# Diversity among agrobacteria isolated from diseased plants of blueberry (*Vaccinium corymbosum*) in Argentina

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Abstract The aim of this study was to isolate, identify and analyze the diversity of the causative agents of crown galls and hairy roots from symptomatic plants of Vaccinium corymbossum by means of biological, biochemical and molecular tools. All the bacteria isolated from blueberries (n=78) were found to be Agrobacterium since they grew on three differential media, provoked cell and/or root proliferation on Kalanchoe, and contained a 730 bp partial sequence that codes for virulence genes within the virC operon found on Ti and/or Ri plasmids. Isolates were highly variable considering the ERIC-PCR patterns as well as biochemical reactions and were all represented by 7 different restriction patterns of the 16SrDNA. While most of the isolates belonged to Agrobacterium bv. 1 (n=33) or Agrobacterium bv. 2 (n=31) only fourteen were Agrobacterium rubi. A representative isolate of each of these three groups was further identified by sequencing the approximately 400 bp 16SrDNA. We concluded that Vaccinium plants are particularly susceptible to Agrobacterium bv. 1, Agrobacterium by. 2, and also to Agrobacterium rubi. To our knowledge this is the first survey of Agrobacterium affecting blueberries in Argentina.

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# Introduction

Agrobacteria are soil-borne Gram-negative bacteria within the alphaproteobacteria subclass that belong to the family Rhizobiaceae. The genus Agrobacterium comprise a diverse group of microorganisms, all of which, when harbouring the appropriate plasmids, provoke uncontrolled cell proliferations on dicotyledoneous plants (Farrand et al. 2003). Depending on the species and strains of Agrobacterium involved, the interaction might result in the development either of galls or hairy roots (Sawada et al. 1995; Moore et al. 2001) or they might not be pathogenic in plants (Llop et al. 2009). Agrobacterium species are particular in that they infect their hosts only through wounds that release compounds such as acetosyringone, that activate the complex and unique mechanism encoded by plasmids. This complex interaction results in the insertion of a Ti or Ri sequence in the plant's genome at the cell nucleus (Gelvin 2010).

Agrobacterium species were first classified by means of a pathogenicity based-taxonomy in *A. tumefaciens*, those strains that induce crown-gall, *A. rhizogenes*, those strains that induce hairy-roots, and *A. radiobacter* that are non-pathogenic strains (Moore et al. 2001). In addition to this, three other species have been described: A. vitis, A. rubi and A. larrymoorei. These species induce galls in Vitis spp.; raspberries, blackberries, blueberries and cranberries; and Ficus benjamina, respectively (Bouzar and Jones 2001; Moore et al. 2001; Young et al. 2005; Alippi et al. 2010). Pulawska and co-workers (2006), based on the phylogenetic analysis of the 23S rRNA gene sequences, made a simpler classification of Agrobacterium in four taxa, Agrobacterium biovar 1, Agrobacterium biovar 2, Agrobacterium vitis (biovar 3), and A. rubi. A. larrymoorei was left as a separate species. However, Young and co-workers (2001), based on phylogenetic analysis of the 16SrDNA, suggested that Agrobacterium species should be renamed Rhizobium radiobacter, Rhizobium rhizogenes, Rhizobium rubi, and Rhizobium vitis. In support of this, researchers recently described the isolation of Rhizobium skierniewicense from tumours of chrysanthemum and Prunus spp. (Pulawska et al. 2012) and Rhizobium pusense from the rhizosphere of chickpea (Panday et al. 2011). Nevertheless, other authors such as Farrand and coworkers (2003) found that as a defined and distinct group within rhizobiaceae, the name should be kept as Agrobacterium. In any case, the phylogenetic-based classification of Agrobacterium, suggests that they can be separated in Agrobacterium biovar 1 that contains at least nine genomic species causing crown gall or hairy roots or are non-pathogenic (Costechareyre et al. 2010; Lasalle et al. 2011); and three closely related species, A. rubi that carries a Ti plasmid and induces galls mostly on Rubus spp. (Anderson and Moore 1979), A. vitis (formerly known as biovar 3) those strains that have a Ti plasmid that provokes galls exclusively on grape (Lim et al. 2009), and A. larrymoorei, carrying a Ti plasmid producing galls on Ficus benjamina (Bouzar and Jones 2001). However, the former Agrobacterium biovar 2 members that also induce galls or hairy roots, whether they have the Ti or Ri plasmid or are nonpathogenic, are considered Rhizobium rhizogenes (Portier et al. 2006). Recently, the Subcommittee on the Taxonomy of Agrobacterium and Rhizobium, confirmed that Rhizobium rhizogenes is a valid designation for biovar 2 (Costecharevre et al. 2010) and also for the A. radiobacter strains K84 and A. tumefaciens AK10 (Velázquez et al. 2010).

Since, the nomenclature and taxonomy of *Agrobacterium* remains obscure and under debate. The Subcommittee on *Agrobacterium* and *Rhizobium* taxonomy suggested that it is up to the authors to

choose the nomenclature to be used (Portier et al. 2006). Here, we decided, mostly based on simplicity, to use the biovar divisions proposed by Pulawska and co-workers (2006) that included: *Agrobacterium* biovar 1 (Syn = *A. tumefaciens*) carrying Ti or Ri plasmids or non-pathogenic; *Agrobacterium* biovar 2 (Syn = A. rhizogenes) carrying Ti or Ri plasmids or non-pathogenic; *Agrobacterium rubi, Agrobacterium vitis* and *Agrobacterium larrymorei*.

An ample array of Agrobacterium strains may share the same soil environment where is most probably that horizontal DNA transfer occurs (Bouzar et al. 1993; Kechris et al. 2006). Considering this, great levels of diversity might be present in soils within isolates of the same species and/or within different species. This diversity might be reflected, among other things, in strain virulence - their capacity to infect and provoke disease on plants (Ryder et al. 1985; Nesme et al. 1987). However, in any plant microbe interaction, the crucial signals are provided by both partners (Kado 1991). Therefore, plants might also differ in their susceptibility to Agrobacterium (Gelvin 2010), i.e. some varieties of grapes, blueberries and raspberries are particularly susceptible to different species or biovars of Agrobacterium (Benjama et al. 2002).

Beginning in the 1990's, the production and culture of blueberries in Argentina has experienced a considerable growth. An expansion of the production area throughout the country as well as an increase in yield, has resulted in an 8,000 t annual export commodity. Blueberries are a group of native species of the northern hemisphere of America that belong to the genus *Vaccinium*, family Ericaceae; among them, a small group of species has commercial value i.e. *Vaccinium corymbosum*.

Nurseries located in different areas spread out within Argentina are providing *V. corymbosum* plants to farmers to initiate production. Symptomatic plants from nurseries as well as from commercial farms were sent to our laboratory between 2006 and 2008. Preliminary analysis showed that almost all of them presented, at the base of the shoot or on the root, crown gall-like structures. Therefore, the aim of this study was to evaluate if these galls were induced by bacteria. If this proved to be true, then we wanted to confirm the identity of the isolates, determine their pathogenic ability and analyze their diversity.

#### Materials and methods

#### Bacterial isolates

The isolates of *Agrobacterium* were obtained from diseased plants of *Vaccinium corymbosum* (n=78) from nurseries (n=44) and also from commercial farms (n=34) that presented galls close to the root crown and/or hairy roots (Table 1); therefore the survey was unevenly distributed across these sites. Three representatives of *Agrobacterium* that were isolated from black raspberries (*Rubus spp.*) and also *Agrobacterium tumefaciens* strains ATCC15955, LBA 958 and K198 and *Agrobacterium radiobacter* (K1026) were included as controls, so the total number of agrobacteria studied was 85. In addition, two strains of *Rhizobium*, two *Ensifer*, one *Bradyrhizobium* and one strain of *Pantoea* were included for comparison (see Table 1).

#### Isolations from symptomatic plants

Bacterial isolation from diseased plants was performed as follows; galls were surface sterilized by immersing them in 1 % sodium hypochlorite for 15-20 min and then, were washed three times with sterile distilled water for 15 min each. Then, galls were cut in  $2 \times$ 2 cm squares by means of a sterile scalpel, mixed with 15 ml of sterile distilled water and homogenized by vortexing for 5 min. The aqueous suspension was incubated at room temperature for 45 min and then, slants of each homogenate were made on the following media: D1 (Moore et al. 2001), D1-M (Perry and Kado 1982) and YEM Congo red (Vincent 1970) supplemented with potassium tellurite, to increase the medium selectivity (Mougel et al. 2001). Then, the plates were incubated in the darkness at 27 °C for 7 days and observations of culture development were made every day. All the colonies that developed in the different media assayed, whose morphological characteristics were similar to those of Agrobacterium, were purified in yeast dextrose carbonate agar (YDC) (Moore et al. 2001). Isolates were cultured on YEM and then aliquots were stored in 20 % glycerol (V/V) at -70 °C.

#### Isolates characterization

All the isolates of *Agrobacterium* from *Vaccinium corymbosum* (n=78) were characterized morphologically, biochemically and physiologically by means of

the following reactions: Gram staining, catalase production, oxidase activity from isolates growing on both Nutrient agar (Ox-NA) and glucose nutrient agar (Ox-GNA) acid production on YDC, utilization of Keto-lactose, and oxidative/fermentative metabolism of glucose (OF-O and OX-F) following standard procedures (Moore et al. 2001). In addition, the isolates were characterized by means of the API 20E strips (Biomerieux,<sup>®</sup>), which included  $\beta$ -galactosidase utilization (ONPG), arginine dihidrolase (ADH), lysine decarboxilase (LDC), ornithine decarboxylase (ODC), citrate utilization (CIT), hydrogen sulfide production (H<sub>2</sub>S), urease production (URE), tryptophane deaminase (TDA), indole production (IND), acetoine production (Voges-Proskauer, VP), gelatinase activity (GEL), utilization of d-Glucose (GLU), d-mannitol (MAN), inositol (INO), d-sorbitol (SOR), 1-rhamnose (RHA), d-sucrose (SAC), d-melibiose (MEL), amygdalin (AMY), 1-arabinose (ARA) and nitrate reduction (NO<sub>3</sub>-NO<sub>2</sub>). In all cases, strains ATCC15955, LBA 958 and K198 and K1026 were included as controls.

#### Pathogenicity tests

Two assays were conducted, one consisted in using a cut leaf of Kalanchoe (Bryophyllum daigremontiana) (Minnemeyer et al. 2006). Briefly, 3-4 cm long young leaves were surface sterilized by immersing them in 2.5 % chlorine bleach solution for 15 min and were subsequently rinsed thoroughly in sterile distilled water. Leaves were injured with a sterile scalpel and 5 µl of 10<sup>8</sup> CFU/ml bacterial suspensions in sterile distilled water made by adjusting the cell concentration of 48 h old YEM cultures were pipette on each wound. Six leaves per strain were inoculated and placed in plates with water agar (2 %) supplemented with 50 µg/ml cicloheximide. They were incubated at 26 °C±2 °C under a 12 h light photoperiod achieved by eight Gro Klux MP100 tubes. In addition, root inducers isolates of Agrobacterium were evaluated by means of carrot discs assay (Ryder et al. 1985). Based on the analysis described above and those described below, we selected five representatives of Agrobacterium rubi (F210, F253, F304, F305 and F315), two of A. tumefaciens bv. 1 (F268 and F293) and two of A. rhizogenes bv. 2 (F288 and F289) and evaluated their pathogenicity on several hosts such as Blueberry (Vaccinium corymbosum cv. Misty) (50-60 cm high), 5-week old pepper (Capsicum annuum cv. California wonder), 5-week

Table 1 Bacterial isolates obtained from diseased blueberries and control strains of Agrobacterium and other strains used in thi	s paper
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Strain <sup>a</sup>	Species <sup>b</sup>	Biovar <sup>c</sup>	Host <sup>d</sup>	Cultivar <sup>e</sup>	Place of isolation <sup>f</sup>	Source <sup>g</sup>	Year <sup>h</sup>	RFLP pattern <sup>i</sup>
F 210	Agrobacterium rubi	rubi	blueberry	U	Buenos Aires	Ν	1997	G
F 253	A. rubi	rubi	blueberry	U	Tucumán	Ν	2006	А
F 254	A. rubi	rubi	blueberry	U	Tucumán	Ν	2006	А
F 255	A. rhizogenes	2	blueberry	Blue Crisp	Tucumán	F	2006	Е
F 256	A. rhizogenes	2	blueberry	Misty	Buenos Aires	Ν	2006	D
F 258	A. rhizogenes	2	blueberry	U	Tucumán	F	2006	В
F 259	A. tumefaciens	1	blueberry	U	Tucuman	F	2006	А
F 260	A. rhizogenes	2	blueberry	Misty	Buenos Aires	Ν	2006	G
F 261	A. rhizogenes	2	blueberry	U	Tucumán	F	2006	В
F 262	A. rhizogenes	2	blueberry	O'Neil	Tucumán	F	2006	В
F 263	A. rhizogenes	2	blueberry	U	Buenos Aires	Ν	2006	А
F 264	A. rhizogenes	2	blueberry	O'Neil	Tucumán	F	2006	G
F 265	A. rhizogenes	2	blueberry	Misty	Tucumán	F	2006	Е
F 266	A. rubi atypical strain	rubi	blueberry	U	Buenos Aires	Ν	2006	Е
F 268	A. tumefaciens	1	blueberry	U	Entre Ríos	F	2006	А
F 269	A. rhizogenes	2	blueberry	U	Entre Ríos	F	2006	В
F 270	A. rubi	rubi	blueberry	U	Entre Ríos	F	2006	В
F 271	A. rubi	rubi	blueberry	U	cvo. in vitro	Ν	2006	В
F 272	A. rubi	rubi	blueberry	O'Neil	E. Rios	Ν	2006	С
F 273	A. rubi	rubi	blueberry	O'Neil	E. Rios	Ν	2006	С
F 274	A. rhizogenes	2	blueberry	O'Neil	Tucumán	F	2006	В
F 276	A. tumefaciens	1	blueberry	Millenia	Tucumán	F	2006	А
F 277	A. rhizogenes	2	blueberry	O'Neil	Tucumán	F	2006	В
F 278	A. rhizogenes	2	blueberry	Blue Crisp	Tucumán	F	2006	В
F 279	A. tumefaciens	1	blueberry	Blue Crisp	Tucumán	F	2006	В
F 280	A. rhizogenes	2	blueberry	U	Entre Ríos	Ν	2006	D
F 281	A. tumefaciens	1	blueberry	U	Entre Ríos	Ν	2006	В
F 282	A. tumefaciens	1	blueberry	Blue Crisp	Tucumán	F	2006	В
F 283	A. rhizogenes	2	blueberry	U	Entre Ríos	Ν	2006	Е
F 284	A. tumefaciens	1	blueberry	U	Entre Ríos	Ν	2006	В
F 285	A. tumefaciens	1	blueberry	U	Tucumán	F	2006	В
F 286	A. rhizogenes	2	blueberry	Misty	Buenos Aires	F	2006	Е
F 287	A. rhizogenes	2	blueberry	Misty	Buenos Aires	F	2006	E
F 288	A. rhizogenes	2	blueberry	Millenia	Tucumán	F	2006	- C
F 289	A. rhizogenes	2	blueberry	O' Neil	Tucuman	F	2006	C
F 291	A. tumefaciens	1	blueberry	Millenia	Tucumán	F	2006	В
F 292	A tumefaciens	1	blueberry	Misty	Buenos Aires	N	2006	Н
F 293	A tumefaciens	1	blueberry	Blue Crisp	Buenos Aires	N	2006	A
F 294	A rhizogenes	2	blueberry	O'Neil	Buenos Aires	N	2006	G
F 295	A tymefaciens	-	blueberry	O'Neil	Buenos Aires	N	2006	B
F 296	A tumefaciens	1	blueberry	O'Neil	Buenos Aires	N	2006	B
F 297	A tumofacions	1	blueberry	Misty	Buenos Aires	N	2000	G
F 298	A tumpfacions	1	hlueberry	Misty	Buenos Aires	N	2000	B
F 200	A tumofacions	1	hlueberry	Blue Crien	Buenos Aires	N	2000	D
1 477	A. iumejuciens	1	oncoenty	Diac Crisp	Ducitos Alics	T.M.	2000	D

#### Table 1 (continued)

Strain <sup>a</sup>	Species <sup>b</sup>	Biovar <sup>c</sup>	Host <sup>d</sup>	Cultivar <sup>e</sup>	Place of isolation $^{\mathrm{f}}$	Source <sup>g</sup>	Year <sup>h</sup>	RFLP pattern <sup>i</sup>
F300	A. tumefaciens	1	blueberry	O' Neil	Buenos Aires	Ν	2006	D
F 301	A. rhizogenes	2	blueberry	O'Neil	Buenos Aires	Ν	2006	D
F 302	A. rhizogenes	2	blueberry	Blue Crisp	Buenos Aires	Ν	2006	Е
F 303	A. rhizogenes	2	blueberry	Blue Crisp	Buenos Aires	Ν	2006	Е
F 304	A. rubi	rubi	blueberry	Misty	Buenos Aires	Ν	2006	А
F 305	A. rubi	rubi	blueberry	Misty	Buenos Aires	Ν	2006	Е
F 306	A. tumefaciens	1	blueberry	Blue Crisp	Buenos Aires	Ν	2006	В
F 309	A. tumefaciens	1	blueberry	Misty	Tucumán	F	2006	В
F 310	A. tumefaciens	1	blueberry	Misty	Buenos Aires	Ν	2006	В
F 311	A. rhizogenes	2	blueberry	Blue Crisp	Tucumán	Ν	2006	G
F312	A. rhizogeness	2	blueberry	Blue Crisp	Buenos Aires	Ν	2006	В
F 313	A. rhizogenes	2	blueberry	Misty	Buenos Aires	Ν	2006	В
F 314	A. tumefaciens	1	blueberry	U	Tucumán	F	2006	В
F 315	A. rubi	rubi	blueberry	U	Tucumán	F	2006	А
F 316	A. tumefaciens	1	blueberry	U	Tucumán	F	2006	В
F 317	A. rhizogenes	2	blueberry	U	Tucumán	F	2006	В
F 319	A. tumefaciens	1	blueberry	Blue Crisp	Buenos Aires	Ν	2006	А
F 320	A. tumefaciens	1	blueberry	U	Tucumán	F	2006	А
F 321	A. tumefaciens	1	blueberry	U	Buenos Aires	F	2006	В
F 332	A. tumefaciens	1	blueberry	U	Corrientes	F	2007	В
F 333	A. tumefaciens	1	blueberry	O' Neill	Buenos Aires	F	2007	B
F 334	A. tumefaciens	1	blueberry	Mistv	Buenos Aires	N	2007	B
F 335	A. rubi	rubi	blueberry	U	Tucumán	N	2007	B
F 336	A. tumefaciens	1	blueberry	U	Tucumán	N	2007	B
F 337	A. rubi	rubi	blueberry	U	Tucumán	N	2007	G
F 338	A. rubi	rubi	blueberry	U	Tucumán	N	2007	В
F 339	A rhizogenes	2	blueberry	U	Tucumán	N	2007	Н
F 340	A rhizogenes	2	blueberry	U	Tucumán	N	2007	Н
F 341	A tumefaciens	1	blueberry	U	Tucumán	N	2007	A
F 346	A tumefaciens	1	blueberry	Mistv	Tucumán	F	2007	B
F 347	A tumefaciens	1	blueberry	Yewell	Tucumán	F	2007	B
F 348	A tumefaciens	1	blueberry	Emerald	Tucumán	N	2007	B
F 349	A rhizogenes	2	blueberry	Gold Coast	Tucumán	N	2007	B
F 400	A rhizogenes	2	blueberry	U	Rs As	F	2007	B
F 239	A tumefaciens	1	raspherry	Heritage	Buenos Aires	F	2000	B
F 240	A tumefaciens	1	raspberry	Rliss	Buenos Aires	F	2004	B
F 241	A tumefaciens	1	raspberry	Rubi	Buenos Aires	F	2004	B
K 1026	A radiobacter	2	I	I	LI	I	11	B
I BA 958	A tumefaciens	1	U	U	U	U	U	B
ATCC 15955	A tumofacions	1	U	U	U	U	U	B
K108	A tumofacions	1	neach	U U	Australia	U U	U	B
K170 Hambi540	л. интеристентs Phizohium galagaa	1	Galaga sp	U	Finland	U	U	٨
CEN4	R atli by -11	_	Galega sp.	U		U	U	A
100	K. etti ov. phaseoli		oean	U	U	U	U	A
109	Ensijer japonicum	-	soybean	U	U	U	U	A

#### Table 1 (continued)

Strain <sup>a</sup>	Species <sup>b</sup>	Biovar <sup>c</sup>	Host <sup>d</sup>	Cultivar <sup>e</sup>	Place of isolation <sup>f</sup>	Source <sup>g</sup>	Year <sup>h</sup>	RFLP pattern <sup>i</sup>
103-1 HH	Ensifer fredii	-	soybean	U	U	U	U	А
USDA 1002T	E. meliloti	-	lucerne	U	U	U	U	Н
F 347	Pantoea ananatis	_	maize	U	Argentina	U	2009	F

<sup>a</sup> dentification number

<sup>b</sup> Species of Agrobacterium isolated

<sup>c</sup> biovar designation according to multiplex PCR

<sup>d</sup> host of isolation

<sup>e</sup> cultivar of Vaccinium corymbosum or U unknown

<sup>f</sup>place of isolation

 $^{g}N$  nursery- F farm; h: year of isolation

<sup>i</sup>Restriction patterns of the 16SrDNA generated by combining the fragments obtained by means of TaqI and HaeIII

old tomato (*Solanum lycopersicum*, formerly *Lycopersicum esculentum* cv. Presto), sunflower (*Helianthus annuum* cv. Dekalb 4040) and 2- to 3- months old tobacco (30–40 cm high) (*Nicotiana tabacum cv.* Xanthi) as described by Peluso and co-workers (2003). Briefly, plants were inoculated by a stabbing a needle containing a bacterial cell suspension ( $10^8$  CFU/ml) of each strain tested or with sterile distilled water (control treatment). Each strain was inoculated on two plants of each species. Inoculated plants were maintained in a greenhouse at  $24\pm5$  °C and the appearance of tumours or hairy roots was visually assessed in herbaceous species 4–6 weeks after inoculation and in blueberries after 3 to 6 months.

Analysis of diversity and identification of the isolates by molecular tools

# DNA preparation

Genomic DNA was isolated by using an interchange ionic resin as described by Alippi et al. (2003). Briefly, all bacterial strains listed in Table 1 were grown in YEM for 24 h at 28 °C. Cells from about one to two single colonies were picked with a sterile toothpick and suspended in 300  $\mu$ l NaCl 1 M. Each sample was vortex-mixed and centrifuged at 16,000 g for 4 min, and once, supernatant was removed and the pellet resuspended in 300  $\mu$ l double-distilled water, vortexmixed and centrifuged at 16,000 g for 3 min. The supernatant was removed and the pellet was resuspended in 150  $\mu$ l of an aqueous suspension of 6 % resin (Chelex<sup>®</sup> 100, 200–400 mesh, Bio-Rad). The mixture of cells and resin was incubated at 56 °C for 20 min and vortex-mixed for 30 s. Finally the mixture was incubated at 99 °C for 8 min and vortex-mixed for 1 min. Bacterial debris and resin were removed by centrifugation, and 5  $\mu$ l of the supernatant containing DNA was used as template for PCR amplifications in a final volume of 25  $\mu$ l.

# PCR detection of virC operon located in Ti and Ri plasmids

A 730 bp fragment representing the partial sequence of *vir*C operon found in Ti and Ri plasmids present on pathogenic strains of *Agrobacterium* species was amplified by means of VCF/VCR primers (Sawada et al. 1995). Five 5  $\mu$ l of genomic DNA isolated as described before was used as template in a PCR reaction with a final volume of 25  $\mu$ l. PCR products were resolved in 1.6 % agarose gels in TBE buffer 0.5 X, and observed under UV light after staining with ethidium bromide. All the isolates listed in Table 1 were compared in the same manner.

# Multiplex PCR

A PCR multiplex reaction based on differences at the 23SrDNA allowed us to discriminate between *Agrobacterium* biovar 1, *Agrobacterium* biovar 2, *A. rubi* and *A. vitis*, respectively. (Pulawska et al. 2006). PCR mixtures and amplifications were performed as described by Pulawska and co-workers (2006), but using

as template 5  $\mu$ l of genomic DNA isolated as specified before. PCR reactions were performed in a thermal cycler (Mastercycler Personal; Eppendorf, Hamburg, Germany) programmed as described (Pulawska et al. 2006). PCR products were resolved in 1.6 % agarose gels in TBE buffer 0.5 X, stain with ethidium bromide and observed under UV light. In addition to all the isolates listed in Table 1, strains K 306 from Australia and S4 from Hungary were included as positive controls of *Agrobacterium vitis*.

# RFLP analysis of PCR-amplified 16S rDNA

We used genomic DNA as template to amplify the full sequence of the 16SrDNA by means of primers fD1 and rD1 (Weisburg et al. 1991). 4- $\mu$ l aliquots of the amplified 16SrDNA were incubated overnight with restriction enzymes *Taq* I (Promega Biotech) and *Hae* III (Promega Biotech) at 65 °C and 37 °C, respectively according to the conditions suggested by the manufacturer. RFLP was resolved by electrophoresis in 2 % agarose gels, the DNA was stained with ethidium bromide and visualized with a UV transiluminator (UVP). All the isolates listed in Table 1 were analyzed.

Analysis of the diversity of isolates by rep-PCR using primers ERIC

Diversity among isolates was assessed based upon fingerprints generated by means of primers associated to Enterobacterial Repetitive Consensus sequences (ERIC) (Versalovic et al. 1994). All PCR reactions were performed in a thermal cycler (Mastercycler Personal; Eppendorf, Hamburg, Germany) as described by López and Alippi (2007). A total of 82 agrobacteria were compared (78 from blueberries and four from Culture Collections) Reactions were resolved in 1.5 % agarose gels that were visualized and analyzed as previously described. Gel images were digitalized and photographed using a digital image capture gel documentation system (Digi Doc-it, UVP, v. 1.1.25) and analyzed by means of Gelcompar II (v 5.1. Applied Math, Kortrijk, Belgium). Cluster analysis was performed using the DICE similarity coefficient and the UPGMA (Unweighted Pair Group Method with Arithmetic Mean) clustering algorithm with a band tolerance of 5 %.

#### Sequence analysis

By sequencing the 16SrDNA (Sanger, et al. 1977) one representative of A. rubi (F266, GU580894), one representative of Agrobacterium rhizogenes bv. 2 carrying Ri plasmid (F289, GU580895) (Alippi et al. 2010) and one representative of Agrobacterium tumefaciens bv.1 carrying Ti plasmid (F268, GU580896) (this paper) were analyzed further. Individual sequences were assembled by means of the GCG Software (University of Wisconsin) and aligned by means of BOXSHADE 3.21 (http:// www.ch.embnet.org/software/BOX form.html). The dendrogram was based on the multiple alignment of the 16SrDNA full sequences by means of the Clustal W version 1.81. This information was processed to build a tree based on the UPGMA by means of Mega 5.0. The support of the groups within the tree was evaluated through bootstrap (Felsenstein 1985), with 1,000 replications. The values obtained are given on the basis of the branches.

#### Results

Bacterial isolations and colony characteristics in different media

All the plants analyzed presented either gall-like lesion on the crown roots or at the base of the shoot or hairy roots. From all these lesions we isolated Gram (-) bacteria, that developed colonies that presented, on the different media tested, morphological characteristics typical of Agrobacterium. On D1 they were of a green-yellowish colour, darker in the centres with creamy white edges. The initial light blue colonies turned to olive green as they aged but always with lighter edges. On D1M colonies were convex, bright, mucoid, of a faint blue colour with darker centres. In YEM-CRT, colonies appeared black, mucous and convex, with entire margins. On YDC, colonies were highly mucous, bright, convex and white cream to light beige in colour, some isolates acidified the medium as revealed by clear halos around the colonies. Based upon this, we decided to confirm by means of biological as well as molecular tools that the identity of the isolates was Agrobacterium and also analyzed their diversity. We obtained 78 Agrobacterium isolates from Vaccinium corymbosum plants showing symptoms,

44 from nurseries and 34 from commercial blueberry crops.

Characterization of bacteria by biochemical and physiological tests

We characterized the Agrobacterium isolates further by means of biochemical reactions. In Fig. 1 we present the data indicating the percentage of isolates of Agrobacterium bv. 1, Agrobacterium bv. 2 and A. rubi that gave a positive response to each of the biochemical reactions tested, including those from the API20E strips on a total of 78 isolates. Among 27 different biochemical reactions there were only four that were positive among all the isolates like glucose (GLU), arabinose (ARA), oxidative metabolism of glucose (OX-O), and oxidase activity of cultures growing in NA (OxNA). Only three other reactions, lysine decarboxilase (LDC), hydrogen sulphide production  $(H_2S)$ , and indole (IND) were negative for all the isolates, except for the utilization of 3-ketolactose (KETO-LAC) that was only positive for Agrobacterium by. 1 isolates (Fig. 1). Analysis of the remaining reactions shows it is obvious that there is no characteristic pattern for Agrobacterium tumefaciens bv.1, Agrobacterium rhizogenes by. 2 or A. rubi isolated from Vaccinium corymbosum (Fig. 1).

# Pathogenicity tests

A virulence assay was performed by inoculating leaves of Kalanchoe with a bacterial suspension of each of the Agrobacterium isolates. All of them provoked cell proliferations, mostly after 10-12 days of incubation at room temperature. On the contrary, Agrobacterium radiobacter K1026 strain, Pantoea ananatis or rhizobia such as Rhizobium galegae, Rhizobium etli bv. phaseoli, Ensifer melilotii, Ensifer japonicum, and Sinorhizobium freedii included as controls, did not induce either galls or hairy roots formation. The symptoms developed by two plants suggested that the causative agents might be representatives of Agrobacterium rhizogenes or Agrobacterium by. 2. Therefore, these two isolates F288 and F289, five A. rubi isolates (F210, F253, F304, F305, F315) and two Agrobacterium bv.1 isolates (F268 and F293) were evaluated by means of additional biological tests. A. rubi and A. tumefaciens by. 1 isolates developed gall like structures on V. corymbosum (highbush blueberry), Bryophyllum daigremontiana (Kalanchoe, whole plants), *Nicotiana tabacum* (Tobacco), *Lycopersicum esculentum* (Tomato), *Capsicum annuum* (Pepper) and *Helianthus annuum* (Sunflower), while isolates F288 and F289 only induced proliferation of roots in all the inoculated hosts which was also observed in the carrot disk assay.

# PCR detection of virC operon

All the Agrobacterium isolates from blueberries and control strains (LBA958, ATCC15955, and K198) proved to have in their genome a 730 bp PCR sequence coding for virulence gene *vir*C (Sawada et al. 1995). On the contrary, *Agrobacterium radiobacter* K1026 strain as well as *Pantoea ananatis* or rhizobia (*Bradyrhizobium japonicum*, *Rhizobium galegae*, *Rhizobium etli* bv. *phaseoli*, *Ensifer melilotii* and *Ensifer freedii*) lacked this sequence.

# Multiplex PCR

The multiplex reaction consisted of amplifying DNA fragments of four different sizes from the 23SrDNA template. Within the Agrobacterium strains isolated from blueberries, 14 generated an amplicon of 1,006 bp, 31 generated an amplicon of 1,066 bp and 33 generated an amplicon of 184 bp, that corresponded to Agrobacterium rubi, Agrobacterium bv. 2 and Agrobacterium bv. 1, respectively (Pulawska et al. 2006). The three isolates from raspberries corresponded to by. 1, reference strains LBA958, ATCC15955 and K 198 were by. 1, while strain K1026 (previously named A. radiobacter) belonged to by. 2, and A. vitis strains S4 and K306 generated an amplicon of 478 bp. Genomic DNA of Bradyrhizobium japonicum, Rhizobium galegae, Rhizobium etli bv. phaseoli, Ensifer melilotii, Ensifer freedii and Pantoea ananatis did not act as templates of any of the described amplicons (Table 1).

# RFLP analysis of PCR-amplified 16S rDNA

Considering that all the isolates belonged to *Agrobacterium* species we amplified the full sequence of the 16SrDNA and made an enzyme restriction analysis. Even though the 16SrDNA is a universal conserved sequence it varied considerably among *Agrobacterium* isolates; seven different restriction patterns were found among the agrobacteria isolated from blueberries (n=78) and those strains of agrobacteria (n=7) and



□ Agrobacterium bv. 1 ■ Agrobacterium bv. 2 SAgrobacterium rubi

Fig. 1 Biochemical characterization of *Agrobacterium* isolates. Percentage of isolates of *A. tumefaciens* biovars 1 and 2 and *A. rubi* that gave a positive response in the analysis. Names of each reaction are listed in Materials and Methods section

rhizobia (n=5) that were used as controls (Table 1 and Fig. 1). Each restriction pattern occurred at different frequency within agrobacteria. Agrobacterium tumefaciens by. 1 strains carrying the Ti plasmid presented the following patterns: A, B, D, G and H; Agrobacterium rhizogenes by.2 carrying the Ti plasmid presented A, B, D, E, G and H. Agrobacterium bv. 2 carrying the Ri plasmid presented only pattern C most probably due to the fact that there were only two isolates and those strains characterized as A. rubi presented the following 16SrDNA patterns: A, B, C, E and G. The most common pattern found within the agrobacteria populations was B with a total of 45 isolates (52 %), which also included all the reference strains, while patterns A, C, D, E, G and H represent 14 %, 5 %, 6 %, 11 % and 8 % respectively (Table 1 and Fig. 2). When analyzing the rhizobial isolates that were used as controls, we found the presence of pattern A in 4 strains (R. galegae, R. etli bv. phaseoli, B. japonicum and E. fredii) and pattern H in one strain of E. meliloti (USDA 1002 T). Additionally, Pantoea ananatis had the F pattern (Table 1).

Analysis of the diversity of isolates by rep-PCR using primers ERIC

The diversity of the isolates from blueberries and Culture Collections (n=82) was analyzed by ERIC-PCR generating 72 different patterns that included four to 13 amplified bands that range in size between 50 and 1,500 bp (Fig. 3). Although the fingerprint patterns were highly diverse, they were clustered at a low similarity level (about 45 %), two main clusters, named I and II were separated at a similarity index of 65 % and 60 % respectively. The rest of the strains (n=10) clustered at a similarity level of 45 % (Fig. 3).

#### Sequence analysis

Identity of three selected isolates was confirmed by sequencing the 16SrDNA of strains F 266, Agrobacterium rubi (GenBank No. GU580894), F 289 Agrobacterium rhizogenes bv. 2 (GU580895) and F268, Agrobacterium tumefaciens bv.1 (GU580896). These sequences were compared with those of Agrobacterium tumefaciens, Agrobacterium rhizogenes, Agrobacterium rubi, Agrobacterium vitis, and Agrobacterium *larrymoorei*, and also with those of the new described species Rhizobium pusense and Rhizobium skierniwicense. In addition, the 16S rDNA sequence of Bradvrhizobium japonicum type strain was used as an outgroup. These sequences were aligned by means of the Clustal W multiple alignment and then a tree was built based on the Maximum Likelihood algorithm. As expected, the sequences were highly related (Fig. 4) and were monophyletic, though grouped in three clusters supported with 1,000 boostraps. Cluster A included four representatives of A. rubi (AY626394.1; X67228.1-type strain-; D14503.1 and EU281314.1), two representatives of A. larrymorei (AY626382.1 and Z30542.1 -type strain-), four representatives of A. tumefaciens (F268 GU580896.1; AY623686.1; GU645017.1 and D14500), R. pusense (FJ969841.2) and R. skierniewicense (HQ 823552.1 -type strain-); cluster B included three representatives of A. vitis (AY636399.1, D14502.1 and AY626401.1) clearly





Fig. 2 Restriction fragment lenght polymorphism (RFLP) patterns of PCR-amplified 16SrDNA found among all the isolates from blueberries tested (n=78). The full sequence of the 16SrDNA was digested with two restriction enzymes *Taq*I and

separated from the rest of the isolates; and cluster C included *A. rubi* F266 (atypical strain) (GU580894.2), one representative of *A. rubi* AY626395.1) and five representatives of *A. rhizogenes* (F289, GU580895.1; AY6263991.1; AY626389.1,D14501.1 and AY945955.1 –type strain-) (Fig. 4).

# Discussion

The 78 bacterial isolates obtained from diseased blueberry plants with symptoms such as root proliferations or galls, grew on selective media and provoked cell proliferation in detached leaves of Kalanchoe. They presented a variable biochemical pattern and even key enzymes were variable, while a few of them were hardly present in a few isolates within *Agrobacterium* bv. 1 and *Agrobacterium* bv. 2 strains and *A. rubi* (Fig. 1). Diversity was confirmed by molecular tools such as ERIC-PCR where there was no correlation between biovars and fingerprint patterns and/or geographical origin of the isolates, since representatives of *Agrobacterium* bv. 1, *Agrobacterium* bv. 2 and *A.* 

HaeIII, respectively. Reference strains of Agrobacterium (n=7)

showed a pattern B and the rhizobia used as controls showed

patterns A (4 strains) and H (1 strain), respectively

Fig. 3 Diversity at the genetic level among 78 strains of *Agrobacterium* isolated from blueberries based on the presence or absence of bands generated by means of ERIC markers. Pairwise comparisons were calculated using the DICE similarity coefficient. A dendrogram was generated from a similarity matrix using unweighted pair group method with arithmetical averages (UPGMA) with a band tolerance of 5 % by means of Gelcompar II v 5.1



Deringer



0.01

Fig. 4 Rooted phylogenetic tree generated based on the 16SrDNA showing the relationship of three *Agrobacterium* isolates each representing *A. tumefaciens* (GU580896), *A. rhizogenes* (GU580895) and *A. rubi* atypical strain (GU580894) provoking symptoms on blueberries, with annotated sequences

rubi, as well as strains from Culture Collections were distributed along the different clusters (Fig. 3). This is in agreement with other authors that also reported high levels of diversity within Agrobacterium representatives (Nesme et al. 1987; Peluso et al. 2003). Therefore, the isolates of Agrobacterium affecting blueberries in Argentina, are like other isolates, quite variable. Some biochemical reaction characteristics of Agrobacterium were negative, which might be related to the methodology used. Based on the phenotypic analysis A. rubi seems to be atypical, though more closely related to Agrobacterium bv.1 (Tighe et al. 2000). Therefore, these results confirmed the variable nature of Agrobacterium isolated from diseased blueberries from Argentina and argue against the reliability of an identification or characterization of Agrobacterium based only on such tools.

of representatives of *Agrobacterium Rhizobium* and *Bradyrrizobium* species. The support of the groups within the tree was evaluated through bootstrap, with 1,000 replications. The values obtained are given on the basis of the branches

Among the nomenclatures of the genus *Agrobacterium*, the traditional pathogenicity-based classification that relies on the pathogenic ability of the isolates, which is coded by genes located on plasmids, might not be the most accurate, considering that plasmids are dispensable and transferable between organisms in their environment. The other nomenclature is the one that refers phenotypic differences to biovars (Holmes and Roberts 1981; Moore et al. 2001; Young et al. 2005), which was repeatedly confirmed by both sequence comparison (Pulawska et al. 2006; Slater et al. 2009; Pulawska et al. 2012) and fatty acid analysis (Tighe et al. 2000).

The multiplex PCR of the blueberries isolates (n=78) indicated that 33 were *Agrobacterium* bv. 1 (Syn. *A. tumefaciens*) (42 %), 31 *Agrobacterium* bv. 2 (40 %) and 14 isolates were *Agrobacterium rubi* (18 %).

Regarding reference strains of *Agrobacterium*, all of them including the three isolates from *Rubus* spp. belonged to bv. 1 while the K 1026 strain of *A. radiobacter* belonged to bv. 2, and as expected, the strains S4 and K306 were *A. vitis*.

Among all the isolated bacteria, we selected five representatives of A. rubi, two of Agrobacterium bv. 1 and two of Agrobacterium by. 2 for further analysis. Five A. rubi and two Agrobacterium by. 1 induced galls on V. corymbossum (blueberry), Bryophyllum daigremontiana (Kalanchoe), Nicotiana tabacum (Tobacco), Lycopersicum esculentum (Tomato), Capsicum annuum (Pepper) and Helianthus annuum (Sunflower) Agrobacterium bv.2 isolates (F288 and F 289) only induced root proliferations in those hosts and in carrot disks. Therefore the identity of the isolates was A. rubi, A. tumefaciens and A. rhizogenes, respectively. Both the biological assays as well as polymorphisms at the 5' sequence of the 23SrDNA (Bautista-Zapanta et al. 2009), support that 33 isolates from blueberries were Agrobacterium bv. 1 (42 %), 31 isolates were Agrobacterium bv. 2 (40 %) and 14 were A. rubi (18 %).

The identity of an Agrobacterium by. 1 (F268) that induced galls (this paper) and one representative of Agrobacterium bv. 2 that provoked root proliferation (F289) (Alippi et al. 2010) were confirmed by the 16SrDNA full sequence. In relation to the strain F266, previously classified as A. rubi based on a multiplex PCR and biochemical tests, the phylogenetic analysis indicated that it is closely related to strain F289 (A. rhizogenes by. 2) and that they both formed part of a cluster that include other representatives of Agrobacterium rhizogenes (n=5) and one A. rubi strains (Fig. 4). However, isolate F 266, with the exception of its ability to metabolize citrate, which was a variable trait among putative A. rubi isolates, showed a biochemical pattern similar to those reported for A. rubi (i.e. positive for β-galactosidase, D-glucose, D-arabinose, n-acetylglucosamine, maltose, mannitol, malonate, urease but negative for L-arabinose, saccharose, lysine decarboxylase, H<sub>2</sub>S production, indole and 3-ketolactose production). Additionally, the sequence of A. tumefaciens isolate F268 was phylogenetically closely related to other A. tumefaciens strains and also to other representatives of A. rubi, A larrymoorei and R. skierniewicense. Young et al. (2004) also found an Agrobacterium rubi strain closely related to Agrobacterium rhizogenes, suggesting that it might be A. rhizogenes rather than A. rubi. On the other hand, Farrand et al. (2003) while analyzing the 16SrDNA sequences stated that *A. rubi* is atypical but more closely related to biovar 1 isolates than to biovar 2 ones. The fact that the phylogenetic analysis based on the 16SrDNA suggest a close association of isolate F266 with *A. rhizogenes* by. 2 while the multiplex PCR and biochemical tests suggest that the isolate is *A. rubi*, raises some questions about the identity of the isolate, that we called atypical, therefore further work should be done to clarify this.

Even though all these organisms are closely related, the phylogenetic analysis confirmed the complexity and variability of the genus *Agrobacterium*. In addition to this, the analysis of Table 1 concomitantly with the identity of the isolates of agrobacteria showed that there was no correlation between the place of origin of sampled plants and the *Agrobacterium* species infecting them. *Agrobacterium* bv. 1, *Agrobacterium* bv. 2 and *A. rubi* were isolated from diseased plants that were sent from nurseries (n=44) and also from field commercial crops (n=34) within four provinces that is Tucumán, Entre Ríos, Corrientes and Buenos Aires.

In Argentina, there is no information about the diversity of agrobacteria affecting *Vaccinium corymbosum* and not much is known about the number and diversity of the *Agrobacterium* population in the soils. Diseased blueberry plants are probably the main way *Agrobacterium* is disseminating into new areas. As many of the analyzed blueberry plants originate in nurseries (n=44) from several places in Argentina; therefore, unknowingly nurseries might be threatening new areas of production by introducing the pathogen through infected plants. The number of isolates of each *Agrobacterium* species suggested that the evaluated nurseries are producing *Vaccinium* plants in infected soils and/or substrates containing mainly *Agrobacterium* by. 1 and by. 2 and low levels of *A. rubi*.

Like other bacterial-plant interactions, the genome of both Agrobacterium and the plant might contribute to the occurrence of the disease. While A. tumefaciens and A. rhizogenes have been found to infect many different hosts, A. rubi, A. vitis and A. larrymoorei have been associated mostly with Rubus spp. and Vaccinium spp., Vitis spp. and Ficus spp., respectively, suggesting that each species has a genetic component that is contributing to infection. It has been proposed that plants differ in their susceptibility to Agrobacterium. Some plants and/or cultivars are particularly susceptible to Agrobacterium (Gelvin 2010); for instance, compatibility of A. tumefaciens and Leuce poplar is controlled at the infraespecific level (Nesme et al. 1987). Regarding this, blueberries seem to be particularly susceptible to Agrobacterium since the frequency of pathogenic populations of Agrobacterium by. 1 and by. 2 isolated from Vaccinium were similar (42 % and 40 %, respectively), with A. rubi represented in a minor proportion (18 %). It is not clear, at this moment, if blueberries are particularly resistant to Agrobacterium isolates carrying the Ri plasmid or if the isolation of only two strains with this characteristic indirectly reflects the frequency of this organism in the soils of those farms. Soil samples from fields cultivated with blueberries (n=15) contained only pathogenic Agrobacterium by. 1 (94 %) and A. rubi (6 %) (data not shown). Future studies should include the isolation of Agrobacterium from soil sampled based on a systematic survey of the area with plant nurseries, which might help to more knowledge about the ecology of these organisms in the soils and also to estimate the risk of spreading the bacteria to other areas.

Agrobacterium is a complex genus and as such its taxonomy is still an issue under discussion (Young et al. 2001; Farrand et al. 2003; Young et al. 2004; Young et al. 2005; Gelvin 2010). In addition to this, it has been demonstrated that the plant mediates instability of A. tumefaciens populations (Dion et al. 1996). It has been demonstrated that non-pathogenic mutants appear in response to cultivation of Agrobacterium in the presence of substances naturally present in plant wounds (Goodner et al. 2001). In this study all the isolates obtained were pathogenic, though it is worth mentioning that we worked with only one colony of each gall, and this is contrary to the proposal of Anderson and Moore (1979) and Nesme and coworkers (1987) that the Agrobacterium population obtained from tumours consists mostly of nonpathogenic strains. Recently, Llop and co-workers (2009) isolated mutants derived from Agrobacterium strains inoculated in pepper and tomato. They found that the appearance of non-pathogenic mutants of Agrobacterium in tumours occurred at low frequency and differed within host species; furthermore, they demonstrated that the genetic changes occurred mostly in large areas of the Ti plasmid. This might explain, at least in part, why all the isolates studied in this work were pathogenic on different hosts.

We conclude that *Vaccinium* plants appear to be highly susceptible to *Agrobacterium* isolates, whether

they are *Agrobacterium* bv. 1, *Agrobacterium* bv. 2 or *A. rubi*. The frequency of each *Agrobacterium* species infecting nurseries and/or field plots of *Vaccinium* might indirectly reflect their presence in the soil. Further work should be done to clarify this.

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#### References

- Alippi, A. M., Dal Bo, E., Ronco, L. B., López, M. V., López, A. C., & Aguilar, O. M. (2003). *Pseudomonas* populations causing pith necrosis of tomato and pepper in Argentina are highly diverse. *Plant Pathology*, 52, 287–302.
- Alippi, A. M., López, A. C., & Balatti, P. A. (2010). First report of Agrobacterium rubi and Agrobacterium rhizogenes, causing crown and root gall and hairy root on blueberry in Argentina. Plant Disease, 94, 1064.
- Anderson, A. R., & Moore, L. W. (1979). Host specificity in the genus Agrobacterium. Phytopathology, 69, 320–323.
- Bautista-Zapanta, J.-N., Arafat, H. H., Tanaka, K., Sawada, H., & Suzuki, K. (2009). Variation of 16 S-23 S internally transcribed spacer sequence and intervening sequence in rDNA among the three major *Agrobacterium* species. *Microbiology Research*, 164, 604–612.
- Benjama, A., Boubaker, H., Khlief, Z., Krimi, Z., Nesme, X., López, M. M., & Zoina, A. (2002). Susceptibility of stonefruit rootstocks, rose and grapevine to Agrobacterium radiobacter var. tumefaciens in Arab Mediterranean countries. OEPP/EPPO Bulletin, 32, 463.
- Bouzar, H., & Jones, J. B. (2001). Agrobacterium larrymoorei sp. nov., a pathogen isolated from aerial tumors of Ficus benjama. International Journal of Systematics and Evolutionary Microbiology, 51, 1023–1026.
- Bouzar, H., Ouadah, D., Krimi, Z., Jones, J. B., Trovato, M., Petit, A., & Dessaux, Y. (1993). Correlative association between resident plasmids and the host chromosome in a diverse Agrobacterium Soil Population. *Applied and Envi*ronmental Microbiology, 59, 1310–1317.
- Costechareyre, D., Rhouma, A., Lavire, C., Portier, P., Chapulliot, D., Bertolla, F., Boubaker, A., Dessaux, Y., & Nesme, X. (2010). Rapid and efficient identification of *Agrobacterium* species by *recA* allele analysis. *Microbial Ecology*, 60, 862–872.
- Dion, P., Belanger, C., Marquis, C., Ream, W., & Gelvin, S. B. (1996). Ecological significance of avirulence in Agrobacterium. In W. Reamand & S. B. Gelvin (Eds.), Crown Gall

advances in understanding interkingdom gene transfer (pp. 44–58). St Paul: APS Press.

- Farrand, S. K., van Berkum, P., & Oger, P. (2003). Agrobacterium is a definable genus of the family Rhizobiaceae. International Journal of Systematics and Evolutionary Microbiology, 53, 1681–1687.
- Felsenstein, J. (1985). Confidence limits on phylogenies: an approach using bootstrap. *Evolution*, *39*, 783–791.
- Gelvin, S. B. (2010). Agrobacterium and plant genes involved in T-DNA transfer and integration. Annual Review of Plant Physiology and Plant Molecular Biology, 51, 223–256.
- Goodner, B., Hinkle, G., Gattung, S., Miller, N., Blanchard, M., Qurollo, B., Goldman, B. S., Cao, Y., Askenazi, M., Halling, C., Mullin, L., Houmiel, K., Gordon, J., Vaudin, M., Iartchouk, O., Epp, A., Liu, F., Wollam, C., Allinger, M., Doughty, D., Scott, C., Lappas, C., Markelz, B., Flanagan, C., Crowell, C., Gurson, J., Lomo, C., Sear, C., Strub, G., Chris Cielo, C., & Slater, S. (2001). Genome sequence of the plant pathogen and biotechnology agent *Agrobacterium tumefaciens* C58. *Science*, 294, 2323–2328.
- Holmes, B., & Roberts, P. (1981). The classification, identification and nomenclature of agrobacteria. *Journal of Applied Bacteriology*, 50, 443–467.
- Kado, C. I. (1991). Molecular mechanisms of crown gall tumorogenesis. *Critical Reviews in Plant Sciences*, 10, 1–32.
- Kechris, K. J., Lin, J. C., Bickel, P. J., & Glazer, A. N. (2006). Quantitative exploration of the occurrence of lateral gene transfer by using nitrogen fixation genes as a case study. *Proceedings of the National Academy of Sciences*, 103, 9584–9589.
- Lasalle, F., Campillo, T., Vial, L., Baude, J., Costechareyre, D., Chapulliot, D., Shams, M., Abrouk, D., Lavire, C., Oger-Desfeux, C., Hommais, F., Guéguen, L., Daubin, V., Muller, D., & Nesme, X. (2011). Genomic species are ecological species as revealed by comparative genomics in *Agrobacterium tumefaciens*. *Genome Biology and Evolution*, *3*, 762–781.
- Lim, S. H., Kim, J. G., & Kang, H. W. (2009). Novel scar primers for specific and sensitive detection of Agrobacterium vitis strains. Microbiology Research, 164, 461–463.
- Llop, P., Murillo, J., Lastra, B., & Lopez, M. M. (2009). Recovery of nonpathogenic mutant bacteria from tumors caused by several *Agrobacterium tumefaciens* strains: a frequent event? *Applied and Environmental Microbiology*, 75, 6504–6514.
- López, A. C., & Alippi, A. M. (2007). Phenotypic and genotypic diversity of *Bacillus cereus* isolates recovered from honey. *International Journal of Food Microbiology*, 117, 175–184.
- Minnemeyer, S. L., Lightfoot, R., & Matthysse, A. G. (2006). A semiquantitative bioassay for relative virulence of Agrobacterium tumefaciens strains on Bryophyllum daigremontiana. Journal of Bacteriology, 173, 7723–7724.
- Moore, L. W., Bouzard, H., & Burr, T. (2001). Agrobacterium. In N. W. Schaad, J. B. Jones, & W. Chun (Eds.), *Labora*tory guide for the identification of plant pathogenic bacteria (pp. 17–35). St. Paul: APS Press.
- Mougel, C., Cournoyer, B., & Nesme, X. (2001). Novel tellurite-amended media and specific chromosomal and Ti plasmid probes for direct analysis of soil populations of

Agrobacterium biovars 1 and 2. Applied and Environmental Microbiology, 67, 65–74.

- Nesme, X., Michel, M.-F., & Digat, B. (1987). Population heterogeneity of Agrobacterium tumefaciens in galls of Populus L. from a single nursery. Applied and Environmental Microbiology, 53, 655–659.
- Panday, D., Schumann, P., & Das, S. K. (2011). *Rhizobium pusense* sp. nov., isolated from the rhizosphere of chickpea (*Cicer arietinum L.*). *International Journal of Systematic and Evolutionary Microbiology*, *61*, 2632–2639.
- Peluso, R., Raio, A., Morra, F., & Zoina, A. (2003). Physiological, biochemical and molecular analyses of an Italian collection of *Agrobacterium tumefaciens* strains. *European Journal of Plant Pathology*, 109, 291–300.
- Perry, K. L., & Kado, C. I. (1982). Characteristics of Ti plasmids from broad-host range and ecologically specific biotype 2 and 3 strains for *Agrobacterium tumefaciens*. *Journal of Bacteriology*, 151, 343–350.
- Portier, P., Le Saux, M. F., Mougel, C., Lerondelle, C., Chapulliot, D., Thiolouse, J., & Nesme, X. (2006). Identification of genomic species in *Agrobacterium* biovar 1 by AFLP genomic markers. *Applied and Environmental Microbiology*, 72, 7123–7131.
- Pulawska, J., Willems, A., & Sobiczewski, P. (2006). Rapid and specific identification of four *Agrobacterium* species and biovars using multiplex PCR. *Systematic and Applied Mi*crobiology, 29, 470–479.
- Pulawska, J., Willems, A., & Sobiczewski, P. (2012). Rhizobium skierniewicense sp. nov. isolated from tumors on chrysanthemum and Prunus in Poland. International Journal of Systematic and Evolutionary Microbiology, 62, 895–899.
- Ryder, M. H., Tate, M. E., & Kerr, A. (1985). Virulence properties of strains of *Agrobacterium* on the apical and basal surfaces of carrot root discs. *Plant Physiol*ogy, 77, 215–221.
- Sanger, F., Nicklen, S., & Coulson, A. R. (1977). DNA sequencing with chain terminating inhibitors. *Proceedings of the National Academy of Sciences USA*, 74, 5463–5467.
- Sawada, H., Leki, H., & Matsuda, I. (1995). PCR detection of Ti and Ri plasmids for phytopathogenic Agrobacterium strains. Applied and Environmental Microbiology, 61, 828–831.
- Slater, S. C., Goldman, B. S., Goodner, B., Setubal, J. C., Farrand, S. K., Nester, E. W., Burr, T. J., Banta, L., Dickerman, A. W., Paulsen, I., Otten, L., Suen, G., Welch, R., Almeida, N. F., Armold, F., Burton, Zijin Du, Z., Ewing, A., Godsy, E., Heisel, S., Houmiel, K. L., Jhaveri, J., Lu, J., Miller, N. M., Norton, S., Chen, Q., Phoolcharoen, W., Ohlin, V., Ondrusek, D., Pride, N., Stricklin, S. L., Sun, J., Wheeler, C., Wilson, L., Zhu, H., & Wood, D. W. (2009). Genome sequences of three *Agrobacterium* biovars help elucidate the evolution of multichromosome genomes in bacteria. *Journal of Bacteriology*, *191*, 2501–2511.
- Tighe, S. W., de Lajudie, P., Dipietro, K., Lindström, K., Nick, G., & Jarvis, B. D. W. (2000). Analysis of cellular fatty acids and phenotypic relationships of *Agrobacterium*, *Bradyrhizobium*, *Mesorhizobium*, *Rhizobium* and *Sinorhizobium* species using the Sherlock microbial identification system. *International Journal of Systematics* and Evolutionary Microbiology, 50, 787–801.

- Velázquez, E., Palomo, J. L., Rivas, R., Guerra, H., Peix, A., Trujillo, M. E., García-Benavídez, P., Mateos, P. F., Wabico, H., & Martínez Molina, E. (2010). Analysis of core genes supports the reclassification of strains Agrobacterium radiobacter K84 and Agrobacterium tumefaciens AKE10 into the species Rhizobium rhizogenes. Systematics and Applied Microbiology, 33, 247–251.
- Versalovic, J., Schneider, M., de Bruijn, F. J., & Lupski, J. R. (1994). Genomic fingerprinting of bacteria using repetitive sequence-based polymerase chain reaction. *Methods in Molecular and Cellular Biology*, 5, 25–40.
- Vincent, J. M. (1970). A manual for the practical study of rootnodule bacteria. Oxford: Blackwell.
- Weisburg, W. G., Barns, S. M., Pelletier, D. A., & Lane, D. J. (1991). 16 S ribosomal DNA amplification for phylogenetic study. *Journal of Bacteriology*, 173, 697–703.
- Young, J. M., Kuykendall, L. D., Martínez-Romero, E., Kerr, A., & Sawada, H. (2001). A revision of *Rhizobium* Frank 1889, with an emended description of the genus, and the inclusion of all species of *Agrobacterium* Conn 1942 and *Allorhizobium undicola* de Lajudie *et al.* 1998 as new combinations: *Rhizobium radiobacter*, *R. rhizogenes*, *R. rubi*, *R. undicola* and *R. vitis. International Journal of Systematics and Evolutionary Microbiology*, *51*, 89–103.
- Young, J. M., Park, D., & Weir, B. S. (2004). Diversity of 16S rDNA sequences of *Rhizobium* spp. implications for species determinations. *FEMS Microbiology Letters*, 238, 125–131.
- Young, J. M., Kerr, A., & Sawada, H. (2005). Genus II. Agrobacterium. In G. M. Garrity, D. J. Brenner, N. R. Krieg, & J. R. Staley (Eds.), *Bergey's manual of systematic bacteriology, volume two: The Proteobacteria, parts A - C* (pp. 340–345). Athens: Springer – Verlag.