

Rhizobia with different symbiotic efficiencies nodulate *Acaciella angustissima* in Mexico, including *Sinorhizobium chiapanecum* sp. nov. which has common symbiotic genes with *Sinorhizobium mexicanum*

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Acaciella angustissima; N₂ fixation; nodulation; rhizobial diversity; legume symbiosis.

Abstract

Bacteria from nodules of the legume Acaciella angustissima native to the south of Mexico were characterized genetically and their nodulation and competitiveness were evaluated. Phylogenetic studies derived from rpoB gene sequences indicated that A. angustissima is nodulated by Sinorhizobium mexicanum, Rhizobium tropici, Mesorhizobium plurifarium and Agrobacterium tumefaciens and by bacteria related to Sinorhizobium americanum, Sinorhizobium terangae, Rhizobium etli and Rhizobium gallicum. A new lineage related to S. terangae is recognized based on the sequences of gyrA, nolR, recA, rpoB and rrs genes, DNA-DNA hybridization and phenotypic characteristics. The name for this new species is Sinorhizobium chiapanecum and its type strain is ITTG S70^T. The symbiotic genes nodA and nifH were similar to those from S. mexicanum strains, which are Acaciella symbionts as well, with nodA gene sequences grouped within a cluster of nod genes from strains that nodulate plants from the Mimosoideae subfamily of the Leguminosae. Sinorhizobium isolates were the most frequently obtained from A. angustissima nodules and were among the best strains to promote plant growth in A. angustissima and to compete in interstrain nodule competition assays. Lateral transfer of symbiotic genes is not evident among the genera that nodulate A. angustissima (Rhizobium, Sinorhizobium and Mesorhizobium) but may occur among the sympatric and closely related sinorhizobia that nodulate Acaciella.

Introduction

Bacteria in the roots or the stems of legumes fix nitrogen and provide the plant with this nutrient. Symbiotic bacteria have been studied from only a small proportion of the extant legume species, and diverse genera such as *Rhizobium*, *Sinorhizobium*, *Mesorhizobium*, *Bradyrhizobium*, *Devosia*, *Methylobacterium*, *Burkholderia* and *Cupriavidus* have been reported to contain nodulating species (Young & Haukka, 1996; Sprent, 2001; Sy *et al.*, 2001; Rivas *et al.*, 2002; Chen *et al.*, 2003; Vandamme & Coenye, 2004; Elliott *et al.*, 2007a, b). Mexico is a very diverse country and occupies the fourth place in plant diversity terms (Rzedowsky, 1978) with many endemic legumes. The genus *Acaciella* is found mainly in Mexico and has well-supported botanical differences to be recognized as a new species different from the *Acacia* genus where it was formerly classified (Rico-Arce & Bachean, 2006).

Nitrogen-fixing trees and shrubs are valuable to maintain forest fertility and N_2 fixation allows their growth in infertile soils while enriching soil nitrogen. In Chiapas, *Acaciella angustissima* shrubs that can grow in poor soils are being used in agroforestry systems, due to their rapid growth rate, high capacity for nitrogen fixation and the quality of the tannins that accumulate in their bark (Rincón-Rosales & Gutiérrez-Miceli, 2008). Interestingly, these shrubs are the preferred hosts of *Llaveia mexicanorum* (Williams & MacVean, 1995), a native homeoptera scale insect, which is used by indigenous people of Chiapas and Mesoamerica to produce a fat for traditional lacquer wood handcrafts (Grillasca, 2007). We established nurseries to propagate *A. angustissima* plants and became aware that inoculants were required to attain good plant development. This prompted us to analyze and select strains for inoculation.

One of the sinorhizobial groups we encountered corresponded to a new species and we proposed the name *Sinorhizobium mexicanum* for this lineage (Lloret *et al.*, 2007). However, modifications in the *Sinorhizobium* genus taxonomy have occurred (Young, 2003). The bacteria belonging to *Sinorhizobium* have been transferred to the genus *Ensifer* (Young, 2003) because according to judicial rules, *Ensifer* has priority over *Sinorhizobium*. This new *Sinorhizobium* species had to be named as *Ensifer mexicanus* (Lloret *et al.*, 2007) instead of *S. mexicanum*. In this work, we chose to use the former name *Sinorhizobium* as used in many recently published papers.

Sinorhizobium mexicanum was not the only symbiont found in *A. angustissima* nodules. The objective of this study was to characterize the other symbionts of *A. angustissima* in Mexico (including a novel sinorhizobial species), their interstrain nodulation competitiveness and their plant growth promotion in *A. angustissima*.

Materials and methods

Sample sites

Isolates used in this study were obtained from root nodules of *A. angustissima* collected from the Sumidero Canyon National Park in Chiapas, Mexico, and from nodulated trap plants grown in pots containing soils collected from an ecological reserve area in Sierra de Huautla in Morelos, Mexico (Supporting Information, Fig. S1). The Chiapas and Morelos collecting sites were *c.* 1000 km apart and both are characterized by deciduous forest vegetation (Lloret *et al.*, 2007).

Bacterial strains

The *A. angustissima* strains analyzed in this study are listed in Table 1. Bacteria were obtained as described by Vincent (1970) using peptone yeast agar (PY) as growth medium (Toledo *et al.*, 2003). Plates were incubated aerobically at 28 °C for 3 days and the isolates were purified by streaking single colonies on fresh PY plates. Single colony formation and morphology were observed in yeast extract mannitol (YEM) and PY media at 28 °C as reported by Toledo *et al.* (2003). The acid/alkaline reaction was verified by spreading the inoculum on YEM plates (pH 7.0) containing 25 µg mL⁻¹ bromothymol blue (Vincent, 1970).

Table 1. Bacteria isolated from nodules of Acaciella angustissima

Species and strains*	Geographical origin
Agrobacterium tumefaciens	
ITTG S2	Chiapas, Mexico
ITTG S6	Chiapas, Mexico
ITTG S9	Chiapas, Mexico
ITTG S10	Chiapas, Mexico
CFN ESH11	Morelos, Mexico
CFN ESH16	Morelos, Mexico
Mesorhizobium plurifarium	
CFN ESH5	Morelos, Mexico
CFN ESH18	Morelos, Mexico
CFN ESH19	Morelos, Mexico
CFN ESH22	Morelos, Mexico
CFN ESH26	Morelos, Mexico
Rhizobium sp. (R. gallicum related)	
ITTG S11	Chiapas, Mexico
Rhizobium sp. (R. leguminosarum/R. etli rel	ated)
CFN ESH6	Morelos, Mexico
CFN ESH7	Morelos, Mexico
CFN ESH34	Morelos, Mexico
Rhizobium tropici	
CFN ESH9	Morelos, Mexico
CFN ESH10	Morelos, Mexico
CFN ESH23	Morelos, Mexico
CFN ESH25	Morelos, Mexico
CFN ESH27	Morelos, Mexico
CFN ESH29	Morelos, Mexico
ITTG S7	Chiapas, Mexico
Sinorhizobium sp. (S. americanum related)	
ITTG S8	Chiapas, Mexico
Sinorhizobium chiapanecum sp. nov.	
ITTG R11	Chiapas, Mexico
ITTG S1	Chiapas, Mexico
ITTG S68	Chiapas, Mexico
ITTG S70 ^T	Chiapas, Mexico
ITTG S71	Chiapas, Mexico
Sinorhizobium mexicanum	
CFN ESH1	Morelos, Mexico
CFN ESH2	Morelos, Mexico
CFN ESH3	Morelos, Mexico
CFN ESH4	Morelos, Mexico
ITTG R4	Chiapas, Mexico
ITTG R7 ^T	Chiapas, Mexico
ITTG S3	Chiapas, Mexico
ITTG S4	Chiapas, Mexico
ITTG S5	Chiapas, Mexico
ITTG S64	Chiapas, Mexico

*Identity according to the sequence analysis of the chromosomal gene rpoB.

Nodulation tests

Acaciella angustissima seeds were scarified with H_2SO_4 for 15 min and surface sterilized with 1% (v/v) sodium hypochlorite for 10 min. Treated seeds were germinated on 0.8% agar–water plates and then placed in glass tubes filled with vermiculite moistened with Fahraeus medium (Fahraeus, 1957). Inoculation tests were also performed with Acacia farnesiana, Leucaena leucocephala and Phaseolus vulgaris cv. Negro Jamapa as described (Lloret et al., 2007). Bacteria for inoculation were grown in individual PY plates and suspended in 1 mL sterile distilled water, serially diluted, and absorbance was determined at A_{600 nm}. Final cell numbers were determined by plating on PY medium to count CFUs. Approximately 10⁶ bacteria mL⁻¹ were added to each plant rootlet and the plants were grown in a plant growth chamber at 28 °C (Räsänen et al., 2001). A negative control with uninoculated seedlings was included. After 30 days, surfacedisinfected nodules were harvested and crushed in PY plates and single colonies were picked up and reinoculated for their authentication. Bacteria were conserved in 65% glycerol-PY broth and stored at -80 °C. Working cultures were maintained on YEM slants at 4 °C (Vincent, 1970).

DNA isolation, genomic fingerprinting and DNA-DNA hybridization

Isolates were grown overnight in 2 mL PY. Total DNA was isolated and purified using the Genomic PrepTM kit (Amersham). Enterobacterial repetitive intergenic consensus (ERIC) genomic fingerprinting was obtained by PCR using primers ERIC1R and ERIC2 as described by Versalovic et al. (1994). The fingerprints were visually analyzed after resolution of PCR products using electrophoresis in 1.5% agarose gels loaded with half the volume of the 25 μ L PCR reaction. ERIC fingerprinting was used only to confirm that the isolates analyzed were not clones or siblings (Ormeño-Orrillo et al., 2006; Lloret et al., 2007). Strains showing different patterns were considered for sequencing and phylogenetic analysis. The DNA relatedness was determined using DNA-DNA hybridization experiments using ³²Plabelled DNA of the newly proposed species (described below) Sinorhizobium chiapanecum ITTG S70^T as a probe. A filter hybridization method described previously was used (Martínez-Romero et al., 1991). The amounts of DNA were standardized using integrating gel fluorescence with the Eagle Eye II system (Stratagene). ANOVA and t-tests were performed to compare the percentage of DNA-DNA hybridization values among species and within species using angular transformation of percentage data (Knudsen & Curtis, 1947; Martínez-Romero & Rosenblueth, 1990).

PCR amplification and gene sequencing

An internal fragment of the chromosomal genes gyrA, nolR, recA, rpoB and 16S rRNA gene (rrs), and the symbiotic genes nifH and nodA were amplified using standard PCRs. Primers and annealing temperatures used for gyrA, nolR, recA, rrs, rpoB and nifH genes were performed as described in Lloret et al. (2007) and by Haukka et al. (1998) for nodA. Before sequencing, the amplification mixture was purified using

the PCR product purification system of RocheTM. The sequences generated were deposited in the GenBank public database and their accession numbers were included in the phylogenetic trees.

Phylogenetic analysis

The protein-coding sequences were aligned using the program CLUSTAL W (Thompson et al., 1994) and then aligned based on codons using DAMBE v4.2.13 (Xia & Xie, 2001). The alignments were edited with BIOEDIT v5 (Hall, 1999). The best-fit evolutive models for each set of sequences were selected by the AKAIKE information criterion implemented in the MODELTEST v3.06 (Posada & Buckley, 2004). rpoB gene sequences were analyzed using the TrN+I+ Γ model of evolution based on an alignment of 642 nucleotides from positions 3262 to 3903; nodA using the GTR+I+ Γ model with 522 nucleotides from positions 67 to 588; nifH with the TrN+I+ Γ model of evolution based on 474 nucleotides from positions 313 to 787; and for the gyrA, recA and nolR genes the model of evolution and alignment positions were as reported by Lloret et al. (2007). These positions were based on the rpoB, nolR, nodA and nifH genes of Sinorhizobium meliloti 1021 and recA and gyrA of Agrobacterium tumefaciens C58. The phylogenetic trees were inferred with the maximum-likelihood (ML) method using the program PHYML v2.4.4 (Guindon & Gascuel, 2003) considering the α -parameter for the Gamma distribution and the proportion of invariable sites estimated by the program. For the inference of the rrs phylogenetic tree, Sinorhizobium type strains were analyzed by the neighbor-joining method (NJ) (Saitou & Nei, 1987) implemented in MEGA v3.1 (Kumar et al., 2004) using the TrN+G model with the α -parameter for the Gamma distribution estimated with MODELTEST. The rrs phylogenetic tree was constructed using an alignment of 1417 nucleotides from positions 28 to 1444 with respect to the rrs gene of S. meliloti 1021. The topology robustness was estimated by a nonparametric bootstrap test using 100 pseudoreplicates for ML and 1000 for NJ.

Competition assays

The nodulation capacity was evaluated in competition assays of *S. mexicanum* ITTG $R7^{T}$ or *S. chiapanecum* ITTG $S70^{T}$ against one randomly selected strain from each of the bacterial groups identified previously by *rpoB* gene sequence analysis. Twenty-one treatments resulted from the 12 combination mixtures plus each of the eight single strains as positive nodulation controls, and the negative control (uninoculated plants). Four replicates of inoculated plants were used per treatment. The plant growth conditions were as mentioned above for nodulation tests. The competitiveness was evaluated by the number of nodules obtained from each member of the mixture with respect to the total number of nodules. The identity of the reisolated strains was determined by plasmid patterns using the Eckhardt procedure (Eckhardt, 1978). The variation in nodule number was analyzed statistically by ANOVA using SAS software (SAS Institute Inc., 1989), followed by comparison of means by Tukey's test (P < 0.05).

Plant inoculation assays

The strains with the best nodulation capacity and high competitiveness were used as inoculants. Germinated seedlings of *A. angustissima* were planted in vermiculite tubes with Fahraeus medium (Fahraeus, 1957) and inoculated as described above. Plants without inoculum, with or without 30 mg KNO₃-N per plant, served as control (Hungria *et al.*, 2001). Six replicate tubes were used per treatment and these were arranged in a completely randomized design. The plants were grown in a climate chamber at 28 °C for 90 days. At harvest, the shoot height, shoot dry weight, root dry weight and nodule number were determined, and total shoot nitrogen was assayed using the Kjeldahl method (Bremner & Mulvaney, 1982). The effect of the inoculation was analyzed statistically by ANOVA, followed by comparison of means using Tukey's test (P < 0.05).

Results

Strain identity, diversity and phylogeny

A total of 94 strains were obtained from *A. angustissima* root nodules in Chiapas and Morelos that were confirmed to form nodules in the original host. Thirty-eight strains that represented the different ERIC-PCR electrophoretic patterns were used for PCR amplification and sequencing. The taxonomic position of the selected strains from *A. angustissima* was determined according to the phylogenetic analysis performed with partial sequences of the chromosomal gene *rpoB*, which encodes the β -subunit of RNA polymerase (Fig. 1).

The largest percentage of isolates found at both sites corresponded to *S. mexicanum* (26.3%) while the lowest corresponded to bacteria related to *S. americanum* and *Rhizobium gallicum*, both with 2.6%. A new lineage related to *S. mexicanum* and *Sinorhizobium terangae* was isolated only in Chiapas while only the strains related to *Rhizobium etli* and *Mesorhizobium plurifarium* were found in Morelos. The largest percentage of the isolates in Chiapas corresponded to *S. mexicanum* (33.3%) and in Morelos *Rhizobium tropici* (30.0%).

The *Sinorhizobium* sp. strain ITTG $S70^{T}$ *rrs* gene was found to be different from all sequences available in the GenBank database and had 99% identity to its closest relative *S. mexicanum*. The phylogenetic tree with the sequences of *rrs* genes (Fig. 2) and the phylogenetic trees with *rpoB* (Fig. 1), *gyrA*, *nolR* and *recA* genes (Fig. 3) were

constructed including all of the type strains of *Sinorhizo-bium* species. The *recA* gene has been used previously in rhizobial phylogenetic studies (Gaunt *et al.*, 2001; Vinuesa *et al.*, 2005); *gyrA*, *recA*, *nolR* and *rpoB* were used previously to describe a new *Sinorhizobium* species (Lloret *et al.*, 2007). *gyrA* encodes the alpha-subunit of DNA gyrase, *nolR* encodes a transcriptional regulator (Chen *et al.*, 2000, 2005) and *recA* encodes the recombination protein RecA. In all phylogenetic trees, the position of strain ITTG S70^T as a different lineage within the *Sinorhizobium* genus was well supported. Strains ITTG R11, ITTG S68 and ITTG S71 had sequences identical to those from ITTG S70^T that was chosen to represent this new lineage.

Total DNA from the strain ITTG S70^T showed low hybridization with the strains belonging to S. mexicanum (< 42%) and S. terangae (< 56%), while hybridization to three strains from its own group, ITTG S68, ITTG S71 and ITTG R11 (> 74%), was higher than the limit proposed for new species (70%, Stackebrandt et al., 2002) (Table S2). DNA-DNA hybridization differences between S. chiapanecum strains and the closest species, S. mexicanum and S. terangae, were statistically significant with P < 0.05 and P < 0.10, respectively. It remains to be established whether similar plasmids in S. terangae and S. chiapanecum account for part of the DNA hybridization obtained. Description of species should be based on chromosomal and not plasmidic characteristics (Martínez-Romero & Jarvis, 1993). Also, phenotypic differences distinguishing S. chiapanecum, S. terangae and S. mexicanum are presented as Table S1.

The DNA–DNA hybridization, the phylogenetic position and phenotypic characteristics support that this *Sinorhizobium* lineage corresponds to a new species within the genus *Sinorhizobium*, and the proposed name is *S. chiapanecum* because it was isolated in Chiapas.

The nifH and nodA phylogenetic trees are shown in Figs 4 and 5, respectively. Symbiotic genes from the different rhizobia isolated from A. angustissima had affiliations with the corresponding genes from species in the genera Rhizobium, Sinorhizobium and Mesorhizobium. The phylogenies of these two symbiotic genes were incongruent with the phylogeny obtained with the chromosomal gene rpoB. The nodA sequences from Rhizobium sp. strains CFN ESH6 and CFN ESH34 (related to R. etli) isolated from A. angustissima were similar to the nodA gene of Rhizobium giardinii H152^T isolated from common bean in France, but with the nifH gene analysis these strains were found to be related to the nifH gene of R. etli bv. mimosae Mim2 isolated from Mimosa affinis in Mexico. The nodA and nifH genes of R. tropici strains CFN ESH23, CFN ESH25, CFN ESH10, CFN ESH29 and CFN ESH9 were related but not identical to nodA and nifH gene sequences from R. tropici CFN299 isolated from P. vulgaris in Mexico. The symbiotic gene sequences of S. chiapanecum and S. mexicanum isolated



Fig. 1. Phylogenetic tree estimated using the ML method with partial sequences of the chromosomal protein encoding gene *rpoB* using the PHYML program. The alignment length was 642 nucleotides from positions 3262 to 3903 of the *rpoB* gene of *Sinorhizobium meliloti* 1021. Only bootstrap values \geq 50% are shown. Type strains are indicated by superscript T. The *Acaciella angustissima* strains are shown in bold. Only haplotypes were included in each terminal branch. The accession numbers for the sequences are indicated within parentheses. Those generated in this work are shown in bold.

from *A. angustissima* clustered together and were related to a different and well-supported group that included mainly sequences from *Sinorhizobium* isolated from American legumes, among them, the strains *Sinorhizobium* sp. BR827

and BR816 from *L. leucocephala* in Brazil and *S. americanum* CFN EI156 isolated from *Acacia acatlensis* in Mexico. The *nodA* and *nifH* gene sequences from *S. terangae*, the closest relative of *S. mexicanum* according to *rpoB* gene sequences,







grouped in a far distant cluster. Mesorhizobium plurifarium isolated from A. angustissima has nodA and nifH gene sequences similar to several Mesorhizobium species isolated from American and African hosts, mainly with the strains Mesorhizobium sp. DWO366 isolated from Acacia polyacantha in Kenya and Mesorhizobium sp. INPA78b isolated from L. leucocephala in Brazil.

Nodulation and nodule occupancy in competition assays

Nodule occupancy evaluated from interstrain competition assays is shown in Table 2. The strains ITTG R7^T, ITTG S70^T, CFN ERSH34, CFN ERSH5 and ITTG S7 showed the best nodulation capacity and high competitiveness.



Fig. 2. Phylogenetic tree estimated by the NJ method with the sequences of *rrs* genes using the Tamura Nei model considering gamma (α -parameter = 0.9271). The alignment length was 1416 nucleotides, from positions 28 to 1444 of the *rrs* gene of *Sinorhizobium meliloti* 1021. Only bootstrap values \geq 50% are shown. Type strains are indicated by superscript T. *Sinorhizobium chiapanecum* is shown in bold.

Sinorhizobium mexicanum strain ITTG R7^T always had a greater occupancy of the nodules than the respective competing strain, ranging from 65% when combined with Sinorhizobium sp. ITTG S8 to 100% when combined with Rhizobium sp. ITTG S11. The S. chiapanecum strain ITTG S70^T did not always have a greater occupancy of the nodules than the competing strain, although this strain and S. mexicanum ITTG R7 were highly effective in inoculation assays with A. angustissima (Table 3). Mesorhizobium plurifarium CFN ESH5 and Rhizobium sp. CFN ESH34 had a greater occupancy than ITTG S70^T (67% and 77%), respectively. Significantly lower numbers of nodules were obtained with M. plurifarium CFN ESH5, R. tropici ITTG S7, Sinorhizobium sp. ITTG S8 (related to S. americanum), Rhizobium sp. ITTG S11 (related to R. gallicum) and A. tumefaciens ITTG S2, with the latter showing the lowest number of nodules and very low nitrogen fixation. All reisolated strains showed colony morphology and plasmid patterns identical to the original inoculated strains (data not shown).

Plant growth, nodulation and nitrogen fixation of *A. angustissima* inoculated with selected strains

The inoculation using the selected rhizobia strains had a significant effect on the growth of *A. angustissima* (Table 3).

Rhizobium sp. CFN ESH34, S. mexicanum ITTG R7^T and S. chiapanecum ITTG $S70^{T}$ had a positive effect on shoot height, shoot dry weight and root dry weight compared with the uninoculated control plants and those with added KNO₃. Plants inoculated with these strains were on average 8.3 cm taller and weighted 109 mg more than noninoculated plants 90 days postinoculation. The number of nodules obtained with S. mexicanum ITTG R7^T and S. chiapanecum ITTG S70^T was significantly different (P < 0.05) compared with the rest of the treatments. None of the noninoculated plants formed nodules. The plants inoculated with ITTG R7^T showed a significantly higher total shoot nitrogen compared with other treatments (P < 0.05). ITTG R7^T was found to be the most effective strain in terms of plant growth promotion as indicated by total plant nitrogen content.

Characteristics of S. chiapanecum sp. nov.

Sinorhizobium chiapanecum (chia.pa.ne'cum. N.L. neut. adj. chiapanecum of Chiapas, the name of a state in Mexico where the bacterium was isolated). Gram-negative, aerobic, motile and nonspore-forming rods. Strains are fast growing and acid producers in YEM medium. The generation time for ITTG S70^T in YEM broth is 2.33 h at 28 °C. Colonies on PY or YEM are circular, pearly, slightly translucent and



Fig. 3. ML phylogenetic trees of partial sequences of the chromosomal protein encoding genes *gyrA*, *recA* and *nolR*. Type strains are indicated by superscript T. Only bootstrap values \geq 50% are shown. The accession numbers for the sequences are indicated within parentheses. Those generated in this work for *Sinorhizobium chiapanecum* are shown in bold. The branches corresponding to the outgroup sequences *Rhizobium leguminosarum* USDA 2370^T and *Rhizobium etli* CFN42^T for the *nolR* tree are not shown because they were too divergent from the *Sinorhizobium* sequences.

produce copious amounts of polysaccharides. Colonies are normally more than 2–4 mm in diameter within 2 days of incubation at 28 °C. The strains are resistant to nalidixic acid ($120 \,\mu g \, m L^{-1}$) but not to carbenicillin ($20 \,\mu g \, m L^{-1}$), ampicillin $(10 \,\mu\text{g mL}^{-1})$ or chloramphenicol $(10 \,\mu\text{g mL}^{-1})$. They grow in media containing 0.5%, 1.0% and 2.0% NaCl but not with 3.0% NaCl. Total DNA from strain ITTG S70^T showed low hybridization values with the strains belonging



Fig. 4. Phylogenetic tree estimated using the ML method with partial sequences of the symbiotic protein encoding *nifH* gene using the PHYML program. The alignment length was 474 nucleotides from positions 313 to 786 of the *nifH* gene with respect to the *nifH* gene encoded on the pSymA of *Sinorhizobium meliloti* 1021. Only bootstrap values \geq 50% are shown. Type strains are indicated by superscript T. The *Acaciella angustissima* strains are shown in bold. The accession numbers for the sequences are indicated within parentheses. Those generated in this work are shown in bold. Host and geographical origin are in parentheses.

to *S. terangae* ORS1009^T (< 48%) and with *S. mexicanum* ITTGR7^T (< 33%). This species can be differentiated from other described *Sinorhizobium* species on the basis of the phylogenetic analysis of the chromosomal genes *rrs*, *gyrA*, *recA*, *rpoB* and *nolR*. The type strain ITTG S70^T was isolated from nodules of *A. angustissima* collected in the Sumidero

Canyon National Park, Chiapas, Mexico. *Sinorhizobium chiapanecum* ITTG S70^T nodulated and fixed nitrogen in *A. angustissima, Acaciella cochliancantha, Acaciella farnesiana, Acaciella pennatula, Dolichos lablab, P. vulgaris, L. leucocephala* and *Lysiloma acapulcensis* and tolerated salinity and acidity (data not shown). ITTG S70^T has characteristics of the species.



0.1

Fig. 5. Phylogenetic tree estimated using the ML method with partial sequences of the symbiotic protein encoding the *nodA* gene using the PHYML program. The alignment length was 522 nucleotides from positions 67 to 588 of the *nodA* gene with respect to the *nodA* encoded on the pSymA of *Sinorhizobium meliloti* 1021. Only bootstrap values \geq 50% are shown. Type strains are indicated by superscript T. The *Acaciella angustissima* strains are shown in bold. The accession numbers for the sequences are indicated within parentheses. Those generated in this work are shown in bold. Host and geographical origin are in parentheses.

		Nodule occupancy (%) by			
Treatments	Nodule number (per plant) (\pm SD)*	First strain of the combination	Second strain of the combination		
Uninoculated	0 ^h				
A. tumefaciens ITTG S2	0.5 (±1.0) ^{gh}				
<i>M. plurifarium</i> CFN ESH5	$8.0 \ (\pm 1.6)^{bcde}$				
Rhizobium sp. ITTG S11	$2.0~(\pm 0.8)^{fgh}$				
Rhizobium sp. CFN ESH34	$10.5 (\pm 2.5)^{abc}$				
R. tropici ITTG S7	$4.0 (\pm 1.6)^{defgh}$				
Sinorhizobium sp. ITTG S8	$2.25 (\pm 2.1)^{fgh}$				
S. chiapanecum ITTG S70 ^T	$10.0 (\pm 1.6)^{abc}$				
S. mexicanum ITTG $R7^{T}$	$14.0(\pm 3.7)^{a}$				
S. chiapanecum ITTG S70 ^T +A. tumefaciens ITTG S2	$4.5~(\pm 1.3)^{defgh}$	61 (18) [†]	39		
S. chiapanecum ITTG S70 ^T +M. plurifarium CFN ESH5	$4.5~(\pm 2.6)^{defgh}$	33 (18)	67		
S. chiapanecum ITTG S70 ^T +Rhizobium sp. ITTG S11	$4.25~(\pm 1.7)^{defgh}$	82 (17)	18		
S. chiapanecum ITTG S70 ^T +Rhizobium sp. CFN ESH34	$3.25~(\pm 1.9)^{efgh}$	23 (13)	77		
S. chiapanecum ITTG S70 ^T +R. tropici ITTG S7	$2.5~(\pm 1.3)^{fgh}$	80 (10)	20		
S. chiapanecum ITTG S70 ^T +Sinorhizobium sp. ITTG S8	$3.75~(\pm 2.1)^{defgh}$	67 (15)	33		
S. mexicanum ITTG R7 ^T +A. tumefaciens ITTG S2	$6.0 \ (\pm 1.4)^{cdef}$	75 (24)	25		
S. mexicanum ITTG R7 ^T +M. plurifarium CFN ESH5	$8.75~(\pm 5.1)^{abcd}$	89 (35)	11		
S. mexicanum ITTG R7 ^T +Rhizobium sp. ITTG S11	5.5 $(\pm 1.9)^{cdefg}$	100 (22)	0		
S. mexicanum ITTG R7 ^T +Rhizobium sp. CFN ESH34	$12.0~(\pm 1.4)^{ab}$	71 (48)	29		
S. mexicanum ITTG R7 T +R. tropici ITTG S7	$3.75~(\pm 1.3)^{defgh}$	67 (15)	33		
S. mexicanum ITTG R7 ^T +Sinorhizobium sp. ITTG S8	$6.5 (\pm 3.4)^{cdef}$	65 (26)	35		

Table 2. Nodule occupancy by strains of *Sinorhizobium mexicanum* ITTG R7^T and *Sinorhizobium chiapanecum* ITTG S70^T and the coinoculated bacteria in competition assays in *Acaciella angustissima*

*Mean values of four replicates. The means followed by the same letter are not significantly different (P < 0.05). [†]In parentheses, total number of nodules analyzed.

Table 3.	Effect of	of inoculatior	n by the	e strains	with high	o competitivity	and	nodulation	capacity	on th	e growth,	nodulation	and	nitrogen	fixation	of
Acaciella	angusti	ssima														

Strains	Shoot height (cm)	Shoot dry weight (mg)	Root dry weight (mg)	Nodule number	Total shoot N (mg per plant)	
Uninoculated	15.0 c*	76.1 b	46.3 b	0 b	30.4 c	
M. plurifarium CFN ESH5	16.5 bc	95.0 b	40.3 a	2.1 b	39.9 c	
Rhizobium sp. CFN ESH34	20.1 b	101.0 b	44.9 a	2.3 b	51.5 c	
R. tropici ITTG S7	17.0 bc	96.4 b	42.9 a	2.3 b	44.3 c	
<i>S. chiapanecum</i> ITTG S70 ^T	24.8 a	112.9 a	44.0 a	5.3 a	101.6 b	
<i>S. mexicanum</i> ITTG R7 ^T	25.1 a	134.7 a	52.6 a	5.8 a	158.9 a	
KNO ₃ -N (30 mg per plant)	15.3 c	56.1 c	25.3 b	0 b	22.4 c	

*Mean values of six replicates. The means followed by the same letter are not significantly different (P < 0.05).

Discussion

Tropical forests in Mexico harbor many endemic plants and a high richness of species (Rzedowsky, 1978). Forests have abiotic and biotic characteristics that allow such diversity to exist. Plant speciation in Mexico seems to be driven by geographical isolation due to the complex topography of the country. The tropics have a large diversity of rhizobia (Wang *et al.*, 1999; Mohamed *et al.*, 2000; Räsänen *et al.*, 2001; Toledo *et al.*, 2003; Wolde-Meskel *et al.*, 2004). Sinorhizobia seem to have radiated in Mexico in relation to the geographical isolation and diversity of climates, conditions and plants (Toledo *et al.*, 2003; Lloret *et al.*, 2007). The sinorhizobia-nodulating legumes in Africa and in the Americas are considered to have had a long period of diverging evolution (Haukka *et al.*, 1998; Toledo *et al.*, 2003; Lloret *et al.*, 2007). Our results showed that *A. angustissima* was preferentially nodulated by closely related members of the *Alphaproteobacteria*, especially sinorhizobia. Differences in symbiotic efficiency and competitiveness were found among the isolates, with *S. mexicanum* and *S. chiapanecum* strains being highly effective symbionts and good competitors. In contrast to acacias, no bradyrhizobia or *Betaproteobacteria* strains were found nodulating this legume. *Acaciella angustissima* was among the legume hosts of Latin American origin that formed nitrogen-fixing nodules with the African sinorhizobial strains *Sinorhizobium arboris* HAMBI 1552^T, *Sinorhizobium kostiense* HAMBI 1489^T and *S. terangae* bv. *acaciae* ORS 1058 (Räsänen *et al.*, 2001). Tropical legumes seem to have a mild specificity when associating with nodulating bacteria (Moreira *et al.*, 1998), although under natural conditions predominant rhizobial species may be preferentially encountered in promiscuous plants (Bala & Giller, 2001; Bala *et al.*, 2003; Martínez-Romero, 2003) as shown here.

Mesorhizobium plurifarium strains originally isolated from Acacia senegal (de Lajudie et al., 1998) encompass a set of diverging strains. In this study, M. plurifarium strains were found in A. angustissima only in Morelos. In Mexico, M. plurifarium were found nodulating Sesbania (Papilioniodeae) trees (Wang et al., 1999) and L. leucocephala plants grown in Morelos soils. Acaciella and Leucaena belong to the Mimosoideae subfamily of the Leguminosae. Plant traps with soils from Morelos were used to collect the bacteria and it has been shown that by doing so a larger diversity of bacteria may be obtained nodulating a single legume (Hungria et al., 2001), and so we predicted that Mesorhizobium strains were the less adapted to nodulate A. angustissima. This turned out to be true.

We found seven isolates of *Rhizobium* similar to *R. tropici* type A, with *nod* genes more closely related (but not identical) to *nodA* of *R. tropici* than to other *nodA* genes. *Rhizobium tropici* strains are common in tropical soils and nodulate some trees from the Mimosoideae subfamily of the *Leguminosae* such as *L. leucocephala* (Martínez-Romero *et al.*, 1991) as well as *A. angustissima* (not shown).

Rhizobium etli is commonly isolated from *P. vulgaris* (Segovia *et al.*, 1993), but has also been isolated from other shrub legumes in Kenya (Odee *et al.*, 2002). Biovars that refer to host specificity have been described in *R. etli*. Nodulation of *Mimosoidea* plants such as *M. affinis* and *Leucaena* spp. is the characteristic of biovar mimosae (Wang *et al.*, 1999). It is probable that the *Rhizobium* sp. strains (related to *R. etli* and *R. leguminosarum*) from *A. angustissima* correspond to biovar mimosae. Strain ITTG S11 was found to be related to *R. gallicum* (Amarger *et al.*, 1997). *Rhizobium gallicum* by. gallicum was isolated from common bean and can also nodulate *L. leucocephala* (Amarger *et al.*, 1997; Silva *et al.*, 2005) and other species from the Mimosoideae subfamily of the *Leguminosae* (Zurdo-Piñeiro *et al.*, 2004) but it was not known that it nodulated *Acaciella*.

In addition, species of *Agrobacterium* were also found in this study. Bala & Giller (2001) reported that the legumes *Acacia auriculiformis*, *L. leucocephala*, *Gliricidia sepium*, *P. vulgaris* and *Sesbania sesban* formed effective nodules with one or more isolates that resembled A. tumefaciens. Agrobacterium strains have been isolated previously from nodules of Acacia mellifera, A. polyacantha, Acacia nilotica and S. sesban (Khbaya et al., 1998; de Lajudie et al., 1999) and shrubs growing in the semi-arid and arid climates of northwestern China (Tan et al., 1999). Odee et al. (2002) indicated that agrobacteria were often found in association with root nodules as a co-occupant with rhizobia. The Agrobacterium strains described here were capable of forming nodules on A. angustissima, but the nitrogen fixation was very low. Agrobacterium sp. ITTG S2 (similar to A. tumefaciens) showed a low level of competitiveness when inoculated in competition assays. Recently, some Agrobacterium strains were found to be capable of forming tumors on plants as well as nodulating (Rivas et al., 2004). In additional experiments, we evaluated the pathogenicity of the strains ITTG S2, ITTG S6 and ITTG S10 (all similar to A. tumefaciens) on sunflower plants (Helianthus annus) and found that these strains are not tumorogenic (not shown).

We showed that the phylogenies of the symbiotic genes were incongruent with the chromosomal genes as has been reported previously (Haukka et al., 1998; Wernegreen & Riley, 1999; Laguerre et al., 2001; Toledo et al., 2003; Lloret et al., 2007). Symbiotic genes on elements such as plasmids and symbiotic islands are prone to lateral gene transfer (Sullivan & Ronson, 1998; Ochman & Moran, 2001) and may be selected by hosts (Ueda et al., 1995; Haukka et al., 1998; Wernegreen & Riley, 1999), as observed here, because the two species S. mexicanum and S. chiapanecum nodulating Acaciella have the same nodA genes. The three main groups described based on nod gene sequences (Haukka et al., 1998) corresponding to African and Latin-American sinorhizobia and some Mesorhizobium spp. were observed in the trees presented here with several more sequences included (Fig. 5). A large group was distinguished that corresponds to nod genes of symbionts with the capacity to nodulate many plants from the Mimosoideae subfamily of the Leguminosae (Fig. 5); it is worth noticing that within this group, R. giardinii and R. gallicum nod gene sequences were included.

Sinorhizobium sp. ITTG S8, a strain related to S. americanum, clustered in the *rpoB* tree with some American strains isolated from L. leucephala in Mexico (Wang et al., 1999) and with strain BR816 (van Rhijn et al., 1994) from Brazil. This group constitutes a sister clade to S. americanum and could have been identified as belonging to the same species, but unpublished DNA–DNA hybridization results from our lab showed that BR816 was not a member of S. americanum. A new biovar has been proposed (mediterranense) (Mnasri et al., 2007) to account for sinorhizobia closely related to Sinorhizobium fredii and with specificity for L. leucocephala and P. vulgaris. This biovar includes strain BR816 and some other strains that, despite being closely related to S. fredii, do not form nodules on soybean. In spite of the close relatedness of the symbiotic genes of bv. mediterranense and *S. americanum* to those from *S. mexicanum* and *S. chiapanecum*, we consider that *A. angustissima* symbionts would not correspond to biovar mediterranense because the isolates that we found to be closely related to biovar mediterranense were not efficient to nodulate *A. angustissima*, comprised only 2.6% of the original isolates and were outcompeted by *S. mexicanum* or *S. chiapanecum*.

Within the enlarged set of sequence data presented here, we observed that the *nodA* gene sequence from *Rhizobium huautlense* (not reported previously) forms a clade with other *Sinorhizobium* species nodulating *Sesbania* (Fig. 5), indicating the strong specificity for *Sesbania* nodulation and evidencing lateral transfer of symbiotic genes between *Rhizobium* and *Sinorhizobium*. The genetic coherence among symbiotic and chromosomal genes has been considered to be characteristic of rhizobia nodulating wild legumes (Wernegreen & Riley, 1999), but *S. chiapanecum* and *S. mexicanum* as well as *R. huautlense* and sinorhizobia from biovar sesbaniae, all from noncultivated hosts, do not follow this observation and show evidence of horizontal transfer of symbiotic genes.

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Authors' contribution

R.R.-R. and L.L. contributed equally to this work.

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Supporting Information

Additional Supporting Information may be found in the online version of this article:

Figure S1. Map of Mexico showing the location of field collection sites in Chiapas and Morelos.

Table S1. Phenotypic characteristics of *Sinorhizobium chiapanecum* strain ITTG S70^T and related reference strains.

Table S2. Levels of total DNA-DNA relatedness as percent of hybridization of *Sinorhizobium chiapanecum* strain ITTG $S70^{T}$ isolated of *A. angustissima* with the *S. mexicanum* and *S. terangae* strains.

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