Interplay between wheat cultivars, biocontrol pseudomonads and soil

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There is a significant potential to improve the plant-beneficial effects of root-colonizing pseudomonads by breeding wheat genotypes with a greater capacity to sustain interactions with these bacteria. However, the interaction between pseudomonads and crop plants at the cultivar level, as well as the conditions which favor the accumulation of beneficial microorganisms in the wheat rhizosphere, are largely unknown. Therefore, we characterized the three Swiss winter wheat cultivars Arina, Zinal and Cimetta for their ability to accumulate naturally occurring plant-beneficial pseudomonads in the rhizosphere. Cultivar performance was measured also by the ability to select for specific genotypes of 2,4-diacetylphloroglucinol (DAPG) producers in two different soils. Cultivar-specific differences were found, however, these were strongly influenced by the soil type. DGGE analysis of fragments of the DAPG biosynthetic gene phlD amplified from natural Pseudomonas spp. rhizosphere populations revealed that phlD diversity substantially varied between the two soils and that there was a cultivar-specific accumulation of certain phlD genotypes in one soil but not in the other. Furthermore the three cultivars were tested for their ability to benefit from Pseudomonas inoculants. Interestingly, Arina which was best protected against Pythium ultimum infection by inoculation with P. fluorescens biocontrol strain CHA0, was the cultivar which profited the least from the bacterial inoculant in terms of plant growth promotion in the absence of the pathogen. Knowledge gained
on the interactions between wheat cultivars, beneficial pseudomonads and soil types allows us to optimize cultivar-soil combinations for the promotion of growth through beneficial pseudomonads. Additionally, this information can be implemented by breeders into a new and unique breeding strategy for low-input and organic conditions.
INTRODUCTION

Improvement of plant fitness and yield by root-colonizing microorganisms is of special value in low-input or organic wheat production. Beneficial soil bacteria, such as certain *Pseudomonas* strains, are known to promote plant growth, which might help to circumvent potential negative consequences of low-input cropping systems such as the limited supply of nutrients and higher disease pressure. A wide range of traits in *Pseudomonas* spp. are responsible for plant-beneficial effects. Many pseudomonads are capable of solubilizing poorly soluble or insoluble mineral phosphates, thereby rendering this element available for the plant and promoting plant growth (25, 43). Root-colonizing pseudomonads are also able to indirectly promote plant growth by providing protection against plant diseases. The most important mechanisms for plant protection against attacking pathogens are the induction of systemic resistance in plants (3) and the direct suppression of soil-borne pathogens through the production of antimicrobial metabolites (16). The protection of wheat plants against *Gaeumannomyces graminis* var. *tritici* by naturally occurring pseudomonads in take-all decline soils is a well described phenomenon and highlights the importance of these bacteria in a successful and environmental friendly wheat production (53). Interestingly, in many naturally disease-suppressive soils a specific group of fluorescent pseudomonads is enriched, which is able to produce the antimicrobial compound 2,4-diacetylphloroglucinol (DAPG) (7, 38, 53). The production of the polyketide DAPG, which has broad-spectrum activity against
bacteria, plants, fungi and nematodes (8, 9, 21, 28, 33, 45), has been shown to be a key factor in the suppression of soil-borne plant diseases by various *Pseudomonas* biocontrol strains (16).

The degree of plant protection and plant growth promotion provided by root-colonizing pseudomonads is highly dependent on different environmental factors. For example, the expression of important biocontrol genes such as DAPG or HCN biosynthetic genes in the rhizosphere, is modulated by biotic factors such as fungi and other bacteria present in the rhizosphere and the secondary metabolites they release (6, 19, 27, 29, 32). Moreover, it has been observed that both the plant species and cultivar as well as the physiological stage of the plant, can influence the expression of biocontrol genes and the production of antimicrobial metabolites (4, 6, 19, 32, 35). In addition to the production of DAPG and other antimicrobial metabolites, efficient colonization of roots is a prerequisite for beneficial plant-*Pseudomonas* interactions. Root colonization is dependent not only upon specific characteristics of the bacterium itself, but also on root morphology and root exudates that vary between host plant species and even between cultivars of the same species (5, 34). The host species/cultivar also influences the abundance and diversity of naturally occurring pseudomonads (13). This has been shown in particular for DAPG-producing populations (4, 5, 26, 30, 36).

Wheat is a crop known to benefit strongly from naturally occurring DAPG-producing pseudomonad populations (52). It has been shown that the size and composition of DAPG-producing populations in the wheat rhizosphere and also the
amount of DAPG produced by these populations may vary substantially between different cultivars (4, 35). However, holistic studies which evaluate specific wheat cultivars for both their ability to benefit from plant growth-promoting pseudomonads and their influence on bacterial populations and production of biocontrol compounds are missing. A comprehensive characterization of different cultivars is needed in order to better understand which cultivars promote beneficial interactions with the pseudomonads. This knowledge has potential in future breeding strategies to be used for selection of new cultivars that optimally attract and respond to these bacteria.

In order to address this gap in knowledge, this study evaluated three Swiss winter wheat cultivars for several characteristics considered important in a successful wheat-pseudomonas interplay: (i) the ability to accumulate pseudomonads and \( \textit{phlD}^+ \) pseudomonads in two different Swiss soils; (ii) the ability to select for individual \( \textit{phlD}^+ \) genotypes in two different soils; (iii) the ability to benefit from the two model biocontrol strains, \textit{Pseudomonas fluorescens} strain CHA0 (a DAPG producer) and \textit{P. putida} KD (a DAPG non producer) in terms of direct plant growth promotion and disease suppression; and finally (iv) the level of biocontrol gene expression (DAPG-biosynthetic gene \( \textit{phlA} \)) in the rhizosphere.
MATERIALS AND METHODS

Wheat cultivars. Three different Swiss winter wheat cultivars were used in this study: Arina, Zinal, and Cimetta. They were bred in the Swiss breeding program at Agroscope ACW in Changins, Switzerland. Arina is one of the most prominent cultivars used in Switzerland. Arina and Zinal have a common ancestor, the cultivar Zenith. Arina is a first generation descendant and Zinal a second generation descendant of Zenith. Cimetta has no near relationship with the other cultivars.

Microorganisms used in this study. *P. fluorescens* CHA0 is a DAPG-producing strain isolated from the rhizosphere of tobacco grown in a Swiss soil naturally suppressive to *Thielaviopsis basicola* (48). *P. fluorescens* KD was isolated in China from the rhizosphere of wheat (46). Strain KD shows an outstanding biocontrol performance against both damping-off disease of cucumber caused by *Pythium ultimum* and crown and root rot of tomato caused by *Fusarium oxysporum* f. sp. *radicis-lycopersici* (FORL) (46), although it does not produce the biocontrol compound DAPG which is often associated with effective biocontrol in pseudomonads. Strain CHA0/pME7100 is a derivative of *P. fluorescens* CHA0 carrying a *phlA-gfp* reporter fusion (2). All strains were routinely cultivated at 27°C on King's B agar (KB) (23) and in Luria broth (LB) (44). For cultivation of CHA0/pME7100, tetracycline hydrochloride was added to the medium at 125 µg/ml.

*Pythium ultimum* strain 67-1 (obtained from Allelix Agriculture, Mississaugua, ON, Canada) was cultured on 1.5% malt agar plates (Oxoid, Basingstoke, England) at
20°C for 7 days. For disease suppression assays, a 0.7-cm plug of an actively growing
*P. ultimum* culture was transferred to a Petri-dish containing 25 g of autoclaved millet
seeds (Biofarm Kleindietwil, Switzerland) and 10 ml of autoclaved double-distilled
water. After 7 days of incubation at 20°C the mycelium-covered millet was sieved
and particles of 1 mm diameter were used to infest soil.

**Quantification of total pseudomonads and phlD+ pseudomonads on the roots**
**of different wheat cultivars grown in two different soils.** Soil samples were taken
in April 2005 from two fields in Switzerland. One site was located near Moudon
(Ecole Cantonale d’Agriculture de Grange-Verney, Moudon, Switzerland, 46°40´43”
N, 6°57´29”E); the other site was located at Delley (Delley Semences et Plantes SA,
Delley, Switzerland, 46°40´34”N, 6°48´22”E). On these fields, the performance of
different wheat cultivars was tested. Soil was collected from three plots each planted
with one of the winter wheat cultivar Arina, Zinal or Cimetta. For experiments
evaluating *Pseudomonas* accumulation and *phlD* diversity, pots (10 cm diameter, 420
cm³ volume) were filled with the collected soils. Into each pot five seeds of the
respective wheat cultivar were sown. Three replicates per cultivar and soil were
prepared. The plants were grown in a growth chamber with 70% relative humidity
and 12 h of light (160 µE/m²/s) at 18°C, followed by a 12-h dark period at 15°C. In
order to accumulate *Pseudomonas* spp. adapted to each cultivar, pots were re-sown
monthly to achieve continuous wheat cultures.

After seven successive cycles of one month each, the *Pseudomonas* populations
in three different locations were investigated: in non-rhizosphere soil, on the root
surface and in the root interior. To this end, 0.1-g samples of non-rhizosphere soil were collected from each pot and suspended in 10 ml 0.9% NaCl solution. One-ml samples of the resulting suspensions were collected for quantification of pseudomonads in the soil. In order to quantify pseudomonads on the roots, plants were harvested carefully, the roots were cut off and washed with tap water. Approximately 500 mg of roots from each sample were placed into a 50 ml Erlenmeyer flask containing 10 ml of a 0.9% sterile NaCl solution. The flasks were vigorously shaken for 20 min at 400 rpm. From the root washes, 1-ml samples were taken to estimate the numbers of *Pseudomonas* spp. closely adhering to the roots. Subsequently, the roots were removed from the flasks, washed with 0.9% NaCl solution, and placed for 45 s into 70% ethanol for surface-sterilization. The roots were washed again and then macerated in 5 ml 0.9% NaCl solution with a homogenizer (Homex 6, Bioreba AG, Reinach, Switzerland) for 45 s. From the resulting suspensions, samples were taken to estimate *Pseudomonas* populations inside the roots. In order to evaluate the efficiency of surface sterilization, subsamples of roots, which had their surface sterilized with ethanol, were not macerated, but shaken again in 0.9% sterile NaCl solution for 20 min at 400 rpm. Afterwards the number of pseudomonads was determined as described below. The obtained results showed that 99.99% of the pseudomonads on the root surface were removed by washing with NaCl solution and subsequent ethanol treatment.

From all samples (soil suspensions, root washes, and macerated root suspensions), aliquots of 20 µl were transferred to 96-well microtiter plates
containing 180 µl of the *Pseudomonas* selective medium KB+++ per well. The KB+++ medium consists of King’s medium B (23), amended with actidione (100 µg/ml), chloramphenicol (13 µg/ml) and ampicillin (40 µg/ml) (37). Tenfold serial dilutions up to a final dilution of 10^-6 were made in the microtiter plates. Four replicates were prepared per sample. The plates were incubated for 4 days at 24°C with continuous shaking at 150 rpm. Bacterial growth was estimated visually, and the highest dilution showing growth was used to calculate total *Pseudomonas* population size of a sample by the most probable number (MPN) technique (1, 14). Sterile glycerol was added to all wells to achieve a final concentration of 50% and microtiter plates were sealed with aluminium tape (Greiner Biotech, Germany) for storage at -80°C.

The proportion of rhizosphere pseudomonads carrying the *phlD* gene was examined using the MPN-PCR approach described by Ramette et al. (40). Briefly, 20 µl of bacterial suspensions from each dilution in the microplate were transferred to a 96-well PCR plate (Simport Plastics, Beloeil, Canada) and incubated for 10 min at 99°C. The heat-lysed suspensions were then frozen at -20°C for 30 min. After thawing, 4-µl aliquots of the suspensions were used as templates for a PCR with forward primer B2BF (25-mer 5’-ACCCACCGCAGCATCGTTTATGAGC -3’) and reverse primer BPR4 (26-mer 5’-CCGCCGGTATGGAAGATGAAAAAGTC -3’) in order to amplify a fragment of *phlD* (31). Primers were synthesized by MWG Biotech, Basel, Switzerland. Amplifications were carried out in 12-µl reaction mixtures containing 4 µl of lysed bacterial suspension, 1 x PCR buffer (Amersham
Pharmacia, Uppsala, Sweden), bovine serum albumin (0.5 g/l; Fluka, Buchs SG, Switzerland), 5% dimethyl sulfoxide (Fluka), 100 µM each of dATP, dCTP, dGTP and dTTP (Amersham Pharmacia), 0.40 µM of each primer and 1.4 U of Taq DNA polymerase (Amersham Pharmacia). The initial denaturation (2 min at 94°C) was followed by 35 cycles of 94°C for 30 s, 60°C for 30 s and 72°C for 60 s with a final extension at 72°C for 10 min. The terminal dilution showing phlD amplification was used to calculate the population size of phlD positive pseudomonads by MPN as described above.

**phlD amplification for DGGE analysis.** phlD diversity was studied in root-surface samples. In order to obtain satisfactory phlD amplification in certain samples, a nested PCR approach was needed. In the first step, phlD was amplified using primers B2BF and BPR4. In the second step, the 10-fold diluted product from the first PCR was used as template to perform a PCR with primers Phl2b1GC (21-mer, 5’-ACC GCA GCA TCG TGT ATG AGC-3’) containing a 40-bp GC clamp (5’-CGCCCCCGCCGCCGCCGCCCAGGCAGGCAGCCCCGCCCCGCCCCG-3’) and Mir1a (21-mer, 5’-GGA GTT CAT GAC CGC CTT GTC-3’).

PCR was carried out in 20-µl reactions containing 1 µl of DNA template, 2 µl of PCR buffer (New England Biolab, Beverly, MA), bovine serum albumine 100x (0.5 g/l (New England Biolab), 5% dimethyl sulfoxide (Fluka), 200 µM of each dATP, dCTP, dGTP, and dTTP (New England Biolab), 0.50 µM of each primer and 1.4 U of Taq DNA polymerase (New England Biolab). The initial denaturation (3 min at 94°C) was followed by 40 cycles of 94°C for 30 s, 60°C for 60 s and 72°C for
60 s with a final extension at 72°C for 10 min. PCR products were analyzed on a 1% agarose gel to check the presence of 670-bp bands for the first and of 571-bp bands for the second PCR.

**DGGE analysis.** DGGE analysis was performed using the DCode Universal Mutation Detection System (Bio-Rad, Hercules, CA). DGGE gels for both type of PCR products were cast using a double gradient ranging from 7% to 12% of acrylamide and from 30% to 60% denaturants (100% denaturant corresponds to 7 M urea and 40% deionized formamide). The marker consisted of individually amplified \( phlD \) fragments from ten reference *Pseudomonas* strains (Table 1, Fig. 1) belonging to different genotypic groups of *phlD*\(^+\) pseudomonads as described by Frapolli et al. (10). The samples were run for 14 h at 140 V in 1 \( \times \) TAE buffer (40 mM Tris base, 20 mM acetic acid, 1 mM EDTA, pH 8) preheated at 60°C. The gels were stained with SYBR Gold (Molecular Probes, Eugene, OR) for 1 h and visualized with a UV trans-illuminator.

**Sequencing of DGGE bands.** Dominant DGGE bands were characterized as follows. The central part of DGGE bands was cut out using sterile pipette tips. The gel pieces were then washed with 100 µl sterile double-distilled water at room temperature for 1 h and subsequently used as template for a 40-µl PCR reaction with primers Mir1a and Phl2b1GC with the conditions described above. The PCR products were run on a DGGE gel to check migration and presence of a single band and then sequenced using the Applied Biosystems 3130 Genetic Analyzer. Sequences were
edited using the Sequencher software (version 4.8, Gene Codes Corporation) and aligned with phlD sequences from known pseudomonad reference strains.

**Analysis of DGGE band patterns.** DGGE pictures were analyzed using the GelCompar II software (version 5.1, Applied Maths). Band patterns of the samples were normalized using the DGGE marker as reference and analyzed using a rolling background subtraction of 10% disk size. For cluster analysis, the Pearson correlation index, which takes the band intensities into account, and the Ward algorithm were used. To test for significant differences ($P \leq 0.05$) between the two soil types and the wheat cultivars, the similarity matrix obtained with the Pearson’s correlation coefficient was submitted to 1’000’000 unrestricted permutations using the Permttest software (24).

**Diversity of phlD+ pseudomonad community.** To compare the phlD+ *Pseudomonas* spp. community diversity between soils and wheat cultivars, the Shannon index ($H'$) was calculated from the band abundance values retrieved from the GelCompar II program. This diversity index takes into account both abundance and distribution (evenness) of band patterns present in a sample and is calculated using the formula $H' = -\sum_{i=1}^{n} p_i \ln p_i$, where $p_i$ is the relative abundance of a given DGGE band expressed in ratio of its intensity on the total DGGE band intensity present in the sample $i$.

**Ability of different wheat cultivars to profit from beneficial *P. fluorescens* strains.** Seeds of the winter wheat cultivars Arina, Zinal and Cimetta were surface-
disinfected for 15 min in 7% sodium hypochlorite (vol/vol), and then thoroughly rinsed with sterile double-distilled water. Seeds were pregerminated for 3 days on 0.85% water agar at 24°C in darkness. Plants were grown in artificial soil containing vermiculite and quartz sand of different fractions (20). The soil was moistened with double-distilled water (100 ml per 1 kg of soil). One-liter Erlenmeyer flasks were part-filled with 300 g of artificial soil, plugged with cotton wool stoppers and autoclaved for 30 min at 121°C. Overnight LB-broth cultures of CHA0 and KD were washed with 0.9% NaCl solution and diluted to give a final concentration of $10^7$ CFU/ml. To each flask, 6 ml of this bacterial suspension was added. Millet powder completely colonized by *P. ultimum* (see above) was mixed into the soil at a rate of 100 mg per flask. Control flasks without the pathogen received the same amount of autoclaved millet powder. Five seedlings were transferred to each flask, covered with soil and incubated in a growth chamber with 16 h of light at 18°C (160 µE/m²/s) followed by 8 h of darkness at 13°C. The flasks were arranged in a randomized complete block design. Three weeks later, plants were removed from the flasks and gently shaken to discard loosely adhering soil. Roots with tightly adhering soil (in this article defined as rhizosphere) were shaken in sterile 0.9% NaCl solution at 450 rpm for 20 min. Rhizosphere colonization with bacteria was determined from the resulting suspension by plating serial dilutions on KB agar. Plants were washed with tap water, blotted dry, and weighed. The experiment was performed twice with four replicate flasks (each flask containing five plants), per treatment. The obtained results are separated in two parts for presentation: Table 3 shows the disease suppression and
Table 4 shows the plant growth promotion in absence of the pathogen. In order to account for the growth differences between the individual wheat cultivars, fresh weights presented in Table 3 and Table 4 are expressed as percentage of the fresh weights of untreated control plants.

**phlA expression in P. fluorescens CHA0 on wheat roots.** The expression of the DAPG biosynthetic gene *phlA* by CHA0 on the roots of the three wheat cultivars was measured using a *phlA-gfp* reporter system and a flow cytometry-based approach as described in detail in De Werra et al. (6). Briefly, surface-sterilized seeds of the wheat cultivars Arina, Zinal, and Cimetta were germinated on soft agar for 2 days and placed into plant growth pouches (Mega International, West St-Paul, MN) at three seedlings per pouch. Growth pouches were amended with 15 ml of modified Knop plant nutrition solution (20). *P. fluorescens* suspensions containing $10^8$ cells/ml were prepared as described above and 1 ml was added to each seedling in the growth pouch. Plants were grown for 10 days in a growth chamber under the same conditions as described above. The experiment consisted of the following treatments: control without added bacteria, inoculation with wild type CHA0 (for setting the background for flow cytometric analysis) and inoculation with CHA0/pME7100 carrying a *phlA-gfp* reporter fusion. Six replications per treatment were made and the experiment was carried out twice. After a 10-day incubation period, plants were removed from the pouches. Roots of each pouch were placed into 7 ml of sterile distilled water and were vigorously shaken at 400 rpm for 20 min. From the resulting suspensions, samples of 2 ml were taken, transferred on ice, and immediately analyzed by
fluorescence-activated cell sorting (FACS) as described by De Werra et al. (6) using a FACSCalibur flow cytometer equipped with a 15 mW, air-cooled argon ion laser excitation light source (488 nm) (Becton Dickinson, San Jose, CA). After subtracting background fluorescence, the green fluorescence measured by FACS corresponding to \( \text{phlA-gfp} \) expression was calculated as mean fluorescence per expressing cell and per gram of analyzed roots (Table 5). Roots were assessed for fresh weight and bacterial root colonization was determined by plating serial dilutions of root suspensions on KB agar.

**Statistical analysis.** Statistical analyses were performed using the statistics program Systat, version 10.0 (Systat Inc., Evanston, IL). Table 2: For each isolation site, the data of the two soils were pooled and soil versus cultivar interactions were analyzed by a two-way analysis of variance (Table 2b). Subsequently means for one soil and one isolation site and also for one cultivar and isolation site were separated using Fisher’s protected least significant difference (LSD) test \((P \leq 0.05)\) (Table 2a). Tables 3-5: Results of independent repetitions over time (disease suppression assay and \( \text{phlA} \) expression assay) were first analyzed by a two-way analysis of variance to examine repetition versus treatment interactions. If this analysis revealed no significant repetition \( \times \) treatment interactions, individual repetitions over time were pooled for statistical analysis and means were separated using Fisher’s protected least significant difference (LSD) test \((P \leq 0.05)\).
RESULTS

Accumulation of total pseudomonads and \textit{phlD}+ pseudomonads by different wheat cultivars in two different soils. To assess the ability of different wheat cultivars to accumulate \textit{Pseudomonas} spp. in distinct soils, wheat plants were grown for seven cycles in pots filled with field soils of two different geographic origins. Pseudomonads were isolated from the soil, the root surface and from inside the roots. Between 5.37 log and 5.98 log CFU/g soil of pseudomonads were detected in the soil originating from Moudon after seven successive cycles (one month each) of wheat growing (Table 2). The \textit{Pseudomonas} spp. population was slightly smaller in the soil originating from Delley. In both soils, no significant differences between the cultivars Arina, Zinal, and Cimetta were found. A much higher \textit{Pseudomonas} concentration was found on the root surface ranging from 6.6 to 8.1 log CFU/g root fresh weight. When the plants were grown in the soil from Moudon no significant differences between cultivars were detected. In the Delley soil, however, the cultivar Arina was able to accumulate 15 to 30 times more pseudomonads on its roots than Cimetta and Zinal. Compared to the root surface, much less pseudomonads were found inside the roots of plants grown in the Moudon soil. Colonization ranged between 4.37 log and 4.52 log CFU/g root fresh weight with no significant differences between cultivars. The situation differed in the soil from Delley. The root interior of plants grown in the Delley soil was generally colonized by more pseudomonads (5.25 log to 7.09 log CFU/g root fresh weight) than that of plants grown in the Moudon soil. In this soil,
the cultivar Cimetta accumulated significantly higher *Pseudomonas* numbers inside the plant roots than the other two cultivars.

In the same experimental set-up, the accumulation of *Pseudomonas* spp. carrying the *phlD* gene was assessed (Table 2). In the Moudon soil the population size of *phlD*+ pseudomonads was below the detection limit of 3.1 log CFU/g soil. In the Delley soil the *phlD*+ population sizes were below the detection limit in two out of three replicates for all cultivars and below 4.3 log CFU/g soil in the third replicate. The numbers of *phlD*+ pseudomonads adhering to the roots differed between the wheat cultivars tested. In the Moudon soil, Cimetta (5.94 log CFU/g root) accumulated significantly higher numbers of *phlD*+ pseudomonads on its roots than Arina (4.40 log CFU/g root). In Delley soil, the colonization of roots by *phlD*+ pseudomonads was significantly superior on the cultivar Arina than on Zinal. The accumulation of *phlD*+ pseudomonads in the root interior differed completely between the two soils. In the Delley soil, wheat plants accumulated 4.99 log to 6.80 log CFU/g root fresh weight inside their roots, with the cultivar Cimetta accumulating significantly more *phlD*+ pseudomonads than the other two cultivars. In the Moudon soil, however, no *phlD*+ pseudomonads were detected inside the roots of two out of three replicates (all cultivars). In the third replicate, the colonization of the root interior was around one log unit above detection limit.

**Resolution power of the new primers used for DGGE analysis.** In a previous study (10), a DGGE approach was developed to analyze communities of *phlD*+ pseudomonads. However, for the soils analyzed in this study the previous approach...
was not effective enough since the amplification of *phlD* was insufficient in many samples (data not shown). In contrast, with the new nested PCR approach, good *phlD* amplification was obtained in all samples. Twelve reference strains (Table 1) that represent the known diversity of DAPG-producing pseudomonads were used to test the resolution power of the new nested method by DGGE. The approach presented here produced ten different migration patterns (Fig. 1, supplemental material) whereas the approach of Frapolli et al. (10) produced 11 different migration patterns.

**phlD** diversity of *Pseudomonas* populations enriched by different wheat cultivars in two different soils. DGGE analysis was performed to study the diversity of *phlD*⁺ pseudomonads in the rhizosphere of the wheat cultivars Arina, Zinal and Cimetua grown in Moudon and Delley soil (Fig. 1A). In the Delley soil, where only four migration patterns were found, the Shannon index was lower (*H*=0.82) than in the Moudon soil (*H*=1.19) where a total of 16 different migration patterns were detected. In each sample of the Delley soil, there were two bands, band 5 corresponding to the migration pattern of Q65c-80 and the more intense band 18 corresponding to the migration pattern of strain CHA0 (Fig. 1A). In Moudon soil samples, there were three prominent bands, i.e. the same two found in Delley soil (Bands 5 and 18) and one (Band 12) with the same migration pattern as strain Q37-87. The Moudon soil was characterized by the presence of other non-dominant bands, in contrast to the Delley soil where only very few additional weak bands were present. When comparing the three different wheat cultivars for the presence of the three dominant bands, it appeared that (i) in the Delley soil there were no differences
between the cultivars; bands 5 and 18 were always present, (ii) in Moudon soil Cimetta also harbored those two bands, (iii) in Moudon soil Zinal was characterized by the presence of band 5 and 12, and (iv) in Moudon soil Arina enriched band 12 and even stronger band 18. The community diversity found on the roots of Zinal was higher than that of the other two cultivars in both types of soil, with H’=1.55 in Moudon and H’=1.03 in Delley soil, whereas Cimetta and Arina, respectively, displayed a diversity of H’=1.19 and H’=0.86 in Moudon soil and of H’=0.79 and H’=0.65 in Delley soil. The DGGE fingerprints (Fig. 1A) were then subjected to Pierson cluster analysis and converted into a dendrogram (Fig. 1B) that shows a clear differentiation between the samples of Moudon and those of Delley soils and displayed two main clusters. The first cluster contained only samples from Moudon soil, whereas the second cluster contained all Delley samples and two Moudon samples (Arina Moudon 1 and Cimetta Moudon 1). The difference between soil types was found to be significant (P≤0.05) when submitted to the permutation test. In the Delley soil, phlD profiles from all three wheat cultivars and replicates were quite similar. In the Moudon soil, DGGE profiles from different wheat cultivars and different replicates of the same cultivar shared lower similarity compared to profiles from the Delley soil. However, as in the Delley soil, no statistically significant differences were found between the different wheat cultivars.

**Sequencing of dominant phlD bands found in Moudon and Delley soils.** The three DGGE bands corresponding to the dominant phlD genotypes were sequenced in at least four samples from different cultivars and soils and sequences were compared
to NCBI sequences of \textit{P. fluorescens} reference strains. Band number 5, with the same migration pattern as Q65c-80, had 100\% identity (based on 460 bp) with the sequences of reference strains S7-42 (GenBank Accession No. EF554345) and Q8r1-96 (GenBank Accession No. AF207693) except samples Cimetta Moudon 3 with 1 bp difference and Cimetta Deley 1 with 2 bp difference. Band number 12, with the same migration pattern as Q37-87, had 100\% identity (based on 472 bp) with the sequence of strain Q37-87 (GenBank Accession No. AY928641) except sample Zinal Moudon 2 (1 bp difference). The most dominant band, number 18, shared the same migration pattern and always had 100\% sequence identity (based on 437 bp) with CHA0 (GenBank Accession No. AJ278806).

\textbf{Ability of different wheat cultivars to profit from beneficial \textit{P. fluorescens} strains.} The potential of the two distinct \textit{P. fluorescens} strains CHA0 and KD to protect three different wheat cultivars against \textit{Pythium ultimum} was assessed in a gnotobiotic system. After 3 wk the total fresh weights of the untreated control plants were 3.04 g (Arina), 2.94 g (Zinal) and 2.75 g (Cimetta) and the root fresh weights were 1.44 g, 1.46 g and 1.42 g, respectively. \textit{P. ultimum} strongly reduced root weights (by 89 - 93\%) and total plant weights (by 66 - 72\%) of all cultivars (Table 3). \textit{P. fluorescens} KD was able to protect the plants only to a limited extent, with no differences between wheat cultivars. The root weights of plants inoculated with KD and \textit{P. ultimum} ranged between 15.9 and 16.9\% of those of non-inoculated control plants and were not significantly different (for two out of three cultivars) from those of plants inoculated with the pathogen alone. Total weights of plants protected by
KD, however, were significantly different from those inoculated with *Pythium* only and ranged between 46.6 and 51.3% of the weights of the respective controls. *P. fluorescens* CHA0 provided significantly better disease suppression than strain KD (Table 3). In presence of strain CHA0, root weights of plants growing in *Pythium*-infested soil were significantly enhanced and reached 41-61% of untreated controls. CHA0-treated pathogen-exposed plants reached between 66 and 83% of the total plant weight of untreated controls. Comparing the three wheat cultivars, strain CHA0 had the best protective effect on the cultivar Arina. In presence of the oomycete pathogen, the total weight of Arina treated with CHA0 was around 25% higher than those of Zinal and Cimetta.

Interestingly, in the absence of the pathogen, the other two cultivars profited more than Arina from the presence of CHA0 (Table 4). Here, *P. fluorescens* KD had no impact on total plant weight of any of the three tested wheat cultivars. Strain CHA0, however, significantly enhanced the growth of all cultivars. The plant growth-promoting effect on Arina, however, was quite small (13%), whereas it was much better for Zinal (30%) and Cimetta (27%).

The effect of the wheat cultivars on root colonization by plant-beneficial pseudomonads was also evaluated. KD proved generally to be a better colonizer than CHA0 (Table 5). Root colonization by strain KD ranged between 7.74 log and 8.30 log CFU/g fresh weight, while colonization by CHA0 ranged between 7.1 log and 7.82 log CFU/g. The roots of Zinal were significantly less colonized by strain KD than the roots of Arina and Cimetta, independent of the presence or absence of
Pythium. This cultivar-specific difference was not observed for root colonization by 
CHA0. In treatments inoculated with CHA0, the presence of Pythium had a 
significant enhancing effect (half a log unit) on the colonization of the roots of Arina 
and Zinal, but not of Cimetta (Table 5).

\textbf{\textit{phlA}}-expression on wheat roots. In order to test the influence of wheat cultivar 
on expression of an important biocontrol gene, Arina, Zinal and Cimetta were grown 
in growth pouches inoculated with CHA0 carrying a \textit{phlA-gfp} reporter fusion. The 
green fluorescence emitted by CHA0 cells, corresponding to the expression of \textit{phlA}, 
was measured by FACS. After 10 days, the relative fluorescence per \textit{phlA} expressing 
CHA0 cell did not differ among the cultivars and ranged between 88.6 on the roots of 
Cimetta and 98.1 on the roots of Arina. Similarly, no differences were found for total 
\textit{phlA}}-expression per root fresh weight. The obtained values were $1.24 \times 10^{11}$ (Arina), 
$9.34 \times 10^{10}$ (Zinal) and $9.02 \times 10^{10}$ (Cimetta) relative fluorescence per g roots.
DISCUSSION

There is great potential for exploitation of host response to beneficial microorganisms (reviewed in 54), which could allow breeders to select for traits that encourage interaction with plant-beneficial pseudomonads. This is of interest especially in breeding programs aimed at low-input agriculture. A quality cultivar should to be able to promote colonization and profit from naturally occurring beneficial organisms in the soil. A prerequisite for breeders to exploit cultivar-driven beneficial interactions with microorganisms is specific knowledge pertaining to the ability of each cultivar to promote or support beneficial interactions. This study represents the first attempt to gather such information for three Swiss winter wheat cultivars currently being used in a Swiss governmental breeding program.

The influence of these three Swiss wheat cultivars on accumulation of naturally occurring pseudomonads and $\textit{phlD}^+$ \textit{Pseudomonas} spp. was assessed in two soils after seven cycles of continuous wheat cultivation in pots. In the Moudon soil, no cultivar-specific differences in the accumulation of total pseudomonads in any of the tested compartments (soil, root surface, root interior) were detected. In contrast, the three cultivars accumulated different numbers of pseudomonads in the Delley soil. In both soils there were cultivar-specific differences in the accumulation of $\textit{phlD}^+$ pseudomonads on the roots. One cultivar, Cimetta, was superior in building up $\textit{phlD}^+$ populations in the Moudon soil, whereas another cultivar, Arina, was found to be superior in the Delley soil. These findings indicate the importance that soil has in the
interaction between the plant and pseudomonads. It has been shown previously that fluorescent pseudomonads play an important role in take-all decline soils or against soil-borne diseases in general. Raaijmakers et al. (39) reported a threshold of $10^5$ CFU/g of root of $phlD^+$ pseudomonads required for a successful suppression of *Gaeumannomyces graminis var. tritici*. In the Moudon soil, colonization of Arina by $phlD^+$ *Pseudomonas* spp. was slightly below this threshold level, in contrast to the Delley soil where Arina displayed the highest root colonization by $phlD^+$ pseudomonads of all three tested cultivars (Table 2). This indicates that a wheat cultivar might in one soil accumulate enough $phlD^+$ pseudomonads to be protected against soil-borne diseases, while the same cultivar in another soil may not reach the threshold level needed for protection. Gu and Mazzola (15) tested different wheat cultivars for the ability to build up suppressiveness against replant disease in apple. They observed that the severity of apple replant disease was reduced after the pre-culture of some wheat cultivars, but not others. As Cimetta proved to be a good attractor of $phlD^+$ pseudomonads in both tested soils, it might be worth to test this cultivar as a pre-culture with apple production or other perennial crops.

Interestingly, the soil type seems to have a greater impact on the accumulation of $phlD^+$ pseudomonads in the root interior than the cultivar. In the Delley soil, all cultivars accumulated high levels of $phlD^+$ pseudomonads inside the roots. In sharp contrast, in the Moudon soil, no $phlD^+$ pseudomonads could be detected inside the roots in two out of three replicates. The importance of soil in determining the population sizes of plant-beneficial pseudomonads has already been shown in
previous studies where soils suppressive to black root rot of tobacco were compared to non-suppressive soils (11, 41).

Soil was also found to be a strong determining factor in the diversity of $\text{phlD}^+$ pseudomonads on the different cultivars. Whereas in the Delley soil the wheat cultivar had no impact on diversity, in the Moudon soil individual cultivars selected for specific bacterial genotypes (Fig. 1). In general, the $\text{phlD}$ diversity was much higher in the Moudon soil (16 bands detected) than in the Delley soil where in the majority of the samples only two bands (number 5 and 18) were detected. The same two bands and additionally a third band (number 12) were the most prominent in the Moudon soil. Sequencing the three major bands 5, 12 and 18 detected by the DGGE analysis showed that they share 99.6 - 100% identity with the well described $P. \text{fluorescens}$ biocontrol reference strains S7-42 (41) and Q8r1-96 (38) (band 5), Q37-87 (22) (band 12) and CHA0 (band 18) which belong to the $\text{phlD}$ genotypes (clusters) 5, 4 and 9 as defined by cluster analysis of $\text{phlD}$ sequences by Frapolli et al. (10), and respectively to $\text{phlD}$ genotypes D, E and A as defined by De la Fuente et al. (5). We therefore suggest that in the Delley soil $\text{phlD}^+$ pseudomonads of genotypes 5 and 9 are predominant. The high intensity of the two bands on DGGE gels made with samples from Delley soil suggests that the population sizes of genotypes 5 and 9 in the Delley soil is higher than that of any genotype present in Moudon soil. The high abundance of genotypes 5 and 9 in the Delley soil may explain the similarity between the two soils in the rhizosphere population level of $\text{phlD}^+$ pseudomonads (Table 2) despite of the large difference in numbers of present genotypes.
Interestingly, the cultivar Arina seems to enrich *phlD* genotype 9 (Band 18) in both soils (Fig. 1A). We speculate that the cultivar’s ability to enrich the most dominant genotype in the Delley soil may have enabled this cultivar to accumulate the highest numbers of *phlD*+ pseudomonads and also of total pseudomonads in this soil (Table 2). We therefore suggest that a wheat cultivar with strong affinity for a *phlD* genotype that is present in abundance in the soil could have an advantage over other cultivars that do not possess the same affinity for that particular genotype. For an optimal cultivar-soil combination with respect to *phlD*+ pseudomonad enrichment it would therefore be necessary to know the preference of different wheat cultivars for individual *phlD*+ genotypes and, additionally, the most prominent *phlD* genotypes present in the soil of interest.

In this study, we demonstrated that the three wheat cultivars differ in their ability to profit from a bacterial inoculant in a gnotobiotic system (Tables 3 and 4). The cultivar Arina was significantly better protected against *Pythium* damping-off by *P. fluorescens* CHA0 than the other cultivars tested. Interestingly, the cultivar Arina enriched pseudomonads with *phlD* sequences identical to strain CHA0 in all samples from both Moudon and Delley soils (Fig. 1), suggesting that the better biocontrol activity of CHA0 on Arina could result from better adaptation of this cultivar to CHA0-like pseudomonads. However, CHA0 displayed neither a higher root colonization level (Table 5) nor a higher expression of the biocontrol gene *phlA* (in absence of the pathogen) on Arina compared to the other varieties. We therefore
suppose that in this simplified system further mechanisms of disease suppression
were responsible for the better protection of Arina against *P. ultimum.*

Although it has been shown, that *P. fluorescens* strain KD is a very good
biocontrol agent in the control of *Pythium* on cress and cucumber (42, 46) this strain
protected all tested wheat cultivars significantly less than CHA0 against *P. ultimum
(Table 3). Since biocontrol activity of strain KD is dependent on the presence of the
Type III secretion system (42), it might be possible that on monocotyledonous plants,
such as wheat, type III secretion system-dependent mechanisms of disease
suppression are of less importance in the control of *Pythium.*

To our best knowledge, we could show for the first time that the plant growth
promotion effects of a *Pseudomonas* inoculant, in absence of a pathogen, can be
wheat cultivar-dependent. Growth promotion in absence of the pathogen was lowest
on Arina. Either this cultivar is less responsive to plant growth promoting factors
secreted by CHA0 or it is more sensitive to potential phytotoxic metabolites produced
by this strain. Some secondary metabolites secreted by CHA0 (eg DAPG and
pyoluteorin) can negatively affect plant growth (28). Direct growth promotion was
lowest on Arina, however protection, in the presence of the pathogen, was found to
be the highest. This indicates a separation between the plant traits which favor plant-
bacteria interactions that protect against disease from those that promote growth. If
genetically controlled, it might thus be possible, through breeding, to combine these
traits in a single cultivar.
For many of the investigated traits, differences between wheat cultivars were found. It is not surprising that the cultivar Cimetta differs from Arina and Zinal since it originates from a distinct genetic background (Dario Fossati, Agroscope ACW Changins, Nyon, Switzerland, personal communication). However, also Zinal and Arina which share a common ancestor differ in several of the investigated traits. This genetic diversity could be exploited to breed a cultivar combining as many beneficial traits as possible. Quantitative trait locus (QTL) analyses in the progeny of crosses between these cultivars could be used to identify some genetic regions responsible for positive interactions with beneficial pseudomonads. Such an approach has been used to explore the genetic basis for interactions of tomato and the disease suppressive bacterium *Bacillus cereus* (47). Smith and coworkers (47) suggest that the discovery of a genetic basis for plant-beneficial interactions with microorganisms, opens up new opportunities to exploit natural genetic variation in host species. Furthermore, studies of this kind will enhance the current understanding of beneficial plant-microbe interactions and promote the development of ecologically sound strategies for disease control in agriculture.

Interactions between cultivars and soil might complicate efforts to exploit the genetic basis of beneficial wheat - bacteria interactions. Nevertheless, our study indicates that there is a large, unexplored potential for better exploitation of plant-bacterial interactions through the breeding of cultivars which are specifically adapted to accumulate and to profit from beneficial microbes.
ACKNOWLEDGMENTS

We gratefully acknowledge Dario Fossatti and Fabio Mascher, Agroscope ACW Changins, Switzerland for helpful discussions and providing seed material. This research was supported by the State Secretariat for Education and Research project C04.0200; COST action 860) and the Swiss National Science Foundation (project 3100A0-105881 and NRP-59 project 405940-115596).
REFERENCES


or conducive to *Thielaviopsis basicola*-mediated black root rot of tobacco. FEMS Microbiol. Ecol. **55**:369-381.


**Figure legends**

**Fig. 1**

A) Comparison of DGGE patterns of *phlD* fragments amplified from *Pseudomonas* populations isolated from the wheat cultivars Arina (A), Zinal (Z), or Cimetta (C) grown in Moudon (m) soil (lanes L1 to L9) or Delley (d) soil (lanes L10 to L18); M = marker composed from *P. fluorescens* strains: PITR2, Q65c-80, K94.37, Q37-87, P97.38, Q2-87, F96.27 and CHA0 belonging to the *phlD* cluster groups 6, 5, 2, 4, 8, 7, 8 and 9, respectively, according to Frapolli et al. (10) as well as F113. The three lanes per cultivar and soil represent DGGE analysis of root samples from three replicate pots. Arrows indicate major bands showing differences between soil type and wheat cultivar.

B) The hierarchical cluster analysis is based on Pearson's correlation index (band intensities) and the Ward algorithm according to Ward (51). A, C, Z = Arina, Cimetta, Zinal. M, m = Moudon soil. D, d = Delley soil. 1, 2, 3 = number of replicate.
Table 1. Fluorescent *Pseudomonas* strains used in this study.

<table>
<thead>
<tr>
<th>Strain</th>
<th>phlD genotype&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>CHA0</td>
<td>9</td>
<td>(48)</td>
</tr>
<tr>
<td>CHA0/pME7100 (<em>phlA-gfp</em>)</td>
<td></td>
<td>(2)</td>
</tr>
<tr>
<td>KD</td>
<td>-</td>
<td>(46)</td>
</tr>
<tr>
<td>Pf-5</td>
<td>9</td>
<td>(18)</td>
</tr>
<tr>
<td>Q2-87</td>
<td>7</td>
<td>(49)</td>
</tr>
<tr>
<td>Q65c-80</td>
<td>5</td>
<td>(17)</td>
</tr>
<tr>
<td>CM1'A2</td>
<td>5</td>
<td>(12)</td>
</tr>
<tr>
<td>F113</td>
<td>-</td>
<td>(9)</td>
</tr>
<tr>
<td>PITR2</td>
<td>6</td>
<td>(22)</td>
</tr>
<tr>
<td>P97.38</td>
<td>8</td>
<td>(50)</td>
</tr>
<tr>
<td>F96.27</td>
<td>8</td>
<td>(50)</td>
</tr>
<tr>
<td>K94.37</td>
<td>2</td>
<td>(50)</td>
</tr>
<tr>
<td>Q37.87</td>
<td>4</td>
<td>(22)</td>
</tr>
<tr>
<td>P12</td>
<td>1</td>
<td>(22)</td>
</tr>
</tbody>
</table>

<sup>a</sup> As determined by cluster analysis of partial *phlD* sequences by Frapolli *et al.*, (10).
Table 2. Accumulation of *Pseudomonas* spp. and *phlD*+ *Pseudomonas* spp. by three Swiss wheat cultivars in two different soils (Moudon and Delley) after seven cycles of wheat growth.

### a) Colonization

<table>
<thead>
<tr>
<th><em>Pseudomonas</em> spp. isolated from</th>
<th>Wheat cultivar</th>
<th>Colonization by</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td><em>Pseudomonas</em> spp.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>log (CFU/g)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Moudon</td>
<td>Delley</td>
<td>Moudon</td>
</tr>
<tr>
<td>Soil</td>
<td>Arina</td>
<td>5.53 a b</td>
<td>4.68 a</td>
<td>b.d.</td>
</tr>
<tr>
<td></td>
<td>Zinal</td>
<td>5.98 a</td>
<td>5.21 a</td>
<td>b.d.</td>
</tr>
<tr>
<td></td>
<td>Cimetta</td>
<td>5.37 a</td>
<td>4.46 a</td>
<td>b.d.</td>
</tr>
<tr>
<td>Root surface</td>
<td>Arina</td>
<td>6.78 a</td>
<td>8.11 a*</td>
<td>4.40 a</td>
</tr>
<tr>
<td></td>
<td>Zinal</td>
<td>6.75 a</td>
<td>6.93 b</td>
<td>5.49 ab</td>
</tr>
<tr>
<td></td>
<td>Cimetta</td>
<td>7.31 a</td>
<td>6.61 b</td>
<td>5.94 b</td>
</tr>
<tr>
<td>Root interior</td>
<td>Arina</td>
<td>4.52 a</td>
<td>6.07 a*</td>
<td>4.43, b.d., b.d.</td>
</tr>
<tr>
<td></td>
<td>Zinal</td>
<td>4.37 a</td>
<td>5.25 a*</td>
<td>3.93, b.d., b.d.</td>
</tr>
<tr>
<td></td>
<td>Cimetta</td>
<td>4.43 a</td>
<td>7.09 b*</td>
<td>4.40, b.d., b.d.</td>
</tr>
</tbody>
</table>

### b) ANOVA P-values

<table>
<thead>
<tr>
<th>Factor</th>
<th>Soil</th>
<th>Root interior</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total <em>Pseudomonas</em> spp.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cultivar</td>
<td>0.231</td>
<td>≤ 0.05</td>
</tr>
<tr>
<td>Soil</td>
<td>≤ 0.05</td>
<td>0.353</td>
</tr>
<tr>
<td>Cultivar × soil</td>
<td>0.984</td>
<td>≤ 0.05</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><em>phlD</em>+-<em>Pseudomonas</em> spp.</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Soil</td>
<td>0.642</td>
<td>0.248</td>
</tr>
<tr>
<td>Cultivar</td>
<td>0.382</td>
<td>0.679</td>
</tr>
<tr>
<td>Cultivar × soil</td>
<td>0.402</td>
<td>≤ 0.05</td>
</tr>
</tbody>
</table>
Wheat cultivars were grown in a growth chamber in pots filled with Moudon or Delley soil in seven successive cycles of one month each. Means of three replicates are presented. Values within the same column for the same isolation site, followed by the same letter are not significantly different according to Fishers protected LSD ($P \leq 0.05$).

In cases where in some of the replicates bacterial populations were below the detection limit (log 3.1 CFU/g for soil and root surface and 2.1 log CFU/g for root interior), results for individual replicates are presented.

Data for the two soils were pooled and analyzed by two-way analysis of variance (Systat, version 10.0, Systat Inc., Evanston, IL).

b.d. = below detection limit.

* = significant differences between soils for the same cultivar and isolation site according to Fishers protected LSD ($P \leq 0.05$).
Table 3. Protection of three Swiss wheat cultivars against the root pathogen *P. ultimum* by *P. fluorescens* strains CHA0 or KD under gnotobiotic conditions

<table>
<thead>
<tr>
<th>Cultivar</th>
<th>Treatment</th>
<th>Root fresh weight&lt;sup&gt;b&lt;/sup&gt; (%)</th>
<th>Plant fresh weight (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arina</td>
<td>Pu</td>
<td>11.5 cd&lt;sup&gt;c&lt;/sup&gt;</td>
<td>27.9 d</td>
</tr>
<tr>
<td></td>
<td>Pu + CHA0</td>
<td>60.7 a</td>
<td>82.9 a</td>
</tr>
<tr>
<td></td>
<td>Pu + KD</td>
<td>15.9 c</td>
<td>46.6 c</td>
</tr>
<tr>
<td>Zinal</td>
<td>Pu</td>
<td>9.6 cd</td>
<td>33.8 d</td>
</tr>
<tr>
<td></td>
<td>Pu + CHA0</td>
<td>40.9 b</td>
<td>65.7 b</td>
</tr>
<tr>
<td></td>
<td>Pu + KD</td>
<td>16.9 c</td>
<td>51.3 c</td>
</tr>
<tr>
<td>Cimetta</td>
<td>Pu</td>
<td>6.9 d</td>
<td>28.7 d</td>
</tr>
<tr>
<td></td>
<td>Pu + CHA0</td>
<td>47.0 b</td>
<td>67.2 b</td>
</tr>
<tr>
<td></td>
<td>Pu + KD</td>
<td>16.3 c</td>
<td>50.4 c</td>
</tr>
</tbody>
</table>

<sup>a</sup>Plants were inoculated with bacteria and on the same day infested with *P. ultimum* (Pu). After 3 wk of growth in artificial soil under sterile conditions plants were harvested and assessed for fresh weights.

<sup>b</sup>Plant fresh weights are expressed as percentage of weights of corresponding non-treated control plants (total plant weights: Arina = 3.04 g, Zinal = 2.94 g, Cimetta = 2.75 g; root weights: Arina = 1.44 g, Zinal = 1.46 g, Cimetta = 1.42 g).

<sup>c</sup>Values are the means of two experiments with four replicates each. Values within the same column followed by the same letter are not significant at the 0.05 level.
significantly different according to Fishers protected LSD ($P \leq 0.05$).

Table 4: Plant fresh weight of three Swiss wheat cultivars grown in presence or absence of *P. fluorescens* strains CHA0 or KD in artificial soil under sterile conditions.

<table>
<thead>
<tr>
<th>Cultivar</th>
<th>None</th>
<th>CHA0</th>
<th>KD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arina</td>
<td>100.00 c</td>
<td>112.84 b</td>
<td>98.12 c</td>
</tr>
<tr>
<td>Zinal</td>
<td>100.00 c</td>
<td>127.43 a</td>
<td>95.50 c</td>
</tr>
<tr>
<td>Cimetta</td>
<td>100.00 c</td>
<td>129.73 a</td>
<td>102.90 c</td>
</tr>
</tbody>
</table>

* Total plant fresh weights are expressed as percentage of weights of corresponding untreated control plants (Arina = 3.04 g, Zinal = 2.94 g, Cimetta = 2.75 g).

* Values are the means of two experiments with four replicates each. Values marked with the same letter are not significantly different according to Fishers protected LSD ($P \leq 0.05$).
Table 5: Root colonization by *P. fluorescens* CHA0 or KD of three Swiss wheat cultivars grown under gnotobiotic conditions in presence or absence of the pathogen *P. ultimum*.

<table>
<thead>
<tr>
<th>Cultivar</th>
<th>CHA0 With <em>Pythium</em></th>
<th>CHA0 Without <em>Pythium</em></th>
<th>KD With <em>Pythium</em></th>
<th>KD Without <em>Pythium</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Arina</td>
<td>7.73 a &lt;sup&gt;b&lt;/sup&gt;</td>
<td>7.10 a&lt;sup&gt;*&lt;/sup&gt;</td>
<td>8.13 a</td>
<td>8.20 a</td>
</tr>
<tr>
<td>Zinal</td>
<td>7.82 a</td>
<td>7.26 a&lt;sup&gt;*&lt;/sup&gt;</td>
<td>7.74 b</td>
<td>7.81 b</td>
</tr>
<tr>
<td>Cimetta</td>
<td>7.68 a</td>
<td>7.50 a</td>
<td>8.30 a</td>
<td>8.08 a</td>
</tr>
</tbody>
</table>

<sup>a</sup>Plants were inoculated with bacteria and on the same day infested with *P. ultimum*. After 3 wk of growth in artificial soil under sterile conditions plants were harvested and assessed for bacterial root colonization.

<sup>b</sup>Values are the means of two experiments with four replicates each. Values in the same column followed by the same letter are not significantly different according to Fishers protected LSD (*P* ≤ 0.05).

<sup>*</sup> = significant differences between treatments with and without *P. ultimum* for the same cultivar and bacterial inoculant according to Fishers protected LSD (*P* ≤ 0.05).