Hubei 430070, China.
2Department of Plant Pathology, Washington State University, Pullman, WA 99164-6430, USA.
3Department of Biological Sciences, The University of Southern Mississippi, Hattiesburg, MS 39406, USA.
4Agricultural Research Service, Root Disease and Biological Control Research Unit, United States Department of Agriculture, Pullman, WA 99164-6430, USA.

Summary

The aim of this study was to inventory the types of plant growth-promoting rhizobacteria (PGPR) present in the rhizosphere of plants grown in soils contaminated with heavy metals, recalcitrant organics, petroleum sewage or salinity in China. We screened 1223 isolates for antifungal activity and about 24% inhibited Rhizoctonia solani or Sclerotinia sclerotiorum. Twenty-four strains inhibitory to R. solani, Gaeumannomyces graminis var. tritici and/or S. sclerotiorum and representing the dominant morphotypes were assayed for PGPR activity. Seven strains contained phiD, prnD, pltC or phzF genes and produced the antibiotics 2,4-diacylphloroglucinol, pyrrolnitrin, pyoluteorin and phenazines respectively. Six strains contained acdS, which encodes 1-aminocyclopropane-1-carboxylic acid deaminase. Phylogenetic analysis of 16S rDNA and phiD, phzF and acdS genes demonstrated that some strains identified as Pseudomonas were similar to model PGPR strains Pseudomonas protegens Pf-5, Pseudomonas chlororaphis subsp. aureofaciens 30–84 and P. brassicacearum Q8r1-96. Pseudomonas protegens- and P. chlororaphis-like strains had the greatest biocontrol activity against Rhizoctonia root rot and take-all of wheat. Pseudomonas protegens and P. brassicacearum-like strains showed the greatest promotion of canola growth. Our results indicate that strains from contaminated soils are similar to well-described PGPR found in agricultural soils worldwide.

Introduction

Soil salinization and contamination by organic compounds (i.e. petroleum products and solvents) and heavy metals (i.e. Cd, Cr, Cu, Hg, Pb) are important environmental problems worldwide and have negative impacts on both human health and agricultural productivity (Jing et al., 2007). Soil pollution is of particular concern in developing countries where rapid industrialization has led to contamination of agricultural lands, especially in small farms that are sources of food for consumption by local populations. In a survey of contamination levels of heavy metals in agricultural soils in China, metal concentrations of Cd were significantly higher than in non-contaminated soils (Wei and Yang, 2010). Similarly, because of the accumulation of mine tailing outfalls, the content of Cu in the wetlands of Daye County, Hubei province, was 18 times higher than the background value (Guo et al., 2011). The build-up of pollutants in soils and their subsequent leaching into the ecosystem has cascading effects, adversely impacting plant growth and yield (Cheng, 2003b; Huang et al., 2005; Parida and Das, 2005). The uptake of soil pollutants by crop plants and the resultant entry of carcinogens, endocrine disruptors and neurotoxins into the food chain are major concerns for human and animal health (Cheng, 2003a; Cai et al., 2009). In addition, soil pollutants can positively or negatively influence crop susceptibility to pathogens and insects (Elmer, 2002; Poschenrieder et al., 2006). For example, salinity can predispose plants to infection by soilborne pathogens or increase the incidence and severity of disease (Triky-Dotan et al., 2005; Roubtsova and Bostock, 2009).
Rhizobacteria are able to colonize and persist on roots and some, known as plant growth-promoting rhizobacteria (PGPR), are able to promote plant growth (Weller, 2007). PGPR improve plant growth directly by supplying nutrients (e.g. phosphorous, nitrogen), producing phytohormones [e.g. indoleacetic acid (IAA)], and/or decreasing the level of ethylene that is produced when the plant is stressed (Li et al., 2007; Lugtenberg and Kamilova, 2009). PGPR also antagonize plant pathogens and/or induce systemic resistance in the plant to disease (Weller et al., 2007; Lugtenberg and Kamilova, 2009). Because plants lack effector-triggered immunity to many soilborne pathogens, they rely on PGPR for defence against pathogen attack (Weller et al., 2007).

Soil pollution, especially by heavy metals, can also decrease the metabolic activity, biomass and diversity of microorganisms in the bulk soil and the rhizosphere (Sanda et al., 1999; Gremion et al., 2004; Jing et al., 2007) and limit the effectiveness of PGPR to protect roots against disease. However, PGPR with metal resistance and the ability to mitigate the toxic effects of soil pollutants are present in polluted soils (Huang et al., 2005; Reed and Glick, 2005; Jing et al., 2007). For example, *Pseudomonas asplenii* AC isolated from polycyclic aromatic hydrocarbon-contaminated soils significantly increased the root and shoot growth of canola in copper and creosote-contaminated soils (Reed and Glick, 2005). Salt-tolerant *Pseudomonas chlororaphis* TSAU13, isolated from the rhizosphere of wheat grown in a salinated soil (Egamberdieva and Kucharova, 2009), improved the shoot and root growth of common bean grown in soil amended with NaCl and protected cucumber and tomato against *Fusarium solani* (Egamberdieva, 2011).

Despite progress in understanding the crucial role of PGPR in helping plants adapt to environmental stress, major gaps remain in our knowledge of how soil pollution affects the population levels, diversity and activity of PGPR. Our study is part of an ongoing cooperative project in China and the United States with the goal of identifying new PGPR and characterizing their role in the promotion of soil and plant health. Our study focused on rhizobacteria isolated from Chinese soils polluted with heavy metals, recalcitrant organic compounds, petroleum sewage or excessive salinization. We have hypothesized that certain groups of PGPR and strains harbouring biocontrol and growth-promoting genes are cosmopolitan and conserved in soil microbiomes in agroecosystems worldwide regardless of the condition of the soil, and that plants enrich and support those PGPR best adapted to help them grow and resist diseases and other pests. Nothing is known about the composition of PGPR in soils from these polluted sites, thus making them ideal to test our hypothesis.

Our specific objectives were to: (i) isolate and identify PGPR, (ii) determine their biocontrol activity against take-all disease and Rhizoctonia root rot on wheat, (iii) determine their growth-promoting activity on rapeseed/canola, and (iv) characterize their growth-promoting traits and genes. Wheat and rapeseed/canola were selected for our study because China is the world’s largest producer of both of these crops and they are often grown in rotation. Rhizoctonia root rot caused by *Rhizoctonia solani* AG-8 and take-all caused by *Gaeumannomyces graminis* var. *tritici* are two important soilborne diseases worldwide, against which wheat has no resistance (Cook et al., 2002; Paulitz et al., 2002; Yang et al., 2011); however, biocontrol with PGPR is a viable approach to controlling them. Poor emergence is a chronic problem in rapeseed/canola production in both China and the United States and application of PGPR as seed treatments can greatly enhance shoot and root growth.

Here we report that strains of well-described PGPR species were among the common morphotypes from the rhizosphere of wheat and rapeseed grown in highly polluted soils in China. Strains of *Lysobacter capsici*, *P. chlororaphis*, *Pseudomonas protegens* and those related to *P. brassicacearum* had some of the greatest biocontrol and/or growth promoting activity. PGPR genes in these morphotypes included *phd*, *phzF*, *pmrD*, *pitC* and *acdS*, and traits included the ability to produce 2,4-diacetylphloroglucinol (2,4-DAPG), phenazines, pyrrolnitrin, pyoluteorin, cellulase, siderophores, protease and IAA.

**Results**

**Population densities and antagonistic activity of rhizobacteria from contaminated soils**

The population sizes of total culturable aerobic, heterotrophic bacteria in the rhizosphere of wheat after three growth cycles in the greenhouse in the contaminated soils from five fields located in Hubei, Jiangxi and Jiangsu provinces of China ranged from $4.2 \times 10^6$ to $3.4 \times 10^6$ colony-forming units (cfu) g$^{-1}$ fresh weight of root (Table 1). Populations recovered from rapeseed grown in the same soils ranged from $3.6 \times 10^6$ to $1.3 \times 10^5$ cfu g$^{-1}$ fresh weight of root (Table 1). From the wheat and rapeseed rhizospheres, 1223 bacterial isolates (605 from wheat and 618 from rapeseed) were randomly selected from dilution plates of $1/5$× King’s medium B (KMB) and $1/10$× tryptic soy agar (TSA). The percentage of isolates from wheat inhibitory to *R. solani* and from rapeseed inhibitory to *Sclerotinia sclerotiorum* varied among soils but averaged 24% and 23% respectively (Table 1). Isolates from the field contaminated with recalcitrant
Crop and native plants and weeds were dug from the contaminated soil at each site, soil attached to the roots was removed, transported to Wuhan, Dongyuan and Yinyang in Jiangsu province; GPS coordinates of these sites are 30°10′E; 29°46′N, 116°15′E; 31°01′N, 112°13′E; 31°58′N, 121°46′E; and 31°44′N, 121°54′E respectively. Crop and native plants and weeds were dug from the contaminated soil at each site, soil attached to the roots was removed, transported to Wuhan, Dongyuan and Yinyang in Jiangsu province; GPS coordinates of these sites are 30°10′E; 29°46′N, 116°15′E; 31°01′N, 112°13′E; 31°58′N, 121°46′E; and 31°44′N, 121°54′E respectively.

Table 1. Population densities of total culturable aerobic, heterotrophic bacteria in the rhizosphere of wheat and rapeseed and ability of the bacteria to inhibit soilborne pathogens in vitro.

| Locationa | Soil contaminant                      | Wheat rhizosphere | Canola rhizosphere |
|-----------|--------------------------------------|-------------------|--------------------|
|           |                                       | cfu g⁻¹ fresh weight root | Inhibitory to Rhizoctonia AG-8 (%) | cfu g⁻¹ fresh weight root | Inhibitory to Sclerotinia sclerotiorum (%) |
| Daye County | Heavy metals                          | 5.48 ± 2.85 × 10⁷ | 27.2               | 1.34 ± 0.44 × 10⁸ | 15.5                |
| Huokou County | Recalcitrant organic compounds       | 4.18 ± 3.0 × 10⁷  | 6.7                | 4.02 ± 1.41 × 10⁸ | 15.1                |
| Jinmen City | Petroleum products                    | 5.54 ± 2.3 × 10⁷  | 28.0               | 8.4 ± 0.35 × 10⁸  | 36.2                |
| Qidong City (Dongyuan) | Salt                                | 2.69 ± 1.4 × 10⁷  | 19.6               | 7.06 ± 0.13 × 10⁸  | 27.5                |
| Qidong City (Yinyang) | Salt                                | 3.39 ± 3.3 × 10⁷  | 37.2               | 3.56 ± 0.66 × 10⁸  | 21.8                |

a. Soil samples were collected in Daye County and Jinmen City in Hubei province, Huokou county in Jiangxi province, and Qidong City near Dongyuan and Yinyang in Jiangsu province; GPS coordinates of these sites are 30°10′E; 29°46′N, 116°15′E; 31°01′N, 112°13′E; 31°58′N, 121°46′E; and 31°44′N, 121°54′E respectively.

Organic compounds in Huokou County, Jiangxi province were the least inhibitory to \( R.\) solani and \( S.\) sclerotiorum (6.7% and 15.1% respectively) as compared with isolates from the other soils.

**Identity of the PGPR and their biocontrol and growth-promoting traits and genes**

Twenty-four strains representative of the dominant morphotypes (Table 2) were screened for growth-promoting and biocontrol-related traits and genes. Of these 24 strains, 15, 11 and 20, respectively, inhibited the growth of \( G.\) graminis var. tritici ARS-A1, \( R.\) solani AG-8 C-1 and \( S.\) sclerotiorum MGSCF180002 (Table 3).

Results of traditional microbiological tests and analysis of partial 16S rDNA gene sequences (Table 2) revealed that five of the 24 strains were Gram-positive and closely related to \( B.\) cereus, \( P.\) putida and \( P.\) protegens and \( D.\) tsuruhatensis. The rest of the strains represented two different clades of Gram-negative bacteria and were closely related to several species within the \( P.\) fluorescens and \( P.\) protegens groups, and to \( L.\) capsici, \( L.\) enymogenes, \( S.\) plymuthica and \( D.\) tauratensis. These 24 strains produced one or more of the extracellular metabolites cellulase, protease and siderophores. Genes for the biosynthesis of the antibiotics 2,4-DAPG (\( phlD\)), phenazines (\( phzF\)), pyrrolnitrin (\( pmD\)) and/or pyoluteorin (\( pltC\)) were detected in strains of \( P.\) protegens, \( P.\) chlororaphis and/or \( S.\) plymuthica (Table 3) and the antibiotics themselves were detected in culture extracts analysed by thin-layer chromatography. These antibiotics are well known to function as major mechanisms of biocontrol of soilborne pathogens.

The amplified fragments of \( phlD\), \( phzF\), \( pmD\) and \( pltC\) from these strains were sequenced and used in phylogenetic analyses together with corresponding alleles previously deposited in GenBank. Phylogenograms inferred from \( phlD\) and \( pltC\) sequences revealed that both of these genes from strains TM1109, TY1502, TY1508 and KY4410 were closely related to their homologues from \( P.\) protegens (Figs S2 and S3). Strains TM1109, TY1502, TY1508 and KY4410 also carried a \( P.\) protegens-like \( pmD\) allele and therefore had a combination of antibiotic biosynthesis pathways similar to those of the model biocontrol agents \( P.\) protegens strains Pf-5 and CHA0 (Fig. S4). The second \( pmD\)-positive \( P.\) protegens strain, KY5406, also carried \( phzF\) (Table 3), a gene required for the production of phenazine antibiotics and highly conserved in all known phenazine producers. The \( pmD\) and \( phzF\) genes of KY5406 clustered with homologues from \( P.\) chlororaphis (Figs S4 and S5). Strain KY5406 also contained \( phzO\) (data not shown), a gene required for the biosynthesis of 2-hydroxylated phenazine compounds, indicating this strain is subsp. \( aureofaciens\). Finally, two more \( pmD\)-positive strains, KM3407 and TM4307-1 (Table 3), carried \( pmD\) alleles similar to those present in pyrrolnitrin-producing strains of \( S.\) sclerotiorum spp. (Fig. S4) and produced Prn in vitro (data not shown). In total, correlation of the antibiotic gene profiling with the taxonomic
placement of the strains by partial 16S rRNA gene sequence analysis identified the antibiotic-producing strains as members of *P. protegens*, *P. chlororaphis* subsp. *aureofaciens* and *S. plymuthica*.

1-aminocyclopropane-1-carboxylic acid (ACC) deaminase and IAA production by PGPR are important determinants in plant–bacterial interactions, and secretion of these compounds can induce plants to produce longer and more vigorous roots (Penrose and Glick, 2003). *Delftia tsuruhatensis* TM3205 and *Pseudomonas* strains KM3113, KY5404, TY3517 and *Serratia plymuthica* KM3407 produced IAA in culture. PCR analysis demonstrated that the aforementioned *Pseudomonas* strains were positive for *acdS*, a gene required for the production of ACC deaminase (Table 4), and the sequences of those strains closely resembled those from other fluorescent pseudomonads (Fig. S6). All *acdS* sequences from the Chinese strains formed a distinct clade within the *Pseudomonas* group and clustered tightly with *acdS* from *Pseudomonas* sp. 2–3 (GenBank Accession No. EU520401). All *acdS*-positive strains grew in defined medium supplemented with ACC as a sole nitrogen source.

**Biocontrol and growth-promoting activity of the rhizobacterial strains**

A subset of strains with the highest antagonistic activity and representative of the dominant morphotypes was selected to determine their biocontrol activity against take-all and Rhizoctonia root rot of wheat under greenhouse conditions in sterile, pasteurized and raw soils from Wuhan, China and Quincy, Washington. Most of the isolates provided some levels of reduction in root disease as compared with the pathogen-inoculated controls (Tables 5–7). However, strains identified as *L. capsici*, *L. enzymogenes*, *P. protegens* and *P. chlororaphis* provided the best and most consistent disease suppression across experiments and pathogens. *Pseudomonas protegens* TM1109 was especially effective in biocontrol of both take-all and Rhizoctonia root rot. These results indicate that indigenous rhizobacteria from the contaminated soil have the capacity to provide protection of plants against soilborne fungal pathogens. Surprisingly, application of *S. plymuthica* TM4307-1, which highly antagonized the pathogens in *vitro* (Table 3), resulted in significantly higher levels of Rhizoctonia root rot (Table 6).
Isolates were tested \((P_{s} < 0.05)\) in at least one of the three experiments (Table 4). \(\text{Pseudomonas protegens}\) and \(\text{P. brassicacearum}\)-like strains demonstrated the most consistent capacity to promote root growth and, on average, increased root length in canola by 41\% to 42\%, respectively, as compared with the non-treated controls. All \(\text{P. brassicacearum}\)-like isolates produced IAA and carried \(\text{acdS}\). In contrast, strains of \(\text{P. protegens}\) had neither trait. Other tested \(\text{Pseudomonas}\) strains (i.e. \(\text{KY3201, TY3101, Ty3116w, Ty5511 and KM2402}\)) increased root length consistently but not always significantly, whereas strains of \(\text{Bacillus, Paenibacillus}\) and \(\text{Lysobacter}\) had no growth promotion activity. These results indicate that rhizobacteria from the contaminated soil have the capacity to directly enhance plant growth.

### Tolerance to abiotic stressors

Based on the results from the biocontrol and growth promotion assays, 10 of the 24 strains were selected and tested for tolerance to two heavy metals and high levels of salt by growing them in LB broth supplemented with \(\text{CdCl}_2\), \(\text{CuCl}_2\) and \(\text{NaCl}\). All 10 strains tolerated high levels of salt and grew well in up to 5–7\% NaCl. Similarly, these 10 strains grew in the presence of heavy metals and demonstrated minimal inhibitory concentrations (MICs) to \(\text{Cu}^2\) of > 4 mM and \(\text{Cd}^{2+}\) of > 1 mM. Overall, \(\text{P. protegens}\) TM1109 was the most tolerant to these stressors (Table 8).

### Discussion

Plants have evolved along with a rhizosphere microbiome that contributes to the growth and health of the plant (Pieterse et al., 2014). Beneficial microbes that directly promote growth or defend roots are recruited by the roots from the bulk soil, making soil type an important determinant of the composition of the rhizosphere microbiome. However, the plant also strongly modulates the shape of the microbial community because both the quality and quantity of root exudates are regulated by the plant genotype (Bais et al., 2006; Hartmann et al., 2009). Our
results support the hypothesis that certain groups of PGPR and biocontrol and growth-promoting genes are cosmopolitan and conserved in soil microbiomes worldwide. Driven by rhizodeposition, the plant enriches and supports PGPR best adapted to colonize the roots, antagonize pathogens, stimulate growth and/or initiate pattern-triggered immunity.

We tested our hypothesis by characterizing cultivable PGPR in the rhizospheres of wheat and rapeseed grown in soils from polluted fields, and assessed their growth-promoting and biocontrol abilities. After the most abundant bacterial morphotypes were isolated, the only other criterion we used for selecting the rhizobacteria was the ability to inhibit G. graminis var. tritici, R. solani or S. sclerotiorum. This was done because in vitro inhibition of plant pathogens is a characteristic commonly found in many PGPR strains (Haas and Défago, 2005) even though antibiosis does not predict biocontrol or growth-promoting activity. The 24 strains described here were shown to be most closely related to species known to

| Treatment                                      | acidS | IAA | Exp 1 Root length | Increase (%) | Exp 2 Root length | Increase (%) | Exp 3 Root length | Increase (%) |
|------------------------------------------------|-------|-----|-------------------|--------------|-------------------|--------------|-------------------|--------------|
| Bacillus sp. TY5107                            | –     | –   | 6.8 ± 0.5 BE      | 20.2         | 4.2 ± 0.4 GH      | –18.5        | 4.8 ± 0.3 I       | –29.4        |
| D. turuhatensis TM3205                         | –     | +   | 8.1 ± 0.4 AD      | 43.3         | 5.7 ± 0.4 DE      | 10.8         | 9.7 ± 0.3 A       | 42.7         |
| L. capsici TM5405                              | –     | –   | 7.0 ± 0.3 BE      | 22.5         | NT                | –           | 5.8 ± 0.4 H       | –13.6        |
| L. enzymogenes TM2502                          | –     | NT  | 4.9 ± 0.4 EG      | –4.5         | 5.9 ± 0.3 H       | –13.5        |
| M. hydrocarbonoxydans TY3211y                  | –     | –   | 7.9 ± 0.3 AD      | 38.8         | 5.5 ± 0.4 DF      | 7.1          | 7.7 ± 0.4 FG      | 13.9         |
| M. hydrocarbonoxydans TY3517                   | –     | +   | 5.8 ± 0.2 DE      | 2            | 5.4 ± 0.3 DF      | 5.2          | 6.5 ± 0.3 H       | –3.5         |
| Pa. polymyxa KM2501                            | –     | –   | 7.4 ± 0.3 AE      | 30.9         | 4.6 ± 0.4 FG      | –11.7        | NT                | –           |
| Pa. polymyxa KM4401                            | –     | –   | 3.1 ± 0.3 E       | –45.8        | 4.9 ± 0.4 EG      | –6.2         | NT                | –           |
| Pseudomonas sp. KM3113                         | +     | +   | 8.1 ± 0.4 AD      | 42.7         | 7.0 ± 0.4 AB      | 35.6         | 9.4 ± 0.4 AB      | 38.4         |
| Pseudomonas sp. KY5404                          | +     | +   | 8.5 ± 0.5 AC      | 50           | 7.9 ± 0.4 A       | 53.4         | 8.2 ± 0.3 DF      | 21.6         |
| Pseudomonas sp. TY1210                          | +     | +   | 9.6 ± 0.4 A       | 68.3         | 7.0 ± 0.3 AB      | 35.7         | 8.4 ± 0.4 CDF     | 23.5         |
| Pseudomonas sp. TY1205                          | +     | +   | 9.4 ± 0.5 A       | 64.8         | 7.0 ± 0.3 AB      | 35.4         | 9.3 ± 0.4 AC      | 37.8         |
| P. chlororaphis KY5406                          | –     | ±   | 6.0 ± 0.4 CE      | 5.9          | 4.4 ± 0.4 GH      | –15.6        | 5.9 ± 0.4 H       | –13         |
| Pseudomonas sp. KY3201                          | +     | –   | 9.7 ± 0.3 A       | 71.4         | 5.9 ± 0.3 CD      | 14.8         | 7.9 ± 0.4 EF      | 17.5         |
| Pseudomonas sp. TY3101                          | +     | +   | 8.4 ± 0.5 AB      | 48.8         | 5.5 ± 0.4 DF      | 6.9          | 9.7 ± 0.4 A       | 43          |
| Pseudomonas sp. TY3116w                         | –     | ±   | 8.1 ± 0.2 AD      | 42.9         | 5.8 ± 0.3 DE      | 12.2         | 7.6 ± 0.4 FG      | 12.7         |
| Pseudomonas sp. KY5511                          | –     | –   | 8.3 ± 0.3 AD      | 45.9         | 5.7 ± 0.3 CD      | 10.6         | 8.8 ± 0.3 AE      | 29.9         |
| Pseudomonas sp. KM2404                          | –     | ±   | 8.4 ± 0.3 AC      | 47.9         | 7.0 ± 0.4 AB      | 36           | 9.7 ± 0.4 DF      | 42.7         |
| P. protegens KY4410                             | –     | –   | 7.9 ± 0.3 AD      | 39.9         | 7.6 ± 0.3 AB      | 47.2         | 8.8 ± 0.4 AE      | 29.8         |
| P. protegens TM1109                             | –     | +   | 8.1 ± 0.4 AB      | 60.6         | 7.6 ± 0.3 AB      | 47.5         | 8.5 ± 0.3 BF      | 25.7         |
| P. protegens TY1502                             | –     | +   | 8.1 ± 0.4 AB      | 60.6         | 6.8 ± 0.3 BC      | 31.7         | 8.6 ± 0.4 BF      | 27.2         |
| P. protegens TY1508                             | –     | –   | 8.7 ± 0.5 AB      | 53.6         | 7.4 ± 0.3 AB      | 42.2         | 9.0 ± 0.4 AD      | 33.2         |
| Ck                                             | –     | –   | 5.7 ± 0.5 CE      | 5.2          | 5.2 ± 0.4 DG      | 6.8          | 5.3 ± 0.3 GH      |             |

a. Surface-sterilized canola seeds were soaked for 1 h in each bacterial suspension. Control (CK), non-treated seeds were soaked in sterile water. Presence (+) or absence (−) of the acidS gene. Bacterial cultures were positive (+) or negative (−) for the presence of IAA.

% increase = [((root length of a treatment - root length of the control)/ root length of the control) × 100%. Means in the same column followed by the same letter are not significantly different at P = 0.05 according to Fisher’s protected least significant difference test (LSD) or Kruskal–Wallis all pairwise comparison test.

Exp, experiment; NT, not tested.

Table 5. Suppression of Rhizoctonia solani AG-8 in sterile soil in greenhouse pot experimentsa.

| Treatment                                      | Disease rating | Shoot length (cm) |
|------------------------------------------------|----------------|------------------|
|                                                | Exp 1 Root length | Exp 2 Root length | Exp 1 | Exp 2 |
| CK1                                           | 0.4 ± 0.1        | 0.3 ± 0.5        | 25.3 ± 4.9 | 36.4 ± 2.7 |
| CK2                                           | 5.0 ± 1.6 A      | 4.6 ± 1.3 AB     | 20.5 ± 5.3 D | 33.6 ± 3.6 BC |
| L. capsici TM5405                              | 3.3 ± 1.6 D      | 3.1 ± 1.7 C      | 23.6 ± 4.6 ABC | 37.1 ± 3.9 A |
| L. enzymogenes TM2502                          | 3.7 ± 1.7 CD     | 3.7 ± 1.7 BC     | 23.1 ± 5.0 ABC | 31.7 ± 4.3 C |
| Pa. polymyxa KM2501                            | 4.1 ± 1.8 ABC    | 3.8 ± 1.5 BC     | 22.6 ± 4.4 BC | 33.9 ± 4.1 B |
| Pa. polymyxa KM4401                            | 3.1 ± 1.7 D      | 4.6 ± 1.3 AB     | 24.4 ± 4.7 A  | 34.8 ± 5.3 B |
| P. protegens TM1109                            | 3.8 ± 1.6 BCD    | 3.8 ± 1.7 BC     | 23.7 ± 4.7 AB | 31.7 ± 3.4 C |

a. CK1, control consisting of sterile soil from Wuhan, China not amended with inoculum of R. solani and sown to non-treated seeds; CK2, control consisting of soil amended with R. solani inoculum and sown to methyl cellulose coated seeds. Severity of Rhizoctonia root rot was evaluated on a scale of 0–8. Means in the same column followed by the same letter are not significantly different at P = 0.05 according to Fisher’s protected least significant difference test (LSD) or Kruskal–Wallis all pairwise comparison test.

Exp, experiment.

© 2014 The Authors. Microbial Biotechnology published by John Wiley & Sons Ltd and Society for Applied Microbiology, Microbial Biotechnology, 8, 404–418
Table 6. Suppression of *Rhizoctonia solani* AG-8 in Quincy virgin soil in a growth chamber tube assays.

| Treatment | Pasteurized soil | Raw soil |
|-----------|-----------------|----------|
|           | Exp 1 | Exp 2 | Exp 3 | Exp 4 |
| CK1       | 0.5 ± 1.5 | 0.1 ± 0.5 | 0.3 ± 0.8 | 0.1 ± 0.3 |
| CK2       | 4.8 ± 0.8 B | 4.9 ± 0.8 B | 4.4 ± 0.6 A | 4.5 ± 0.7 AB |
| CK3       | 4.7 ± 0.9 BC | 5.2 ± 0.8 B | 4.3 ± 0.6 AB | 4.6 ± 0.6 A |
| *D. tsuruhatensis* TM3205 | NE | NE | 3.9 ± 0.4 ABC | 4.1 ± 0.4 BC |
| *L. capsici* TM5405 | 4.3 ± 1.1 BCD | 4.8 ± 0.8 B | 3.8 ± 0.5 BC | 4.1 ± 0.7 C |
| *L. enzymogenes* TM2502 | NE | NE | 3.9 ± 0.9 ABC | 4.0 ± 0.5 C |
| *Pa. polymyxa* KM4410 | 4.4 ± 1.7 BCD | 4.9 ± 0.9 B | 3.9 ± 0.5 BC | 4.0 ± 0.4 C |
| *Pseudomonas* sp. KM3113 | NE | NE | 4.2 ± 0.7 AB | 4.1 ± 0.6 BC |
| *P. chlororaphis* KY5406 | 4.3 ± 0.8 CD | 3.9 ± 1.0 C | 3.4 ± 1.6 C | 3.7 ± 1.8 CD |
| *Pseudomonas* sp. KM2404 | 4.4 ± 0.9 BCD | 4.8 ± 0.8 B | NE | NE |
| *P. protegens* KY4410 | NE | NE | 2.4 ± 1.2 D | 3.2 ± 1.1 D |
| *P. protegens* TM1109 | 4.2 ± 0.8 D | 4.2 ± 0.8 C | 2.0 ± 1.2 D | 3.1 ± 1.2 D |
| *S. plymuthica* TM4307-1 | 6.6 ± 0.9 A | 6.3 ± 1.0 A | NE | NE |

a. CK1, soil not amended with *R. solani* inoculum and sown to non-treated seed; CK2, soil amended with *R. solani* inoculum and sown to methyl cellulose coated seed; and CK3, soil amended with oat-kernel inoculum and sown to non-treated seed.

Severity of Rhizoctonia root rot was evaluated on a scale of 0–8. Experiments 1 and 2 were conducted in Quincy virgin soil that had been pasteurized (60°C, 30 min); experiments 3 and 4 were conducted in raw Quincy virgin soil. Means in the same column followed by the same letter are not significantly different at *P* = 0.05 according to Fisher’s protected least significant difference test (LSD) or Kruskal–Wallis all pairwise comparison test.

Exp, experiment; NE, not evaluated.

Table 7. Suppression of *G. graminis var. tritici* ARS-A1 in Quincy virgin soil in a growth chamber tube assays.

| Treatment | Pasteurized soil | Raw soil |
|-----------|-----------------|----------|
|           | Exp 1 | Exp 2 | Exp 3 | Exp 4 |
| CK1       | 0.5 ± 1.4 | 0.0 ± 0.3 | 0.5 ± 1.4 | 0.0 ± 0.2 |
| CK2       | 5.2 ± 1.2 AB | 6.7 ± 0.8 A | 4.5 ± 0.9 ABC | 6.1 ± 1.0 AB |
| CK3       | 5.3 ± 1.0 AB | 6.6 ± 0.8 A | 4.7 ± 1.0 AB | 6.1 ± 1.0 A |
| *L. capsici* TM5405 | 3.8 ± 0.8 D | 5.9 ± 1.0 BC | 3.3 ± 0.7 E | 5.3 ± 1.4 BCD |
| *Pa. polymyxa* KM4410 | 5.0 ± 1.3 AB | 6.6 ± 0.9 A | 4.2 ± 0.9 BC | 5.9 ± 1.2 ABC |
| *P. chlororaphis* KY5406 | 4.0 ± 1.2 CD | 4.8 ± 1.2 D | 3.7 ± 1.2 DE | 4.4 ± 1.0 E |
| *Pseudomonas* sp. KM2404 | 3.9 ± 1.1 D | 5.4 ± 0.9 CD | 3.2 ± 0.8 E | 4.6 ± 1.3 DE |
| *P. protegens* TM1109 | 5.9 ± 1.1 A | 5.6 ± 0.9 A | 5.3 ± 1.6 A | 6.1 ± 1.4 AB |
| *S. plymuthica* TM4307-1 | 5.8 ± 1.3 A | 6.6 ± 0.9 A | 5.3 ± 1.6 A | 6.1 ± 1.4 AB |

a. CK1, Quincy virgin soil not amended with inoculum and sown to non-treated seed; CK2, soil amended with *G. graminis var. tritici* inoculum and sown to methyl cellulose coated seed; and CK3, soil amended with *G. graminis var. tritici* inoculum and sown to non-treated seed.

Severity of take-all was evaluated on a scale of 0–8. Experiments 1 and 2 were conducted in pasteurized Quincy virgin soil (60°C, 30 min); experiments 3–4 were conducted in raw Quincy virgin soil. Means in the same column followed by the same letter are not significantly different at *P* = 0.05 according to Fisher’s protected least significant difference test (LSD) or Kruskal–Wallis all pairwise comparison test.

Exp, experiment.

© 2014 The Authors. *Microbial Biotechnology* published by John Wiley & Sons Ltd and Society for Applied Microbiology.
biosynthesis of the broad-spectrum antibiotic 2,4-DAPG in *Pseudomonas* spp. and 2,4-DAPG producers are responsible for the natural suppression of take-all of wheat in the United States and black root rot of tobacco in soils of the Morens region of Switzerland (Weller *et al.*, 2007). 2,4-DAPG also induces systemic resistance in plants to pathogens (Weller *et al.*, 2012). *prnD* and *pltC* are genes that encode for the biosynthesis of pyrrolnitrin and pyoluteorin, antibiotics that are highly active against *Rhizoctonia* and *Pythium* species respectively (Loper *et al.*, 2007). The gene *acdS* encodes ACC deaminase, which promotes plant growth and development by decreasing ethylene levels in the plant. ACC deaminase-producing PGPR convert the ethylene precursor ACC to α-ketobutyrate and ammonia and can relieve stress resulting from pathogens, polyaromatic hydrocarbons, heavy metals, salt and drought (Glick *et al.*, 2007).

To validate the biocontrol activity of our strains, we used the *Rhizoctonia* root rot- and take-all-wheat pathosystems. Strains of *P. protegens*, *P. chlororaphis* subsp. *aureofaciens* and *L. capsici* provided the most consistent disease suppression and members of all three species are biocontrol agents of a broad range of diseases. For example, *P. protegens* PI-5 produces a wide arsenal of antibiotics and toxins and approximately 6% of its genome is devoted to the production of secondary metabolites (Paulsen *et al.*, 2005; Loper *et al.*, 2007). Strain CHA0, isolated from a tobacco black root rot suppressive soil, is another notable *P. protegens* with exceptional biocontrol activity (Ramette *et al.*, 2011). Our four *P. protegens* strains, TM1109, TY1502, TY1508 and KY4410, contained the three antibiotic biosynthesis genes, *phID*, *prnD* and *pltC*, and produced the antibiotics that are characteristic of Pf-5 and CHA0.

*Pseudomonas chlororaphis* subsp. *aureofaciens* contains *phzO* (Mavrodi *et al.*, 2010), and besides PCA, it also produces hydroxypHENazines, which have characteristic orange and red colours and differ from PCA in antibiotic activity (Mavrodi *et al.*, 2010). Notable examples of *P. chlororaphis* subsp. *aureofaciens* with biocontrol activity include strains 30–84 (Pierson and Thomashow, 1992) and TX-1 (Powell *et al.*, 2000). In *P. chlororaphis*, phenazine production also contributes to rhizosphere competence (Mavrodi *et al.*, 2006). *Lysobacter* sp. strains also have broad antifungal and antibacterial activities.

For example, Park and colleagues (2008) reported that *L. capsici* YC5194 

\[ \text{Inhibited Pythium ultimum, Colletotrichum gloeosporioides, Fusarium oxysporum, Botrytis cinerea, R. solani, Botryosphaeria dothidea and Bacillus subtilis.} \]

*Lysobacter* sp. SB-K88 (formerly *Stenotrophomonas*) (Nakayama *et al.*, 1999) produced the antibiotics xanthobaccins A, B and C, and xanthobaccin A was involved in the suppression of Pythium damping-off of sugar beet (Islam *et al.*, 2005).

To test our strains for the ability to directly promote plant growth, we used the canola root elongation assay, a model system commonly employed to demonstrate direct growth-promoting activity by PGPR (Patten and Glick, 2002). The most consistent and effective growth promoters were strains of *Delftia* and *Pseudomonas* that produced IAA and, in the case of *P. fluorescens* and *P. brassicacearum* strain Am3, with ACC deaminase activity, promoted root elongation of *Arabidopsis thaliana*, *Brassica napus* (Belimov *et al.*, 2007; Glick *et al.*, 2007). Members of these two species also are biocontrol agents (Loper *et al.*, 2012). *Pseudomonas brassicacearum* has commonly been isolated from *Arabidopsis thaliana*, *Brassica napus* (Achouak *et al.*, 2000) and wheat (Ross *et al.*, 2000). *Pseudomonas brassicacearum* strain Am3, with ACC deaminase activity, promoted root elongation of Indian mustard and increased root and shoot biomass of rape and pea (Belimov *et al.*, 2001; Safronova *et al.*, 2006). A mutation in *acdS* resulted in a loss of growth-promoting activity in *P. brassicacearum* (Belimov *et al.*, 2007).

We tested 10 strains for tolerance to heavy metals and high levels of salt by growing them in broth supplemented

### Table 8. Minimum inhibitory concentrations (MICs) of heavy metals and salt for selected strains.*

| Isolate          | Type of soil contamination strain from | Cd²⁺ (mM) | Cu²⁺ (mM) | NaCl (%) |
|------------------|----------------------------------------|-----------|-----------|----------|
| *D. tsuruhatensis* TM3205 | Petroleum products                      | 1.3       | 4.4       | 7        |
| *Pseudomonas* sp. KM3113 | Petroleum products                      | 1         | 4.4       | 6        |
| *Pseudomonas* sp. TY1205 | Heavy metals                            | 1         | 4.8       | 5.5      |
| *Pseudomonas* sp. KY5404 | Excessive salinization                   | 1         | 5         | 5.5      |
| *Pseudomonas* sp. TY1210 | Heavy metals                            | 1         | 4         | 6        |
| *P. chlororaphis* KYS406 | Excessive salinization                   | 1.5       | 5         | 5.5      |
| *Pseudomonas* sp. TY3101 | Petroleum products                      | 1.5       | 4         | 5.5      |
| *Pseudomonas* sp. KY3201 | Petroleum products                      | 1         | 4         | 7.5      |
| *P. protegens* TM1109 | Heavy metals                            | 1.5       | 5         | 7        |
| *P. protegens* KY4410 | Excessive salinization                   | 1.5       | 4.8       | 7        |

* MICs were determined in Luria–Bertani (LB) broth amended with CuCl₂ to give a range of concentrations from 3 to 6 mM; CdCl₂ to give a range of concentrations from 0.4 to 2.0 mM; and NaCl to give concentrations ranging from 0% to 8%.

© 2014 The Authors. *Microbial Biotechnology* published by John Wiley & Sons Ltd and Society for Applied Microbiology, *Microbial Biotechnology, 8*, 404–418
with CdCl₂, CuCl₂ and NaCl. Unfortunately, there are no records of how long the sampled fields have been polluted but discussions with local inhabitants indicated that some fields may have been in that condition for decades. Not surprising, all of the PGPR strains tested tolerated high levels of salt and demonstrated MICs to Cu²⁺ and Cd²⁺ on par or higher than those previously reported for strains stored in 60% glycerol at -80°C. The fungal wheat root pathogen S. sclerotiorum (Lib.) de Bary isolate MGSCF180002 (from rapeseed) were grown at room temperature on one-fifth-strength homemade potato dextrose agar (1/5 PDA) (Yang et al., 2011). In vitro inhibition assays of pathogens by PGPR were conducted on full-strength PDA.

Sample collection

Soil was sampled from five polluted sites in China: farmland contaminated with heavy metals in Daye County, Hubei Province; a seaside field at a recalcitrant organic sewage outlet in Gold Sand Bay Industrial Park, Hukou County, Jiangxi Province; farmland beside a petroleum products sewage outlet in Jinmen City, Hubei Province; and saline-alkali fields on two separate seaside farms in Qidong City (near Dongyuan and Yinyang respectively), Jiangsu Province. At each site, plants were dug to a depth of 30 cm from five random locations. Two kilograms of soil surrounding the roots of plants from each location was placed in plastic bags, transported to the laboratory and stored at 4°C until processed as described below.

Wheat and rapeseed greenhouse cycling assays in contaminated soils

Soil from each location was sieved, put into three square pots (6.5 cm high × 7 cm wide) and 10 ml of a metalaxyl (Syngenta, Wilmington, DE, USA) solution (75 mg l⁻¹) was added to prevent damping-off disease caused by indigenous Pythium spp. Each pot served as a replicate. Six surface-sterilized seeds of wheat (cv. Zhengmai 9023) or rapeseed (cv. Xiangyou 571) were pre-germinated for 48 h, sown in each pot and then covered with a 1.5 cm-thick layer of sterile vermiculite. Each pot was then given 30 ml of water. The pots were covered with a plastic bag for 4 days and incubated in a greenhouse (15–18°C; dark/light cycle, 12 h). Each pot received 30 ml of water two times weekly and diluted (1:3, vol/vol) Hoagland’s solution once weekly. After 3 weeks, the seedlings were removed from the pots, roots were excised and soil and roots from all of the pots of the same treatment were mixed and added back to the pots and sown as before with wheat or rapeseed to begin a second growth cycle. At the end of the third growth cycle, roots from two randomly chosen plants from each pot were selected for enumeration of populations of total cultivable aerobic, heterotrophic rhizobacteria (total bacteria) by the end-point dilution assay as described below.

Experimental procedures

Bacterial strains, soilborne pathogens and growth media

Bacterial strains used in this study are listed in Table 2. Pseudomonas isolates were grown in KMB or one-third-strength KMB (1/3 × KMB) agar or broth at 28°C as described by Mavrodi and colleagues (2012). Other strains were grown in TSA or tryptic soy broth (TSB) or these media at one-tenth strength (1/10 × TSB or TSA) (Mavrodi et al., 2012) at 28°C. When needed to inhibit fungi, these media were supplemented with cycloheximide (100 μg ml⁻¹). All strains were stored in 60% glycerol at -80°C. The fungal wheat root pathogen Gaeumannomyces graminis (Sacc.) von Arx and Olivier var. tritici Walker isolate ARS-A1 (Yang et al., 2011) and R. solani Kühn [teleomorph Thanatephorus cucumeris (Frank) Donk] AG-8 isolate C-1 (Huang et al., 2004), and the rapeseed/canola pathogen S. sclerotiorum (Lib.) de Bary isolate MGSCF180002 (from rapeseed) were grown at room temperature on one-fifth-strength homemade potato dextrose agar (1/5 × PDA) (Yang et al., 2011). In vitro inhibition assays of pathogens by PGPR were conducted on full-strength PDA.

Enumeration and isolation of rhizosphere bacteria

Roots with adhering rhizosphere soil of two plants from the same pot were excised and placed in a 50 ml screw-cap tube with 10 ml of sterile distilled water, vortexed (1 min) and sonicated in an ultrasonic cleaner (1 min) to dislodge rhizobacteria (Mavrodi et al., 2012). The dilution endpoint assay with 1/10 × TSB plus cycloheximide was used to determine the population density of total bacteria as described by
Mavrodi and colleagues (2012). The 96-well microplates were incubated at room temperature in the dark, and after 72 h, each well was scored for growth. At the end of the 3rd growth cycle, rhizobacteria were isolated by dilution plating of the root washings onto \( \frac{1}{10} \times \) KMB agar and \( \frac{1}{10} \times \) TSA, each with cycloheximide. Plates were incubated at room temperature for 2 to 3 days or until colonies were visible. Colonies of different morphotypes were re-streaked onto \( \frac{1}{10} \times \) KMB and \( \frac{1}{10} \times \) TSA plates and stored in 60% glycerol at \(-80^\circ C\).

**Pathogen inhibition in vitro**

Inhibition of soilborne pathogens by rhizobacteria was tested on PDA by two approaches. In the first, aliquots (20 \( \mu l \)) from overnight broth cultures were introduced into a hole in the agar cut with a 5 mm cork borer, 1 cm from the edge of the Petri dish. A 5 mm plug from a culture of *R. solani* AG-8 or *S. sclerotiorum* grown on \( \frac{1}{10} \times \) PDA was placed in the centre of the plate. Plates were incubated at 28°C and scored after 4 days by measuring the distance between the edges of the bacteria and the fungal mycelium. Four isolates were tested on each plate and each isolate was tested three times. In a second approach, aliquots (2 \( \mu l \)) from overnight cultures were spotted twice, 1 cm from the edge of a plate of PDA, and a plug of *R. solani* AG-8 was placed in the centre. For tests against *G. graminis var. tritici*, the plug of the fungus was placed in the centre of the plate 24 h before the bacteria were spotted. The zone of inhibition was measured 5 days later. Each isolate was tested four times.

**Morphological and physiological characteristics of isolates**

Selected isolates were characterized based on morphological and biochemical tests as described by Zhao and He (2012). Morphological characteristics included colony morphology, pigmentation, cell shape and Gram stain reaction. Biochemical and physiological characterization included tests for starch hydrolysis, gelatin liquefaction, catalase production, pectin decomposition, indole production and production of phenazine-1-carboxylic acid, 2,4-DAPG, pyrrolnitrin and pyoluteorin. Cellulase activity was determined on carboxymethylcellulose (CMC) medium (per 1 l: peptone, 10 g; yeast extract, 10 g; sodium CMC, 10 g; NaCl, 5 g; KH\(_2\)PO\(_4\), 1 g; agar, 15 g; pH 7.0). A 10 \( \mu l \) aliquot of an overnight bacterial culture was spotted on sterile filter paper (5 mm diameter, 0.5 mm thick) on the surface of CMC agar. To visualize the culture was spotted on sterile filter paper (5 mm diameter, 15 ml) were macerated with a mortar and pestle in an equal volume of acetone. The acetone was evaporated in a hood overnight and the residual aqueous phase was extracted twice with equal volumes of ethyl acetate. The ethyl acetate was recovered, dried and the residue was dissolved in 50 \( \mu l \) of methanol. Samples (5 \( \mu l \)) of the extracts, standards of the four antibiotics, and positive control extracts from strains *P. protegens* Pf-5 and *P. brassicaevarum* Q8r1-96 were spotted on silica gel (GHLF) thin layer chromatography plates (Anateltech, Newark, DE, USA). The plates were developed with chloroform/acetone (9:1) and observed under UV light (254 nm).

**Amplification and sequencing of 16S rDNA, antibiotic biosynthesis and ACC deaminase genes**

Genomic DNA was extracted from strains using a QIAamp DNA Mini Kit (Qiagen, Valencia, CA, USA) and quantified by fluorometry using a DNA quantitation kit (Bio-Rad, Hercules, CA, USA). Genes encoding 16S rDNA, subunit B of DNA gyrase, sigma 70 factor of RNA polymerase and key antibiotic biosynthesis enzymes (i.e. *phzF*, *phlD*, *prnD* and *pltC*) were amplified by using published oligonucleotide primer sets (McSpadden Gardener et al., 2001; De Souza and Raaijmakers, 2003; Mulet et al., 2009; Mavrodi et al., 2010) and cycling conditions (Table 9). The ACC deaminase gene (*acds*) was targeted with primers F1936f-tail and F1938-tail. These primers were derived from primers F1936 and F1938 (Blaha et al., 2006) by adding to their 5′-ends sequences GCTCCCTACTCTGTACCTATC and CTGTGCG TCTGGGCTTGC, respectively, for direct sequencing of *acds* amplicons with primers Tail1 and Tail2 (Table 9). Amplification conditions for *acds* included initial denaturation at 94°C for 1 min, followed by 10 cycles of 94°C for 30 s, 53°C for 20 s and 72°C for 45 s, followed by 25 more cycles with primer annealing temperature of 60°C. All amplifications were performed with a PTC-200 gradient thermal cycler (Bio-Rad) using GoTaq DNA polymerase (Promega, Madison, WI, USA). The amplifications were carried out in 25 \( \mu l \) reactions that contained 100 ng of DNA; 1 \( \times \) GoTaq Flexi buffer, 200 \( \mu M \) dNTPs, 1.5 mM MgCl\(_2\), 20 pmol of each primer pair and 1.2 U of GoTaq DNA polymerase (Promega). Positive controls included DNA from *P. chlororaphis* subsp. aureofaciens TX-1 (*phzF*), *P. brassicaevarum* Q8r1-96 (*phlD*, *acds*) and *P. protegens* Pf-5 (*phlD*, *prnD*, *pltC*). Amplification products were cleaned with QIAquick PCR purification spin columns (Qiagen) and sequenced directly with a BigDye Terminator v. 3.1 cycle sequencing kit (Applied Biosystems, Foster City, CA, USA) at ELIM Biopharmaceuticals (Hayward, CA, USA).

**Sequence and phylogenetic analysis**

Sequence data were assembled and analysed using Geneious Pro v. 7.1.5 (Biomatters, Auckland, New Zealand). The 16S rDNA similarity searches and taxa assignments were carried out using the Classifier and SeqMatch tools available through the Ribosomal Database Project (http:// rdp.cme.msu.edu). For antibiotic, ACC deaminase, *gyrB* and *ropD* gene sequences, database searches for similar DNA and protein sequences were carried out with NCBI’s BLAST network service (http://www.ncbi.nlm.nih.gov/BLAST). Sequences were aligned and neighbour-joining
The Tail1/Tail2 primer set was used for direct sequencing of plt acids phz phl.

Table 9. Target genes and PCR and primers used in this study.

| Gene  | Primer          | Sequence (5’-3’)                | Annealing temp (°C) | Putative gene function          | Amplicon (bp) | References                     |
|-------|-----------------|---------------------------------|---------------------|---------------------------------|---------------|--------------------------------|
| 16S   | 8F              | AGAGTTTTGATCTGGCTGTCAG          | 55                  | Small-subunit rRNA              | 1500          | Weisburg et al., 1991          |
|       | 1492R           | TACGGHTACCTGTGATTACCTT          |                     |                                 |               |                                |
| gyrB  | Up-1G-          | YGGCAGGGCGGATAGCTGAGGA          | 60                  | DNA gyrase subunit B            | 1208          | Mavrodi et al., 2010           |
|       | Up-2G-          | CCGCCGGATGCTGAAAGTGAAGAAGAAGTC |                     |                                 |               |                                |
| gyrBCh1| F1936f-tail     | CCGCCGGATGCTGAAAGTGAAGAAGAAGTC | 57                  | DNA gyrase subunit B            | 1208          | This study                     |
| rpoD  | PsEG30F         | CTGTCGCTCTGGCTGTC              | 60                  | Sigma 70 factor of RNA polymerase| 760           | Mulet et al., 2009             |
|       | PsEG790R        | CCGCCGGATGCTGAAAGTGAAGAAGAAGTC |                     |                                 |               |                                |
| phD   | B2BF            | ACCACCGGAGCAGCTTGTTTTAGGCC      | 66.5                | 2,4-DAPG-specific type III PKS  | 629           | McSpadden Gardener et al., 2001|
|       | BPR4            | CCGCCGGATGCTGAAAGAAGAAGAAGAAGTC|                     |                                 |               |                                |
| phF   | Ps_up 1         | ATCTTCAACCGGTCGGCTACCG         | 57                  | Phenazine biosynthesis enzyme    | 427           | Mavrodi et al., 2010           |
|       | Ps_low 1        | CCRTAGGCGGTCGGAAC              |                     |                                 |               |                                |
| pmD   | PRND1           | GGCGGGCGGCGGTTGATGGA           | 65                  | Pyrrolnitrin biosynthesis enzyme| 786           | De Souza and Raaijmakers, 2003 |
|       | PRND2           | YCCCCGCGGCGGCGGCTGTCGGTG       |                     |                                 | 438           | De Souza and Raaijmakers, 2003 |
| plC   | PLTC1           | AACAAGTGCGGCGGCTACGAGBGCC      | 58                  | Type I PKS                      | 438           | De Souza and Raaijmakers, 2003 |
|       | PLTC2           | AAGCGCGGACATCGAAGAAGCAAGCG     |                     |                                 |               |                                |
| acdS  | F1936f-tail     | GCTCCTAATGTCGATCTAGCGTAAGC     | 50                  | ACC deaminase-encoding gene      | 792           | This study                     |
|       | F1938r-tail     | CAGATGCCAGGAGTTCAGATTGCAGC     |                     |                                 |               |                                |
|       | Tail1           | GCTCCTAATGTCGATCTAGCGTAAGC     | 60                  | See footnote                    |               | Nikolic et al., 2011           |
|       | Tail2           | CAGATGCCAGGAGTTCAGATTGCAGC     |                     |                                 |               |                                |

The Tail1/Tail2 primer set was used for direct sequencing of acdS amplicons generated with primers F1936f-tail/F1938r-tail.

Preparation of barley and oat grain inoculum

Inocula of G. graminis var. tritici and R. solani AG-8 were prepared as described by Yang and colleagues (2011). Briefly, oat or barley grains (250 g) and water (350 ml) were combined in a flask and autoclaved on each of two consecutive days. Sterilized grains were inoculated with pieces of agar cut from 1/4 × PDA plates of R. solani or G. graminis var. tritici. After 21 days at room temperature, the colonized grains were tested for contamination, dried under a stream of sterile air and stored at 4°C. Prior to use, inoculum was fragmented and sieved into sizes, with particles of 0.25–0.5 mm added to soil.

Bacterial treatment of wheat seed

For biocontrol studies, seeds were coated with bacteria by methods similar to those of Yang and colleagues (2011). Strains were inoculated onto plates of KMB or TSA and incubated at room temperature for 24–48 h. Bacteria were scraped into 1.0 ml H2O, washed by centrifugation (twice for 3 min at 14 000 r.p.m.) and then suspended in H2O. The concentration of the cell suspension was adjusted based on optical density (600 nm), and an aliquot was mixed with a 2% solution of methylene blue (MC) and deionized H2O. Wheat (130 seeds) was added into the mixture, shaken for 3 min and seeds were dried under a stream of sterile air. To determine the final cfus per seed, 10 seeds were placed in a tube with 10 ml of sterile water, vortexed (1 min) and sonicated (1 min). Population size as determined by the end-point dilution assay ranged from 10^4 to 10^7 cfu seed^-1.

Biocontrol activity of rhizobacteria

The biocontrol activity of strains was determined by several approaches, and against both Rhizoctonia root rot and take-all, in autoclaved, pasteurized (60°C, 30 min) and raw soil. For pot experiments, soil from a farm at Huazhong Agricultural University, Wuhan, China, was sieved, autoclaved and amended with barley grain inoculum of R. solani (1%, w/w). Each plastic pot (10 cm deep × 6 cm wide) was filled with a 5 cm-thick layer of sterile vermiculite followed by 8 g of infested soil. Four treated wheat seeds (cv. Yangmai 158) were sown on the surface of the soil and covered with a 1.5 cm layer of sterile vermiculite. Controls included: control 1 (CK1), soil not amended with inoculum and sown to non-treated seeds; control 2 (CK2), soil amended with inoculum and sown to seed coated with MC; and control 3 (CK3), soil amended with inoculum and sown to non-treated seed. Each pot was given 10 ml of 2.5 mg ml^-1 metalaxyl (Syngenta Wilmington, DE, USA) to prevent Pythium damping-off. Pots were covered with a plastic sheet to reduce evaporation, incubated at room temperature (22°C) overnight and then transferred to a greenhouse (17 ± 5°C). The plastic was removed when the tips of the shoots were visible. Each pot was watered twice weekly with water and once with 1/4 × Hoagland’s solution (macro-elements only). Treatments were replicated five times and arranged in a randomized complete block design; each pot served as a replicate. After 3 to 4 weeks, the seedlings were removed from the pots.

© 2014 The Authors. Microbial Biotechnology published by John Wiley & Sons Ltd and Society for Applied Microbiology, Microbial Biotechnology, 8, 404–418
washed free of soil, and the plants were evaluated for disease severity on a scale of 0 to 8, where 0 = no lesions evident and 8 = seedling dead or nearly so (Huang et al., 2004). The length of the seedling shoot was measured.

Biocontrol studies of Rhizoctonia root rot and take-all were also conducted using the tube assay as described by Yang and colleagues (2011) with pasteurized or raw Shano sandy loam (Quincy virgin) from a non-cropped site near Quincy, WA, USA. Plastic tubes (2.5 cm diameter, 16.5 cm long) with a hole in the bottom were hung in plastic racks (200 per rack). Each tube had a cotton ball placed in the bottom, followed by a 6.5 cm-thick column of sterile vermiculite, and then soil (10 g), with or without inoculum of R. solani at 1.0 % (w/w) or G. graminis var. tritici at 0.7% (w/w). A 1.5 cm-thick layer of vermiculite was placed over the infested soil, followed by three bacteria-treated wheat seeds (cv. Louise) and finally another 1.5 cm-thick topping of vermiculite. Each tube then received 10 ml of water with metalaxyl. Racks of tubes were covered with plastic sheets until shoots emerged and were incubated in a growth chamber (15–18°C; dark/light cycle 12 h). Each cone was watered twice weekly with water and once weekly with 1/3 Hoagland's solution. Seedlings were harvested after 3–4 weeks and washed under a stream of water. The severity of Rhizoctonia root rot and take-all was evaluated on a scale of 0–8 (Yang et al., 2011).

Detection of IAA

IAA production was determined colorimetrically (Patten and Glick, 2002; Egamberdieva et al., 2008). Strains were incubated with shaking for 48 h at 28°C in KMB broth alone or supplemented with 500 μg per milliliter of tryptophan. Cultures were centrifuged at 13 000 r.p.m. for 10 min and a 1 ml aliquot of the supernatant was mixed vigorously with 4 ml of Saikowski reagent. The mixture was incubated at room temperature for 25 min and the absorbance of the pink color was measured spectrophotometrically at 540 nm. The concentration of IAA was determined by using a calibration curve of pure IAA (0, 20, 40, 60, 80 and 100 μg ml−1) as a standard. Readings of < 1 μg ml−1 per OD600 unit were considered as negative for IAA production.

Growth on ACC as a sole nitrogen source

Strains were tested for ability to grow in a defined medium supplemented with ACC as a sole nitrogen source as described by Penrose and Glick (2003). Strains were first cultured in 3 ml of sterile DF salts minimal medium (pH 7.2): per litre, KH2PO4, 4.0 g; Na2HPO4, 6.0 g; MgSO4 7H2O, 0.2 g; glucose, 2.0 g; gluconic acid, 2.0 g; trace elements (FeSO4 7H2O, 1 mg; H3BO3, 10 μg; MnSO4 H2O, 11.19 μg; ZnSO4 7H2O, 124.6 μg; CuSO4 5H2O, 78.22 μg; MoO3, 10 μg); and (NH4)2 SO4, 2.0 g as a nitrogen source. After 72 h of shaking at 28°C, a 5 μl aliquot was transferred to 3 ml sterile DF salts medium with 3.0 mM ACC instead of (NH4)2 SO4 as the source of nitrogen. Pseudomonas brassicacearum QBr1-96 was used as a positive control.

Canola root elongation assay

A gnotobiotic canola root elongation assay was used to determine the ability of strains to promote growth (Egamberdieva et al., 2008). An aliquot of an overnight culture was centrifuged, the cell pellet was washed twice and suspended in 1 ml of sterile 0.03 M MgSO4. The suspension was adjusted to an OD600 of 0.15. Surface-sterilized spring canola seeds (cv. InVigor 8440) were soaked for 1 h in 700 μl of a bacterial suspension. Depending on the isolate, the concentration ranged from 104 to 106 cfu ml−1. Growth pouches were made using 1-gallon Ziploc plastic bags and sterile germination paper soaked with 50 ml of 1/3 Hoagland's solution. Twenty-five seeds of a single bacterial treatment were distributed along the germination paper and the paper was placed inside of the bag. Individual bags were then hung vertically, covered with aluminium foil for 1–2 days and incubated at 20°C in a dark/light cycle of 12 h. The length of the primary roots was measured 5 days after sowing. Roots of seeds that failed to germinate 2 days after sowing were not measured. This study was conducted five times with three representative experiments shown.

Tolerance of bacteria to abiotic stress

The tolerance of selected strains to copper, cadmium and NaCl was determined in Luria–Bertani (LB) broth amended with CuCl2 to give concentrations from 3 to 6 mM; CdCl2 to give concentrations from 0.4 to 2.0 mM; and NaCl to give concentrations from 0% to 8%. Each well of a 96-well microplate was filled with 200 μl of LB supplemented with the heavy metal or salt solutions and inoculated with 2 μl of an overnight culture of each isolate (OD600 = 0.1). Each isolate at each concentration of heavy metal or salt was replicated eight times. The microplates were incubated at room temperature in the dark for 72 h and assayed spectrophotometrically at 600 nm using a model 680 microplate reader (Bio-Rad). An OD600 > 0.1 was scored as positive for growth. The MIC was defined as the lowest concentration that completely inhibited the growth of the isolate.

Statistical analysis

Comparisons of means of root disease ratings were performed using standard analysis of variance (ANOVA) followed by Kruskal–Wallis all pairwise comparisons (P ≤ 0.05) (Statistix 8.1, Analytical Software, Tallahassee, FL, USA). Mean comparisons of plant height and root length among treatments were determined by ANOVA followed by using either the Fisher’s protected least significant difference test (P = 0.05) or the Kruskal–Wallis all pairwise test (P = 0.05). Bacterial population sizes were log transformed before analysis.

Accession numbers

Nucleotide sequences were deposited in GeneBank under the following accession numbers: prIC, KC357589 through KC357592; pmD, KC693004 through KC693007; phiD, KC693000 through KC693003; acoD, KC430107 through KC430112; gyrB, KM030038 through KM030046; rpoD, KM030047 through KM030054; 16S rDNA, KM030056 through KM030064. The phzF sequence of strain KY5406 was deposited under accession number KC572125.
Acknowledgements

Mention of trade names or commercial products in this publication is solely for the purpose of providing specific information and does not imply recommendation or endorsement by the U.S. Department of Agriculture. USDA is an equal opportunity provider and employer.

Conflict of interest
None declared.

References

Achouak, W., Sutra, L., Heulin, T., Meyer, J.M., Fromin, N., Degraeve, S., et al. (2000) Pseudomonas brassicacearum sp. nov., and Pseudomonas thivervalensis sp. nov., two root-associated bacteria isolated from Brassica napus and Arabidopsis thaliana. Int J Syst Evol Micr 50: 9–18.

Altimira, F., Yanez, C., Bravo, G., Gonzalez, M., Rojas, L.A., and Seeger, M. (2012) Characterization of copper-resistant bacteria and bacterial communities from copper-polluted agricultural soils of central Chile. BMC Microbiol 12: 193.

Bais, H.P., Weir, T.L., Perry, L.G., Gilroy, S., and Vivanco, J.M. (2006) The role of root exudates in rhizosphere interactions with plants and other organisms. Annu Rev Plant Biol 57: 233–266.

Belimov, A.A., Safronova, V.I., Sergeyeva, T.A., Egorova, T.N., Matveyeva, V.A., Tsyganov, V.E., et al. (2001) Characterization of plant growth-promoting rhizobacteria isolated from polluted soils and containing 1-aminocyclopropane-1-carboxylate deaminase. Can Microbiol 47: 642–652.

Belimov, A.A., Dodd, I.C., Safronova, V.I., Hontzeas, N., and Davies, W.J. (2007) Pseudomonas brassicacearum strain Am3 containing 1-aminocyclopropane-1-carboxylate deaminase can show both pathogenic and growth-promoting properties in its interaction with tomato. J Exp Bot 58: 1485–1495.

Blaha, D., Prigent-Combaret, C., Mirza, M.S., and Moenne-Loccoz, Y. (2006) Phylogeny of the 1-aminocyclopropane-1-carboxylic acid deaminase-encoding gene accD in phyto-beneficial and pathogenic Proteobacteria and relation with strain biogeography. FEMS Microbiol Ecol 56: 455–470.

Cai, Q., Long, M., Zhu, M., Zhou, Q., Zhang, L., and Liu, J. (2009) Food chain transfer of cadmium and lead to cattle in a lead-zinc smelter in Guizhou, China. Environ Pollut 157: 3078–3082.

Cheng, S. (2003a) Effect of heavy metal on plants and resistance mechanisms. Environ Sci Pollut R 10: 256–264.

Cheng, S. (2003b) Heavy metal pollution in China: origin, pattern and control. Environ Sci Pollut R 10: 192–198.

Cook, R.J., Schilling, W.F., and Christensen, N.W. (2002) Rhizoctonia root rot and take-all of wheat in diverse direct-seed spring cropping systems. Can J Plant Pathol 24: 349–358.

De Souza, J.T., and Raaijmakers, J.M. (2003) Polymorphisms in the pmD and pltC genes from pyrrolnitrin and pyoluteorin-producing Pseudomonas and Burkholderia spp. FEMS Microbiol Ecol 43: 21–34.

Egamberdieva, D. (2011) Pseudomonas chlororaphis: a salt-tolerant bacterial inoculant for plant growth stimulation under saline soil condition. Acta Physiol Plant 34: 751–756.

Egamberdieva, D., and Kucharova, Z. (2009) Selection for root colonizing bacteria stimulating wheat growth in saline soils. Biol Fert Soils 45: 563–571.

Egamberdieva, D., Kamilova, F., Validov, S., Gafurova, L., Kucharova, Z., and Lugtenberg, B. (2008) High incidence of plant growth-stimulating bacteria associated with the rhizosphere of wheat grown on salinized soil in Uzbekistan. Environ Microbiol 10: 1–9.

Elmer, W.H. (2002) Influence of formononetin and NaCl on mycorrhizal colonization and fusarium crown and root rot of asparagus. Plant Dis 86: 1318–1324.

Ghose, T.K. (1987) Measurement of cellulase activities. Pure Appl Chem 59: 257–268.

Glick, B.R., Cheng, Z., Czarzy, J., and Duan, J. (2007) Promotion of plant growth by ACC deaminase-producing soil bacteria. Eur J Plant Pathol 119: 329–339.

Gremion, F., Chatzinotas, A., Kaufmann, K., von Sigler, W., and Harms, H. (2004) Impacts of heavy metal contamination and phytoremediation on a microbial community during a twelve-month microcosm experiment. FEMS Microbiol Ecol 48: 273–283.

Guo, Y., Gao, Z.Y., Deng, Y.M., Ma, Z.Z., and Yan, S. (2011) Environmental geochemistry of abandoned flotation tailing reservoir from the Tonglvshan Fe-Cu sulfide mine in Daye, Central China. B Environ Contam Tox 87: 91–95.

Haas, D., and Défago, G. (2005) Biological control of soil-borne pathogens by fluorescent pseudomonads. Nat Rev Microbiol 3: 307–319.

Hartmann, A., Schmid, M., Tuinen, D., and Berg, G. (2009) Plant-driven selection of microbes. Plant Soil 321: 235–257.

Huang, X.D., El-Alawai, W., Gurska, J., Glick, B.R., and Greenberg, B.M. (2005) A multi-process phytoremediation system for decontamination of persistent total petroleum hydro-carbons (TPHs) from soils. Microchem J 81: 139–147.

Huang, Z., Bonsall, R.F., Mavrodi, D.V., Weller, D.M., and Thomashow, L.S. (2004) Transformation of Pseudomonas fluorescens with genes for biosynthesis of phenazine-1-carboxylic acid improves biocontrol of Rhizoctonia root rot and in situ antibiotic production. FEMS Microbiol Ecol 49: 243–251.

Islam, M.T., Hashidoko, Y., Deora, A., Ito, T., and Tahara, S. (2005) Suppression of damping-off disease in host plants by the rhizoplane bacterium Lysobacter sp. strain SB-K88 is linked to plant colonization and antibiotics against soilborne Peronosporomycetes. Appl Environ Microbiol 71: 3786–3796.

Jing, Y., He, Z., and Yang, X. (2007) Role of soil rhizobacteria in phytoremediation of heavy metal contaminated soils. J Zhejiang Univ Sci B 8: 192–207.

Jones, D.T., Taylor, W.R., and Thornton, J.M. (1992) The rapid generation of mutation data matrices from protein sequences. Computer Appl Biosci 8: 275–282.

Kimura, M. (1980) A simple method for estimating evolutionary rates of base substitutions through comparative studies of nucleotide sequences. J Mol Evol 16: 111–120.

© 2014 The Authors. Microbial Biotechnology published by John Wiley & Sons Ltd and Society for Applied Microbiology, Microbial Biotechnology, 8, 404–418
Kraus, J., and Loper, J.E. (1992) Lack of evidence for a role of antifungal metabolite production by Pseudomonas fluorescens Pf-5 in biological control of Pythium damping-off of cucumber. *Phytopathology* **82**:264–271.

Li, W.C., Ye, Z.H., and Wong, M.H. (2007) Effects of bacteria on enhanced metal uptake of the Cd/Zn-hyperaccumulating plant Sedum alfredii. *J Exp Bot* **58**:4173–4182.

Loper, J.E., Kobayashi, D.Y., and Paulsen, I.T. (2007) The genomic sequence of *Pseudomonas fluorescens* Pf-5: insights into biological control. *Phytopathology* **97**:233–238.

Loper, J.E., Hassan, K.A., Mavrodi, D.V., Davis, E.W., Lim, C.K., Shaffer, B.T., et al. (2012) Comparative genomics of plant-associated *Pseudomonas* spp.: insights into diversity and inheritance of traits involved in multitrophic interactions. *PLoS Genet* **8**:e1002784.

Lugtenberg, B., and Kamilova, F. (2009) Plant-growth-promoting rhizobacteria. *Annu Rev Microbiol* **63**:541–556.

McSpadden Gardner, B.B.M., Mavrodi, D.V., Thomashow, L.S., and Weller, D.M. (2001) A rapid polymerase chain reaction-based assay characterizing rhizosphere population of 2,4-diacetylphloroglucinol-producing bacteria. *Phytopathology* **91**:44–54.

Mavrodi, D.V., Blankenfeldt, W., and Thomashow, L.S. (2006) Phenazine compounds in fluorescent *Pseudomonas* spp.: biosynthesis and regulation. *Annu Rev Phytopathol* **44**:417–445.

Mavrodi, D.V., Peever, T.L., Mavrodi, O.V., Parejko, J.A., Raaijmakers, J.M., Lemanceau, P., et al. (2010) Diversity and evolution of the phenazine biosynthesis pathway. *Appl Environ Microbiol* **76**:866–879.

Mavrodi, D.V., Mavrodi, O.V., Parejko, J.A., Bonsall, R.F., Kwak, Y.S., Paulitz, T.C., et al. (2012) Accumulation of the antibiotic phenazine-1-carboxylic acid in the rhizosphere of dryland cereals. *Appl Environ Microbiol* **78**:804–812.

Mazurier, S., Corberand, T., Lemanceau, P., and Raaijmakers, J.M. (2009) Phenazine antibiotics produced by fluorescent pseudomonads contribute to natural soil suppressiveness to Fusarium wilt. *ISME J* **3**:977–991.

Mitra, B., Ghosh, P., Henry, S.L., Mishra, J., Das, T.K., Ghosh, S., et al. (2004) Novel mode of resistance to *Fusarium* infection by a mild dose pre-exposure of cadmium in wheat. *Plant Physiol Bioch* **42**:781–787.

Mulet, M., Bennasar, A., Lalucat, J., and García-Valdés, E. (2009) An rpoD-based PCR procedure for the identification of *Pseudomonas* species and for their detection in environmental samples. *Mol Cell Probes* **23**:140–147.

Mulet, M., Lalucat, J., and García-Valdés, E. (2010) DNA sequence-based analysis of the *Pseudomonas* species. *Environ Microbiol* **12**:1513–1530.

Nakayama, T., Homma, Y., Hashidoko, Y., Mizutani, J., and Tahara, S. (1999) Possible role of xanthobactin produced by *Stenotrophomonas* sp. strain SB-K88 in suppression of sugar beet damping-off disease. *Appl Environ Microbiol* **65**:4334–4339.

Nikolic, B., Schwab, H., and Sessitsch, A. (2011) Metagenomic analysis of the 1-aminocyclopropane-1-carboxylate deaminase gene (acdS) operon of an uncultured bacterial endophyte colonizing *Solanum tuberosum* L. *Arch Microbiol* **193**:665–676.

Parida, A.K., and Das, A.B. (2005) Salt tolerance and salinity effects on plants: a review. *Ecotox Environ Safe* **60**:324–349.

Park, J.H., Kim, K., Aslam, Z., Jeon, C.O., and Chung, Y.R. (2008) *Lysobacter capsici* sp. nov., with antimicrobial activity, isolated from the rhizosphere of pepper, and emended description of the genus *Lysobacter*. *Int J Syst Evol Microb* **58**:387–392.

Patten, C.L., and Glick, B.R. (2002) Role of *Pseudomonas putida* indoleacetic acid in development of the host plant root system. *Appl Environ Microbiol* **68**:3795–3801.

Paulitz, T.C., Smiley, R.W., and Cook R.J. (2002) Insights into the prevalence and management of soliborne cereal pathogens under direct seeding in the Pacific Northwest, U.S.A. *Can J Plant Path* **24**:416–428.

Paulsen, I.T., Press, C., Ravel, J., Kobayashi, D.Y., Myers, G.S.A., Mavrodi, D.V., et al. (2005) Complete genome sequence of the plant commensal *Pseudomonas fluorescens* Pf-5: insights into the biological control of plant disease. *Nat Biotechnol* **23**:873–878.

Penrose, D.M., and Glick, B.R. (2003) Methods for isolating and characterizing ACC deaminase-containing plant growth-promoting rhizobacteria. *Physiol Plantarum* **118**:10–15.

Pierson, L.S., and Thomashow, L.S. (1992) Cloning and heterologous expression of the phenazine biosynthetic locus from *Pseudomonas aureofaciens* 30–84. *Mol Plant-Microbe In* **5**:330–339.

Pieterse, C.M.J., Zamhioudis, C., Berendsen, R.L., Weller, D.M., Van Wees, S.C.M., and Bakker, P.A.H.M. (2014) Induced systemic resistance by beneficial microbes. *Annu Rev Microbiol* **52**:347–375.

Poschenrieder, C., Tolrà, R., and Barceló, J. (2006) Can metals defend plants against biotic stress? *Trends Plant Sci* **11**:288–295.

Powell, J.F., Vargas, J.M., Nair, M.G., Detweiler, A.R., and Chandra, A. (2000) Management of dollar spot on creeping bentgrass with metabolites of *Pseudomonas aureofaciens* (TX-1). *Plant Dis* **84**:19–24.

Ramette, A., Frapolli, M., Fischer-Le Saux, M., Gruftaz, C., Meyer, J.-M., Défago, G., et al. (2011) *Pseudomonas protegens* sp. nov., widespread plant-protecting bacteria producing the biocontrol compounds 2,4-diacetylphloroglucinol and pyoluteorin. *Syst Appl Microbiol* **34**:180–188.

Reed, M.L.E., and Glick, B.R. (2005) Growth of canola (*Brassica napus*) in the presence of plant growth-promoting bacteria and either copper or polycyclic aromatic hydrocarbons. *Can J Microbiol* **51**:1061–1069.

Ross, I.L., Alami, Y., Harvey, P.R., Achouak, W., and Ryder, M.H. (2000) Genetic diversity and biological control activity of novel species of closely related pseudomonads isolated from wheat field soils in South Australia. *Appl Environ Microbiol* **66**:1609–1616.

Roubisova, T.V., and Bostock, R.M. (2009) Episodic abiotic stress as a potential contributing factor to onset and severity of disease caused by *Phytophthora ramorum* in *Rhododendron* and *Viburnum*. *Plant Dis* **93**:912–918.

Safrovnova, V.I., Stepanov, V.V., Engqvist, G.L., Aleksyev, Y.V., and Belimov, A.A. (2006) Root-associated bacteria containing 1-aminocyclopropane-1-carboxylate deaminase

© 2014 The Authors. *Microbial Biotechnology* published by John Wiley & Sons Ltd and Society for Applied Microbiology, *Microbial Biotechnology*, 8, 404–418
improve growth and nutrient uptake by pea genotypes cultivated in cadmium supplemented soil. Biol Fert Soils 42: 267–272.

Sandraa, R.A., Torvik, V., Enger, Ø., Daae, F.L., Castberg, T., and Hahn, D. (1999) Analysis of bacterial communities in heavy metal-contaminated soils at different levels of resolution. FEMS Microbiol Ecol 30: 237–251.

Shin, S.H., Lim, Y., Lee, S.E., Yang, N.W., and Rhee, J.H. (2011) CAS agar diffusion assay for the measurement of siderophores in biological fluids. Microbiol Meth 44: 89–95.

Tamura, K., Dudley, J., Nei, M., and Kumar, S. (2007) MEGA4: Molecular Evolutionary Genetics Analysis (MEGA) software version 4.0. Mol Biol Evol 24: 1569–1599.

Thomas, R., Mohamed, L., and Merroun, S.S. (2008) Interactions of Paenibacillus sp. and Sulfolobus acidocaldarius strains with U(VI). In Uranium, Mining and Hydrogeology. Merkel, B.J., and Haschne-Berger, A. (eds). Berlin, Germany: Springer. pp. 703–710.

Thomashow, L.S., and Weller, D.M. (1988) Role of a phenazine antibiotic from Pseudomonas fluorescens in biological control of Gaeumannomyces graminis var. tritici J Bacteriol 170: 3499–3508.

Triky-Dotan, S., Yermiyahu, U., Katan, J., and Gamliel, A. (2005) Development of crown and root rot disease of tomato under irrigation with saline water. Phytopathology 95: 1438–1444.

Wei, B., and Yang, L. (2010) A review of heavy metal contaminations in urban soils, urban road dusts and agricultural soils from China. Microchem J 94: 99–107.

Weisburg, W.G., Barns, S.M., Pelletier, D.A. and Lane, D.J. (1991) 16S ribosomal DNA amplification for phylogenetic study. J Bact 173: 697–703.

Weller, D.M. (2007) Pseudomonas biocontrol agents of soilborne pathogens: Looking back over 30 years. Phytopathology 97: 250–256.

Weller, D.M., Landa, B.B., Mavrodi, O.V., Schroeder, K.L., De La Fuente, L., Blouin Bankhead, S., et al. (2007) Role of 2,4-diacetylphloroglucinol-producing fluorescent Pseudomonas spp. in the defense of plant roots. Plant Biol 9: 4–20.

Weller, D.M., Mavrodi, D.V., van Pelt, J.A., Pieterse, C.M., van Loon, L.C., and Bakker, P.A. (2012) Induced systemic resistance in Arabidopsis thaliana against Pseudomonas syringae pv. tomato by 2,4-diacetylphloroglucinol-producing Pseudomonas fluorescens. Phytopathology 102: 403–412.

Yang, M., Mavrodi, D.V., Mavrodi, O.V., Bonsall, R.F., Parejko, J.A., Paulitz, T.C., et al. (2011) Biological control of take-all by fluorescent Pseudomonas spp. from Chinese wheat fields. Phytopathology 101: 1481–1491.

Zhao, B., and He, S. (2012) Microbiological Experiment. Beijing, China: Beijing Science Press.

Zhou, Y. (2013) Soil heavy metals and flourine pollution and health. In Soil Pollution and Physical Health. Zhou, Y. (ed.). Beijing, China: China Environmental Science Press, pp. 132–194.

Supporting information

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site:

Fig. S1. Neighbour-joining phylogeny inferred from concatenated housekeeping genes 16S, gyrB and rpoD. The concatenated housekeeping gene data set contained 2911 characters. Evolutionary distances were estimated using the Kimura two-parameter model of nucleotide substitution. Indels were ignored in the analysis and reproducibility of clades was assessed by bootstrap resampling with 1000 pseudoreplicates. The branch lengths are proportional to the amount of evolutionary change. The scale bars indicate substitution per site. Subgroups within the P. fluorescens lineage are defined according to Mulet and colleagues (2010).

Fig. S2. Neighbour-joining phylogeny inferred from data for aligned 578–bp fragments of phlD sequences of fluorescent Pseudomonas spp. Indels were ignored in the analysis, and evolutionary distances were estimated using the Kimura two-parameter model of nucleotide substitution. Bootstrap values greater than 60% are indicated at the nodes, and the branch lengths are proportional to the amount of evolutionary change. Clades indicated by capital letters correspond to BOX-PCR genotypes of 2,4-diacytethylphloroglucinol-producing Pseudomonas spp. as defined by De La Fuente and colleagues (2006).

Fig. S3. Neighbour-joining phylogeny inferred from data for aligned 390–bp fragments of pltC sequences of fluorescent Pseudomonas spp. Indels were ignored in the analysis, and evolutionary distances were estimated using the Kimura two-parameter model of nucleotide substitution. Bootstrap values greater than 60% are indicated at the nodes, and the branch lengths are proportional to the amount of evolutionary change.

Fig. S4. Neighbour-joining phylogeny inferred from data for aligned 745–bp fragments of prnD sequences of pyrrolnitrin-producing species. Indels were ignored in the analysis, and evolutionary distances were estimated using the Kimura two-parameter model of nucleotide substitution. Bootstrap values greater than 60% are indicated at the nodes, and the branch lengths are proportional to the amount of evolutionary change. Clades C1 through C2.4 correspond to distinct groups of pyrrolnitrin-producing bacteria as defined by Costa and colleagues (2009).

Fig. S5. Neighbour-joining phylogeny inferred from data for aligned 391–bp fragments of phzF sequences of fluorescent Pseudomonas spp. Indels were ignored in the analysis, and evolutionary distances were estimated using the Kimura two-parameter model of nucleotide substitution. Bootstrap values greater than 60% are indicated at the nodes, and the branch lengths are proportional to the amount of evolutionary change. Clades A1 through A5 correspond to distinct groups of phenazine-producing bacteria as defined by Mavrodi and colleagues (2010).

Fig. S6. Neighbour-joining phylogeny inferred from data for aligned 714–bp fragments of acdS sequences of ACC-deaminase-producing bacteria. Indels were ignored in the analysis, and evolutionary distances were estimated using the Kimura two-parameter model of nucleotide substitution. Bootstrap values greater than 60% are indicated at the nodes, and the branch lengths are proportional to the amount of evolutionary change. Clades A through C correspond to distinct groups of ACC-deaminase-producing bacteria as defined by Blaha and colleagues (2006).