Identification and Specific Detection of a Novel Pseudomonadaceae Cluster Associated with Soils from Winter Wheat Plots of a Long-Term Agricultural Field Experiment

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The genus Pseudomonas (sensu stricto) represents a group of microorganisms directly involved in functions conferring plant health. We performed a study in the DOK long-term agricultural field experiment on the basis of previously published Pseudomonas-selective PCR primers in order to investigate the community structure of the microbial groups defined by the target range of these primers. Three different agricultural management systems, i.e., conventional, biodynamic, and bio-organic, along with mineral and unfertilized controls were investigated, with each system planted with either winter wheat or a grass-clover ley. Amplified small-subunit rRNA gene fragments were analyzed using the genetic profiling techniques restriction fragment length polymorphism (RFLP) and denaturing gradient gel electrophoresis (DGGE), revealing distinct differences between soils planted with winter wheat and grass clover but only minor differences between the management systems. Phylogenetic analyses of 59 clone sequences retrieved from bio-organic and unfertilized systems identified sequences related to Pseudomonas fluorescens and a novel cluster termed Cellvibrio-related Pseudomonadaceae (CRP). The CRP clones were exclusively isolated from winter wheat soil samples and were responsible for the crop-specific differences observed in RFLP and DGGE profiles. New primers were designed for the amplification of CRP targets directly from soil DNA, yielding strong signals exclusively for winter wheat soils. We concluded that crop-associated CRP exist in agricultural soils and that genetic profiling followed by specific probe design represents a valuable approach for identification as well as sensitive and rapid monitoring of novel microbial groups in the environment.

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In order to maintain the fertility of agricultural soils, it is important to learn more about the structures and functions of the indigenous microbial communities. A promising strategy may be to gain detailed information about microbial community structures in relation to various agricultural practices. For bacteria and fungi, a number of studies have been performed to investigate the relative influences of factors such as soil properties, agricultural practices, or plant types on their community structures. Soil type has consistently been reported as one of the main determinants of microbial community structures (15, 47). Furthermore, agricultural practices, including types of fertilizer, the addition of lime, and pesticide application, have been shown to significantly affect microbial community structures as well (14, 19, 48). The influence of plants on soil microbial communities has been assessed intensively in rhizosphere soil (7, 20, 41) as well as in bulk soil (15, 19, 48). Whereas clear differences between bulk and rhizosphere soil and also between different plant species have been detected, the influence of plants on microbial communities in bulk soil has often been reported as less significant (15, 19, 48).

In the DOK field experiment, located in northwestern Switzerland, three different farming systems, i.e., biodynamic (BIODYN), bio-organic (BIOORG), and conventional (CONFYM) management, and two control treatments with mineral fertilization (CONFYM) or no fertilization (NOFERT) have been established since 1978 (25). A study of soils from winter wheat plots from the DOK field experiment based on Biolog profiles revealed differences between the farming systems, but variation among field replications was relatively high (10). Recently, these differences have been confirmed with Biolog and bacterial terminal restriction fragment length polymorphism (T-RFLP) analyses performed on soils from winter wheat and grass-clover ley plots (48). These studies have shown that long-term farmyard manure (FYM) application has a highly significant effect on soil bacterial community structures, while short-term crop effects were weaker but still significant. Differences among BIODYN, BIOORG, and CONFYM were not significant. However, analyses of soil organic matter in the DOK system have indicated that distinct differences exist in the differently managed systems (9).

While the detection of entire bacterial or fungal communities has proven to be a successful approach for assessing changes in dominant community components, the resolution provided by current community-level profiling methods may not be sufficient to obtain information on the structure of specific microbial subgroups. This limitation can be overcome by directly targeting specific subgroups of particular interest.

The bacterial genus Pseudomonas represents one such subgroup that plays important roles in agricultural ecosystems. Specifically, it is involved in disease suppression in many crops (23, 46), exhibits plant growth-promoting properties (16, 40),
and is able to degrade a variety of pesticides used in agriculture for crop protection purposes (30, 39). Besides these beneficial traits, some members of the genus *Pseudomonas* also represent plant pathogens (35, 37, 50) and cause reductions in crop yields. Limited information is available on environmental or anthropogenic factors that determine or influence the occurrence, distribution, and activity of indigenous *Pseudomonas* populations in soil (1, 3, 21). The majority of studies on soil *Pseudomonas* communities have relied on cultivation-dependent techniques (1, 21, 33). Although selective cultivation of *Pseudomonas* spp. has been successfully applied, only a small fraction of the entire soil *Pseudomonas* community can be retrieved by cultivation (24, 34). On the other hand, few studies have been performed to analyze soil *Pseudomonas* communities by using less biased molecular biological techniques (12, 28, 32, 34, 49). The agronomic importance and the saprophytic nature of *Pseudomonas* spp. make them an important microbial group for investigation in agricultural management systems such as the long-term DOK field experiment.

Our hypothesis was that differences in soil organic matter quality among BIODYN, BIOORG, and CONFYM derived from system-specific (FYM) qualities influence saprophytic *Pseudomonas* populations in soil. In addition, different crops may also have an influence on different components of soil *Pseudomonas* communities. In order to address these issues, we first performed a theoretical reevaluation of the *Pseudomonas*-selective PCR primers published by Widmer et al. (49). Second, the structure of the bacterial group defined by the *Pseudomonas*-selective PCR primers was examined in soils of the different farming systems of the DOK field experiment planted with either winter wheat or a grass-clover ley.

**MATERIALS AND METHODS**

Site description and sample collection. The DOK field experiment was established in 1978 near Basel, in northwestern Switzerland, to compare three different agricultural management systems regarding soil fertility and productivity, nutrient dynamics, and impact on the environment (26). The soil at the site is classified as haplic luvisol (sandy loam) on deep deposits of alluvial loess. The experiment consists of BIODYN, BIOORG, and CONFYM farming systems, which all receive FYM as fertilizer. Two controls include a system with exclusive mineral fertilization (CONMIN) and an unfertilized system (NOFERT). The 7-year crop rotation includes potato, winter wheat 1, soy, maize, and winter wheat 2 followed by 2 years of grass-clover ley and is implemented in all treatments in three temporally shifted parallels. Each combination of management system and crop is replicated four times in the field (25). Altogether, 96 single plots with a size of 5 by 20 m are arranged in a split-split plot design on an area of 1.4 ha.

Soil samples (40 total) were obtained in March 2000 from each replicate plot for all five treatments planted with either winter wheat 1 (preceding crop, potato) or the grass-clover ley in its second year (for details, see references 15, 19, and 48). Briefly, 16 soil cores per plot were sampled to a depth of 20 cm, sieved to 2 mm, and stored on ice until further processing. DNA was extracted from the soil of each replicate plot, but for efficient screening of treatment-dependent population changes, equal amounts of DNA from the four corresponding replicates were pooled. This strategy allows for efficient screening and, if necessary, reanalysis of individual replicates.

DNA extraction and quantification. DNA was extracted following the protocol of Burgmann et al. (4), using a bead-beating procedure with slight modifications. Briefly, 0.5 g of soil (dry weight) was subjected to three repeated extractions using a bead beater (FP 120; Savant Instruments Inc., NY). Supernatants from all three extractions were pooled and subsequently purified with one chloroform/isoamyl alcohol (24/1) extraction. DNA concentrations were determined using a fluorometric assay with PicoGreen (Molecular Probes, Eugene, OR) as described previously (4).

Reevaluation of primers Ps-for and Ps-rev. Theoretical reevaluation of the Ps-for and Ps-rev primer sequences (49) was performed by calculating the coverage (percentage of *Pseudomonas* small-subunit (SSU) rRNA gene sequences that contain the primer sequence) and specificity (ratio of *Pseudomonas* SSU rRNA gene sequences that contain the primer sequence to the number of bacterial SSU rRNA gene sequences [including *Pseudomonas*] that contain the primer sequence) of the primers. For these analyses, exclusively bacterial SSU rRNA sequences longer than 1,200 bp and deposited in RDP-II release 9.24 (January 2005; 44,890 sequences) were used. The Primer_Match algorithm in RDP-II does not distinguish between sequences that lack one primer site due to missing sequence information and those that display mismatches with primer sequences.

PCR amplification, RFLP, and denaturing gradient gel electrophoresis (DGGE) analyses. *Pseudomonas*-selective PCR was performed with primers Ps-for (5'-GGTCTGAGGAGTAGATCATG-3') and Ps-rev (5'-TTAGCTCCCACTGCGCCG-3') (49). Reaction cocktails (50 µl) contained 2 mM MgCl₂, 0.2 µM each primer, 0.4 mM each deoxyribonucleoside, 0.6 mg/ml bovine serum albumin, 2 µl of HotStar Taq polymerase (QIAGEN GmbH, Hilden, Germany), 1× PCR buffer (QIAGEN), and 40 ng soil DNA. The cycling conditions consisted of an initial denaturation step for 15 min at 95°C to activate the HotStar polymerase, followed by 40 cycles of denaturation for 45 s at 94°C, primer annealing for 45 s at 65°C, and DNA synthesis for 45 s at 72°C. A final synthesis step for 5 min at 72°C was performed to complete the reaction.

Prior to RFLP analyses, PCR products were mixed with concentration conversion buffer (18) to convert the PCR mixture into a solution suitable for restriction digestion. For digestion, 3 U of the restriction endonuclease HaeIII (Promega, Madison, WI) was added and incubated overnight at 37°C. Digestion products were cleaned using a Montage SEQ® reagent purification system (Millipore Corporation, Billerica, MA) and resuspended in 30 µl Tris-EDTA buffer. Restriction fragments were electrophoresed in 12% polyacrylamide gels (bisacrylamide/acrylamide, 1/37.5) for 6 h at 150 V and 35°C (Dcode system; Bio-Rad Laboratories Ltd., Hercules, CA). Gels were stained for 30 min with SYBR green I nucleic acid gel stain (1:5,000 in 1× Tris-acetate-EDTA buffer; Molecular Probes, Eugene, OR) and were subsequently analyzed using the Gel Doc system and Quantity One software (Bio-Rad).

For preparation of PCR products suitable for DGGE analyses, a nested PCR approach for Ps-for/Ps-rev PCR products was developed. Two hundred picograms of Ps-for/Ps-rev PCR product or 1 ng of plasmid DNA was used as the template for PCR with the primers UNI-b-for (5'-TGGCCAGMCGCAGGTT ATC-3' [modified from reference 13]; positions 516 to 532 according to *Escherichia coli* numbering) and 520-rev (5'-CGTGGCCTACTACGGGTATC-3') [49]; positions 791 to 809, including a 40-bp GC clamp at the 5' end [29]. Nested PCR yielded products of approximately 300 bp. Reaction cocktails (50 µl) and cycling conditions were the same as those described above, except for the annealing temperature, which was set to 55°C. DGGE analyses were performed in 16- by 16-cm polyacrylamide gels (10%) containing 1× Tris-acetate-EDTA buffer, 2% glycerol, with a denaturing gradient of formamide and urea of 35 to 65% (29). Separation was performed with the Dcode system (Bio-Rad) for 14 h at 70 V and 60°C. Gel staining and analysis were the same as those described above for RFLP analysis. A mixture of SSU rRNA gene fragments from the following organisms was used as a DGGE migration standard: (i) *Pseudomonas fluorescens* ATCC 26663, (ii) *Bacillus subtilis* ATCC 14893, (iii) *Rizobium meliloti* DSM 1, (iv) *Flavobacterium capsulatum* DSM 1, and (v) *Arthrobacter globiformis* DSM 1.

DNA cloning and sequencing. Ps-for/Ps-rev PCR products were cloned using a pGEM-T Easy cloning kit (Promega, Madison, WI). The approximately 990-bp PCR products were sequenced on both strands according to the method of Widmer et al. (49), using an ABI3100 genetic analyzer (Applied Biosystems, Foster City, CA).

DNA sequence analyses. Sequences were subjected to theoretical RFLP analyses using the online program RestrictionMapper (http://www.restrictionmapper.org). Clones were classified into restriction types labeled A, B, C, etc., according to the calculated RFLP patterns. The 59 clone sequences from this study and 26 control sequences retrieved from RDP-II (6) were aligned using BioEdit, version 7.0.1 (17). Pairwise Jukes-Cantor distance determinations and cluster analysis by the unweighted-pair group method with mathematical averages (UPGMA) with 100 bootstrap resamplings were performed with Treecon, version 1.3b (44). Maximum likelihood analysis was performed with the fastDNAml routine in BioEdit.

**Primer design, testing, and application.** Primer sequences for the detection of *Cellulobiose*-related *Pseudomonas aeruginosa* (CRP) were derived from the sequence alignment established for phylogenetic analyses. The determined primer sequences were evaluated using the Probe Match tool from RDP-II (5) as well as
the BLASTn tool from GenBank (2). The sequence of the forward primer, CRP 149-for (positions 440 to 460), was 5'-CAGTGGGAGAAARCTG-3', and the sequence of the reverse primer, CRP 565-rev (positions 837 to 856), was 5'-CCACTAAAGCTCAGGACT-3'. The PCR conditions for these primers were optimized using clones and soil DNA extracts from this study. Reaction cocktails and cycling conditions were the same as those described above, except for the annealing temperature, which was set to 57°C.

**Nucleotide sequence accession numbers.** The nucleotide sequences of the 59 clones retrieved in this study are deposited in GenBank under accession numbers AY854436 to AY854494.

**RESULTS**

**Reevaluation of primers Ps-for and Ps-rev.** Primer Ps-for covered 1,370 (96.8%) and primer Ps-rev covered 1,338 (94.6%) of 1,415 *Pseudomonas* (sensu stricto) SSU rRNA sequences deposited in RDP-II. The combination of both primers detected 1,299 of 1,415 *Pseudomonas* sequences, corresponding to 91.8% coverage. Primer Ps-for revealed an exact match with 364 non-*Pseudomonas* bacterial SSU rRNA sequences in RDP-II (79.0% specificity), while primer Ps-rev matched 588 non-*Pseudomonas* sequences (69.5% specificity). For the combination of both primers, only 12 non-*Pseudomonas* sequences were identified, leading to a specificity of 99.1%. Among these nontarget sequences, five were associated with the genus *Cellvibrio* (sequence accession numbers AF006505, AI289162, AF452103, AY212574, and AY038046), which is the closest known relative to *Pseudomonas*. The remaining seven sequences were classified as *Oceanospirillales* (two *Zooshikella* sp. sequences [AB021682 and AF195410] and one unclassified sequence [AB020600]), unclassified *γ-Proteobacteria* (three sequences [AY676463, AF006502, and AF006506]), and unclassified *β-Proteobacteria* (one sequence [AF006507]).

**Pseudomonas-selective RFLP analysis.** Distinct changes in HaeIII RFLP patterns were observed between winter wheat and grass-clover bulk soils across all five treatments in the DOK field experiment (Fig. 1). Prominent bands with calculated fragment sizes of 512, 279, 126, and 72 bp were detected in bulk soil from winter wheat plots but were absent or only faint for soil samples from grass-clover plots. Band intensities and fragment sizes observed in RFLP patterns from winter wheat or grass-clover bulk soil appeared highly similar across the five management systems.

**Pseudomonas-selective DGGE analyses.** The DGGE banding patterns for winter wheat plot soils of all five treatments revealed two distinct additional bands in the higher denaturing region of the gel compared to the corresponding soils from grass-clover plots (Fig. 2, region II, open arrowheads). For soils from grass-clover plots of the farming systems BIOYDN, BIOOR, and NOFERT, these bands were not detectable, and for soils from the systems CONFYM and CONMIN, they were only weakly detectable. DGGE patterns derived from soils of winter wheat plots showed high similarity among BIOYDN, BIOOR, CONFYM, and CONMIN, while NOFERT soils planted with winter wheat displayed two additional bands that were not detected in the other soils (Fig. 2, closed arrowheads).

**Cloning of Pseudomonas-selective PCR products.** Four gene libraries of cloned Ps-for/Ps-rev PCR products were constructed from the BIOOR and NOFERT systems for both crops. The libraries were screened by HaeIII RFLP analysis, and a total of 59 randomly selected clones were isolated and sequenced (Table 1). Theoretical restriction analysis of individual clone sequences with the restriction endonuclease HaeIII revealed eight different RFLP patterns (Table 1). Pattern A, which yielded the dominant fragment sizes observed in

![FIG. 1. HaeIII RFLP analysis of PCR-amplified SSU rRNA gene fragments from bulk soil DNA extracts with Pseudomonas-selective primers. Samples represent five different treatments of the DOK field experiment (lanes BIOYDN, BIOOR, CONFYM, CONMIN, and NOFERT) planted with winter wheat (lanes WW) or grass-clover (lanes GC). Bands referred to in the text are identified with arrows and DNA fragment sizes (bp). Underlined fragment size values indicate bands that revealed clear intensity changes across different soils and crops.](image1)

![FIG. 2. DGGE analysis of PCR-amplified SSU rRNA gene fragments based on Pseudomonas-selective PCR products from bulk soil DNA extracts. Samples represent five different treatments of the DOK field experiment (BIODYN, BIOOR, CONFYM, CONMIN, and NOFERT) planted with winter wheat (WW) or grass-clover (GC). Migration standard (S) bands 1 and 3 define the migration region for sequences associated with *Pseudomonas* (sensu stricto) (lower denaturing region [I]), and bands 3 and 4 define the migration region for sequences of CRP (higher denaturing region [II]). Open arrowheads, dominant bands detected in all winter wheat soils; closed arrowheads, bands exclusively detected in NOFERT system planted with winter wheat.](image2)
the whole community patterns for grass-clover soils (Fig. 1, size labels without underlining), was also dominant (100% and 81.3%) in the two libraries derived from grass-clover soils (Table 1). Accordingly, pattern B, which yielded the additional bands in the community patterns for winter wheat soils (Fig. 1, underlined size labels), was most frequently (23.1 and 48%) found in the libraries from winter wheat soils.

Of the 59 isolated clones, 45 differing within the 300 bp amplified by nested PCR were used for DGGE classification into lower and higher denaturing groups (Fig. 2). Clones characterized by HaeIII RFLP types A, C, E, G, and H were exclusively assigned to the lower denaturing group (Table 1). Clones with RFLP types B, D, and F were assigned to the higher denaturing group, except for clone DOK_N_GC 60 (RFLP type B), which migrated in the lower denaturing group. Five clones (DOK_N_WW 07, 45, 21, 13, and 52) from the higher denaturing group migrated at exactly the positions of the two additional bands observed for the NOFERT system planted with winter wheat (data not shown).

**Phylogenetic analysis.** Cluster analysis revealed that all 59 sequences matched the target range of the *Pseudomonas*-selective primers and were associated with three branches within the *Pseudomonadaceae* (Fig. 3). Forty sequences (68%) grouped within the *P. fluorescens* branch (Fig. 3, branch Ia) of the whole community patterns for grass-clover soils (Fig. 1, size labels without underlining), was also dominant (100% and 81.3%) in the two libraries derived from grass-clover soils (Table 1). Accordingly, pattern B, which yielded the additional bands in the community patterns for winter wheat soils (Fig. 1, underlined size labels), was most frequently (23.1 and 48%) found in the libraries from winter wheat soils.

**TABLE 1.** HaeIII RFLP types and frequencies among Ps-for/Ps-rev PCR product clone libraries from BIOORG and NOFERT plot soils planted with either winter wheat or grass-clover

<table>
<thead>
<tr>
<th>RFLP type</th>
<th>BIOORG</th>
<th>NOFERT</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Grass-clover</td>
<td>Winter wheat</td>
</tr>
<tr>
<td>A Low</td>
<td>5 (100)</td>
<td>5 (38.5)</td>
</tr>
<tr>
<td>B High</td>
<td>0</td>
<td>3 (23.1)</td>
</tr>
<tr>
<td>C Low</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>D High</td>
<td>0</td>
<td>1 (7.7)</td>
</tr>
<tr>
<td>E Low</td>
<td>0</td>
<td>1 (7.7)</td>
</tr>
<tr>
<td>F High</td>
<td>0</td>
<td>1 (7.7)</td>
</tr>
<tr>
<td>G Low</td>
<td>0</td>
<td>1 (7.7)</td>
</tr>
<tr>
<td>H Low</td>
<td>0</td>
<td>1 (7.7)</td>
</tr>
<tr>
<td>Total</td>
<td>5 (100)</td>
<td>13 (100)</td>
</tr>
</tbody>
</table>

a Clone libraries labeled according to their sources, i.e., BIOORG grass-clover, DOK_O_GC; BIOORG winter wheat, DOK_O_WW; NOFERT grass-clover, DOK_N_GC; and NOFERT winter wheat, DOK_N_WW.

b Clones were divided into a lower and a higher denaturing group according to their migration positions in DGGE gels relative to the standard (Fig. 2).

c One clone from the DOK_N_GC library migrated with the lower DGGE group.

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**FIG. 3.** Phylogenetic tree of SSU rRNA gene sequences amplified with *Pseudomonas*-selective PCR primers. Sequences from this study are printed in bold and are annotated with their RFLP types (A to H; see Table 1). The arrowhead marks the branch including the family *Pseudomonadaceae*. Cluster I includes the genus *Pseudomonas* (sensu stricto); collapsed subclusters Ia, Ib, and Ic are defined by the marker species *P. fluorescens*, *P. aeruginosa*, and *P. putida*, respectively (49). Cluster II includes the genus *Cellvibrio* (IIa) and CRP (IIb). The proteobacterial sequence SCB11 represents the outgroup, as originally defined for the design of the Ps-for/Ps-rev primers (49). Maximum likelihood calculation supported the definition of these clusters and the locations of all retrieved clones within these clusters.
the *Pseudomonas* (sensu stricto) cluster (Fig. 3, cluster I). In DGGGE profiles, these sequences corresponded to the lower denaturing group, i.e., region I in Fig. 2. Two sequences (3%), one in the lower (DOK _N_ GC 60) and the other in the higher (DOK _N_ GC 49) denaturing group, were associated with the genus *Cellvibrio* (Fig. 3, branch IIa). Seventeen sequences (29%) formed a novel cluster (Fig. 3, branch IIb) located close to *Cellvibrio* and therefore termed *Cellvibrio*-related *Pseudomonadaceae* (CRP). In DGGGE analysis, these sequences migrated in the higher denaturing group, i.e., region II in Fig. 2. Clones DOK _N_ WW 07, 13, 21, 45, and 52 associated with the CRP cluster and accounted for the two unique bands in the soil from winter wheat plots in the NOFERT system (Fig. 2, closed arrowheads).

The sequences, which grouped in the *P. fluorescens* cluster (Fig. 3, branch Ia), originated from all four clone libraries. The sequences associated with the *Cellvibrio* cluster (Fig. 3, branch IIa) derived from NOFERT soils planted with grass-clover. The sequences from the CRP cluster (Fig. 3, branch IIb) derived exclusively from soils planted with winter wheat. No sequences were retrieved which clustered within the *Pseudomonas aeruginosa* (Fig. 3, branch Ib) and the *Pseudomonas putida* (Fig. 3, branch Ic) branches. Phylogenetic analyses using maximum likelihood revealed an identical basic tree topology to that by UPGMA cluster analysis (data not shown).

**Selective detection of winter wheat-associated CRP.** Theoretical analyses in RDP-II revealed that in combination, the newly designed primers CRP 149-for and CRP 565-rev discriminated all published nontarget sequences by at least 3 bp, except for one uncultured γ-proteobacterium (GenBank accession number AJ518787), which contained only two mismatches within the forward and none within the reverse primer. Experimental testing of the primers was performed, with the 45 different clones as templates. Positive PCR signals were exclusively observed for clones associated with the CRP cluster (data not shown). Clones belonging to the *Pseudomonas* (sensu stricto) and *Cellvibrio* clusters (Fig. 3) yielded no PCR products (data not shown).

PCR analyses of soil DNA extracts from all farming systems, controls, and crops resulted in strong products of the expected size from soils planted with winter wheat, while no or faint signals were detected in soil samples from grass-clover plots (Fig. 4a). With DGGGE analysis, two dominant bands were resolved (Fig. 4b). These bands corresponded to those detected by *Pseudomonas*-selective PCR in soils planted with winter wheat (Fig. 2 and 4b, open arrowheads). For soils from grass-clover plots, only very weak bands were detected in DGGGE patterns from the CONFYM and CONMIN systems, whereas no signals were detected for the BIODYN, BIOORG, and NOFERT systems. This finding was in perfect agreement with the *Pseudomonas*-selective DGGGE profiles (Fig. 2).

**DISCUSSION**

Data from RFLP and DGGGE profiling were directly comparable because the analyses were based on identical PCR products generated with the primer pair Ps-for and Ps-rev (Fig. 1 and 2) (49). Both techniques produced consistent results by clearly discriminating between target populations in soils planted with winter wheat and grass-clover. DGGGE analysis yielded higher resolution within the *Pseudomonas* (sensu stricto) and CRP clusters but required a nested PCR step, potentially leading to increased bias by preferential amplification of certain sequence types (42). On the other hand, the limited resolution power of the applied RFLP approach for analysis of *Pseudomonas* spp. could be overcome by using multiple enzymes for the digestion of PCR products (32). Furthermore, both profiling techniques yielded objectively analyzable information for detection of and discrimination between *Pseudomonas* spp. and CRP. In RFLP analysis and DGGGE analysis, fragment sizes of particular bands and lower and higher denaturing regions, respectively, represented the diagnostic criteria (Fig. 1 and 2). Although performed on pooled DNA samples, our analyses allowed us to determine that both fingerprinting techniques do not have the sensitivity to resolve possible differences within the soil *Pseudomonas* (sensu stricto) communities. However, we could clearly demonstrate a pronounced crop effect on the CRP community.

The majority of SSU rRNA gene sequences retrieved, i.e., 55% for winter wheat plots and 90% for grass-clover plots, were derived from *Pseudomonas* (sensu stricto) (Table 1; Fig. 3). These sequences exclusively associated with the *P. fluorescens* branch, although our theoretical analyses and previous experimental testing (49) revealed that the Ps primers also amplify SSU rRNA genes from the *P. aeruginosa* and *P. putida* clusters (49). The results from the present study underscore the importance of repeated theoretical and experimental reevaluations of primer specificities as new sequences are continuously added to databases. In silico reevaluation of primers Ps-for and Ps-rev (49) in RDP-II revealed high coverage.
(91.8%) and specificity (99.1%) for the target group *Pseudomonas* (sensu stricto) and fully justified the use of these primers in our analysis. Subsequent phylogenetic analysis of 59 clone sequences demonstrated that the primers were highly selective for the originally defined target range characterized by the outgroup sequence SCB11 (Fig. 3) (49).

A group of sequences isolated from soils planted with winter wheat clustered apart from *Pseudomonas* (sensu stricto) and formed the novel CRP cluster (Fig. 3, branch Iib). It is premature to define whether CRP constitute a new species of the genus *Cellvibrio* or a new genus of the *Pseudomonadaceae*. The isolation of CRP and detailed analyses will be required for this definition. It was surprising that close relatives of *Pseudomonas* (sensu stricto) have only rarely been detected in soil and that the genus *Cellvibrio* has only been recently isolated by selective cultivation from bulk soil (22, 45). These isolates were reported to display agarolytic or cellulolytic properties. A few *Cellvibrio* spp. have been isolated from maize rhizospheres (36), and potato endospheres (38). The detection of *Cellvibrio* spp. in plant rhizo- and endospheres suggested that they may be preferentially associated with specific plants. This would be in agreement with our finding that the presence of CRP was highly correlated with winter wheat.

The novel CRP primers, i.e., CRP 149-for and CRP 565-rev, allowed us to confirm the results obtained with RFLP and DGGE analyses based on *Pseudomonas*-selective PCR. Forty PCR cycles were used for the detection of *Pseudomonas* spp. and CRP, suggesting a comparable abundance of CRP and *Pseudomonas* spp. in soils planted with winter wheat. The same conclusion could be drawn from the comparable band intensities observed for *Pseudomonas* spp. and CRP in the RFLP and DGGE analyses (Fig. 1 and 2). *Pseudomonas* spp. have been reported to account for approximately 1% of the bacterial communities in soil (24), which may also represent the approximate abundance of CRP in the winter wheat plots. If more detailed quantitative analyses of these microbial groups are required, specific quantitative PCR approaches will have to be developed (43). The independent analysis of CRP also provided strong evidence that the observed differences in RFLP and DGGE profiles (Fig. 1 and 2) reflected actual differences in bacterial community structures and were not related to a PCR bias. The identification of changing components in genetic profiles by cloning and sequencing has been frequently performed in microbial community analysis (7, 41). However, the gained information has rarely been confirmed by designing specific primers for independent analysis of the changing community components (31). It has previously been shown that such a strategy can also provide tools with increased detection sensitivities (31). We strongly believe that, if consistently applied, this approach will provide an increasing set of tools for indicator detection of high value for research and soil quality analyses.

The main agricultural factor determining CRP community structure in the DOK field soils appeared to be the plant type. Currently, it is unknown whether CRP are promoted in the presence of winter wheat or whether they are excluded by the grass-clover ley. Applying the developed detection tools to soil samples from different crops at different time points during growth and at different locations will provide further information regarding the mechanisms involved in enriching CRP in soils and their possible functions. Previous analyses of the same DOK field samples have focused on the bacterial domain by using T-RFLP analyses (48). These data revealed that the type of fertilizer, i.e., organic versus mineral or no fertilization, dominated plant effects (48), suggesting that bacterial subgroups which responded to the fertilizer regimen were present. In agreement with these results, Kennedy et al. (19) reported that bacterial community structures were influenced by lime and nitrogen additions, and to a much smaller extent, by the plant species. This apparent difference compared to our findings may be explained by the fact that a bacterial community profile represents a composite of various bacterial subgroups, which may display different responses. Relatively small groups may either not be detected in bacterial community profiles (8, 29) or be outweighed by other components in statistical analysis (11, 18), thus emphasizing the necessity of genus-level analyses in microbial ecology investigations.

**Conclusions.** With RFLP and DGGE analyses, no clear differences were detected in the *Pseudomonas* soil communities in the DOK field experiment. Therefore, we had to reject our initial hypothesis that *Pseudomonas* communities respond to the different treatments in the DOK long-term field experiment. However, the novel group of soil CRP identified in the present study appeared to be highly responsive to the crop type. The isolation and detailed characterization of CRP will allow us to taxonomically and functionally define this interesting novel soil bacterial group.

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