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Abstract	In this work we analysed dit Lupinus albus (white lupin) 16S-23S intergenic spacers genus Bradyrhizobium. Mor some strains isolated on mai divided into two ITS subgro Ornithopus. The remaining (American continent) formit ISLU207 isolated from the C close to different species of gene showed that all strains four different nodC lineages isolated from different conti	fferent chromosomal and symbiotic markers in rhizobial strains nodulating in several continents. Collectively the analysis of their <i>rrs</i> and <i>atpD</i> genes, and (ITS), showed that they belong to at least four chromosomal lineages within the st isolates from the Canary Islands (near to the African continent) grouped with nland Spain and were identified as <i>Bradyrhizobium canariense</i> . These strains are oups coincident with those previously described from isolates nodulating strains isolated on mainland Spain grouped with most isolates from Chile ng a new lineage related to <i>Bradyrhizobium japonicum</i> . The strains BLUT2 and Canary Islands and Chile, respectively, formed two new lineages phylogenetically <i>Bradyrhizobium</i> depending on the marker analyzed. The analysis of the <i>nodC</i> nodulating <i>L. albus</i> belong to the biovar genistearum; nevertheless they form s of which lineage C is at present exclusively formed by <i>L. albus</i> endosymbionts inents.
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ORIGINAL PAPER

Strains nodulating *Lupinus albus* on different continents belong to several new chromosomal and symbiotic lineages within *Bradyrhizobium*

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6 Álvaro Peix · Inne Gantois · José M. Igual · Milagros León-Barrios ·
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10 Abstract In this work we analysed different chro-11 mosomal and symbiotic markers in rhizobial strains nodulating Lupinus albus (white lupin) in several 12 13 continents. Collectively the analysis of their rrs and 14 atpD genes, and 16S-23S intergenic spacers (ITS), 15 showed that they belong to at least four chromosomal 16 lineages within the genus Bradyrhizobium. Most 17 isolates from the Canary Islands (near to the African 18 continent) grouped with some strains isolated on 19 mainland Spain and were identified as Bradyrhizobi-20 um canariense. These strains are divided into two ITS 21 subgroups coincident with those previously described from isolates nodulating Ornithopus. The remaining 22

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strains isolated on mainland Spain grouped with most 23 isolates from Chile (American continent) forming a 24 new lineage related to Bradyrhizobium japonicum. 25 The strains BLUT2 and ISLU207 isolated from the 26 Canary Islands and Chile, respectively, formed two 27 new lineages phylogenetically close to different 28 species of Bradyrhizobium depending on the marker 29 analyzed. The analysis of the *nodC* gene showed that 30 all strains nodulating L. albus belong to the biovar 31 genistearum; nevertheless they form four different 32 nodC lineages of which lineage C is at present 33 exclusively formed by L. albus endosymbionts iso-34 lated from different continents. 35

Keywords Bradyrhizobium · Lupinus · Phylogeny 36

Introduction

Lupinus albus (white lupin) is a legume which has been 40 cultivated in Europe for the last 2000 years, used in 41 human and animal feeding, as green manure in 42 agriculture (Rosolem et al. 2002; Jensen et al. 2004) 43 and in soil stabilization (Clapham 1997). This species is 44 currently considered a good alternative as an animal 45 foodstuff due to the high quality of its proteins (Erbas 46 et al. 2005). Therefore there is increasing interest in this 47 plant to be used in sustainable agriculture due to its high 48 potential to provide protein without nitrogen fertiliza-49 tion (Robinson et al. 2000; Dijkstra et al. 2003), 50 estimated at 150-200 kg of nitrogen per ha in symbiosis 51

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,	Journal : Medium 10482	Dispatch : 28-1-2010	Pages : 14	
	Article No. : 9415	□ LE	□ TYPESET	
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52 with Bradyrhizobium (Robinson et al. 2000). Despite 53 the interest of this symbiosis there are few studies about the identity of strains nodulating L. albus in different 54 55 continents whereas several studies have been carried 56 out with other Lupinus species (Barrera et al. 1997; Stepkowski et al. 2005; Andam and Parker 2007; 57 Stepkowski et al. 2007). Although L. albus may be also 58 59 nodulated by strains of Ochrobactrum this symbiosis 60 was not very effective (Trujillo et al. 2005). The strains isolated to date from effective nodules of L. albus in 61 62 different countries belong to the genus Bradyrhizobium 63 (Barrera et al. 1997; Jarabo-Lorenzo et al. 2003; 64 Stepkowski et al. 2007; Rivas et al. 2009). However 65 these strains have not been extensively analysed and only for a few strains have the same genes been studied. 66 For example, the ARDRA profiles of Canary Island 67 68 isolates were previously analysed but not their rrs gene 69 sequences. Of the Chilean strains, this sequence is only 70 known in the strains ISLU227 and ISLU207 (Jarabo-71 Lorenzo et al. 2003). Several housekeeping genes have 72 been analysed for the strains isolated in mainland Spain 73 (Stepkowski et al. 2007; Rivas et al. 2009) but the rrs 74 gene and the 16S-23S intergenic spacer (ITS) have not 75 been previously analysed. Finally, the nodC gene has been only analysed in the strain ISLU207 (Jarabo-76 77 Lorenzo et al. 2003).

78 Therefore the aim of this study was to analyse the 79 phylogenetic relationships of L. albus strains isolated on 80 three different continents using three chromosomal markers with different rates of evolutionary divergence 81 82 (the rrs and atpD genes and the ITS spacer) and a 83 symbiotic marker, the nodC gene, related with the host range of legumes (Roche et al. 1996; Perret et al. 2000, 84 Rivas et al. 2006; Laranjo et al. 2008; Laranjo et al. 85 86 2009). The results showed that the L. albus endos-87 ymbionts belong to several chromosomal lineages within 88 the genus Bradyrhizobium that could represent new 89 species of this genus. A group of these endosymbionts 90 constitute a *nodC* lineage that could represent an allelic 91 group present up to date only in L. albus bradyrhizobia.

92 Materials and methods

93 Strains and nodulation experiments

94 The reference and the L. albus strains analysed in this 95 study are listed in Table 1. These strains were isolated from L. albus nodules in previous studies according to 96

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the method of Vincent (1970). For nodulation exper-97 iments L. albus plants were inoculated with represen-98 tative isolates, under growth chamber conditions, in 99 modified Leonard jars using vermiculite as substrate 100 and nitrogen-free nutrient solutions. Non-inoculated 101 nitrogen-free and nitrogen-supplemented plants were 102 used as controls. Five replicates were set per treatment 103 and plants were harvested 6 weeks after planting. Shoot 104 dry weight and number of nodules were the parameters 105 measured. Symbiotic efficiency was determined as 106 described by Somasegaran and Hoben (1994). Data 107 were analyzed by one-way analysis of variance, and 108 mean values compared by Fisher's Protected LSD test 109 (Least Significant Differences) ($P \le 0.05$). 110

RAPD fingerprinting

RAPD patterns were obtained using the primer M13 112 (5'-GAGGGTGGCGGTTCT-3') according to Rivas 113 et al. (2006) in the following PCR conditions: 114 preheating at 95°C for 9 min; 35 cycles of denaturing 115 at 95°C for 1 min; annealing at 45°C for 1 min and 116 extension at 75°C for 2 min, and a final extension at 117 72°C for 7 min. The PCR products were electropho-118 resed on 1.5% agarose gel in TBE buffer (100 mM 119 Tris, 83 mM boric acid, 1 mM EDTA, pH 8.5) at 120 6 V/cm, stained in a solution containing 0.5 µg/ml 121 ethidium bromide, and photographed under UV light. 122 Standard VI (Roche, USA) was used as molecular 123 weight marker. An 8 µl aliquot of loading solution 124 (40% sucrose and 0.25% bromophenol blue) was 125 added to each sample. The bands present in each 126 profile were coded for input into a database including 127 all the strains studied and Jaccard's similarity coeffi-128 cient was calculated to construct the distance matrix. A 129 dendrogram was constructed from the distance matrix 130 using the unweighted pair group with arithmetic mean 131 (UPGMA) using the GelCompar II program from 132 Bionumerics platform. 133

Analysis of <i>rrs</i> , <i>atpD</i> and <i>nodC</i> genes	134
and 16S-23S intergenic spacer (ITS)	135

The rrs was amplified and sequenced according to Rivas 136 et al. (2007a), the *atpD* gene according to Gaunt et al. 137 (2001) and the ITS as described by Willems et al. 138 (2003). A partial sequence of the nodC gene was 139 obtained by using the primers designed in this study 140 NodCBradyF (5'-CGCAAGGCGCAG(AT)TCGC-3') 141

2	Journal : Medium 10482	Dispatch : 28-1-2010	Pages : 14
	Article No. : 9415	🗆 LE	□ TYPESET
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Strain	Geographical location	Source or nodulated hosts	Reference	RAPD pattern	ITS group	nodC group
<i>B. japonicum</i> bv glycinearum LMG 6138 ^T	China	Glycine max	Vinuesa et al. (2005a)	А		NA
<i>B. japonicum</i> by genistearum BGA-1	Canary Islands (Spain)	Teline stenopetala	Vinuesa et al. (2005a)	В	III	А
B. canariense BTA-1 ^T	Canary Islands (Spain)	Chamaecytisus proliferus	Vinuesa et al. (2005a)	С	I*	D
Bradyrhizobium genosp. αBC-C1	Canary Islands (Spain)	Chamaecytisus proliferus	Vinuesa et al. (2005a)	D	IV	А
Bradyrhizobium genosp. βBRE-1	Canary Islands (Spain)	Teline canariense	Jarabo-Lorenzo et al. (2003)	Е		В
RLA08	Léon (Spain)	L. albus	Rivas et al. (2009)	F	III	А
RLA09	Léon (Spain)	L. albus	Rivas et al. (2009)	G	III	А
RLA10	Léon (Spain)	L. albus	Rivas et al. (2009)	Н	III	А
RLA11	Léon (Spain)	L. albus	Rivas et al. (2009)	Ι	III	А
MCLA07	Salamanca (Spain)	L. albus	Rivas et al. (2009)	J	I*	D
MCLA12	Salamanca (Spain)	L. albus	Rivas et al. (2009)	Κ	II*	D
MCLA22	Salamanca (Spain)	L. albus	Rivas et al. (2009)	L	I*	С
MCLA23	Salamanca (Spain)	L. albus	Rivas et al. (2009)	М	II*	D
BLUT1	Canary Islands (Spain)	L. albus	Jarabo-Lorenzo et al. (2003)	Ν	I*	D
BLUT2	Canary Islands (Spain)	L. albus	Jarabo-Lorenzo et al. (2003)	0		В
BLUT3	Canary Islands (Spain)	L. albus	Jarabo-Lorenzo et al. (2003)	Р		С
BLUT5	Canary Islands (Spain)	L. albus	Jarabo-Lorenzo et al. (2003)	Q	I*	С
BLUT6	Canary Islands (Spain)	L. albus	Jarabo-Lorenzo et al. (2003)	R	I*	С
BLUT8	Canary Islands (Spain)	L. albus	Jarabo-Lorenzo et al. (2003)	S	I*	D
ISLU203	Cautín (Chile)	L. albus	Jarabo-Lorenzo et al. (2003)	Т	III	С
ISLU207	Cautín (Chile)	L. albus	Jarabo-Lorenzo et al. (2003)	U	IV	В
ISLU213	Cautín (Chile)	L. albus	Jarabo-Lorenzo et al. (2003)	v	III	А
ISLU220	Valdivia (Chile)	L. albus	Jarabo-Lorenzo et al. (2003)	W	III	А
ISLU227	Valdivia (Chile)	L. albus	Jarabo-Lorenzo et al. (2003)	Х	III	А

Table 1 Representative characteristics of reference strains and strains isolated from Lupinus albus

* According Safronova et al. (2007)

and NodCBradyR (5'-GG(GT)GTG(AGC)AGCG(AC) 142 GAAGCCG-3') in the following PCR conditions: pre-143 heating at 95°C for 9 min; 35 cycles of denaturing at 144 145 95°C for 1 min; annealing at 45°C for 1:30 min and extension at 72°C for 1 min, and a final extension at 146 72°C for 7 min. The sequences were obtained in an 147 148 ABI377 sequencer (Applied Biosystems Inc.) using a BigDye terminator v3.0 cycle sequencing kit as 149 supplied by the manufacturer. The sequences obtained 150 151 were compared with those from GenBank using the BLASTN program (Altschul et al. 1990). Sequences 152 153 were aligned using the Clustal W software (Thompson 154 et al. 1987). The distances were calculated according to Kimura's two-parameter method (Kimura 1980). 155

Phylogenetic trees were inferred using the neighbour-
joining method (Saitou and Nei 1987). Bootstrap
analysis was based on 1,000 resamplings. The MEGA1564 package (Tamura et al. 2007) was used for all analyses.
Genbank Accession numbers for the sequences deter-
mined in this study are provided in Table 2.161

Results and discussion

RAPD fingerprinting is a useful tool for genetic 164 diversity analysis of rhizobia allowing the selection 165

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•	Journal : Medium 10482	Dispatch : 28-1-2010	Pages : 14	
	Article No. : 9415	🗆 LE	□ TYPESET	
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Strains	rrs	ITS	atpD	nodC
MCLA07	EF694770	EF694745	FM253158, AM168301	EF694751
MCLA12	EF694741	EF694746	FM253159	EF694752
MCLA22	EF694742	EF694747	FM253160	EF604753
MCLA23	EF694743	EF694748	FM253161	EF694754
BLUT1	EU333379	EU333383	AM168276	EU333389
BLUT2	GQ863574	GQ863558	GU338032	GQ863566
BLUT3	GQ863575	GQ863559	GU338033	GQ863567
BLUT5	GQ863568	GQ863552	GU338034	GQ863560
BLUT6	GQ863569	GQ863553	*	GQ863561
BLUT8	GQ863570	GQ863554	*	GQ863562
RLA08	EF694744	EF694749	FM253166, AM168303	EF694755
RLA09	EF694744	EU333386	FM253167 ^a	EU333391
RLA11	EU333381	EU333387	FM253169 ^a	EU333392
ISLU203	GQ863571	GQ863555	GU338035	GQ863563
ISLU207	AJ558028	GQ863557	GU338036	AJ560652
ISLU213	GQ863572	GQ863556	*	GQ863564
ISLU220	GQ863573	EF990556	*	GQ863565
ISLU227	AJ558032	EU333385	*	EU333390

Table 2 Sequence accession numbers of genes analysed in this study corresponding to the strains isolated from L. albus nodules

* *atpD* gene sequences of strains BLUT6 and BLUT8 were not deposited in databases because they are identical to that of strain BLUT5. Also, the sequences of strains ISLU213, ISLU207 and ISLU227 were identical to that of strain ISLU203

^a atpD gene sequences of the strains RLA09 and RLA11 were obtained by other authors in a previous work and were not included in the phylogenetic tree since they are identical to that of strain RLA08

166 of strains for gene sequencing (Valverde et al. 2006, Iglesias et al. 2007; Santillana et al. 2008; Álvarez-167 168 Martínez et al. 2009; Ramírez-Bahena et al. 2009). The 169 results of this analysis in the strains isolated from L. albus nodules are shown in Fig. 1 and Table 1. They 170 171 presented 19 different patterns (Fig. 1, lanes 6 to 24) 172 that were also different to those of the reference strains 173 (Fig. 1, lanes 1 to 5). The results of the mathematical 174 analysis showed low similarity coefficients among 175 most of L. albus strains and with respect to the reference strains. These results imply a high genetic diversity of 176 177 Lupinus strains and confirmed the usefulness of RAPD 178 patterns to analyze the genetic diversity of rhizobial 179 populations. Considering the low similarity values 180 found after the mathematical analysis (lower than 90%) 181 we sequenced the rrs gene and the ITS fragment in all 182 L. albus strains.

183 Analysis of rrs gene

184 Considering that the current phylogenetic classifica-185 tion of rhizobia is predominantly based on *rrs* gene

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sequences (Kuykendall 2005) the classification of 186 nodule isolates should be performed on the basis of the 187 results of the analysis of this gene. According to the rrs 188 gene analyses previously published the strain BLUT1 189 was identified as Bradyrhizobium canariense (Step-190 kowski et al. 2007 and the strain ISLU207 was 191 classified in the phylogenetic group of Bradyrhizobium 192 japonicum and Bradyrhizobium genospecies alpha 193 (Jarabo-Lorenzo et al. 2003). These results are con-194 firmed in the case of strains BLUT1 and ISLU207, 195 whereas strain ISLU227 belongs to an independent 196 lineage formed by several strains isolated in different 197 continents (Fig. 2). To this lineage (with 100% iden-198 tity) belong the strains ISLU227, ISLU213 and 199 ISLU220 isolated from different geographical loca-200 tions in Chile (South America), the strain FN13 201 isolated from Lupinus montanus in Mexico (North 202 America) and the strains isolated in León (mainland 203 Spain, Europe). In spite of the proximity of León and 204 Salamanca (mainland Spain, Europe), the strains 205 isolated in Salamanca belong to a different lineage 206 that contains most of the isolates from L. albus in the 207





Fig. 1 Results of the mathematical analysis of RAPD patterns from strains isolated in this study. *B. japonicum* by glycinearum LMG 6138^{T} (*lane 1*), *B. japonicum* by genistearum BGA-1 (*lane 2*), *B. canariense* BTA-1^T (*lane 3*), *Bradyrhizobium* genosp. α BC-C1 (*lane 4*), *Bradyrhizobium* genosp. β BRE-1 (*lane 5*), RLA08 (*lane 6*), RLA09 (*lane 7*), RLA10 (*lane 8*), RLA11 (*lane 8*),

208 Canary Islands and coincides with the species 209 B. canariense (identities higher than 99.8%). Two strains isolated on the Canary Islands, BLUT2 and 210 211 BLUT3, clustered with the genospecies alpha of 212 Bradyrhizobium represented by the strain BC-C1 isolated in the same location but from different hosts 213 (identities higher than 99.6%). Finally, the strain 214 215 ISLU203 isolated from L. albus in Chile belongs to the 216 same lineage as the strain BGA-1 isolated from Teline stenopetala in Canary Islands (100% identity). 217

218 From the rrs analysis it was concluded that the 219 strains isolated from L. albus in different continents 220 belong to several different lineages that in some cases 221 presented high identity to already known species of 222 genus Bradyrhizobium. Nevertheless, the resolution 223 of rrs analysis is too low for Bradyrhizobium species 224 assignment and other more variable molecules must 225 be studied (Willems et al. 2003; Rivas et al. 2004).

9), MCLA07 (lane 10), MCLA12 (lane 11), MCLA22 (lane 12), MCLA23 (lane 13), BLUT1 (lane 14), BLUT2 (lane 15), BLUT3 (lane 16), BLUT5 (lane 17), BLUT6 (lane 18), BLUT8 (lane 19), ISLU203 (lane 20), ISLU207 (lane 21), ISLU213 (lane 22), ISLU220 (lane 23), ISLU227 (lane 24). MW: Molecular weight standard VI (Roche, Germany)

Analysis of 16S-23S intergenic spacer (ITS)

The ITS sequence analysis has been reported as a better 227 tool than 16S rDNA analysis for species delineation 228 within the genus Bradyrhizobium, in which ITS 229 sequence similarities higher than 95.5% indicate a 230 genospecies level relatedness (Willems et al. 2003, 231 Rivas et al., 2004). The results of the phylogenetic 232 analysis of ITS sequences (Fig. 3) confirmed that the 233 strains isolated from Salamanca (mainland Spain) and 234 most of the strains isolated from the Canary Islands 235 belong to the species B. canariense. These strains were 236 divided into two close ITS groups with identities 237 higher than 98% (gaps not considered) together with 238 several strains isolated in different geographical 239 regions from different Genisteae legumes and Ornith-240 opus. These two ITS groups coincide with those 241 designed as ITS-I and ITS-II by Safronova et al. (2007) 242

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•	Journal : Medium 10482	Dispatch : 28-1-2010	Pages : 14
Į	Article No. : 9415	□ LE	□ TYPESET
	MS Code : ANTO-09-147	🗹 СР	🖌 disk





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Fig. 2 Neighbour-joining phylogenetic tree based on *rrs* gene sequences (1,480 nt) showing the position of representative strains from each RAPD group. Bootstrap values calculated for 1,000 replications are indicated. *Bar*, 0.2 nt substitution per

243 within B. canariense, which can be differentiated by 244 the presence of an insert in the strains from ITS-I 245 group. Nevertheless, we found that the insert is smaller 246 (15 nucleotides) than that reported by Safronova et al. (2007) comprising only the sequence "TAG-247 248 AGACTTAGGTTT" (located from 731 to 745 in the 249 ITS of the strain Oc9 isolated from Ornithopus sp.). All 250 Canary Island isolates belong to the ITS-I group, 251 whereas the Salamanca isolates were distributed in the ITS groups I and II. 252

The strains isolated in León (mainland Spain,Europe), most strains isolated in Chile (South

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•	Journal : Medium 10482	Dispatch : 28-1-2010	Pages : 14
	Article No. : 9415	🗆 LE	□ TYPESET
	MS Code : ANTO-09-147	🗹 СР	🖌 disk

100 nt. Strains isolated from *L. albus* nodules are in *bold*. *Asterisks* indicate the strains identified as a previously described species of genus *Bradyrhizobium*

America) and the strain BLup-MR1 isolated from 255 Lupinus polyphylus in Germany (Europe) have iden-256 tical ITS sequences and grouped in a cluster (III) 257 258 phylogenetically divergent from that formed by the strains identified as B. canariense (identities lower 259 than 98%). They presented high identity values (higher 260 than 99%) with respect to B. japonicum by genistearum 261 strain BGA-1, which presented near 100% identity 262 with the strain ISLU256 isolated from Ornithopus in 263 mainland Spain (Jarabo-Lorenzo et al. 2003) and strain 264 FN13 isolated from L. montanus in Mexico (North 265 America) (Barrera et al. 1997). This high identity value 266



Fig. 3 Neighbour-joining phylogenetic tree based on 16S-23S rDNA intergenic sequences (725 nt) showing the position of representative strains from each RAPD group. Bootstrap values calculated for 1,000 replications are indicated. *Bar*, 1 nt

showed that the strains from these two groups probably
belong to the same species but this species could not be *B. japonicum*. The low identity (about 95%) found
between the strains of this group, that includes *B. japonicum* by genistearum BGA-1, and *B. japon- icum* by glycinearum LMG 6138^T suggested they
could belong to different species and emphasises the

substitution per 100 nt. Strains isolated from *L. albus* nodules are in bold. *Asterisks* indicate the strains identified as a previously described species of genus *Bradyrhizobium*

need for a revision of the taxonomic status of the 274 current species *B. japonicum*. 275

The high similarity (99%) found between the strain276ISLU207 isolated in Chile (South America) and277the strain BC-C1 (group IV) suggested that both278strains belong to the genospecies alpha of genus279Bradyrhizobium.280

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2	Journal : Medium 10482	Dispatch : 28-1-2010	Pages : 14
	Article No. : 9415	□ LE	□ TYPESET
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281 Finally, and in spite of their high *rrs* gene identity, 282 the strains BLUT2 and BLUT3 isolated in Canary 283 Islands have 96% identity in their ITS sequences, the 284 limit of ITS similarity for species differentiation in 285 genus Bradyrhizobium (Willems et al. 2003). How-286 ever more strains of these groups are necessary to 287 define new species according to the recommendations 288 of the Subcommittee on the taxonomy of Agrobac-289 terium and Rhizobium (Lindström and Young 2009).

290 Analysis of *atpD* gene

291 In recent years, analyses of housekeeping genes have 292 been performed in Bradyrhizobium showing their 293 usefulness in taxonomic and phylogenetic studies (Vinuesa et al. 2005a; Stepkowski et al. 2007; 294 295 Ramírez-Bahena et al. 2009; Rivas et al. 2009). 296 From these genes, the *atpD* gene has been previously 297 analysed in strains from Lupinus isolated in different 298 geographical locations (Stepkowski et al. 2007; Rivas et al. 2009). In these previous studies, some strains 299 300 isolated from L. albus in Spain and the Canary 301 Islands were already included, with the conclusion 302 that they belong to different phylogenetic groups (Stepkowski et al. 2007; Rivas et al. 2009). In this 303 304 work we have sequenced the *atpD* genes of strains 305 that had not been previously analysed. As the strains presenting identical ITS sequences also have identi-306 307 cal *atpD* genes, only a representative strain from each ITS group and geographical location has been 308 309 included in the phylogenetic analysis (Fig. 4). Nev-310 ertheless, in some cases the phylogenies based on 311 *atpD* genes and ITS fragments were not completely congruent. For example the strain BLUT3 has a 312 313 distant ITS sequence but close *atpD* gene with respect 314 to the strain BLUT1. Also, the strain BRE-1 from the 315 genospecies beta of Bradyrhizobium has a distant ITS 316 sequence but a close *atpD* gene with respect to the strain BGA-1. Finally, the strain ISLU207 has a close 317 318 ITS sequence but a divergent *atpD* gene with respect 319 to the strain BC-C1. The significance of these 320 findings should be further studied since strains 321 isolated from the Canary Islands are involved in all 322 these exceptions. The special characteristics of rhi-323 zobia present in these Islands have been recently 324 reported for strains nodulating Phaseolus having 325 common characteristics with strains isolated in North 326 Africa nodulating this legume and in mainland Spain nodulating Medicago (Zurdo-Piñeiro et al. 2009). 327

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The representative strains of ITS groups I and II, 328 isolated in Salamanca and Canary Islands, belong to 329 five different subgroups with *atpD* gene identities 330 higher than 97% among them and with respect to the 331 strain *B. canariense* by genistearum BTA-1^T. These 332 high identity values are in agreement with those 333 obtained after the ITS analysis and confirmed the 334 identification of strains from ITS groups I and II as B. 335 canariense. Several strains isolated from other Lupi-336 nus species in European countries (Poland and Iceland) 337 and several strains isolated from diverse Genisteae 338 legumes in Africa (Morocco) and nearby geographical 339 zones such as Canary Islands also cluster in this group 340 (Vinuesa et al. 2005a; Stepkowski et al. 2007). 341

The representative strains from ITS group III, 342 isolated in León and Chile, form a cluster phylogenet-343 ically divergent from B. canariense that also includes 344 strains isolated from other Lupinus species mostly in 345 American countries. These strains were close to the 346 strain B. japonicum by genistearum BGA-1 (99.8%) 347 identity), that itself presented about 97% identity with 348 respect to the type strain of B. japonicum by glycinea-349 rum LMG 6138^T. This identity value is at the limit for 350 species differentiation in the genus Bradyrhizobium 351 (Ramírez-Bahena et al. 2009) and thus confirms that 352 the taxonomic status of the strains currently included in 353 B. japonicum by genistearum should be revised. 354

Related to the B. canariense group (about 96% 355 identity), the strains BLUT2 and ISLU207 represent 356 two independent lineages, congruent with the ITS 357 analysis. Nevertheless, in disagreement with this 358 analysis, the high identity found in the *atpD* gene 359 (near 98%) suggested that they could belong to the 360 same species. Moreover the results of the *atpD* gene 361 analysis are in disagreement with the identification of 362 the strain ISLU207 as belonging to the genospecies 363 alpha since it has less than 94% identity with respect to 364 the strain BC-C1. These discordant results depending 365 on the phylogenetic marker analysed showed that it is 366 necessary to have more strains from these groups in 367 order to establish their taxonomic status as well as that 368 369 of other phylogenetic lineages also formed by single strains isolated from other Lupinus species (Stepkow-370 ski et al. 2007). 371

Although according our results and those from372Stepkowski et al. (2007), the strains nodulating373different Lupinus species in Europe mostly belong to374B. canariense. Our results showed that strains isolated375on mainland Spain belong to both B. canariense and376

•	Journal : Medium 10482	Dispatch : 28-1-2010	Pages : 14	
ĺ	Article No. : 9415	□ LE	□ TYPESET	
	MS Code : ANTO-09-147	🖌 СР	🖌 disk	



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Fig. 4 Neighbour-joining phylogenetic tree based on *atpD* gene (460 nt) showing the position of representative strains compared with strains isolated from other legume hosts mainly from *Lupinus* and other *Genisteae* and *Loteae* isolates. Bootstrap values calculated for 1,000 replications are

indicated. *Bar*, 5 nt substitution per 100 nt. Strains isolated from *L. albus* nodules are in *bold. Asterisks* indicate the strains identified as a previously described species of genus *Bradyrhizobium*

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Journal : Medium 10482	Dispatch : 28-1-2010	Pages : 14
Article No. : 9415	□ LE	□ TYPESET
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B. japonicum clusters. This finding suggests that the
distribution of these two species in *Lupinus* nodules
could be more related with the local ecological
conditions than with its geographic (continent) location. Nevertheless, further studies on biodiversity and
biogeography of strains nodulating *Lupinus* are necessary to establish more reliable conclusions.

384 Analysis of the *nodC* gene

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385 The *nodC* gene determines the host range of rhizobia 386 and it is therefore related with the host promiscuity (Roche et al. 1996; Perret et al. 2000; Laguerre et al. 387 388 2001; Rivas et al. 2007b; Iglesias et al. 2007). It has 389 been reported that restrictive hosts such as Cicer are 390 nodulated by species bearing almost identical nodC 391 genes (Rivas et al. 2007b; Laranjo et al. 2009) whereas 392 promiscuous hosts such as Phaseolus or Prosopis are 393 nodulated by rhizobial species carrying divergent 394 nodC genes (Laguerre et al. 2001; Iglesias et al. 395 2007). Previous analyses of some isolates from 396 nodules of legumes belonging to the Genisteae Tribe 397 showed the high conservation degree of this gene 398 (Jarabo-Lorenzo et al. 2003; Kalita et al. 2006) and led 399 to the definition of the biovar genistearum within 400 B. canariense and B. japonicum (Vinuesa et al. 2005a). 401 However, the nodC genes of B. japonicum by genis-402 tearum strains clustered separately (identity lower than 403 75%) from those of Bradyrhizobium japonicum by 404 glycinearum (Fig. 5). The strains from biovar genis-405 tearum nodulating Genisteae, including Lupinus, and 406 Ornithopus (Tribe Loteae), belong to a wide phyloge-407 netic *nodC* clade supported by a bootstrap value of 99% and with an internal similarity of about 90%. 408 409 Although the rhizobia nodulating the Genisteae and 410 Ornithopus belong to the same cross-inoculation 411 group, the relatively low identity level of their nodC 412 genes (near to 92% in some cases) indicated that these 413 legumes are less restrictive than other hosts such as 414 Cicer whose endosymbionts have almost 100% iden-415 tity in their nodC genes (Rivas et al. 2006). Within the 416 clade formed by the biovar genistearum strains, those 417 nodulating L. albus are located in four groups (A-D) 418 that also contain other strains isolated from Genisteae 419 except in the case of group C that is formed exclusively 420 by L. albus endosymbionts. The results from nodC 421 analysis basically agree with those found after the 422 analysis of the *nodA* gene since the strains from biovar genistearum are included in the nodA clade II including 423

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Journal : Medium 10482	Dispatch : 28-1-2010	Pages : 14
Article No. : 9415	□ LE	□ TYPESET
MS Code : ANTO-09-147	🛃 СР	🖌 DISK

the strains BLUT1, MCLA07 and RLA08 that belong 424 to three different subgroups of nodA (Stepkowski et al. 425 2007). On the basis of the *nodA* gene analysis strain 426 BLUT1 belongs to a group that also contains isolates 427 from other Lupinus species in Europe and America. 428 However according to the results of the *nodC* gene, 429 strains BLUT1 and MCLA07 isolated from L. albus in 430 Canary Islands (geographically near to Africa) and 431 Salamanca (mainland Spain, Europe), respectively, 432 belong to the same group (group D) which also 433 grouped other strains isolated from these two locations 434 (Fig. 5). The León strains (mainland Spain, Europe) 435 and several isolates from Chile (South America) 436 belong to a phylogenetically distant group (group A). 437 The strains WM9 and RLA08 (Poland) isolated in 438 Europe from L. luteus and L. albus, respectively, 439 belong to the same group after both nodC and nodA 440 gene analyses (Stepkowski et al. 2007). Furthermore, 441 the *nodC* group A includes the strain BGA-1 (that was 442 also close to the same strains on the basis of the ITS 443 sequences) but also the strain BC-C1, a representative 444 strain of the genospecies alpha. 445

The group B was constituted by miscellaneous 446 strains isolated from different hosts and geographical 447 origins including the strain BRE-1 representative of 448 the genospecies beta. To this group belong the strain 449 ISLU207, isolated in Chile (South America) and 450 strains BLUT2 and BLUH1, isolated from L. albus 451 and L. angustifolius, respectively, in Canary Islands. 452 This last strain belonged to the same group as strain 453 MCLA07 when the nodA gene was analysed (Step-454 kowski et al., 2007). Therefore, some differences in the 455 phylogenetic arrangement of L. albus strains BLUT1 456 and MCLA07 were found depending on the gene 457 analyzed. Nevertheless, in agreement with the results 458 found on the basis of the nodA gene (Stepkowski et al. 459 2007), the strains from nodC groups B and D are 460 dominant in Europe and Africa. 461

Finally, it must be highlighted that the group C was 462 exclusively formed by strains nodulating L. albus on 463 different continents i.e. strains BLUT3, BLUT5 and 464 BLUT6 isolated on the Canary Islands (near to Africa), 465 strain MCLA22 isolated on mainland Spain (Europe) 466 and strain ISLU203 isolated in Chile (South America). 467 Although this finding could suggest a closer coevolu-468 tion among the strains from group C and their host, no 469 significant differences in the number of nodules per 470 plant (ranging from 20 to 30) and shoot dry weight 471 (ranging from 300 to 400 mg/plant) were found 472



Fig. 5 Neighbour-joining phylogenetic tree based on *nodC* gene sequences (610 nt) showing the position of representative strains from each RAPD group. Bootstrap values calculated for

1,000 replications are indicated. *Bar*, 5 nt substitution per 100 nt. Strains isolated from *L. albus* nodules are in *bold*

	Journal : Medium 10482	Dispatch : 28-1-2010	Pages : 14
	Article No. : 9415	□ LE	□ TYPESET
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Table 3	Symbiotic	characteristics	of	strains	isolated	from
L. albus	nodules repr	resentative of ea	ch g	group of	nodC ger	e and
geograph	ical origin					

Strains	nodC subgroup	NN*	SDW ^a (g)	SE ^b (%)
ISLU213	А	30	0.40	66.7
RLA08	А	25	0.30	50.0
ISLU207	В	21	0.36	60.3
BLUT2	В	21	0.30	50.0
ISLU203	С	20	0.35	58.3
BLUT3	С	22	0.38	63.3
MCLA22	С	20	0.36	60.0
BLUT1	D	25	0.35	58.3
MCLA23	D	20	0.40	66.7

Not significant differences were found at P = 0.05 according to Fisher's protected LSD (Least Significant Differences)

* NN Number of nodules per plant

^a *SDW* Shoot Dry Weigth per plant, *SDW* inoculated plants/ SDW non-inoculated control plants (140 ppm nitrogen as NH₄NO₃). SDW-average shoot dry weight from five replicates ^b *SE* Symbiotic efficiency

473 between the strains from group C and the remaining 474 *nodC* groups (Table 3). The presence in the groups A 475 and C of strains isolated from Spanish and South 476 American soils suggests that L. albus endosymbionts could have been dispersed from Europe to American 477 478 countries together with the legume seeds. Neverthe-479 less, as was mentioned in the case of the chromosomal genes, more strains isolated from different countries 480 481 and from different cultivated and wild Lupinus species 482 should be analyzed to establish the geographical 483 distribution patterns of lupine endosymbionts.

484 In summary, the results of chromosomal and 485 symbiotic markers analysis from this study showed that the L. albus endosymbionts isolated in different 486 487 continents belong to at least four genetic lineages 488 within the genus Bradyrhizobium. However, the exis-489 tence of some discordant results among these markers 490 showed that a revision of genus Bradyrhizobium 491 through a polyphasic study is necessary to establish 492 the taxonomic status of several phylogenetic groups 493 within this genus. For this, additional strains isolated 494 from different legumes of tribes Genisteae and Loteae 495 should be analysed in order to describe the potential 496 new species detected in this and previous works and to 497 establish more reliable conclusions about the bioge-498 ography and host range of these species. The results of 499 this work also showed a certain degree of coevolution between chromosomes and the *nodC* gene in *L. albus* 500 isolates since the strains of *B. canariense* carry *nodC* 501 genes phylogenetically related among them (groups C 502 and D) and distant to those carried by the remaining 503 strains. Also the strains from the chromosomal cluster 504 of B. japonicum by genistearum BGA-1, with the 505 exception of strain ISLU203, carry phylogenetically 506 related nodC genes (cluster A). Therefore a tripar-507 tite coevolution could be occurring among chromo-508 somes, symbiotic elements and hosts at least in the case 509 of L. albus nodulating strains. 510

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References

- Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ (1990) Basic local alignment search tool. J Mol Biol 215: 403–410 Andam CP, Parker MA (2007) Novel alphaproteobacterial root 521
- Andam CP, Parker MA (2007) Novel alphaproteobacterial root nodule symbiont associated with *Lupinus texensis*. Appl Environ Microbiol 73:5687–5691
- Barrera LL, Trujillo ME, Goodfellow M, García FJ, Hernández-Lucas I, Dávila G, van Berkum P, Martínez-Romero E (1997) Biodiversity of bradyrhizobia nodulating *Lupinus* spp. Int J Syst Bacteriol 47:1086–1091
- Clapham WM (1997) Lupin development. Field Crops Res 52:283–284
- Dijkstra DS, Linnemann AR, van Boekel TA (2003) Towards sustainable production of protein-rich foods: appraisal of eight crops for Western Europe. Part II: Analysis of the technological aspects of the production chain. Crit Rev Food Sci Nut 43:481–506
- Erbas M, Certel M, Uslu MK (2005) Some chemical properties of white lupin seeds (*Lupinus albus* L.). Food Chem 89:341–345
- Gaunt MW, Turner SL, Rigottier-Gois L, Lloyd-Macgilp SA, Young JWP (2001) Phylogenies of *atpD* and *recA* support the small subunit rRNA-based classification of rhizobia. Int J Syst Evol Microbiol 51:2037–2048
- Iglesias O, Rivas R, García-Fraile P, Abril A, Mateos PF, Martinez-Molina E, Velázquez E (2007) Genetic characterization of fast-growing rhizobia able to nodulate *Prosopis alba* in North Spain. FEMS Microbiol Lett 277:210–216
- Jarabo-Lorenzo A, Pérez-Galdona R, Donate-Correa J, Rivas R, Velázquez E, Hernández M, Temprano F, Martínez-Molina E, Ruiz-Argüeso T, León-Barrios M (2003) Genetic diversity of bradyrhizobial populations from diverse geographic origins that nodulate *Lupinus* spp. and Ornithopus spp. Syst Appl Microbiol 26:611–626 552

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Journal : Medium 10482	Dispatch : 28-1-2010	Pages : 14
Article No. : 9415	🗆 LE	□ TYPESET
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- Jensen CR, Joernsgaard B, Andersen MN, Christiansen JL, Mogensen VO, Friis P, Petersen CT (2004) The effect of lupins as compared with peas and oats on the yield of the subsequent winter barley crop. Eur J Agron 20:405–418
- Kalita M, Stepkowski T, Lotocka B, Malek W (2006) Phylogeny of nodulation genes and symbiotic properties of *Genista tinctoria* bradyrhizobia. Arch Microbiol 186:87–97
- Kimura M (1980) A simple method for estimating evolutionary rates of base substitutions through comparative studies of nucleotide-sequences. J Mol Evol 16:111–120
- Kuykendall LD (2005) Order VI. Rhizobiales ord. nov. In: Brenner DJ, Krieg NR, Staley JT, Garrity GM (eds) Bergey's manual of systematic bacteriology, (The Proteobacteria), part C (The Alpha-, Beta-, Delta-, and Epsilonproteobacteria), vol 2, 2nd edn. Springer, New York, pp 324–340
- Kuykendall LD, Saxena B, Devine TE, Udell SE (1992) Genetic diversity in *Bradyrhizobium japonicum* Jordan 1982 and a proposal for *Bradyrhizobium elkanii* sp. nov. Can J Microbiol 38:501–505
- Laguerre G, Nour SM, Macheret V, Sanjuan J, Drouin P, Amarger N (2001) Classification of rhizobia based on nodC and nifH gene analysis reveals a close phylogenetic relationship among *Phaseolus vulgaris* symbionts. Microbiology 147:981–993
- Laranjo M, Alexandre A, Rivas R, Velázquez E, Young JP,
 Oliveira S (2008) Chickpea rhizobia symbiosis genes are
 highly conserved across multiple *Mesorhizobium* species.
 FEMS Microbiol Ecol 66:391–400
- Lindström K, Young JPW (2009) International committee on systematics of prokaryotes; subcommittee on the taxonomy of *Agrobacterium* and *Rhizobium*: minutes of the meetings, 31 August 2008, Gent, Belgium. Int J Syst Evol Microbiol 59:921–922
- 587 Perret X, Staehelin C, Broughton WJ (2000) Molecular basis
 588 of symbiotic promiscuity. Microbiol Mol Biol Rev 64:
 589 180–201
- Ramírez-Bahena MH, Peix A, Rivas R, Rodríguez-Navarro
 DN, Camacho M, Mateos PF, Martínez-Molina E, Willems A, Velázquez E (2009) *Bradyrhizobium pachyrhizi*sp. nov. and Bradyrhizobium jicamae sp. nov., isolated
 from effective nodules of Pachyrhizus erosus. Int J Syst
 Evol Microbiol 59:1929–1934
- 596 Rivas R, Velázquez E, Valverde A, Mateos PF, Martínez597 Molina E (2001) A two primers random amplified poly598 morphic DNA procedure to obtain polymerase chain
 599 reaction fingerprints of bacterial species. Electrophoresis
 600 22:1086–1089
- Rivas R, Willems A, Palomo JL, García-Benavides P, Mateos
 PF, Martínez-Molina E, Gillis M, Velázquez E (2004) *Bradyrhizobium betae* sp. nov., isolated from roots of
 Beta vulgaris affected by tumour-like deformations. Int J
 Syst Evol Microbiol 54:1271–1275
- Rivas R, Peix A, Mateos PF, Trujillo ME, Martínez-Molina E, Velázquez E (2006) Biodiversity of populations of phosphate solubilizing rhizobia that nodulate chickpea in different Spanish soils. Plant Soil 287:23–33
- 610 Rivas R, García-Fraile P, Mateos PF, Martínez-Molina E,
 611 Velázquez E (2007a) Characterization of xylanolytic
 612 bacteria present in the bract phyllosphere of the date palm
 613 Phoenix dactylifera. Lett Appl Microbiol 44:181–187

- Rivas R, Laranjo M, Velázquez E, Mateos PF, Oliveira S, Martínez-Molina E (2007b) Strains of *Mesorhizobium amorphae* and *M*. tianshanense carrying symbiotic genes of common chickpea endosymbiotic species constitute a novel biovar (ciceri) able to nodulate Cicer arietinum. Lett Appl Microbiol 44:412–418
 Roche P, Maillet F, Plazanet C, Debellé F, Ferro M, Truchet G,
- Roche P, Maillet F, Plazanet C, Debellé F, Ferro M, Truchet G, Prome JC, Dénarié J (1996) The common *nodABC* genes of *Rhizobium meliloti* are host-range determinants. Proc Natl Acad Sci USA 93:15305–15310

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- Rosolem CA, Foloni JSS, Tiritan CS (2002) Root growth and nutrient accumulation in cover crops as affected by soil compaction. Soil Tillage Res 65:109–115
- Safronova V, Chizhevskaya E, Bullitta S, Andronov E, Belimov A, Charles TC, Lindstrom K (2007) Presence of a novel 16S-23S rRNA gene intergenic spacer insert in Bradyrhizobium canariense strains. FEMS Microbiol Lett 269:207–212
- Saitou N, Nei M (1987) A neighbour-joining method: a new method for reconstructing phylogenetic trees. Mol Biol Evol 4:406–425
- Somasegaran P, Hoben HJ (1994) Handbook for rhizobia methods in legume-rhizobium technology. Springer, New York
- Stepkowski T, Moulin L, Krzyzanska A, McInnes A, Law IJ, Howieson J (2005) European origin of *Bradyrhizobium* populations infecting lupins and serradella in soils of Western Australia and South Africa. Appl Environ Microbiol 71:7041–7052
- Stepkowski T, Hughes CE, Law IJ, Markiewicz L, Gurda D, Chlebicka A, Moulin L (2007) Diversification of lupine *Bradyrhizobium* strains: evidence from nodulation gene trees. Appl Environ Microbiol 73:3254–3264
- Tamura K, Dudley J, Nei M, Kumar S (2007) MEGA4: Molecular Evolutionary Genetics Analysis (MEGA) software version 4.0. Mol Biol Evol 24:1596– 1599
- Thompson JD, Gibson TJ, Plewniak F, Jeanmougin F, Higgins DG (1987) The clustalX windows interface: flexible strategies for multiple sequence alignment aided by quality analysis tools. Nucleic Acid Res 25:4876–4882
- Trujillo ME, Willems A, Abril A, Planchuelo AM, Rivas R, Ludeña D, Mateos PF, Martínez-Molina E, Velázquez E (2005) Nodulation of *Lupinus albus* by strains of *Ochrobactrum lupini* sp. nov. Appl Environ Microbiol 71: 1318–1327
- Valverde A, Igual JM, Peix A, Cervantes E, Velázquez E (2006) *Rhizobium lusitanum* sp. nov. a bacterium that nodulates Phaseolus vulgaris. Int J Syst Evol Microbiol 56:2631–2637
- Vincent JM (1970) The cultivation, isolation and maintenance of rhizobia. In: Vincent JM (ed) A manual for the practical study of root-nodule. Blackwell Scientific Publications, Oxford, pp 1–13
- Vinuesa P, León-Barrios M, Silva C, Willems A, Jarabo-Lorenzo A, Pérez-Galdona R, Werner D, Martínez-Romero E (2005a) *Bradyrhizobium canariense* sp. nov., an acid-tolerant endosymbiont that nodulates endemic genistoid legumes (Papilionoideae: Genisteae) from the Canary Islands, along with Bradyrhizobium japonicum bv. genistearum, Bradyrhizobium genospecies alpha and 674

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- 676 677 678 679 680 681 682 683 684 685 686 687 688 689
- 675Bradyrhizobium genospecies beta. Int J Syst Evol676Microbiol 55:569–575
 - Vinuesa P, Silva C, Werner D, Martínez-Romero E (2005b)
 Population genetics and phylogenetic inference in bacterial molecular systematics: the roles of migration and
 recombination in *Bradyrhizobium* species cohesion and
 delineation. Mol Phylogenet Evol 34:29–54
 - Willems A, Munive A, de Lajudie P, Gillis M (2003) In most Bradyrhizobium groups sequence comparison of 16S-23S rDNA internal transcribed spacer regions corroborates DNA–DNA hybridizations. Syst Appl Microbiol 26: 203–210
 - Xu LM, Ge C, Cui Z, Li J, Fan H (1995) Bradyrhizobium liaoningense sp. nov., isolated from the root nodules of soybeans. Int J Syst Bacteriol 45:706–711

- Yao ZY, Kan FL, Wang ET, Wei GH, Chen WX (2002)
 Characterization of rhizobia that nodulate legume species of the genus *Lespedeza* and description of *Bradyrhizobi um yuanmingense* sp. nov. Int J Syst Evol Microbiol 52:2219–2230
 690
 691
 692
 693
 694
- Zurdo-Piñeiro JL, García-Fraile P, Rivas R, Peix A, León-Barrios M, Willems A, Mateos PF, Martínez-Molina E, Velázquez E, van Berkum P (2009) Rhizobia from Lanzarote, the Canary Islands, that nodulate *Phaseolus vulgaris* have characteristics in common with *Sinorhizobium meliloti* isolates from mainland Spain. Appl Environ Microbiol 75:2354–2359
 695
 695
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 697
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