

## Effects of Bacterial Antibiotic Production on Rhizosphere Microbial Communities from a Culture-Independent Perspective

EDUARDO A. ROBLETO, JAMES BORNEMAN,<sup>†</sup> AND ERIC W. TRIPLETT\*

Department of Agronomy, Center for the Study of Nitrogen Fixation,  
University of Wisconsin-Madison, Madison, Wisconsin 53706

Received 29 June 1998/Accepted 13 September 1998

**The effects of antibiotic production on rhizosphere microbial communities of field-grown *Phaseolus vulgaris* were assessed by using ribosomal intergenic spacer analysis. Inoculum strains of *Rhizobium etli* CE3 differing only in trifolixitin production were used. Trifolixitin production dramatically reduced the diversity of trifolixitin-sensitive members of the  $\alpha$  subdivision of the class *Proteobacteria* with little apparent effect on most microbes.**

The great majority of microorganisms in the environment remain unidentified because they are not culturable with standard techniques (2, 14, 15, 23). The phylogenetic diversity of soil bacteria observed by culture-independent methods is enormous (4, 5, 17, 22). Ribosomal intergenic spacer analysis (RISA) is used to assess microbial diversity in complex systems, as well as estimate effects of disturbance on diversity (5), such as the introduction of an antibiotic-producing bacterium into soil. To date, nothing is known about the ability of antibiotic production to affect microbial diversity in the environment from a culture-independent perspective. In this study, we assessed microbial diversity changes in the rhizosphere of bean plants following inoculation with bacterial strains that differ only in the ability to produce a narrow-spectrum peptide antibiotic, trifolixitin (TFX).

The *Rhizobium* TFX production phenotype reduces the number of TFX-sensitive rhizobia in bean rhizospheres and enhances the ability of a strain to limit root nodulation by TFX-sensitive *Rhizobium* strains (12, 13, 18–20). By using the nearly isogenic strains of *Rhizobium etli* available that differ only in the presence or absence of the TFX production phenotype, we assessed the ability of the TFX production phenotype to alter microbial diversity in the rhizosphere by using RISA. As the taxonomic range of in vitro TFX sensitivity is restricted to a specific group of members of the  $\alpha$  subdivision of the class *Proteobacteria* ( $\alpha$ -*Proteobacteria*) (21), the specificity of the TFX action on rhizosphere microbes was estimated.

The inoculum strains used are derivatives of *R. etli* CE3. CE3(pT2TFXK) and CE3(pT2TX3K) are nearly isogenic, TFX-producing and -nonproducing strains, respectively (13). Culture conditions, inoculation methods, antibiotic concentrations, seed coating, bean cultivar, soil properties, and experimental design and location were described previously (12).

To amplify RISA PCR products specific to the taxonomic group of bacteria sensitive to TFX, a forward 16S ribosomal DNA (rDNA) primer referred to as RB1 (5'-TGGTGACAG

TGGGCAGCG-3') was designed. The specificity of RB1 for the TFX-sensitive bacteria was confirmed by PCR analysis with template DNA of organisms within and outside this group by using TR8 as the reverse primer (3, 11). Further evidence of

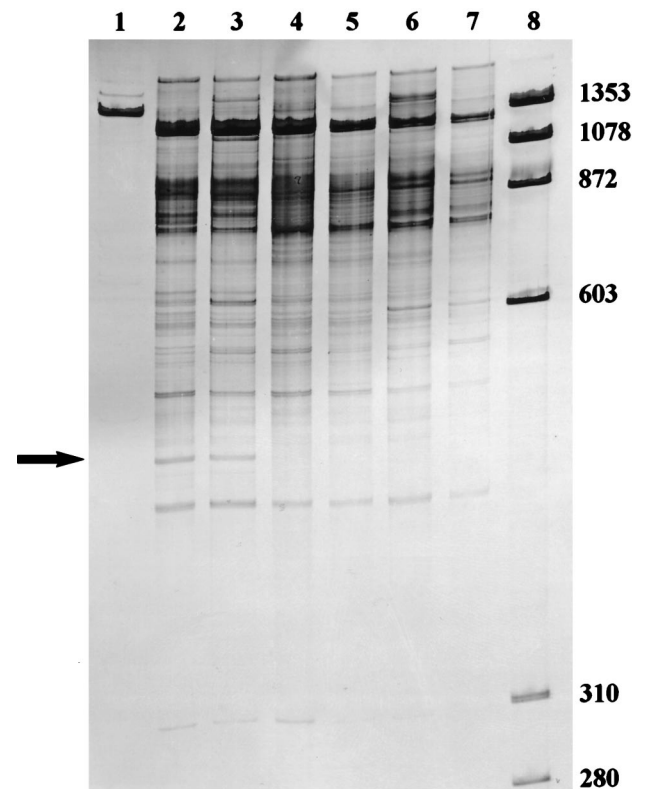


FIG. 1. Intergenic 16S to 23S rDNA patterns of dominant microbial populations in the rhizosphere of beans inoculated with *R. etli* CE3 strains that differ in TFX production in Arlington, Wis., 1997. Lanes: 1, pure culture of *R. etli* CE3; 2 and 3, rDNA bands from rhizosphere of uninoculated plants; 4 and 5, rDNA bands from rhizospheres inoculated with CE3(pT2TFXK) (the TFX-producing strain); 6 and 7, rDNA bands from rhizospheres inoculated with CE3(pT2TX3K) (the TFX-nonproducing strain); 8, molecular size markers. The arrow indicates a 450-bp fragment absent in *R. etli* treatments. Each lane represents an independent replicate. The values on the right are molecular sizes in base pairs.

\* Corresponding author. Mailing address: University of Wisconsin-Madison, Department of Agronomy, 1575 Linden Drive, Madison, WI 53706. Phone: (608) 262-9824. Fax: (608) 262-5217. E-mail: triplett@facstaff.wisc.edu.

<sup>†</sup> Present address: University of California-Riverside, Department of Plant Pathology, Riverside, CA 92521.

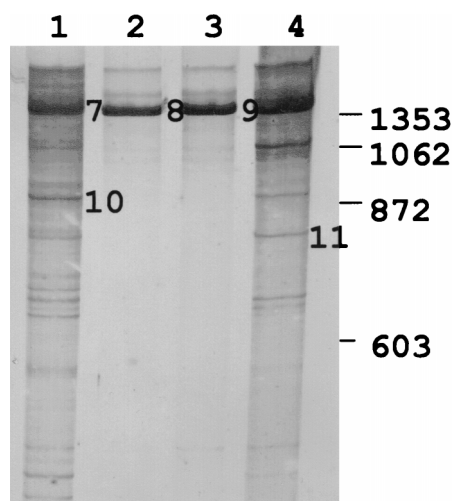


FIG. 2. Intergenic 16S to 23S rDNA patterns of dominant  $\alpha$ -proteobacterial populations in the rhizosphere of beans inoculated with *R. etli* CE3 strains that differ in TFX production in Arlington, Wis., 1997. Lanes: 1, rDNA bands from rhizosphere of uninoculated plants; 2 and 3, rDNA bands from rhizosphere of plants inoculated with CE3(pT2TFXK) (the TFX-producing strain); 4, rDNA bands from rhizospheres of plants inoculated with CE3(pT2TX3K) (the TFX-nonproducing strain). Numbers inside the gel, to the right of bands, refer to fragments that were excised and sequenced corresponding to TFX-resistant or TFX-sensitive populations. Bands corresponding to TFX-resistant populations were identified by their presence in rhizosphere samples. Bands corresponding to TFX-sensitive populations were identified by their absence in the rhizosphere samples from the treatment containing the TFX-producing strain. Each lane represents an independent replicate. Rhizosphere DNA was isolated 63 days after inoculation. The values on the right are molecular sizes in base pairs.

RB1 specificity was obtained by using the Check Probe program from the Ribosomal Database Project (9).

Sixty-three days after planting, root sections from four plants per plot were removed from soil, placed in 40 ml of sterile

water, and shaken for 30 min at 250 rpm at room temperature. Rhizosphere DNA was extracted from 1 ml of root washes by using a soil DNA extraction kit (Bio 101) and then further purified with Wizard columns (Promega). RISA-PCR was performed as described previously (5). To amplify rDNA from  $\alpha$ -Proteobacteria sensitive to TFX, the RB1 forward primer was used. The PCR conditions used to amplify rDNA from TFX-sensitive  $\alpha$ -Proteobacteria were 94°C for 60 s; 30 cycles of 94°C for 15 s, 58°C for 15 s, and 72°C for 25 s; and extension at 72°C for 60 s. Fragments common or specific to treatments were excised from the gel, extracted in water, and reamplified by using the RB1 and TR8 primers. The resulting 230 to 250-bp fragments were sequenced by dye termination (5). Sequences from the RISA bands were compared with all sequences in the databases by using BLAST (1).

The RISA products obtained by using universal rDNA primers suggest that the microbial diversity of rhizosphere soil is immense (5) (Fig. 1). Many bands were observed in the size range of 300 to 1,400 bp. Very few differences in the overall microbial diversity were observed, regardless of the inoculum strain used (Fig. 1). However, a significant difference between the RISA patterns obtained with the uninoculated treatment and the two inoculated treatments was observed. The RISA pattern from the uninoculated rhizosphere shows a 450-bp fragment that is not present in patterns from the rhizospheres inoculated with either CE3(pT2TFXK) or CE3(pT2TX3K). This suggests that inoculation with *R. etli* reduces the population of the microorganism(s) represented by the 450-bp fragment, while the TFX production phenotype has little detectable effect on the overall microbial diversity.

When RB1 was used for amplification, the RISA pattern was considerably less complex than that obtained with 1406F (5) (Fig. 2). The complexity decreased even more dramatically when the rhizosphere was inoculated with CE3(pT2TFXK). There was very little difference between the patterns obtained

TABLE 1. Results of sequence analysis of selected RISA bands that are labeled in Fig. 2

rrn DNA band	Accession no. of RISA band sequence	Closest relatives <sup>a</sup>	% Identity	Accession no. of closest relative
7TR	AF073476	<i>Rhizobium leguminosarum</i> <i>R. etli</i> TAL 182 <i>R. tropici</i> IIB <i>Agrobacterium tumefaciens</i>	99	D14513 U28939 X67234 D14505
8TR	AF073477	<i>Rhizobium leguminosarum</i> bv. trifolii <i>R. leguminosarum</i> bv. viciae <i>R. mongolense</i> <i>R. tropici</i> IIB	99	U31074 U89829 U89822 X67234
9TR	AF073478	<i>Rhizobium</i> sp. <i>Rhizobium</i> sp. <i>R. etli</i> TAL182 <i>Rhizobium leguminosarum</i>	100	AF041443 Y10174 U28939 X67233
10TS	AF073479	<i>Agrobacterium tumefaciens</i> bv. 2 <i>R. leguminosarum</i> <i>R. leguminosarum</i> bv. viciae <i>R. mongolense</i>	99	D14501 X67233 U29386 U89822
11TS	AF073480	<i>Agrobacterium tumefaciens</i> bv. 2 <i>R. leguminosarum</i> bv. viciae <i>R. tropici</i> IIB <i>R. gallicum</i>	98	D14501 U89829 X67234 AF008128

<sup>a</sup> The closest relatives of these partial rDNA sequences were identified by BLAST.

with the uninoculated treatment and the treatment inoculated with CE3(pT2TX3K) (Fig. 2).

A few bands from the  $\alpha$ -proteobacterial RISA gel were excised and sequenced. Some bands were chosen that were common to all of the treatments. Others were chosen because they were not found in the rhizosphere of plants inoculated with CE3(pT2TFXK). All sequences showed very high homology to the TFX-sensitive  $\alpha$ -*Proteobacteria*, consistent with the specificity of the forward primer used to amplify these fragments (Table 1).

Previous studies have examined the culturable microbial community in the rhizosphere upon inoculation with an antibiotic-producing strain (7, 8, 10). All of these studies examined only the culturable microorganisms, which meant that only a small portion of the rhizosphere microbial community was examined. In addition, the effects observed on the cultured communities could not be attributed to antibiotic production, since undefined chemical mutants were used for comparison with the wild type (7, 8).

Thus, a culture-independent approach was used here to examine changes in the bean rhizosphere after inoculation with two *Rhizobium* strains that differ only in antibiotic production. An uninoculated treatment was also included to determine the effects of *Rhizobium* inoculation on the rhizosphere microbial community. This approach also examined the effects on the taxonomically defined group of TFX-sensitive microorganisms. Our results are consistent with the taxonomic range of activity of TFX, since the  $\alpha$ -proteobacterial RISA profile was considerably altered by the TFX-producing strain, while the total microorganism RISA profile was not substantially affected (Fig. 1 and 2).

Thus, TFX production has a significant effect on TFX-sensitive bacteria in the rhizosphere of beans under agricultural conditions. This effect is not noticeable with the total microbial RISA profile, as expected, given the inability of TFX to inhibit most bacteria and given that the  $\alpha$ -*Proteobacteria* are not common in soil (4, 5). Although the culture bias has been removed from this work, PCR and primer biases remain (6, 16). However, despite these potential biases, this report strongly suggests that a reduction in the diversity of  $\alpha$ -*Proteobacteria* has occurred with TFX production.

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