## **CURRENT REVIEW**

# Legume-Nodulating Betaproteobacteria: Diversity, Host Range, and Future Prospects

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Rhizobia form specialized nodules on the roots of legumes (family Fabaceae) and fix nitrogen in exchange for carbon from the host plant. Although the majority of legumes form symbioses with members of genus Rhizobium and its relatives in class Alphaproteobacteria, some legumes, such as those in the large genus Mimosa, are nodulated predominantly by betaproteobacteria in the genera Burkholderia and Cupriavidus. The principal centers of diversity of these bacteria are in central Brazil and South Africa. Molecular phylogenetic studies have shown that betaproteobacteria have existed as legume symbionts for approximately 50 million years, and that, although they have a common origin, the symbiosis genes in both subclasses have evolved separately since then. Additionally, some species of genus Burkholderia, such as B. phymatum, are highly promiscuous, effectively nodulating several important legumes, including common bean (Phaseolus vulgaris). In contrast to genus Burkholderia, only one species of genus Cupriavidus (C. taiwanensis) has so far been shown to nodulate legumes. The recent availability of the genome sequences of C. taiwanensis, B. phymatum, and B. tuberum has paved the way for a more detailed analysis of the evolutionary and mechanistic differences between nodulating strains of alpha- and betaproteobacteria. Initial analyses of genome sequences have suggested that plant-associated Burkholderia spp. have lower G+C contents than Burkholderia spp. that are opportunistic human pathogens, thus supporting previous suggestions that the plant- and human-associated groups of Burkholderia actually belong in separate genera.

#### History and taxonomy of legume-nodulating bacteria.

The Fabaceae (Leguminosae) family consists of over 19,000 species that are divided among three subfamilies, the Papilionoideae, the Mimosoideae, and the polyphyletic Caesalpinioideae (Lewis et al. 2005). There have been intensive efforts over the last 50 years to determine the full range of legumes that can nodulate, especially in the tropics, where legume diversity is very high (Doyle 2011; Sprent 2001, 2009). It is now known that the vast majority of species (assuming that nodulation is a generic character) in the Papilionoideae (96%) and the Mimosoideae (96%) form nodules, but relatively few of the Caesalpinioideae (22%) that have been examined are nodulated (Sprent 2009; J. I. Sprent unpublished). However, until 20 years ago, with the exception of a few crop species, we knew relatively little about the microorganisms that were involved in nodulating the majority of these legumes, with most being classified as types of genus Rhizobium, a bacterial genus that had been first described in the nineteenth century by Frank (1889). This situation changed after a revolution in bacterial phylogenetics based on sequences of the conserved small subunit ribosomal 16S rRNA gene (Young and Haukka 1996), which led to the division of rhizobia into the currently accepted genera Azorhizobium, Bradyrhizobium, Rhizobium, Mesorhizobium, and Ensifer (syn. Sinorhizobium) (Graham 2008; MacLean et al. 2007; Velázquez et al. 2010; Willems 2006), which so far comprise approximately 100 defined species, with many more remaining to be classified to species level (Velázquez et al. 2010).

Interestingly, the legume-nodulating genera are not monophyletic but are widely dispersed within four families in class Alphaproteobacteria, in which they are intermingled among genera not regarded as nodulating, such as *Afipia*, *Brucella*, and *Zoogloea* (Velázquez et al. 2010; Willems 2006; Young and Haukka 1996). Another consequence of the molecular phylogenetical revolution combined with the intensification of interest in native legumes from many parts of the world is an

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increasing number of reports of nodulation by bacteria belonging to "nonrhizobial" genera. For example, nodulation has so far been confirmed in Neptunia natans by Devosia spp. (Rivas et al. 2002), Lupinus albus by Ochrobactrum lupini (Trujillo et al. 2005), L. albus and Trifolium repens by Phyllobacterium trifolii (Valverde et al. 2005), members of genera Crotalaria and Lotononis by Methylobacterium spp. (Sy et al. 2001; Yates et al. 2007), and members of genus Lotononis by Microvirga spp. (Ardley et al. in press). This list continues to grow annually, and the reader is encouraged to consult the websites maintained by B. S. Weir (Auckland, New Zealand) and the Center for Genomic Sciences (CCG, Cuernavaca, Mexico) for comprehensive and updated lists. However, although these bacteria appear to be quite "exotic" in rhizobial terms, all of them belong to the class Alphaproteobacteria, and so their ability to harbor nodulation genes is not actually as problematic as it might first appear. Indeed, some, such as Phyllobacterium trifolii (Valverde et al. 2005) and Ochrobactrum lupini (Trujillo et al. 2005), are actually more closely related to Rhizobium spp. than are the "traditional" rhizobial genera Bradyrhizobium and Azorhizobium (Willems 2006), and given the plasticity of the nodulation trait, which, in genera Rhizobium and Sinorhi*zobium*, is largely based upon easily transferred plasmid-borne genes (Barnett et al. 2001; Cummings et al. 2009; González et al. 2006; Young et al. 2006) and, in genus Mesorhizobium, by relatively mobile "symbiosis islands" in the chromosome (Sullivan and Ronson 1998), it is, thus, not surprising that the known diversity of legume-nodulating strains has increased greatly beyond genus Rhizobium and its immediate relatives in the family Rhizobiaceae (Velázquez et al. 2010).

#### The discovery of legume-nodulating betaproteobacteria.

Originally it was doubted that legumes could be nodulated by bacteria in other classes of the proteobacteria, such as the betaproteobacteria ( $\beta$ -rhizobia) or gammaproteobacteria. This was because, although N<sub>2</sub> fixation is common in these classes, the bacteria within them had long been considered to be exclusively free-living or loosely associated with plants, or both, and not as nodulating symbionts. Indeed, in his famous manual, Vincent (1970) cautioned against the possibility of isolating nonrhizobial "contaminants" from nodules (particularly if they were fast-growing). Nodules, with their ample supply of nutrients and their enclosed protective environment, are an attractive niche for a wide variety of nonsymbiotic bacteria that have the capability to colonize plants opportunistically (Sprent 2009). Recent published examples of nonnodulating contaminants (which are more correctly termed endophytes) that have been isolated from nodules include Agrobacterium strains (alphaproteobacteria) from various legumes in China (Wang et al. 2006), Labrys neptuniae (alphaproteobacteria) from Neptunia (Chou et al. 2007), Herbaspirillum lusitanum (betaproteobacteria) from Phaseolus vulgaris (Valverde et al. 2003), and various Enterobacter isolates (gammaproteobacteria) from genus Hedysarum (Muresu et al. 2008). In all of these examples, attempts to fulfill Koch's postulates (i.e., to see if they can renodulate their hosts) have failed. Therefore, it is not surprising that reports of symbiotic bacteria in so-called nonrhizobial genera being isolated from nodules were generally greeted with some degree of skepticism. Indeed, when the first claims of legume-nodulating β-rhizobia emerged in 2001 (Chen et al. 2001; Moulin et al. 2001), they were not unconditionally and universally accepted, especially as they were based largely on 16S rRNA sequences (van Berkum et al. 2003). Nevertheless, both studies contained intriguing results suggesting that legume nodulation may not be restricted to alphaproteobacteria. For example, Moulin and associates (2001) presented evidence that two Burkholderia strains, STM678 and

STM815, which had been respectively isolated from nodules on the papilionoid legumes *Aspalathus carnosa* (South Africa) and *Machaerium lunatum* (French Guiana), possessed nodulation genes (*nodA* in this case) and, thus, might be nodulating symbionts. However, neither strain was tested for symbiosis on its original host, although it was demonstrated that they could form ineffective (i.e., non-N<sub>2</sub>-fixing) nodules on the promiscuous host siratro (*Macroptilium atropurpureum*) and that STM678 required common nodulation genes to form these nodules. In parallel to this study, Chen and associates (2001) reported that *Ralstonia taiwanensis* (later renamed *Cupriavidus taiwanensis*), isolated from the nodules of *Mimosa pudica* and *M. diplotricha* in Taiwan, could form nodules on these legumes, but no nodulation data were presented.

Conclusive evidence for nodule formation and effective nitrogen fixation by a betaproteobacterium came from the study by Chen and associates (2003a) in which they used a green fluorescent protein (GFP)-marked strain of C. taiwanensis to perform a detailed microscopic analysis of the nodulation process on its Mimosa hosts. This study showed that the infection of Mimosa spp. was through root hairs, and that the development of classically indeterminate N<sub>2</sub>-fixing nodules was essentially the same as that described for many other legumes (Sprent 2009). Effective nodulation by Burkholderia was finally demonstrated in two studies published in 2005, in which GFPmarked strains of the Brazilian isolates B. nodosa Br3461 and B. mimosarum MAP3-5 (Chen et al. 2005a), and the Taiwanese isolate *B. mimosarum* PAS44<sup>T</sup> (Chen et al. 2005b), were used to confirm nodulation of various Brazilian native and invasive Mimosa spp.

The two Burkholderia strains in the study of Moulin and associates (2001), which were subsequently described as two new species, B. tuberum STM678<sup>T</sup> and B. phymatum STM815<sup>T</sup> (Vandamme et al. 2002), were also later shown to be effective nodulators of legumes, but surprisingly, not of their original hosts. Elliott and associates (2007a) examined the nodA gene sequences of these two strains, and observing that the sequence from B. phymatum STM815 was very similar to that of *Mimosa*-nodulating  $\beta$ -rhizobia, such as *B. mimosarum* and C. taiwanensis, inoculated B. phymatum STM815 onto 30 different Mimosa species, and obtained nodules on 29 of them, including 20 that could fix N2 effectively. Elliott and associates (2007a) thus demonstrated for the first time that B. phymatum is, indeed, a symbiotic bacterium, at least with Mimosa spp. They failed to nodulate a Machaerium species with this strain, so there is no evidence that B. phymatum actually nodulates legumes in this genus. Burkholderia tuberum STM678 has a very different *nodA* sequence from all the other (all *Mimosa-*) nodulating Burkholderia spp. so-far described (Chen et al. 2005a), and thus, it was predicted that it would not nodulate Mimosa spp. This was, indeed, found to be the case by Elliott and associates (2007b), who showed that it could, however, nodulate members of the genus Cyclopia, which is a papilionoid genus native to South Africa.

The aforementioned studies have shown beyond doubt that betaproteobacteria in the genera *Burkholderia* and *Cupriavidus* can both nodulate legumes and fix N<sub>2</sub> within the nodules to the benefit of their plant hosts. These nodulating betaproteobacteria can thus be considered essentially rhizobial in nature and have been termed  $\beta$ -rhizobia to distinguish them from *Rhizobium* and relatives in the Alphaproteobacteria subgroup, which, accordingly, are termed  $\alpha$ -rhizobia. This terminology, which was first used by Moulin and associates (2002), is not universally accepted by all researchers, some of whom prefer 'legume-nodulating bacteria' or 'root-nodulating bacteria' (Lima et al. 2009; Yates et al. 2007), but as they are simple in concept, and now in general use (Graham 2008; Lee and Hirsch 2006; Sprent 2009), we shall use the terms  $\alpha$ - and  $\beta$ -rhizobia for the remainder of this review.

#### Nodulation of *Mimosa* by $\beta$ -proteobacteria.

Nodulation by Burkholderia and Cupriavidus has been further confirmed by several other studies, but it is found most particularly associated with species of Mimosa, and all the newly described  $\beta$ -rhizobial species, with the exception of South African strains of B. tuberum (Elliott et al. 2007b; Kock 2004), either come from or are capable of nodulating Mimosa (Fig. 1; Table 1). For example, C. taiwanensis is frequently isolated from nodules on the pan-tropical weeds M. pudica and M. diplotricha in Taiwan (Chen et al. 2001, 2003b), India (H. S. Gehlot and P. Gyaneshwar unpublished; Verma et al. 2004) and China (Liu et al. 2011), and Burkholderia spp. such as B. mimosarum and B. phymatum are also often isolated from these two Mimosa species, as well as from a third species, M. pigra, which is an extremely aggressive invader in South East Asia (Chen et al. 2005b; Liu et al. 2011) and Australia (Parker et al. 2007), and is rated in the top 10 world's worst weeds (Lowe et al. 2000). With further regard to  $\beta$ -rhizobia from southeast Asia, it is interesting to note that Trinick (1980) isolated many strains of so-called Rhizobium from invasive Mimosa spp. in Papua New Guinea in the early 1960s, and many of these isolates were later identified as Burkholderia and Cupriavidus spp. (Elliott et al. 2007a, 2009).

But what nodulates Mimosa spp. in its native environments, and what is the nature of the relationship between Mimosa spp. and  $\beta$ -rhizobia? Genus *Mimosa* is a large genus of more than 500 species, most of which are native to the New World (Simon et al. 2011). Its principal centers of radiation are in Brazil (about 300 species) and Mexico (about 100 species), where there are many endemic species, particularly in highland regions (Simon and Proença 2000; Simon et al. 2011). There are also several widespread species that are found throughout South and Central America, and these include the three invasive species *M. diplotricha*, *M. pigra*, and *M. pudica*, which are known to be nodulated by  $\beta$ -rhizobia in southeast Asia and Australia. So, are invasive Mimosa spp. also nodulated by  $\beta$ -rhizobia in their native ranges? The answer is very much affirmative, as shown by studies in Brazil and Venezuela (Bontemps et al. 2010; Chen et al. 2005a), Panama (Barrett and Parker 2005), Costa Rica (Barrett and Parker 2006), and Texas (Andam et al. 2007). Indeed, molecular analysis based on 16S rRNA, *nifH*, and *nodA* sequences has suggested that  $\beta$ rhizobial strains isolated from the nodules of plants growing in nonnative environments are very similar, if not identical, to the ones isolated from the native regions, e.g., genetically very similar strains of B. mimosarum nodulate M. pigra in both Taiwan and South America (Chen et al. 2005a and b), and this also appears to be the case in Australia, where M. pigra Burkholderia isolates are also very similar to those from plants in their native range (Parker et al. 2007). This suggests that these invasive species have somehow taken their symbionts with them when they were transported (accidentally or deliberately) from their native environments to nonnative parts of the tropics.

The relationship between  $\beta$ -rhizobia and Mimosa spp. was investigated in more depth in a large-scale study reported by Bontemps and associates (2010) and dos Reis and associates (2010) in the Cerrado and Caatinga biomes of central and northeast Brazil, where the genus Mimosa has evolved and diversified into more than 200 native and endemic species (Fig. 2A and B). Bontemps and associates (2010) isolated rhizobia from 47 Mimosa spp. and all were found to be nodulated by Burkholderia spp., with one widespread species, M. xanthocentra, also nodulated by Rhizobium spp. The phylogenies obtained from the concatenated 16S rDNA and recA sequences of the isolates showed that, although they were related to the known legume-nodulating Burkholderia species, particularly to B. nodosa and B. tuberum, they were in seven deep and distinct clades that were sufficiently distant from the established species that they could probably be new species. Indeed, three of these clades (Fig. 1) are currently in the process of being formally described as new species using a polyphasic approach (W.-M. Chen et al. unpublished).

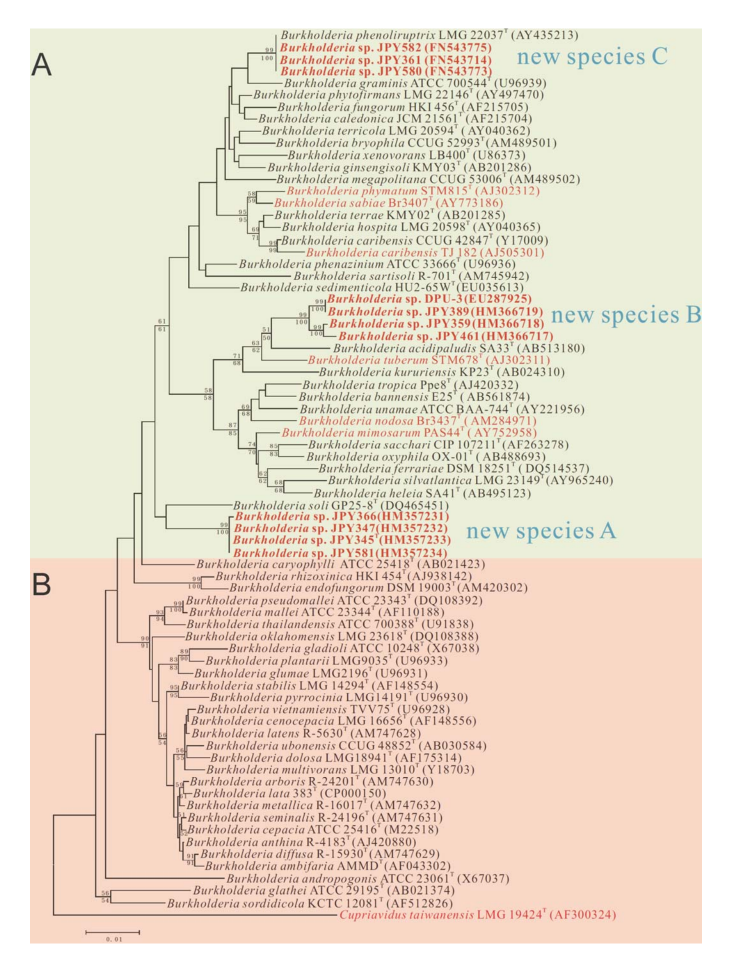
In parallel to the study of Bontemps and associates (2010), dos Reis and associates (2010) performed in situ immunolocalization on the nodules of 67 *Mimosa* species growing in the same biomes, using antibodies specific to *Burkholderia* spp., *C. taiwanensis*, and the *nifH* protein of nitrogenase, and confirmed the overwhelming predominance of *Burkholderia* spp. as symbionts of *Mimosa* spp. in Brazil (Fig. 2C and D). Additionally, dos Reis and associates (2010) showed, using the <sup>15</sup>N natural abundance technique, that the *Burkholderia* symbionts were capable of providing at least 60% of the total plant N in *Mimosa* spp. endemic to Brazil.

# Conditions favoring nodulation of *Mimosa* spp. by $\beta$ -rhizobia.

Given how frequently it is isolated from *Mimosa* spp. (both native and nonnative) in other regions, an unexpected finding by both studies was the absence of C. taiwanensis in Mimosa nodules from central Brazil (Bontemps et al. 2010; dos Reis et al. 2010). This curious observation might be partly explained by the results of inoculation studies that have shown that most Brazilian Mimosa spp. appear incapable of being nodulated effectively (if at all) by C. taiwanensis, whereas many of the same species are nodulated effectively by the promiscuous Mimosa-nodulating Burkholderia strain B. phymatum STM815 (dos Reis et al. 2010) (Fig. 2E). It would, thus, appear that most Mimosa spp. are inherently incapable of being nodulated by C. taiwanensis and that this bacterium is largely confined to nodulating either a few, widespread and invasive Mimosa spp. at the edges of the native range of the genus (Andam et al. 2007), invasive Mimosa spp. outside their native range (Chen et al. 2001, 2003a and b, 2005b; Elliott et al. 2007a, 2009), or both.

Although most *Mimosa* spp. are nodulated by *Burkholderia* spp. and some by *C. taiwanensis*, some of them can also form effective symbioses with  $\alpha$ -rhizobia (Chen et al. 2003b; Elliott et al. 2009; Wang et al. 1999), which suggests the existence of genetic and physiological factors that determine their apparent preference for nodulation by  $\beta$ -rhizobia. To determine the conditions that favor nodulation by  $\beta$ -rhizobia, Elliott and associ-

**Fig. 1.** 16S rRNA gene phylogeny of all known *Burkholderia* spp. (betaproteobacteria species marked in red), including three proposed new *Burkholderia* species (in bold red) consisting of strains isolated from *Mimosa* nodules collected in central and northeast Brazil by Bontemps and associates (2010). Neighbor-joining bootstrap percentages are given above nodes and those for maximum likelihood below the nodes. The genus can be divided into two groups, group A (in the green box), which contains most of the plant-associated species and all of the currently known legume-nodulating species, and group B (in the pink box), which contains all the phytopathogenic species as well as the species that may cause disease in humans and animals. P. Estrada-de los Santos, P. Vinuesa, L. Martinez-Aguilar, A.M., Hirsch, and J. Caballero-Mellado (*unpublished*) propose that group A should be moved to the new genus *Caballeronia*.



ates (2009) set up paired competition experiments between defined strains of Burkholderia, C. taiwanensis, and Rhizobium spp. and showed that the Burkholderia strains (B. mimosarum PAS44, B. phymatum STM815) outcompeted both C. taiwanensis LMG19424 and all the Mimosa-nodulating  $\alpha$ -rhizobial strains (R. etli TJ167, R. tropici NGR181, and R. tropici UPRM8021) for nodulation of three invasive Mimosa species (M. diplotricha, M. pigra, M. pudica) in all the conditions tested. However, the competitive domination of B. mimosarum over C. taiwanensis was reduced in the presence of nitrate for the three plant hosts, with the largest significant effect on M. pudica, in which C. taiwanensis formed 57% of the nodules in the presence of 0.5 mM KNO<sub>3</sub>. Further study of Brazilian legumes suggests that physical environment, rather than the host species, largely determines the distribution of Burkholderia species (Bontemps et al 2010; dos Reis et al. 2010). These data indicate that, in acidic soils containing very low levels of inorganic nitrogen, Burkholderia spp. will most likely be the preferred symbionts of endemic Mimosa spp. (and possibly other legumes that have evolved to live in low-N, acidic soils [Garau et al. 2009]).

### Host range of $\beta$ -rhizobia.

Although published data at the time of writing this review indicate that nodulation by C. taiwanensis is confined to the genus Mimosa, Burkholderia spp. appear to be able to nodulate a much wider range of legumes. This was indicated by Barrett and Parker (2005), who, in addition to isolating several Burkholderia strains from Mimosa spp. in Panama, also isolated strains from the mimosoid legume Abarema macademia (tribe Ingaeae). However, although these strains were capable of nodulating *Mimosa* spp., they were not tested on their original host, owing to lack of available seeds. More direct evidence comes from host-range studies with B. phymatum STM815, a strain that nodulates more than 40 Mimosa species from various parts of the world (dos Reis et al. 2010; Elliott et al. 2007a). These studies have shown this strain to be highly promiscuous outside the genus Mimosa, nodulating several legumes in the subfamily Mimosoideae (E. Gross, C. Bontemps, and E. K. James unpublished). Burkholderia phymatum STM815 has a particular affinity for species in the tribe Mimoseae, such as those in the "sister" group to Mimosa, Piptadenia (sensu stricto), and both of the other two clades in the polyphyletic genus Piptadenia (Jobson and Luckow 2007) (Fig. 2F and G). It also nodulates some species of genera *Anadenanthera*, *Leucaena*, and *Prosopis* (all in tribe Mimoseae) as well as several *Acacia* spp., but only those in the subgenus *Acacia*, which is phylogenetically close to the Mimoseae (Fig. 2H). A clue to the ability of *B. phymatum* STM815 to nodulate legumes in this tribe lies in its nodulation genes, as *Burkholderia* strains isolated from *Piptadenia* spp. in Brazil have very similar *nodC* sequences to *B. phymatum* and they are capable of nodulating invasive *Mimosa* spp. (e.g., *M. pudica*) (E. Gross, C. Bontemps, and E. K. James *unpublished*). It, thus, appears likely that there has been some coevolution between the plants in tribe Mimoseae and their *Burkholderia* symbionts, at least in a Brazilian context, and that this coevolution is not confined to the genus *Mimosa* (Bontemps et al. 2010).

Recent studies have extended the nodulation of legumes by Burkholderia spp. to the subfamily Papilionoideae, but there are distinct preferences in terms of the host range of Papilionoideae-nodulating Burkholderia spp., and these are based upon the nodulation genes. For example, the B. tuberum STM678 nodA gene sequence is distant from those of the Mimosa-nodulating Burkholderia spp. (Chen et al. 2003b, 2005a), and it does not nodulate Mimosa spp. or other members of the Mimosoideae (Elliott et al. 2007b). Interestingly, as well as Cyclopia (Elliott et al. 2007b; Kock, 2004), B. tuberum nodulates other South African genera in the same tribe, the Podalyrieae, such as Podalyria and Virgilia (Fig. 2I; E. K. James, J. I. Sprent, and W.-M. Chen unpublished). However, B. tuberum has not so far been demonstrated to nodulate species in the genus Aspalathus (tribe Crotalarieae) (Elliott et al. 2007b), which, although not closely related to the Podalyrieae, grow in the same acidic soils of the South African Cape Fynbos biome, and thus will be exposed to the same nodulating microflora, including burkholderias. Burkholderia tuberum STM678 also effectively nodulates siratro (Elliott et al. 2007b) and Phaseolus vulgaris (A. Angus and A. M. Hirsch unpublished), which are in the tribe Phaseoleae. Garau and associates (2009), in their study of Rhynchosia ferufolia, also in the tribe Phaseoleae but native to the acidic soils of the South African Cape, have shown that this plant is nodulated by Burkholderia strains with nodA genes similar to that of B. tuberum STM678. Indeed, the South African Cape Fynbos biome would appear to be a major reservoir of Papilionoideae-nodulating burkhold-

Table 1. Species of  $\beta$ -proteobacteria that have been confirmed to nodulate and fix N<sub>2</sub> in symbiosis with legumes

Species	Original host	Host-range	Geographical location	References
Burkholderia caribensis	M. pudica, M. diplotricha	M. pudica, M. diplotricha	Taiwan	Chen et al. 2003a; G. N. Elliott (unpublished)
B. sabiae	M. caesalpiniifolia	M. caesalpiniifolia, M. pudica	Brazil	Chen et al. 2005a, 2008
B. mimosarum	M. pigra, M.pudica	M pigra, M pudica	Taiwan, Brazil, Venezuela, China	Chen et al. 2005a and b, 2006; Liu et al. 2010
B. nodosa	M. bimucronata, M. scabrella	M. bimucronata, M. scabrella, M pudica	Brazil	Chen et al. 2005a, 2007
B. phymatum	Not known <sup>a</sup>	<i>Mimosa</i> spp., and other members of the tribe Mimoseae, <i>Phaseolus vulgaris</i> .	French Guiana, Papua New Guinea, Morocco, China	Trinick 1980; Moulin et al. 2001; Elliott et al 2007a; Talbi et al. 2010; Liu et al. 2011; E. Gross, C. Bontemps, and E. K. James, ( <i>unpublished</i> )
B. tuberum	Aspalathus carnosa (Crotalarieae, Papilionoideae) but not yet shown to nodulate it; Cyclopia (Podalyrieae, Papilionoideae).	Cyclopia, and other members of the tribe Podalyrieae, Macroptilium atropurpureum, Phaseolus vulgaris	South Africa	Moulin et al. 2001; Elliott et al. 2007b; E. K. James ( <i>unpublished</i> )
C. taiwanensis	M. pudica, M. diplotricha, M. pigra	Mimosa spp.	Taiwan, India, Panama, Costa Rica	Chen et al. 2003a and b, 2005b; Verma et al. 2004; Barrett and Parker 2005, 2006; Elliott et al. 2007a, 2009

<sup>a</sup> Reported to be isolated from the nodules of *Machaerium lunatum* but not confirmed to nodulate this legume (Moulin et al. 2001; Elliott et al. 2007a).

erias, as another native legume, *Lebeckia* (Crotalarieae), is also nodulated by *Burkholderia* spp. (Ardley et al. in press).

It would thus appear that there are at least two distinct groups of legume-nodulating burkholderias defined and separated neither by their core genomes nor by their nif genes (which are closely related [Bontemps et al. 2010]) but by geography and by their nodulation genes. These are the mimosoid-nodulating burkholderias centered on South America (e.g., B. mimosarum, B. nodosa, B. phymatum, B. sabiae) and the papilionoid-nodulating burkholderias centered on South Africa (e.g., B. tuberum STM678). Interestingly, however, although the papilionoid-nodulating B. tuberum-type symbionts appear incapable of nodulating mimosoids (Elliott et al. 2007b; Garau et al. 2009), the opposite is not the case. For example, Martinez-Romero (2009) has described partially effective nodulation of common bean (Phaseolus vulgaris) by the highly promiscuous strain, B. phymatum STM815. In addition, other B. phymatum strains have recently been isolated from nodules on common bean in Morocco by Talbi and associates (2010). These strains have a nodC gene almost identical to that of B. phymatum strains STM815 and NGR195A and, besides nodulating Mimosa spp., they are also apparently capable of fully effective nodulation of common bean.

Data emerging from Australia suggest that it may house a third center for  $\beta$ -rhizobia, because two *Burkholderia* strains (WSM2230 and WSM2232) have been demonstrated to nodulate and fix nitrogen with the Australian native papilionoid legumes *Kennedia coccinea* (Phaseoleae) and *Gastrolobium capitatum* (Mirbelieae). Strains WSM2230 and WSM2232 were isolated from *K. coccinea* and *G. capitatum* trap plants grown in soil taken from Karijini National Park in western Australia, which has a semiarid climate and nutrient-poor soils. However, these soils differ from the South African Fynbos and Brazilian Cerrado soils in that they are alkaline (pH 8.0). Initial phylogenetic analyses suggest that the *Burkholderia* strains isolated from the *Burkholderia* strains that are microsymbionts of the invasive *M. pigra* in Australia (Ardley et al. in press).

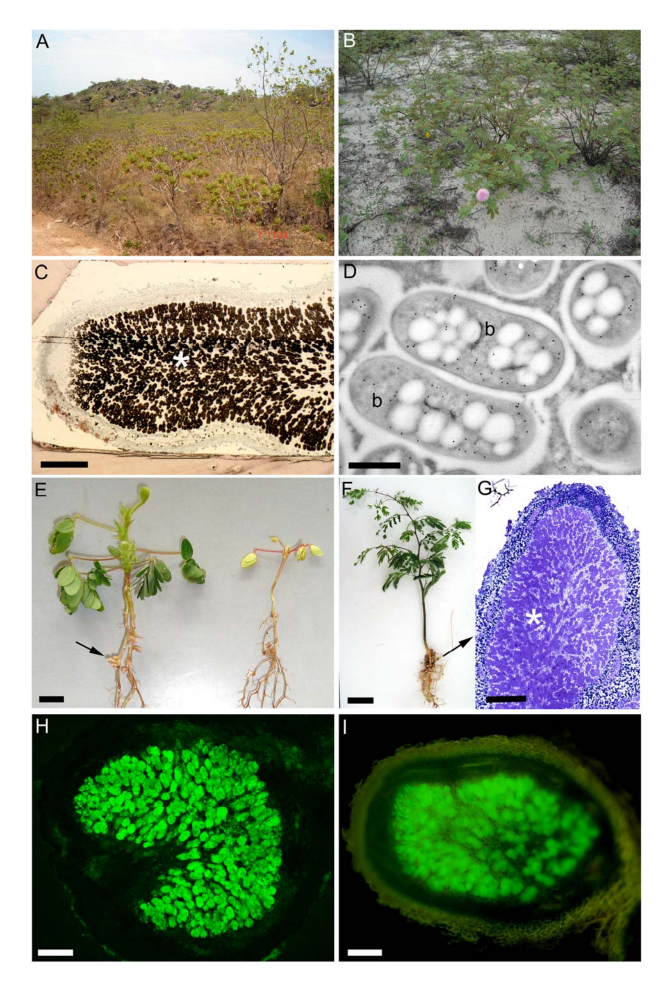
#### Evolution of symbiosis in β-proteobacteria.

The recent availability of whole-genome sequences of both  $\alpha$ - and  $\beta$ -rhizobia can provide a better understanding of the mechanisms and evolution of nodulation and nitrogen fixation by members of genera Burkholderia and Cupriavidus. Analysis of the DNA sequences of genes essential for nitrogen fixation (nifH) has shown that nif genes of the nodulating strains are closely related to nifH of nonnodulating Burkholderia strains and show some homology to nifH genes from genera Bradyrhizobium and Azorhizobium but are only distantly related to other alphaproteobacterial rhizobial strains (Bontemps et al. 2010; Chen et al. 2003b, 2005a and b). Indeed, genera Bradyrhizobium and Azorhizobium are postulated to have acquired nif genes from betaproteobacteria (Young 2005). It has been suggested that symbiotic Burkholderia strains may have contained nif genes before acquiring the essential nodulation genes (Bontemps et al. 2010). Unlike most  $\alpha$ -rhizobial strains, which do not exhibit free-living diazotrophy, both B. phymatum STM815 and B. tuberum STM678 have significant nitrogenase (acetylene reduction) activity in free-living conditions (Elliott et al. 2007a; Wong-Villarreal and Caballero-Mellado 2010), but this activity is dependent upon the presence of some fixed nitrogen in the growth medium (e.g., yeast extract) and is significantly lower than acetylene reduction by nonsymbiotic Burkholderia strains (Elliott et al. 2007a). In contrast to B. phymatum and B. tuberum, C. taiwanensis strains did not reduce acetylene in free-living conditions (Elliott et al. 2007a), although Verma and associates (2004) have induced expression

of the *nifH* protein of nitrogenase by *C. taiwanensis* growing in a semisolid N-free medium that is usually used to detect diazotrophic *Azospirillum* strains. These results indicate physiological differences for the regulation of nitrogen fixation between nonsymbiotic and symbiotic betaproteobacteria, as well as between  $\beta$ - and  $\alpha$ -rhizobia, but the mechanisms underlying these differences remain to be determined.

In addition to nif genes, all legume-nodulating bacteria, with the exception of some photosynthetic stem-nodulating bradyrhizobia (Giraud et al. 2007), contain nodulation (nod) genes involved in the synthesis and transport of Nod factors. These are signaling molecules that are recognized by the plant hosts, and they confer various degrees of specificity in terms of which host the bacterium can nodulate (Sprent 2009). The nod genes are sufficiently conserved to use for determining the evolution of rhizobial-legume interactions and nodulation. Recent analysis using nodC sequences of 143 Burkholderia strains isolated from 47 native Mimosa species in central Brazil showed a monophyletic origin, suggesting a single acquisition of these genes (Bontemps et al. 2010). The almost complete congruence of the *nodC* sequence tree with those of other phylogenetic markers (16S rRNA, recA, and nifH) suggests that they have evolved together with little horizontal gene transfer. Additionally, the branch depth of nodC groups in genus Burkholderia was found to be similar to that observed in  $\alpha$ -rhizobial groups. These data indicate that nodulation in Burkholderia spp. is an ancient and stable ecological trait, with a possible age of at least 50 million years (Angus and Hirsch 2010; Bontemps et al. 2010). Legumes evolved about 60 million years ago, and nodulation in some groups shortly after that (Doyle 2011; Sprent 2009). The tribes and genera that are nodulated by Burkholderia spp. are more recent, for example, Mimosa first appeared 28 million years ago (Simon et al. 2011).

So far, with the exception of the Rhynchosia-nodulating strains from the South African Cape Fynbos environment (Garau et al. 2009), no  $\beta$ -rhizobia have been reported to have nod gene sequences closely related to those of B. tuberum STM678, and this most likely explains its very different host range compared with the Mimosa-nodulating strains (discussed above). The possible origins of the nod genes in B. tuberum STM678 are hinted at by Kock (2004), who described more than 120 authenticated nodulating rhizobial strains isolated from 14 Cyclopia spp. The vast majority of these, according to their 16S rRNA sequences, were B. tuberum and related Burkholderia strains. Just as Bontemps and associates (2010) found with Brazilian Mimosa-nodulating Burkholderia symbionts, a very strong relationship is apparent between the genus Cyclopia and B. tuberum STM678-like strains, and a long history of coevolution between them is suggested. Interestingly, a few of the strains nodulating Cyclopia spp. were Bradyrhizobium spp. or R. tropici, and these had very similar nodA sequences to the B. tuberum strains, so it is highly likely that there had been horizontal gene transfer between the  $\alpha$ - and  $\beta$ rhizobia (Kock 2004). Although the direction of this gene transfer is not clear, it seems most likely that Burkholderia was the donor, given its preponderance as a Cyclopia symbiont. Interestingly, this situation is very different from the Mimosa symbiosis, in which the nodulation genes of  $\alpha$ -rhizobia are not closely related to those of  $\beta$ -rhizobia and the host specificity must have evolved independently (Bontemps et al. 2010). Certainly the fascinating data of Kock (2004) are worthy of much further analysis, because they hint that the Cape region of South Africa and the acidic soils of the Fynbos biome, in particular (discussed above), might be another center of  $\beta$ -rhizobial evolution and diversity to rival that of the Cerrado and Caatinga biomes of Brazil (Bontemps et al. 2010).



# Genomic analysis of nodulation and nitrogen fixation by $\beta$ -rhizobia.

The complete genome sequence of C. taiwanensis has been published (Amadou et al. 2008) and consists of a chromosome of 3.4 Mb, a chromid of 2.5 Mb (Harrison et al. 2010), and a large symbiosis plasmid of 0.56 Mb (pRalta). The genes required for nodulation and nitrogen fixation are present on a symbiosis island of only 38 kb, which is the most compact known so far for any rhizobial species. Cupriavidus taiwanensis has one copy of nodD and makes a Nod factor that is sulfated at its reducing end and acetylated at its nonreducing end. Interestingly, cytochrome cbb3, which has a high affinity for oxygen and is critical for nitrogen fixation in the nodules, is localized in the chromosome, in contrast to its presence in the symbiosis plasmids in  $\alpha$ -rhizobia. In addition, the genome lacks homologs of the  $\alpha$ -rhizobial FixLJ two-component regulatory system that is required for nitrogen fixation in the nodules. This indicates that, in C. taiwanensis, the expression of genes required for efficient nitrogen fixation (nif and fix) might be regulated in a different way from the  $\alpha$ -rhizobia, and this needs to be explored further.

Like that of many Burkholderia species (Harrison et al. 2010), the genome of *B. phymatum* STM815 (National Center for Biotechnology Information Bioproject website) comprises a chromosome (3.5 Mb), two chromids ('chromosome 2', 2.7 Mb and pBPHY01, 1.9 Mb), and a plasmid (0.59 Mb). The genome is in the process of being published together with its symbiotic transcriptome (L. Moulin unpublished). The organization of the nod and nif gene operons in the pSym of C. taiwanensis LMG19424 and B. phymatum STM815 are very similar, indicating that these β-rhizobia have acquired the symbiotic functions either from each other or from a common donor. It is interesting that, even though both C. taiwanensis and B. phymatum contain the same nod genes and, thus, probably make similar Nod factors, B. phymatum forms an effective symbiosis with many mimosoid legumes, whereas C. taiwanensis forms mostly ineffective symbioses with them (Fig. 2E; discussed above; Elliott et al. 2007a). This suggests that the broad host range of B. phymatum might involve genes other than those on the symbiosis plasmid and that other components besides Nod factors are needed for effective nodulation. It is also possible that C. taiwanensis contains protein secretion systems that are recognized as pathogenic by the legume hosts. Deletion of the type III secretion system in a Ralstonia solanacearum strain containing the C. taiwanensis symbiosis plasmid (pRalta) led to nodule formation on M. pudica (Marchetti et al. 2010).

The genome of *B. tuberum* STM678 has been sequenced and is still in the process of being annotated (A. M. Hirsch *unpublished*). With regard to nitrogenase genes, *B. tuberum* STM678 exhibits similar gene organization to that of other nitrogen-fixing *Burkholderia* species, including *B. unamae* MTI641, *B. xenovorans* LB400 (Chain et al. 2006), and *B. vietnamiensis* G4.

However, unlike these but similar to *B. phymatum* STM815, *nifA* and *nifB* are not next to one other but, rather, are separated by a large number of genes. Also, in B. phymatum STM815, the nifEN genes are located at some distance from nifHDK, whereas, for B. tuberum STM678, the nifHDK and nifEN genes show a similar organization to the comparable genes in other nitrogen-fixing strains (A. M. Hirsch unpublished). Although the nif genes are similar to other nodulating Burkholderia spp., such as B. phymatum STM815, the nod genes of B. tuberum STM678 are very different from those of the other  $\beta$ -rhizobia. This has already been shown for *nodA* and *nodC* (Bontemps et al. 2010; Chen et al. 2003b, 2005a and b; Kock 2004), and the genome has provided deeper insight into these differences. For example, a second *nodC* gene exists in the *B. tuberum* genome, but it is a partial copy, as previously reported by Moulin and associates (2001). In terms of Nod factors, it is likely that those purified by Boone and associates (1999) from one of three "Bradyrhizobium aspalati" strains were actually from the Cyclopia-nodulating strain B. tuberum STM678. Elliott and associates (2007b) provides a detailed history of the relationship between Aspalathus and Cyclopia-nodulating rhizobia. Analysis of the nod genes in the genome of B. tuberum STM678 shows that these match the Nod factor structure reported by Boone and associates (1999), who identified two major Nod factors, each with a backbone consisting of three to five glucosamine residues, which is typical of the Nod factors of  $\alpha$ -rhizobial species. However, the *B. as*palati Nod factors are substituted on the nonreducing end sugar with an N-methyl and two carbamoyl groups but have no substitutions at the reducing end. Nod factors of  $\alpha$ -rhizobia typically have substitutions on the reducing end, making B. aspalati Nod factors significantly different.

Numerous  $\beta$ -rhizobial genome-sequencing projects are currently under way, as listed in Table 2. Future analyses of these genomes in comparison with those of  $\alpha$ -rhizobia should shed light on the adaptations that  $\beta$ -rhizobia possess that predisposes them to form symbioses with particular legumes in certain environments.

# Relationship between plant-associated *Burkholderia* and human pathogens.

Apart from the nodulating symbiotic strains, many species of genus *Burkholderia* have been described that associate with plants such as maize, tomato, rice, sorghum, sugarcane, pine-apple, and coffee (Caballero-Mellado et al. 2004; Castro-González et al. 2011; Estrada de los Santos et al. 2001; Gillis et al. 1995; Martinez-Aguilar et al. 2008; Perin et al. 2006; Reis et al. 2004), and several of these fix nitrogen and increase plant growth. On the other hand, a number of other *Burkholderia* species are phytopathogens, such as *B. caryophylli*, *B. gladioli*, and *B. glumae*. These are related to *B. cepacia*, which was originally isolated from rotting onions, as the name '*cepacia*' implies (Burkholder 1950). *Burkholderia cepacia* and some close relatives, the *B. cepacia* complex

**Fig. 2.** Some legumes that are commonly nodulated by betaproteobacteria ( $\beta$ -rhizobia). **A**, *Mimosa setosissima*, a species endemic to the Pirenopolis municipality in the Cerrado biome in central Brazil. The Cerrado is a tropical savannah characterized by highly seasonal rainfall (500 to 1,000 mm per annum and very acidic soils (pH 3 to 5) with low fertility and is regularly subjected to burning during the dry season. **B**, *Mimosa cordistipula*, a Caatinga endemic, growing in sand close to the Rio São Francisco in northeast Brazil. The Caatinga is a xeric shrub land with a climate classed as semiarid (<500 mm per annum); it has Fe-rich soils, many of which are as acidic as those in the Cerrado. **C**, Light micrograph of a *Mimosa velloziana* nodule from the Brazilian Cerrado that has been immunogold-labeled with an antibody against *Burkholderia phymatum* (the N<sub>2</sub>-fixing *Burkholderia*-infected zone in the nodule section is marked by an asterisk). **D**, Transmission electron micrograph of bacteroids (b) from a *Mimosa caesalpinijfolia* nodule collected in central Brazil that has been immunogold-labeled with an antibody against *Burkholderia phymatum* (the N<sub>2</sub>-fixing *Burkholderia*-infected zone in the nodule section is marked by an asterisk). **D**, Transmission electron micrograph of bacteroids (b) from a *Mimosa caesalpinijfolia* nodule collected in central Brazil that has been immunogold-labeled with an antibody against nitrogenase. **E**, *Mimosa ursina* nodulated by *B*. *phymatum* STM815 (left, nodules indicated by an arrow) and *C. taiwanensis* LMG19424 (right). **F**, *Piptadenia viridiflora* (Mimosoideae, tribe Mimoseae) nodulated by *B*. *phymatum* STM815. **G**, The inset shows a section through the nodules marked with an arrow in F. An asterisk marks the N<sub>2</sub>-fixing infected zone in the nodule section. **H**, Section through a nodule on *Acacia pennatula* (Mimosoideae, tribe Mimoseae) infected with *B*. *tuberum* STM815GFP. **I**, Section through a nodule on *Podalyria canescens* (Papilionoideae, tribe Podal

Table 2. Beta-rhizobia genome sequencing projects (as of June 22, 2011).

Taxonomy	Strain name	Plant host	Country of origin	Sequencing center <sup>a</sup>	GOLD identity <sup>b</sup>	
Finished						
Burkholderia phymatum STM815		Machaerium lunatum <sup>c</sup>	French Guiana	JGI	Gc00775	
Cupriavidus taiwanensis LMG19424		Mimosa pudica Taiwan		Genoscope	Gc00754	
Current		•		-		
Burkholderia tuberum	STM678	Aspalathus carnosa <sup>c</sup>	South Africa	Washington University	Gi07505	
Burkholderia tuberum	STM3649	Mimosa pudica	French Guiana	Genoscope, France	Gi09600	
Burkholderia mimosarum	LMG23256T	Mimosa pigra	Taiwan	GEBA-RNB, JGI	Gi08823	
	STM3621	Mimosa pudica	French Guiana	GEBA-RNB, JGI,	Gi08839	
Burkholderia sp.	WSM3937	Rhynchosia ferulifolia	South Africa	GEBA-RNB, JGI	Gi08878	
-	JPY580	Mimosa cordistipula	Brazil, Bahia	GEBA-RNB, JGI	Gi08877	
	CCGE1002	Mimosa occidentalis	Mexico, Tepic	DOE-JGI	Gc01354	
	JPY366	Mimosa misera	Brazil, Bahia	GEBA-RNB, JGI	Gi08876	
	JPY347	Mimosa cordistipula	Brazil, Bahia	GEBA-RNB, JGI	Gi08875	
	JPY251	Mimosa velloziana	Brazil	GEBA-RNB, JGI	Gi08874	
	WSM4176	Lebeckia ambigua	South Africa,	GEBA-RNB, JGI	Gi08873	
	WSM3556	Lebeckia ambigua	South Africa	GEBA-RNB, JGI	Gi08872	
	Mcas7.1	Mimosa casta	Panama	GEBA-RNB, JGI	Gi08846	
	MP20	Mimosa pudica	India, Bokaro	GEBA-RNB, JGI	Gi08833	
	WSM2232	Nemcia capitata	Australia	GEBA-RNB, JGI	Gi08832	
	WSM2230	Kennedia coccinea	Australia	GEBA-RNB, JGI	Gi08831	
	4.13	Parapiptadenia rigida	Uruguay	GEBA-RNB, JGI	Gi08829	
Cupriavidus taiwanensis	STM6018	Mimosa pudica	French Guiana	GEBA-RNB, JGI	Gi08840	
-	STM6070	Mimosa pudica	N.Caledonia	GEBA-RNB, JGI	Gi08841	
Cupriavidus sp.	Amp6	Mimosa asperata	U.S.A. (Texas)	GEBA-RNB, JGI	Gi08845	
- •	5v12	Parapiptadenia rigida	Uruguay	GEBA-RNB, JGI	Gi08830	

<sup>a</sup> JGI = United States Department of Energy (DOE) Joint Genome Institute; GEBA-RNB:Genome encyclopedia of bacteria and archaea on root nodulating bacteria, a specific program of JGI.

<sup>b</sup> GOLD = Genome On-Line database.

<sup>c</sup> Strain STM815 has a broad host range on many *Mimosa* species, but was never proved to nodulate its original host *Machaerium lunatum*. Strain STM678 nodulates *Cyclopia* species but was also not proven to nodulate *Aspalathus* spp. Only the *C. taiwanensis* LMG19424 genome sequence has been published so far (Amadou et al. 2008).

Table 3. Characteristics of genomes from several sequenced and annotated Burkholderia species of varying life styles<sup>a</sup>

		GC% Size <sup>b</sup>	% Genes for					
	GC%		Metabolism					
Species and strain			Xenobiotic	Secondary	Lipid	Signal transduction	Lifestyle	nifH <sup>c</sup>
B. ambifaria MC40-6	66	7.64	11.66	9.4	10.78	0.22	Biocontrol, human pathogen	No
B. cenocepacia HI2424	67	7.7	12.50	10.22	10.54	0.27	Human pathogen	No
B. glumae BGR1	68	7.28	10.68	9.35	9.6	0.19	Plant pathogen	No
B. graminis C4D1M	63	7.48	13.98	9.05	10.75	0.37	Biocontrol	No
B mallei ATCC 23344	68	5.84	10.68	9.35	10.08	0.27	Human pathogen	No
B. multivorans ATCC 17616	67	7.01	12.28	9.74	10.02	0.18	Human pathogen	No
B. phymatum STM815	62	8.68	13.72	9.95	10.45	0.30	Mutualist	Yes
B. phytofirmans PsJN	62	8.21	13.67	9.74	10.83	0.20	Biocontrol	No
B. pseudomallei 1106a	68	7.09	10.53	9.45	10.19	0.17	Human pathogen	No
B. tuberum STM678	63	8.10	14.73	10.49	12.04	0.31	Mutualist	Yes
B. unamae MT1-641	65	9.64	16.52	11.34	14.79	0.17	Mutualist	Yes
B. vietnamiensis G4	66	8.39	11.28	9.44	9.88	0.22	Mutualist, human pathogen	Yes
B. xenovorans LB400	63	9.73	16.97	10.94	13.35	0.16	Mutualist	Yes

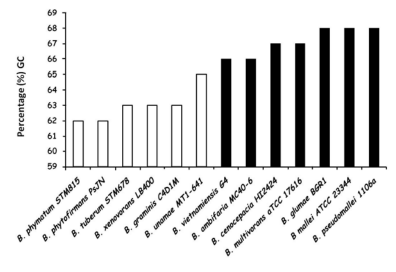
<sup>a</sup> All the genomes listed are finished, except for *B. graminis* C4D1M, *B. tuberum* STM678, and *B. unamae* MTI-641, which are still in draft form. The information above comes from IMG/ER (Markowitz et al. 2009) and A. M. Hirsch *unpublished*.

<sup>b</sup> Genome size in Mbp.

<sup>c</sup> Evidence for *nifH* in the sequenced genome.

(Bcc), are found in hospital environments and are opportunistic human pathogens (Mahenthiralingam et al. 2008). Interestingly, strains belonging to the Bcc are also frequently isolated from legume nodules (Rasolomampianina et al. 2005; Vandamme et al. 2002), although none have yet been confirmed as effective symbionts (Vandamme et al. 2007). Nevertheless, their presence within nodules well illustrates the diversity of niches that can be occupied by Bcc strains that are outside the clinical and phytopathogenic environments.

Because of the potential use of these plant growth-promoting, biocontrol-affecting,  $N_2$ -fixing, and nodulating *Burkholderia* strains as agricultural inoculants (Compant et al. 2005) as well as the perception (based on their generic name) that they may cause disease, various studies have compared the phylogenetic relationships between beneficial and nodulating plant–associated *Burkholderia* strains and those in the Bcc. Two subclades were evident from early studies of 16S rRNA sequences (Caballero-Mellado et al. 2004, Reis et al. 2004), and deeper analysis resulted in a distinct separation between *Burkholderia* spp. that are plant-associated and generally beneficial and those species that are either plant pathogens or opportunistic mammalian pathogens (Perin et al. 2006). This phylogenetic separation into two separate lineages was also observed when 16S rRNA and *recA* gene sequences of many nodulating strains of *Burkholderia* were analyzed (Bontemps et al. 2010). The separation based on 16S rRNA sequences is depicted in Figure 1, showing the same dichotomy between the plant- and animal-associated sublineages of *Burkholderia* spp.



Burkholderia species

**Fig. 3.** The mean G+C content of the genomes of several *Burkholderia* species is displayed in graph form. The species with the lowest G+C content are members of the plant-associated clade (white bars) and are used for biocontrol, bioremediation, or to provide fixed nitrogen to plants. The species with the highest G+C content are phytopathogens or opportunistic mammalian pathogens (black bars).

as do phylogenetic schemes that are based upon the sequences of other genes, such as *acdS*, which encodes for 1-aminocyclopropane-1-carboxylate (ACC) deaminase (Onofre-Lemus et al. 2009) and on the analysis of whole genomes (Vanlaere et al. 2009). A recent multilocus sequence analysis of five housekeeping genes from 67 type and reference *Burkholderia* strains confirmed the two distinct evolutionary lineages within the genus (P. Estrada-de los Santos, P. Vinuesa, L. Martinez-Aguilar, A.M., Hirsch, and J. Caballero-Mellado *unpublished*). These authors plan to propose that the plant- and animal-associated strains be split into two distinct genera, with *Caballeronia* being suggested as the name for a new genus containing the plant-associated but nonpathogenic strains, in honor of the late J. Caballero-Mellado, a Mexican microbiologist who pioneered many of the studies on the plant-associated strains.

Analysis of 22 Burkholderia species with completely or almost completely sequenced genomes available through The Integrated Microbial Genome IMG/ER system website (Markowitz et al. 2009) shows that the majority of plant-associated Burkholderia species have a lower G+C content than those in the Bcc (Table 3; Fig. 3). Expanding the analysis to 34 strains recapitulated the difference in G+C content in the two sublineages (A. M. Hirsch unpublished). Furthermore, the plant-associated species lack some genes that are important for the virulence of animal pathogens in the Bcc. Perin and associates (2006) could not detect the Burkholderia virulence genes cblA and esmR in nitrogen-fixing Burkholderia spp. associated with maize and sugarcane. By analyzing the entire genome sequence of four different plant-associated Burkholderia strains, including B. tuberum STM678, we have found that various secretion systems that are indicators of pathogenesis are missing (A. M. Hirsch unpublished), although additional studies are required to establish the effect of these Burkholderia strains on animal models.

#### Concluding remarks and future work.

 $\beta$ -Rhizobia are highly effective symbionts of legumes and have coevolved with their hosts for up to 50 million years, with their principal centers of diversity in South America and South Africa and, possibly, another center of diversity in Australia. It is only a decade since their existence was first recognized, and many more will undoubtedly be discovered in the coming years. There is much to learn about the genetic, taxonomic, and geographical factors underlying the ability (or preference) of particular legumes to nodulate with  $\beta$ -rhizobia and, after such a long period of separate evolution of the symbiosis genes in very different genomic backgrounds, there could be substantial differences from the way that  $\alpha$ -rhizobia interact with their hosts. Although  $\beta$ -rhizobia are particularly associated with genus Mimosa and some related genera, they also nodulate several agriculturally important papilionoid legumes, including common bean (Phaseolus vulgaris) and honeybush tea (*Cyclopia* spp.), thus raising the possibility that they could be used as agricultural inoculants when their particular characteristics (e.g., tolerance to pH extremes, high salt tolerance) make them more suited to specific environments, such as in Morocco (Talbi et al. 2010) and the South African Cape (Elliott et al. 2007b). However, before developing these inocula, we need to know whether their relationship to human, animal, and plant pathogens would preclude their use in agriculture. Current fears about their use should be at least partly allayed by the evidence for divergence that underlies the proposed transfer of the plant-associated and nonpathogenic Burkholderia spp. to the new genus Caballeronia.

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## AUTHOR-RECOMMENDED INTERNET RESOURCES

- ICSP–Subcommittee on the taxonomy of Rhizobium and Agrobacterium website: edzna.ccg.unam.mx/rhizobial-taxonomy
- GOLD: Genomes Online database: www.genomesonline.org The Integrated Microbial Genome (IMG) family systems website:
- genomebiology.jgi-psf.org/Content/IMG\_system.htm National Center for Biotechnology Information's Bioproject *B. phymatum* STM815 page: www.ncbi.nlm.nih.gov/bioproject/58699
- B. S. Weir's taxonomy of rhizobia webpage: www.rhizobia.co.nz/taxonomy/rhizobia.html