Differences in ‘Candidatus Phytoplasma cynodontis’ Based on 16S rRNA and groEL Genes and Identification of a New Subgroup, 16SrXIV-C

J. Mitrović and M. Smiljković, Laboratory of Applied Phytopathology, Institute of Pesticides and Environmental Protection, Belgrade, Serbia; Erich Seemüller, Julius Kuehn Institute, Federal Research Centre for Cultivated Plants, Institute for Plant Protection in Fruit Crops and Viticulture, Dossenheim, Germany; Richard Reinhardt and Bruno Hüttel, Max Planck Genome Centre Cologne, Köln, Germany; Carmen Büttner, Department of Crop and Animal Sciences, Humboldt-Universität zu Berlin, Berlin, Germany; Assunta Bertaccini, DipSA, Plant Pathology, Alma Mater Studiorum, University of Bologna, Bologna, Italy; Michael Kube, Department of Crop and Animal Sciences, Humboldt-Universität zu Berlin, Berlin, Germany; and Bojan Duduk, Laboratory of Applied Phytopathology, Institute of Pesticides and Environmental Protection, Belgrade, Serbia

Abstract


‘Candidatus Phytoplasma cynodontis’ is widespread in bermudagrass and has only been found in monocotyledonous plants. Molecular studies carried out on strains collected in Italy, Serbia, and Albania confirmed the ability to identify molecular variability in the 16S ribosomal RNA (rRNA) gene. Based on restriction fragment length polymorphism and sequence analyses, the strains from Serbia were clearly differentiated from all others and assigned to a new ribosomal DNA (rDNA) subgroup designated as 16SrXIV-C. A system for amplification of fragments containing the ‘Ca. P. cynodontis’ groEL gene was developed to enable study of its variability in related strains belonging to different 16SrXIV subgroups. Despite the fact that the groEL gene exhibited a greater sequence variation than 16S rRNA, the phylogenetic tree based on groEL gene sequence analysis was highly congruent with the 16S rDNA-based tree. The groEL gene analyses supported differentiation of the Serbian strains and definition of the new subgroup 16SrXIV-C. Phylogenetic analyses of both genes confirmed distinct phylogenetic lineages for strains belonging to 16SrXIV subgroups. Furthermore, groEL is the only nonribosomal marker developed for characterization of ‘Ca. P. cynodontis’ thus far, and its application in molecular surveys should provide better insight into the relationships among these phytoplasmas and correlation between strain differentiation and their geographical distribution.

Bermudagrass white leaf (BGWL) is a disease of bermudagrass (Cynodon dactylon (L.) Pers.) associated with ‘Candidatus Phytoplasma cynodontis’ and transmitted by the leafhopper Exitianus capricola (Marcone et al. 2004; Salehi et al. 2009). Typical symptoms of BGWL include extensive chlorosis, proliferation of axillary shoots, bushy growing habit, small leaves, shortened stolons and rhizomes, stunting, and plant death (Marcone et al. 2004). ‘Ca. P. cynodontis’ (BGWL phytoplasma) has been reported to have the smallest genome, with a size of 530 kbp, among all studied self-replicating organisms, estimated by pulsed-field gel electrophoresis, and is specifically associated with infection of monocotyledonous plants (Marcone et al. 2004). Therefore, its characterization may be important to studies of phytoplasma genome evolution and gene variability. ‘Ca. P. cynodontis’ belongs to the BGWL group (16SrXIV), which consists of subgroups 16SrXIV-A (Lee et al. 1998) and 16SrXIV-B, recently described in Iran (Salehi et al. 2009). The disease was reported for the first time in Taiwan in 1972 and has been reported since then throughout Asia, Africa, Australia, and Europe, and in Cuba (Arocha et al. 2005; Chen et al. 1972; Dafalla and Cousin 1988; Marcone et al. 1997; Salehi et al. 2009; Sdoodee et al. 1999; Viswanathan 1997; Zahoor et al. 1995). Although BGWL is a widespread disease, little is known about sequence variations in ‘Ca. P. cynodontis’ genes other than 16S ribosomal DNA (rDNA), which has shown some variability. Phylogenetic analyses of 16S rDNA sequences have indicated that BGWL phytoplasma strains, together with Brachiaria and Cynodon white leaf phytoplasmas, form a discrete subclade within the phytoplasma clade, having some distinct lineages also within the BGWL subclade (Çağlar et al. 2013; Marcone et al. 2004; Salehi et al. 2009). Phytoplasma classification was established using the properties of the 16S rDNA but it does not always provide molecular distinction of closely related strains. Because of the conserved nature of the 16S ribosomal RNA (rRNA) gene, more variable single-copy genes, such as ribosomal protein (rpl22 and rps3), secY, tuf, and groEL have been employed for finer differentiation of other phytoplasmas (Lee et al. 2004, 2006, 2010; Marcone et al. 2000; Martini et al. 2007; Mitrović et al. 2011); however, corresponding information is not available for ‘Ca. P. cynodontis’. The 16S–23S rDNA spacer region has been found highly conserved among different BGWL strains, while the secA gene has been used only for diagnostic purposes (Bekele et al. 2011; Marcone et al. 2004).

In the present study, we identified a new subgroup (16SrXIV-C) of ‘Ca. P. cynodontis’ and developed a groEL gene polymerase chain reaction (PCR) assay, which we used, in addition to 16S rDNA and the 16S–23S rDNA spacer regions, for differentiation of ‘Ca. P. cynodontis’-related strains from three European countries.

Materials and Methods

Sample collection and nucleic acid extraction. Eight bermudagrass samples exhibiting BGWL symptoms (leaf whitening, small leaves, bushy growing, and stunting) were collected from the following locations: two samples from mowed lawns on two locations in Italy; one sample from a park in an urban area in Tirana, Albania; and five samples from mowed lawns on three locations in public parks in Serbia (Table 1). Total nucleic acid was extracted from 0.5 g of symptomatic leaves following a previously described protocol (Doyle and Doyle 1990), then
resuspended in Tris-EDTA buffer and stored at −20°C. Nucleic acids were diluted 25- to 100-fold in sterile distilled water to obtain a DNA concentration of 20 ng/μl, before performing PCR assays.

Primer design. Two sequences obtained in a genomic survey and available online (GenBank accession numbers KF437623 and KF437624), containing partial groEL sequences of BGWL phytoplasma, were used for primer design. The sequence KF437623 containing the 3’ end of the groES, groES-groEL spacer region and the 5’ end of the groEL gene, and the sequence KF437624 containing the 3’ end of the groEL gene, together enabled the design of primers that specifically amplify the DNA fragment containing the groEL gene of BGWL phytoplasma. A forward primer was designed inside the groES gene while two reverse primers were designed on and near the 3’ end of the groEL gene. The forward primer CYNgroesF2 (5’-ACTATTCCGGAACTCAAT-3’) was based on a sequence starting 89 bp upstream of the 3’ end of groES, while the reverse primers CYNgroelR (5’-TATAAAGAAGAAG...

Table 1. ‘Candidatus Phytoplasma cynodontis’ strains and accession numbers of their 16S ribosomal DNA (rDNA) and groEL gene sequences

<table>
<thead>
<tr>
<th>Locality</th>
<th>Sampling year</th>
<th>Strain</th>
<th>Accession numbers for 16S rDNA, groEL*</th>
<th>Ribosomal subgroup</th>
</tr>
</thead>
<tbody>
<tr>
<td>Montemassi, Italy</td>
<td>2010</td>
<td>IT71/10</td>
<td>KF383979 (1,768), KF383983 (1,687)</td>
<td>16SrXIV-A</td>
</tr>
<tr>
<td>Belgrade-Zvezdara, Serbia</td>
<td>2011</td>
<td>RS59/11</td>
<td>KF383981 (1,755), KF383985 (1,748)</td>
<td>16SrXIV-C</td>
</tr>
<tr>
<td>Tirana, Albania</td>
<td>2011</td>
<td>AL85/11</td>
<td>KF383980 (1,764), KF383984 (1,725)</td>
<td>16SrXIV-A</td>
</tr>
<tr>
<td>Tormancina, Italy</td>
<td>2012</td>
<td>IT35/12</td>
<td>KF383978 (1,749), KF383982 (1,718)</td>
<td>16SrXIV-A</td>
</tr>
<tr>
<td>Jarak, Serbia</td>
<td>2013</td>
<td>RS123/13</td>
<td>KJ000024 (1,748), KJ000022 (1,748)</td>
<td>16SrXIV-C</td>
</tr>
<tr>
<td>Belgrade-Zemun, Serbia</td>
<td>2013</td>
<td>RS306/13</td>
<td>KJ000021 (1,729), KJ000023 (1,748)</td>
<td>16SrXIV-C</td>
</tr>
<tr>
<td>Belgrade-Zemun, Serbia</td>
<td>2013</td>
<td>RS305/13</td>
<td>KP019340 (1,755), KP019342 (1,709)</td>
<td>16SrXIV-C</td>
</tr>
<tr>
<td>Belgrade-Zemun, Serbia</td>
<td>2013</td>
<td>RS304/13</td>
<td>KP019339 (1,763), KP019341 (1,709)</td>
<td>16SrXIV-C</td>
</tr>
</tbody>
</table>

* Length in base pairs of obtained sequences is given in parentheses

Table 2. Bermudagrass white leaf (BGWL) phytoplasmas and related strains used as representatives for phylogenetic analyses

<table>
<thead>
<tr>
<th>Straina</th>
<th>Geographic origin</th>
<th>GenBankb</th>
<th>Host</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>BGWL-C1</td>
<td>Italy</td>
<td>A550984</td>
<td>Cynodon dactylon</td>
<td>Marcone et al. 2004</td>
</tr>
<tr>
<td>M4</td>
<td>Turkey</td>
<td>HE599391</td>
<td>C. dactylon</td>
<td>Çağlar et al. 2013</td>
</tr>
<tr>
<td>FBGWL</td>
<td>Iran</td>
<td>EF444485</td>
<td>C. dactylon</td>
<td>Salehi et al. 2009</td>
</tr>
<tr>
<td>JBGWL</td>
<td>Iran</td>
<td>EF444486</td>
<td>C. dactylon</td>
<td>Salehi et al. 2009</td>
</tr>
<tr>
<td>BGWL3</td>
<td>Thailand</td>
<td>AF248961</td>
<td>C. dactylon</td>
<td>Davis and Dally 2001</td>
</tr>
<tr>
<td>BGWL-D.s.a</td>
<td>India</td>
<td>GQ403689</td>
<td>Digitaria sanguinalis</td>
<td>Rao et al. 2010</td>
</tr>
<tr>
<td>BGWL-Op.bu</td>
<td>India</td>
<td>GQ403690</td>
<td>Oplismenus burmannii</td>
<td>Rao et al. 2010</td>
</tr>
<tr>
<td>CWL</td>
<td>Australia</td>
<td>AF509321</td>
<td>C. dactylon</td>
<td>Blanche et al. 2003</td>
</tr>
<tr>
<td>GBGWLE</td>
<td>Myanmar</td>
<td>AB741630</td>
<td>C. dactylon</td>
<td>Win et al. 2013</td>
</tr>
<tr>
<td>GBGWLE2</td>
<td>Myanmar</td>
<td>AB642601</td>
<td>Chrysopogon aciculatus</td>
<td>Win and Jung 2012</td>
</tr>
<tr>
<td>BraWL-KK</td>
<td>Thailand</td>
<td>AB052872</td>
<td>Brachiaria sp.</td>
<td>Jung et al. 2003</td>
</tr>
<tr>
<td>Gorakhpur</td>
<td>India</td>
<td>EU032485</td>
<td>Cynodon dactylon</td>
<td>Snehi et al. 2008</td>
</tr>
<tr>
<td>KF-C1</td>
<td>China</td>
<td>EU377477</td>
<td>C. dactylon</td>
<td>Unpublished</td>
</tr>
</tbody>
</table>

* CWL = Cynodon white leaf phytoplasma and BraWL = Brachiaria white leaf phytoplasma.  
GenBank accession numbers for 16S ribosomal DNA.

Fig. 1. Restriction maps showing positions of restriction sites in 16S ribosomal DNA (marked by black bar) and 16S-23S spacer region sequences of strains BGWL-C1, RS59/11, and JBGWL, the reference strains of 16SrXIV-A, -C, and –B ribosomal subgroups, respectively. MseI and TruI are isoschizomers of each other.
GATCG-3') and CYNgroelR2 (5'-ACACTACTGCTGCTCCT-3') were based on a sequence ending 1 and 56 bp upstream, respectively, of the 3' end of the groEL gene. The expected PCR product sizes were approximately 1,800 bp for CYNgroesF2/CYNgroelR and 1.7 kbp for CYNgroesF2/CYNgroelR2 primer pairs.

**PCR and restriction fragment length polymorphism analyses.**

PCR assays with the universal phytoplasma primer pair P1/P7 (Deng and Hiruki 1991; Schneider et al. 1995) and the primer pairs developed in this study, CYNgroesF2/CYNgroelR and CYNgroesF2/CYNgroelR2, were performed to amplify the BGWL phytoplasma 16S rDNA and partial groEL gene, respectively. Each 25-μl PCR reaction mix contained 1 μl of diluted template DNA, 1× PCR Master Mix (Fermentas, Vilnius, Lithuania), and 0.4 μM each primer. Asymptomatic bermudagrass samples and reaction mixes lacking DNA were employed as negative controls. Thirty-five amplification cycles were performed under described conditions (Deng and Hiruki 1991), except that a 56°C annealing temperature was used for the CYNgroesF2/CYNgroelR2 primer pair. PCR products (6 μl) were separated by electrophoresis in 1% agarose gel, stained with ethidium bromide, and visualized with a UV transilluminator.

Restriction fragment length polymorphism (RFLP) analyses of P1/P7 amplicons were performed using the *HindIII, TaqI, MboI,* Tsp509I, and TruII restriction enzymes (Fermentas) under conditions described by the manufacturer. All restriction products were separated by electrophoresis in 8% polyacrylamide gel and stained and visualized as described above.

**Sequence analyses.**

Direct P1/P7-amplified products of 16S rDNA and CYNgroesF2/CYNgroelR, or CYNgroesF2/CYNgroelR2 of groEL, were purified using the mi-PCR purification kit (Metabion International AG, Martinsried, Germany). The P1/P7 amplicons were sequenced in both directions with primers P1, R16F2n (Gundersen and Lee 1996), P7, and 350R (Namba et al. 1993), while the CYNgroesF2/CYNgroelR amplicons were sequenced with the same primers used for amplification. For sequencing of the CYNgroesF2/CYNgroelR2 amplicons, in addition to the primers used for amplification, two primers, forward CYNgelFseq (5'-TACAAAGTGTTAATTCAGAC-3') and reverse CYNgelRseq (5'-GCTCCCATATTGTGATAAGG-3'), were used for sequencing to cover the complete amplicon. Sequencing was performed by a commercial service (Macrogen Inc.). The obtained sequences were assembled using Pregap4 from the Staden program package (Staden et al. 1999) and deposited in the National Center for Biotechnology Information (NCBI) under accession numbers listed in Table 1.

---

**Fig. 2.** Phylogenetic tree constructed by parsimony analyses of 16S ribosomal DNA sequences of 21 *Candidatus Phytoplasma cynodontis* and 7 other *Ca. Phytoplasma* strains, employing *Acholeplasma laidlawii* as outgroup. One of nine equally parsimonious trees is shown. Numbers on the branches are bootstrap values obtained for 1,000 replicates (only values above 70% are shown). Strains obtained and analyzed in this work are in bold.

---

1580 Plant Disease / Vol. 99 No. 11
The 16S rDNA sequences were aligned using Clustal W (Larkin et al. 2007) from the Molecular Evolutionary Genetics Analysis program MEGA6 (Tamura et al. 2013) and a search for single-nucleotide polymorphisms (SNP) was performed in the Bi edit program (Hall 1999). The 16S rDNA sequences of strains BGWL-C1 and JBGWL (GenBank accession numbers AJ550984 and EF444486, respectively), representing the 16SrXIV-A and -B subgroups, respectively, were added to the alignment, and putative restriction site maps were generated using MapDraw (DNASTAR Inc.).

Evolutionary history was inferred based on the 16S rDNA of BGWL strains sequenced in this work and 12 other representatives of BGWL phytoplasma available in the NCBI using the maximum parsimony (MP) method (MEGA6). The 12 representatives of BGWL and related phytoplasmas were selected according to their geographical origin and are listed in Table 2. Additional sequences BGWL and related phytoplasmas were selected according to their parsimony (MP) method (MEGA6). The 12 representatives of BGWL strains sequenced in this work and 12 other representatives generated using MapDraw (DNASTAR Inc.) were added to the alignment, and putative restriction site maps were generated using MapDraw (DNASTAR Inc.).

The phylogenetic tree derived by MP analysis of the 16S rDNA gene sequences showed that the analyzed ‘Ca. P. cynodontis’ strains formed a well-supported (bootstrap value of 96%) phylogenetic

Results
Analyses of 16S rDNA. The 16S rDNA and the partial spacer region of all tested BGWL phytoplasma strains were successfully amplified using the P1/P7 primer pair, confirming the presence of phytoplasma DNA. The sequences obtained from P1/P7 amplicons ranged from 1,729 to 1,768 bp and contained the 16S rDNA and 16S-23S rDNA spacer region. The search for SNP revealed no differences at any nucleotide position among the strains from Italy and Albania and showed the highest homology with the 16S rDNA sequence of ‘Ca. P. cynodontis’ reference strain BGWL-C1 (GenBank accession number AJ550984), confirming their identification as ‘Ca. P. cynodontis’, ribosomal subgroup 16SrXIV-A. In contrast, the five sequenced Serbian strains were identical to each other but different from all others. The Serbian strains showed 11 nucleotide differences from the other European strains, of which 8 were in the 16S rDNA and 3 in the 16S-23S rDNA spacer region. Compared with JBGWL (GenBank accession number EF444486), the reference strain of 16SrXIV-B, the Serbian strain and other European BGWL strains showed 10 and 8 SNP in 16S rDNA, respectively. The 16S rDNA sequence of ribosomal subgroup 16SrXIV-C (strain RS 59/11) was 99.37% and 99.33% similar to the corresponding sequences of subgroups 16SrXIV-A (strain IT71/10) and 16SrXIV-B (strain JBGWL), respectively. The 16S rDNA sequences of ribosomal subgroups 16SrXIV-A and 16SrXIV-B (strains IT71/10 and JBGWL) were 99.46% similar.

Virtual RFLP analyses indicated that five restriction enzymes (HinfI, TaqI, MboII, Tsp509I, and TruII) can differentiate the Serbian strains from all other European strains tested that belong to subgroup 16SrXIV-A, while the Serbian strains were also differentiated from strains in subgroup 16SrXIV-B (Fig. 1). The differential restriction sites for TaqI, MboII, and TruII restriction enzymes are located in the 16S rRNA gene, fulfilling conditions for description of the new ribosomal subgroup 16SrXIV-C. For confirmation, the obtained P1/P7 amplicons of the Serbian strains and IT71/10 (representative of subgroup 16SrXIV-A) were subjected to RFLP analyses with the five restriction enzymes. The Serbian strains showed identical profiles with all restriction enzymes used (data not shown). Three restriction profiles (HinfI, TaqI, and MboII) of the Serbian strain were different when compared with the corresponding restriction profiles of 16SrXIV-A BGWL strain, while the differences in Tsp509I and TruII restriction profiles, visible in virtual RFLP analyses, were hardly or not at all visible in the polyacrylamide gel run after RFLP reaction.

Fig. 3. Phylogenetic tree constructed by parsimony analyses of the groEL gene sequences of eight Candidatus Phytoplasma cynodontis and other available Ca. Phytoplasma strains, employing Acholeplasma laidlawii as outgroup. One of six equally parsimonious trees is shown. Numbers on the branches are bootstrap values obtained for 1,000 replicates (only values above 70% are shown).
group closely related to ‘Ca. P. oryzae’ (Fig. 2). Interestingly, the Serbian strains formed a well-supported (bootstrap value of 99%) separate lineage from the other ‘Ca. P. cynodontis’ strains.

**Analyses of groEL gene.** Using the CYNgroesF2/CYNgroelR primer pair in direct PCR assays, amplicons of approximately 1.8 kb were obtained with the two Italian and one Albanian BWGL phytoplasma strains employed, while no amplification with Serbian strains was obtained (data not shown). When the CYNgroesF2/CYNgroelR2 primer pair was used in direct PCR, amplicons of approximately 1.7 kb were obtained with all eight BWGL phytoplasma strains (data not shown). The three sequences obtained from CYNgroesF2/CYNgroelR and five sequences from CYNgroesF2/CYNgroelR2 amplicons ranged from 1,687 to 1,748 bp containing the 3′ end of groES gene, groES-groEL spacer region, and most of the groEL gene. The search for SNP revealed no differences at any nucleotide position between strains IT71/10 and AL85/11 from Italy and Albania, respectively. Strain IT71/10, the other one from Italy, showed one nucleotide difference (position 1,236 of groEL [C/T]), synonymous when translated, from the other two strains from Italy and Albania. Serbian strains RS304/13, RS305/13, and RS306/13 were identical in the groEL gene and showed one nucleotide difference (position 687 of groEL [C/T]), synonymous when translated, from the other two Serbian strains (RS59/11 and RS123/13), which were identical. The Serbian and Italian or Albanian strains showed 3.0 to 3.1% differences in the groEL nucleotide and 1.7% of deduced amino acid sequences.

The phylogenetic tree derived by MP analysis of the groEL gene sequences showed that ‘Ca. P. cynodontis’ strains formed a monophyletic group within the phytoplasma clade (Fig. 3). The ‘Ca. P. cynodontis’ subclade position within the phytoplasma clade is congruent with its position according to the 16SrDNA-derived phylogenetic tree (data not shown). The Serbian BWGL strains tested, as in case of 16SrDNA, clustered separately from the other European ‘Ca. P. cynodontis’ strains analyzed.

**Discussion**

The obtained results confirm earlier reports that had associated the symptoms observed on bermudagrass plants in Europe with the presence of ‘Ca. P. cynodontis’ (Marcone et al. 1997), and present the first molecular characterization of ‘Ca. P. cynodontis’ with a molecular marker other than 16SrDNA.

Sequence analyses of 16SrDNA confirmed that the 16SrXIV ribosomal group is highly supported, with a bootstrap value of 96%, and forms subclades within the ‘Ca. P. cynodontis’ clade. Strains from Serbia formed a highly supported (bootstrap value of 99%) distant lineage from all other BWGL phytoplasma strains analyzed in this study. The other cluster contains all other European strains, one strain each from Turkey and Iran, and the remaining BWGL strains from the Far East (China, India, Myanmar, Thailand, and Australia). The two Italian and Albanian strains clustered closest to the reference strain from Italy (BWGL-C1), previously described by Marcone et al. (2004). Based on the RFLP analyses of the 16S rDNA gene, the intergenic spacer region, and beginning of the 23S ribosomal gene, specific primers for groEL gene, specific primers for groEL gene amplification of all tested BWGL phytoplasmas were successfully developed and employed. The results suggest that, in agreement with previous work on other phytoplasma groups (Mitrovic et al. 2011), groEL also may be used as an additional marker gene for BWGL phytoplasmas after additional testing has been performed on other BWGL strains, including strains in the subgroup 16SrXIV-B.

The comparative sequence analyses conducted on groEL and 16SrDNA sequences substantiated the data obtained through RFLP and sequence analyses of the 16SrDNA gene. Furthermore, the groEL gene exhibited a greater sequence variation than 16SrDNA among the tested BWGL phytoplasma strains. In particular, the strains which were identical regarding the 16SrDNA, groEL revealed one nucleotide difference (synonymous when translated) in sequences of three strains from the Serbian group (RS304/13, RS305/13, and RS306/13) and one from Italy (IT71/10), whereas the Serbian strains showed more differences from other European strains in the groEL gene nucleotide sequence than in 16SrDNA (3.0 to 3.1% compared with 0.54%). These results are in agreement with data already reported for the groEL gene in asteys yellow phytoplasma strain differentiation (Mitrovic et al. 2011). The phylogenetic tree based on groEL gene sequence analysis was highly congruent with the 16SrDNA-based tree and confirmed distinct phylogenetic lineages among the BWGL phytoplasma strains analyzed. One lineage contains the strains from Italy and Albania (IT71/10, IT35/12, and AL85/11), while strains from Serbia (RS59/11, RS123/13, RS304/13, RS305/13, and RS306/13) formed a separate lineage. This separation of the two groups of strains is strongly supported by high bootstrap values. Because groEL is the only nonribosomal marker developed for BWGL phytoplasmas thus far, its further application in molecular surveys of non-European strains, including those assigned to subgroup 16SrXIV-B, should give a better insight into the relationships among BWGL phytoplasmas and correlation between differentiation and geographic distribution of their strains.

**Acknowledgments**

This work was funded as a grant from the project TR31043 of the Serbian Ministry of Education, Science, and Technological Development and supported by the German Research Foundation (DFG projects KU 2679/2-1 and BU 890/21-1), Max-Planck Society, and bilateral project of the German Academic Exchange Service (DAAD) 53626384 and the Serbian Ministry of Education, Science, and Technological Development 451-02-00446/2013-09/11.

**Literature Cited**


