APPLIED GENETICS AND MOLECULAR BIOTECHNOLOGY

Ribosomal protein biomarkers provide root nodule bacterial identification by MALDI-TOF MS

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Received: 12 January 2015 / Revised: 19 February 2015 / Accepted: 28 February 2015 / Published online: 18 March 2015 © Springer-Verlag Berlin Heidelberg 2015

Abstract Accurate identification of soil bacteria that form nitrogen-fixing associations with legume crops is challenging given the phylogenetic diversity of root nodule bacteria (RNB). The labor-intensive and time-consuming 16S ribosomal RNA (rRNA) sequencing and/or multilocus sequence analysis (MLSA) of conserved genes so far remain the favored molecular tools to characterize symbiotic bacteria. With the development of mass spectrometry (MS) as an alternative method to rapidly identify bacterial isolates, we recently showed that matrix-assisted laser desorption ionization (MALDI) time-of-flight (TOF) can accurately characterize RNB found inside plant nodules or grown in cultures. Here,

Electronic supplementary material The online version of this article (doi:10.1007/s00253-015-6515-3) contains supplementary material, which is available to authorized users.

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Laboratory of Applied Microbiology, Department of Environment Construction and Design, University of Applied Sciences of Southern Switzerland, via Mirasole 22a, CH-6500 Bellinzona, Switzerland we report on the development of a MALDI-TOF RNBspecific spectral database built on whole cell MS fingerprints of 116 strains representing the major rhizobial genera. In addition to this RNB-specific module, which was successfully tested on unknown field isolates, a subset of 13 ribosomal proteins extracted from genome data was found to be sufficient for the reliable identification of nodule isolates to rhizobial species as shown in the putatively ascribed ribosomal protein masses (PARPM) database. These results reveal that data gathered from genome sequences can be used to expand spectral libraries to aid the accurate identification of bacterial species by MALDI-TOF MS.

Keywords Bacterial fingerprints · Phylogeny · Cluster analysis · Rhizobia · Legume nodules · GEBA-RNB

Introduction

Soil bacteria, collectively known as Rhizobia or root nodule bacteria (RNB), form beneficial nitrogen-fixing associations with most leguminous plants, including important crops such as soybean (Glycine max), bean (Phaseolus vulgaris), pea (Pisum sativum), chickpea (Cicer arietinum), pigeon pea (Cajanus cajan), and pastures such as clovers (Trifolium spp.) or Medicago spp. These associations provide significant sustainable inputs of reduced nitrogen (N) to agricultural systems that are estimated to be ca. 50-70 Tg/year (Herridge et al. 2008). A key aspect of symbiotic nitrogen fixation is the development of specialized structures called nodules on the roots (or more rarely stems) of host plants. Within the nodule cells, RNB differentiate into bacteroids, which reduce atmospheric nitrogen (N_2) to ammonia (Oldroyd and Downie 2008; Udvardi and Poole 2013). Rhizobial infection and nodule organogenesis require a coordinated exchange of molecular signals between the host and its symbiotic bacteria (Gibson et al. 2008; Perret et al. 2000).

Although most of the approximately 20,000 species of the Leguminosae are able to nodulate (Doyle 2011), only a small fraction of their rhizobial associations have been characterized. In contrast to legumes, which are monophyletic (Werner et al. 2014), molecular techniques have demonstrated that rhizobia are polyphyletic, belonging to diverse bacterial genera of the Alphaproteobacteria and Betaproteobacteria (Masson-Boivin et al. 2009). In RNB, symbiotic ability is conferred by genes for nodulation and nitrogen fixation that can be acquired via lateral transfer of plasmids or genomic islands (Broughton 2003; Freiberg et al. 1997; Moulin et al. 2004; Nandasena et al. 2007; Sullivan and Ronson 1998). To date, 16 rhizobial genera and over 100 species have been validly described (ICSP Subcommittee on the taxonomy of Rhizobium and Agrobacterium-diversity, phylogeny, and systematics; http://edzna.ccg.unam.mx/rhizobial-taxonomy/ node/4), and these numbers are expected to increase as more symbioses are studied.

Phenotypic methods of identifying and typing bacteria, such as colony morphology and serotyping, have largely been superseded by molecular markers. Sequence analysis of the 16S ribosomal RNA (rRNA) gene allows the construction of valid bacterial phylogenies (Schleifer 2009), but lacks resolving power at and below the species level, due to the extent of gene conservation (Willems 2006). Multilocus sequence analysis (MLSA) of conserved core genes was proposed as a robust and reliable alternative method for classifying bacteria to species level (Gevers et al. 2005; Hanage et al. 2006) and has been successfully used to delineate species within a number of genera including Ensifer, formerly Sinorhizobium (Martens et al. 2008), and Bradyrhizobium (Rivas et al. 2009). However, incongruent phylogenies have been reported among several genes commonly used in MLSA, along with evidence of lateral gene transfer or intergenic recombination, resulting in unclear taxonomic resolution (Tian et al. 2012; Zhang et al. 2012). No single genealogical reconstruction or typing method currently encompasses all levels of bacterial diversity, from domain to strain. Moreover, molecular marker analysis is costly and laborious, making it unsuitable for use in high throughput, broad-scale studies of rhizobial taxonomy, phylogeny, biogeography, or population genetics.

Alternatively, matrix-assisted laser desorption ionization (MALDI) time-of-flight (TOF) mass spectrometry (MS) has increasingly replaced DNA-based procedures for the reliable identification of bacteria, in particular, in diagnostic laboratories responsible for characterizing clinical isolates (Seng et al. 2009). MALDI-TOF MS offers the possibility of accurately resolving bacterial identity to the genus, species, and subspecies levels in some taxa and has successfully been applied to diverse rhizobial strains grown on solid media (Ferreira et al. 2011; Sánchez-Juanes et al. 2013; Ziegler et al. 2012) and

found inside legume nodules (Ziegler et al. 2012). In most direct MALDI-TOF MS analyses, bacterial cells are lysed to release the intracellular proteins, which are then ionized and separated according to their mass-to-charge ratio (m/z) and recorded as distinct peaks, generating a complex spectrum, or fingerprint that is characteristic of a bacterial sample. In general, microbial MALDI-TOF MS signatures can be used to identify unknown bacterial samples down to the subspecies level, providing they can be matched against comprehensive databases of empirical spectra obtained for sets of wellcharacterized (but not necessarily type) strains selected to represent the microbial genera of interest. For example, the enlarged Spectral ARchive And Microbial Identifications System (SARAMIS[™] ver. 4.10, system ver. 3.4.1.2; Mabritec, Riehen, Switzerland) database contains characteristic signatures for more than 650 bacterial species, mostly of clinical importance.

MALDI-TOF MS biomarkers do not need to be linked to specific proteins to ensure correct identification of bacterial strains. However, many of the individual peaks that form a MALDI-TOF mass spectrum correspond to ribosomal proteins (RPs), which are encoded by more than 50 genes scattered in mostly chromosomal loci. RPs are ubiquitous and among the most abundant cytosolic proteins, regardless of the cell life stage or growth conditions (Ishihama et al. 2008), and have masses mostly within the 4 to 30 kDa range that is examined with MALDI-TOF MS. Ribosomal multilocus sequence typing (rMLST) has been shown to be consistent with current bacterial nomenclature schemes and to provide accurate phylogenies for diverse taxonomic groups (Jolley et al. 2012; Maiden et al. 2013; Yutin et al. 2012). These characteristics make RPs reliable biomarkers in MALDI-TOF MS analyses to obtain species and subspecies identification of diverse bacteria including bifidobacteria (Sato et al. 2011), lactobacilli (Sato et al. 2012; Teramoto et al. 2007a), rhodococci (Teramoto et al. 2009), members of the Sphingomonadaceae (Hotta et al. 2012), and strains of Staphylococcus aureus (Josten et al. 2013) and Neisseria meningitidis (Suarez et al. 2013).

Additionally, the calculated masses of RPs deduced from partial or whole genome sequences were shown to be sufficient for correctly identifying unknown bacteria, even in the absence of empirical mass spectral fingerprints from a corresponding reference microorganism (Pineda et al. 2003). Together, these results suggested that empirical spectra of strains assembled into a reference database, such as SARAMIS[™], could be supplemented by lists of biomarkers corresponding to masses of RPs calculated from the sequenced genomes of reference strains.

Given the potential for MALDI-TOF MS to facilitate and expedite the exploration of rhizobial diversity in agricultural and natural ecosystems (Ziegler et al. 2012), our first objective was to establish a comprehensive empirical

 Table 1
 List of 114 Rhizobia and two tumorigenic strains used to generate the reference database of MALDI-TOF MS spectra for the identification of nodule bacteria

#	Genus	Species	Strain	Isolated from	GOLD ID	Source
1	Agrobacterium	A. tumefaciens	C58-UWash	Cherry tree tumor	Gc00073	J.F. Pothier
2		A. vitis	S4	Vitis vinifera	Gc00943	J.F. Pothier
3	Azorhizobium	A. caulinodans	ORS 571 ^T	Sesbania rostrata	Gc00669	S. Goormachtig
ŀ		A. doebereinerae	UFLA1-100 ^T	Sesbania virgate	Gi08849	W.G. Reeve
	Bradyrhizobium	B. canariense	WSM4349	Syrmatium glabrum	Gi08886	W.G. Reeve
		B. elkanii	USDA 76 ^T	Glycine max	Gi08850	W.G. Reeve
		B. elkanii	USDA 94	G. max	Gi10733	W.G. Reeve
;		Bradyrhizobium sp.	WSM2783	Leobordea carinata	Gi08855	W.G. Reeve
)		B. elkanii	USDA 3259	Phaseolus lunatus	Gi22151	W.G. Reeve
0		B. elkanii	USDA 3254	Phaseolus acutifolius	Gi08852	W.G. Reeve
1		Bradyrhizobium sp.	WSM1741	Rhynchosia minima	Gi08854	W.G. Reeve
2		B. japonicum	USDA 6^{T}	G. max	Gc02045	W.G. Reeve
3		B. japonicum (B. diazoefficiens)	USDA 110	G. max	Gc00116	X. Perret
4		B. japonicum (B. diazoefficiens)	USDA 122	G. max	Gi08856	W.G. Reeve
5		Bradyrhizobium sp.	USDA 135	G. max	Gi08859	W.G. Reeve
6		Bradyrhizobium sp.	USDA 38	G. max	Gi08860	W.G. Reeve
7		Bradyrhizobium sp.	USDA 4	G. max	Gi08861	W.G. Reeve
8		Bradyrhizobium sp.	WSM1743	Indigofera spp.	Gi08863	W.G. Reeve
9		Bradyrhizobium sp.	WSM2793	Rhynchosia totta	Gi08908	W.G. Reeve
0		Bradyrhizobium sp.	ARR65	Stylosanthes viscosa	Gi08868	W.G. Reeve
1		Bradyrhizobium sp.	BTAi1	Aeschynomene indica	Gc00561	E. Giraud
2		Bradyrhizobium sp.	ORS285	Aeschynomene spp.	Gi09732	E. Giraud
3		Bradyrhizobium sp.	ORS278	Aeschynomene spp.	Gc00552	E. Giraud
4		Bradyrhizobium sp.	USDA 3384	Crotalaria paulina	Gi10733	W.G. Reeve
5		B. canariense	WSM1253	Omithopus compressus	Gi06492	W.G. Reeve
6		B. canariense	WSM1417	Lupinus spp.	Gi06490	W.G. Reeve
7		Bradyrhizobium sp.	WSM2254	Acacia spp.	Gi14039	W.G. Reeve
8		Bradyrhizobium sp.	WSM3983	Kennedia coccinea	Gi08871	W.G. Reeve
9		B. canariense	WSM471	Lupinus angustifolius	Gi06491	W.G. Reeve
0	Burkholderia	B. mimosarum	LMG 23256 ^T	Mimosa pigra	Gi08823	W.G. Reeve
1		Burkholderia sp.	WSM2230	K. coccinea	Gi08831	W.G. Reeve
2		B. dilworthii	WSM3556 ^T	Lebeckia ambigua	Gi08872	W.G. Reeve
3		B. tuberum	WSM4176	L. ambigua	Gi08873	W.G. Reeve
4		Burkholderia sp.	JPY347	Mimosa cordistipula	Gi08875	W.G. Reeve
5		Burkholderia sp.	JPY251	Mimosa velloziana	Gi08874	W.G. Reeve
6		Burkholderia sp.	UYPR1.413	Parapiptadenia rigida	Gi08829	W.G. Reeve
7		B. sprentiae	WSM5005 ^T	L. ambigua	Gi06497	W.G. Reeve
8	Cupriavidus	Cupriavidus sp.	UYPR2.512	P. rigida	Gi08830	W.G. Reeve
9		C. taiwanensis	LMG 19424 ^T	Mimosa pudica	Gc00754	M. M. Saad
0		C. taiwanensis	STM6070	M. pudica	Gi08841	W.G. Reeve
1	Ensifer	E. arboris	LMG 14919 ^T	Prosopis chilensis	Gi08822	W.G. Reeve
2		E. fredii	HH103	G. max	Gc02113	J.E. Ruiz-Sainz
3		E. fredii	USDA257	G. max	Gc02300	W.G. Reeve
4		E. fredii	NGR234	Lablab purpureus	Gc00990	X. Perret
5		E. medicae	WSM419	Medicago murex	Gc00590	W.G. Reeve
6		E. medicae	Di28	Medicago arabica	Gi08905	W.G. Reeve
7		E. medicae	WSM1115	Medicago polymorpha	Gi08906	W.G. Reeve
8		E. medicae	WSM1369	Medicago sphaerocarpos	Gi08907	W.G. Reeve

Table 1 (continued)

ŧ	Genus	Species	Strain	Isolated from	GOLD ID	Source
19		E. medicae	WSM244	M. polymorpha	Gi08908	W.G. Reeve
0		E. medicae	WSM4191	Melilotus siculus	Gi08903	W.G. Reeve
1		E. meliloti	AK83	Medicago falcata	Gc01810	A. Mengon
2		E. meliloti	Mlalz-1	Medicago laciniata	Gi08913	W.G. Reeve
3		E. meliloti	CIAM1775	Medicago lupulina	Gi08844	W.G. Reeve
4		E. meliloti	WSM1022	Medicago orbicularis	Gi08916	W.G. Reeve
5		E. meliloti	1021	Medicago sativa	Gc00059	X. Perret
5		E. meliloti	BL225C	M. sativa	Gc01811	A. Mengon
7		E. meliloti	MVII-I	M. sativa	Gi08914	W.G. Reev
3		E. meliloti	RR1128	Medicago truncatula	Gi08915	W.G. Reev
)		E. meliloti	4H41	Phaseolus vulgaris	Gi08911	W.G. Reev
)		E. meliloti	GVPV12	P. vulgaris	Gi08912	W.G. Reev
		<i>Ensifer</i> sp.	TW10	Tephrosia wallichii	Gi08835	W.G. Reev
		<i>Ensifer</i> sp.	WSM1721	Indigofera spp.	Gi08904	W.G. Reev
	Mesorhizobium	M. australicum	WSM2073 ^T	Biserrula pelecinus	Gc02468	W.G. Reev
		M. ciceri	WSM4083	Bituminaria bituminosa	Gi08880	W.G. Reev
		M. ciceri	CMG6	Cicer arietinum	Gi08879	W.G. Reev
		M. ciceri bv. Biserrulae	WSM1271	B. pelecinus	Gc01578	W.G. Reev
		M. loti	USDA 3471 ^T	Lotus corniculatus	Gi08881	W.G. Reev
		M. loti	CJ3sym	L. corniculatus	Gi08828	W.G. Reev
		M. loti	MAFF303099	Lotus japonicus	Gc00040	X. Perret
		M. loti	NZP2037	Lotus divaricatus	Gi08826	W.G. Reev
		M. loti	R7A	L. corniculatus	Gi08825	W.G. Reev
		M. loti	R88b	L. corniculatus	Gi08827	W.G. Reev
		M. loti	WSM1293	Lotus spp.	Gi08882	W.G. Reev
Ļ		M. opportunistum	WSM2075 ^T	B. pelecinus	Gc01853	W.G. Reev
		Mesorhizobium sp.	WSM2561	Lessertia diffusa	Gi08883	W.G. Reev
		Mesorhizobium sp.	WSM3626	L. diffusa	Gi08885	W.G. Reev
7		Mesorhizobium sp.	WSM3224	Otholobium candicans	Gi08884	W.G. Reev
	Methylobacterium	M. nodulans	ORS 2060^{T}	Crotalaria podocarpa	Gc00935	E. Giraud
	menyloouelerium	Methylobacterium sp.	WSM2598	Listia bainesii	Gi08887	W.G. Reev
	Microvirga	M. lotononidis	WSM3557 ^T	Listia angolensis	Gi06493	W.G. Reev
	merovirgu	M. lupini	Lut6 ^T	Lupinus texensis	Gi06478	W.G. Reev
	Rhizobium	R. etli	CFN 42 ^T	P. vulgaris	Gc00342	S. Brom
	101120010111	R. phaseoli	CIAT 652	P. vulgaris	Gc00823	S. Brom
		R. giardinii bv. giardinii	$H152^{T}$	P. vulgaris	Gi08897	W.G. Reev
		R. leguminosarum bv. phaseoli	4292	P. vulgaris	Gi08895	W.G. Reev
, ,		R. leguminosarum bv. phaseoli R. leguminosarum bv. phaseoli	FA23	P. vulgaris	Gi08821	W.G. Reev
		R. leguminosarum bv. trifolii	TA1	Trifolium spurmosum	Gi06488	W.G. Reev
		R. leguminosarum bv. trifolii	WSM2297	Trifolium africanum	Gi06477	W.G. Reev
		R. leguminosarum bv. trifolii R. leguminosarum bv. trifolii	CC283b	Trifolium ambiguum	Gi06484	W.G. Reev W.G. Reev
		R. leguminosarum bv. trifolii R. leguminosarum bv. trifolii	CC2830 CC278f	Trifolium amolguum Trifolium nannum	Gi06479	W.G. Reev W.G. Reev
		R. leguminosarum bv. trifolii R. leguminosarum bv. trifolii	WSM597	Trifolium nannum Trifolium pallidum	Gi06479 Gi06486	W.G. Reev W.G. Reev
		R. leguminosarum bv. trifolii R. leguminosarum bv. trifolii	WSM2304	Trifolium paluaum Trifolium polymorpha	G106486 Gc00870	W.G. Reev W.G. Reev
						W.G. Reev W.G. Reev
		R. leguminosarum by. trifolii R. leguminosarum by. trifolii	WSM2012	Trifolium ruepellianum Trifolium spp	Gi06480	
ļ		R. leguminosarum bv. trifolii	WSM1325	Trifolium spp.	Gc01039	W.G. Reev
5		R. leguminosarum bv. trifolii R. leguminosarum bv. trifolii	SRDI565 SRDI943	Trifolium subterraneum T. subterraneum	Gi08843 Gi08842	W.G. Reev W.G. Reev
5						

Table 1 (continued)

# Ge	enus	Species	Strain	Isolated from	GOLD ID	Source
98		R. leguminosarum bv. trifolii	CB782	Trifolium semipilosum	Gi06498	W.G. Reeve
99		R. leguminosarum bv. viciae	128C53	Pisum sativum	Gi08894	W.G. Reeve
100		R. leguminosarum bv. viciae	3841	P. sativum	Gc00385	P.S. Poole
101		R. leguminosarum bv. viciae	GB30	P. sativum	Gi08820	W.G. Reeve
102		R. leguminosarum bv. viciae	Ps8	P. sativum	Gi08888	W.G. Reeve
103		R. leguminosarum bv. viciae	UPM1131	P. sativum	Gi08837	W.G. Reeve
104		R. leguminosarum bv. viciae	UPM1137	P. sativum	Gi08836	W.G. Reeve
105		R. leguminosarum bv. viciae	TOM	P. sativum	Gi08893	W.G. Reeve
106		R. leguminosarum bv. viciae	Vc2	Vicia cracca	Gi08889	W.G. Reeve
107		R. leguminosarum bv. viciae	248	Vicia faba	Gi08896	W.G. Reeve
108		R. leguminosarum bv. viciae	VF39	V. faba	Gi08892	W.G. Reeve
109		R. leguminosarum bv. viciae	WSM1455	V. faba	Gi06482	W.G. Reeve
110		R. leguminosarum bv. viciae	WSM1481	V. faba	Gi08891	W.G. Reeve
111		R. leguminosarum bv. viciae	Vh3	Vicia hirsuta	Gi08890	W.G. Reeve
112		R. mongolense	USDA 1844 ^T	Medicago ruthenica	Gi08900	W.G. Reeve
113		Ensifer ^a sp.	BR816	Leucaena leucocephala	Gi08910	W.G. Reeve
114		R. mesoamericanum	STM6155	M. pudica	Gi08838	W.G. Reeve
115		Rhizobium sp.	OR 191	M. sativa	Gi08824	W.G. Reeve
116		R. sullae	WSM1592	Hedysarum coronarium	Gi08899	W.G. Reeve

^T type strains. Reference strains were obtained from Wayne G. Reeve (Center for *Rhizobium* Studies, Murdoch University, Perth, Australia); Maged M. Saad (Microbiology Unit, University of Geneva, Geneva, Switzerland); Eric Giraud (Laboratoire des Symbioses Tropicales et Méditéranéennes, Montpellier, France); Julio E. Ruiz-Sainz (Departamento de Microbiología, Universidad de Sevilla, Spain); Susana Brom (Center for Genomic Sciences, Universidad Nacional Autonoma de Mexico, Cuernavaca, Mexico); Sofie Goormachtig (Department of Plant Biotechnology and Genetics, Ghent University, Ghent, Belgium); Alessio Mengoni (Department of Animal Biology and Genetics, Florence University, Florence, Italy); Philip S. Poole (John Innes Center, Norwich, UK); and Joël F. Pothier (ZHAW Wädenswil, Wädenswil, Switzerland). Sequenced genomes were accessed as genome online database identifiers (GOLD ID; https://gold.jgi-psf.org).

^a Due to changes in taxonomy, strain BR816 has previously been identified as *Rhizobium*, *Sinorhizobium*, and *Ensifer*. The following two strains were recently reclassified as new type strains: *Bradyrhizobium diazoefficiens* USDA110^T (Delamuta et al. 2013) and *Burkholderia dilworthii* WSM3556^T (De Meyer et al. 2014)

reference database for unambiguously assigning RNB to known rhizobial genera. To this end, two strains of planttumorigenic and 114 strains of plant-symbiotic bacteria that were representative of the major rhizobial genera were used to construct the RNB-specific module extension to the SARAMISTM database. As the genomes of these 116 selected strains had been sequenced, mostly as part of the Genomic Encyclopedia of Bacteria and Archaea-Root Nodule Bacteria (GEBA-RNB) project, we expanded the empirical RNB-specific module with additional SARA MISTM-compatible databases of biomarkers that corresponded to (a) the deduced ribosomal protein masses (DRPM) calculated from genomic data and (b) a subset of putatively ascribed ribosomal protein masses (PARPM) that matched mass peaks in the empirical spectra. The DRPM and PARPM databases could be used to provide accurate identification of rhizobial isolates, suggesting that lists of biomarker masses deduced from genomic data may complement the databases of empirical spectra obtained from pure culture of bacteria.

Materials and methods

Acquisition of root nodules and bacteria

Reference strains used in this study are listed in Table 1 together with some of their major characteristics including the name of the host plant they were isolated from and the genome online database identifier (GOLD ID) number under which sequence data was stored. Root-nodulating and tumorigenic bacteria were grown in/on TY (Beringer 1974) or YM (Vincent 1970) media and incubated at 28 °C as growth temperature. Nodules were collected from field-grown legumes at locations in Western Australia (Brookton and Esperance) and Ivory Coast (Bondoukou and Kossou-Bouafla) and kept desiccated until further use. For analysis of nodule bacteria, desiccated nodules were rehydrated 30 min prior to surface sterilization with 70 % EtOH and 4 % bleach, with respective incubation times adjusted for nodule size. Once thoroughly rinsed with sterile H₂O, surface sterilized nodules were then processed as described by Ziegler et al. (2012).

Determining symbiotic properties of nodule isolates

Nodulation assays were carried out in Magenta jars containing vermiculite (Lewin et al. 1987) with B&D nitrogen-free nutritive solution (Broughton and Dilworth 1971). Once surfacesterilized, seeds of *C. cajan* (pigeon pea) and *Vigna unguiculata* cv. Red Caloona (cowpea) were incubated several days in the dark at 27 °C for germination. Seedlings were then planted in autoclaved Magenta jars, and 2–3 days later each plantlet was inoculated with 200 μ l of a solution containing 2×10^8 bacteria. Once inoculated, plants were grown for 6 weeks at a day temperature of 27 °C, a night temperature of 20 °C and a light phase of 12 h prior to harvesting of nodules.

Preparation of samples for MALDI-TOF MS analyses

For generating spectra for strain identification and/or construction of the Rhizobia-specific module of SARAMISTM, free-living bacteria were harvested after 1, 2, 3, 5, and 7 days growth on TY (Beringer 1974) or YM (Vincent 1970) plates incubated at 28 °C. Bacterial samples were spotted in quadruplicate on MALDI steel target plates. Spots were overlaid with 1 µl of 25 % formic acid, air-dried, and again overlaid with 1 µl of matrix solution consisting of saturated solution of alpha-cyano-4 hydroxycinnamic acid (CHCA; Sigma-Aldrich, Buchs, Switzerland) in 33 % acetonitrile (Sigma-Aldrich), 33 % ethanol and supplemented with 3 % trifluoroacetic acid (TFA). Alternatively, sinapinic acid (SA) diluted in 60 % acetonitrile and 0.3 % TFA was used as matrix. Finally, all samples processed in these ways were air-dried for a few minutes at room temperature. For ascribing ribosomal proteins to specific mass peaks in empirical spectra, cell lysates were used instead of whole bacteria. Cell lysates were obtained as follows: after growth on the appropriate medium cells were collected and washed three times in TMA-I buffer (10 mM Tris-HCl pH 7.8, 30 mM NH₄Cl, 10 mM MgCl₂, and 6 mM 2-mercaptoethanol). Once resuspended in 500 µl TMA-I buffer, cells were ground three times for 20 s at 600 rpm using 0.1 mm glass beads in a FastPrep FP120 bead beater cell disrupter (Farmingdale, NY, USA). Glass beads and cell debris were removed by centrifugation at $5,800 \times g$ for 10 min and aliquots of the supernatant spotted for MALDI-TOF MS measurements as described above and using SA as matrix.

Acquisition of MALDI-TOF mass spectra and data processing

Empirical spectra were obtained using a MALDI-TOF Mass Spectrometer Axima[™] Confidence machine (Shimadzu-Biotech, Kyoto, Japan), with detection in the linear positive mode at a laser frequency of 50 Hz and within a mass range of 3 to 20 kDa or 4 to 30 kDa when CHCA or SA were used as matrices, respectively. Acceleration voltage was 20 kV, and the extraction delay time was 200 ns. A minimum of ten laser shots per sample was used to generate each ion spectrum. For each bacterial sample, a total of 50 to 100 protein mass fingerprints were averaged and processed using the LaunchpadTM 2.8 software (Shimadzu-Biotech). For peak processing of raw spectra, the following settings were selected for LaunchpadTM 2.8: (i) the advanced scenario from the Parent Peak Cleanup menu, (ii) a peak width set at 80 chans, (iii) the smoothing filter width set at 50 chans, (iv) a baseline filter width of 500 chans, and (v) a threshold apex method set as dynamic. The threshold offset was set at 0.020 mV with a response factor of 1.2. Each target plate was externally calibrated using spectra of the reference strain *Escherichia coli* DH5 α (Invitrogen, Carlsbad, USA).

Constructing the databases of deduced and putatively ascribed ribosomal protein masses (DRPM and PARPM)

Genomes of the 116 selected RNB or tumorigenic bacteria were accessed at the US Department of Energy's (DoE) Joint Genome Institute (JGI) or the National Center for Biotechnology Information (NCBI) using the strain-specific GOLD ID listed in Table 1. DNA sequences annotated as coding for RPs were collected, aligned, and checked for correct start and stop codons. Genes coding for RPs that were either misannotated or missing from a given genome were manually curated using corresponding sequences from closely related strains as in silico probes. DNA sequences verified in this way and coding for a complete set of 54 ribosomal proteins for each of the 116 selected strains were translated into corresponding amino acid sequences. Masses were then deduced taking into account the most frequent posttranslational modifications in bacteria (Arnold and Reilly 1999). Thus, 131 Da corresponding to the loss of the N-terminal methionine was subtracted from the deduced ribosomal protein mass when the penultimate amino acid was found to be either Gly, Ala, Pro, Ser, Thr, Val, or Cys. With the exception of Burkholderia strains for which empirical spectra showed no such modifications, ribosomal protein L33 was always considered to be methylated. For each strain, a text file compatible with the SARAMISTM database and containing the relevant taxonomical information and deduced masses for all 54 RPs together with arbitrary intensity values was created and stored into a separate DRPM database. Using a possible error interval of 800 ppm, this dataset was matched against empirical spectra to ultimately identify among mass peaks those that corresponded to RPs. Mass peaks putatively ascribed to RPs L20, L24, L28, L29, L31, L32, L33, L36, S10, S15, S16, S17, and S20 were found to be consistently present in empirical spectra generated for each of the 116 reference strains. For each strain, mass peaks ascribed to each of these 13 marker RPs and intensity values were compiled into a SARAMISTM-compatible text file that was stored into a

distinct subdatabase called PARPM for putatively ascribed ribosomal proteins masses.

Genetic analyses

Sequences of 16S rRNA, recA, gyrB, rpoB, dnaJ, and atpD genes or ribosomal protein genes (rplT, rplX, rpmB, rpmC, rpmE, rpmF, rpmG, rpmJ, rpsJ, rpsO, rpsP, rpsO, and rpsT) were accessed using the NCBI (www.ncbi.nlm.nih.gov) and JGI (genome.jgi.doe.gov) databases. As full 16S rRNA gene sequences were missing from the draft genomes of Rhizobium leguminosarum by. viciae strains 128C53, Ps8, and Vh3, as well as strain Rhizobium sullae WSM1592, this gene was amplified and sequenced. Genomic DNA was extracted using the InstaGene[™] kit (Bio-Rad Laboratories, Hercules, CA, USA). Primers nucleotide sequence 5'-ATTCTAGAGT TTGATCATGGCTCA-3' (UNI-L) and nucleotide sequence 5'-ATGGTACCGTGTGACGGGGGGGGTGTGTA-3' (UNI-R) were then used to amplify and sequence a fragment of ca. 1,400 bp (Brosius et al. 1978). Sequencing reactions were carried out using Big DyeTerminator v3.1 Cycle Sequencing Kit and results were analyzed using the 3500 Data Collection software v.1.00 (Applied Biosystems, Foster City, CA, USA).

Phylogenetic analyses

For each locus of interest, DNA sequences were trimmed to the same length and aligned using the MEGA 5 software package (Tamura et al. 2011). Phylogenetic trees were reconstructed using the maximum composite likelihood method and default settings. For hierarchical clustering of protein masses, the presence/absence binary matrices were generated using the SARAMIS Superspectra tool. Results were then imported into the PAleontological STatistics (PAST) software (Natural History Museum, Oslo University, Norway). Multivariate neighbor joining cluster analyses with similarity distances were calculated using Dice algorithm in PAST software (Feltens et al. 2010). Figure 1 was generated and colored using the R software (http://www.r-project.org/).

Results

A comprehensive database for the identification of nodule bacteria via MALDI-TOF MS

In order to establish a database of MALDI-TOF MS spectra that could capture the existing diversity of rhizobia, we sourced representative strains of diverse rhizobial species (see Table 1). Most of the selected RNB were part of the DoE Joint Genome Institute's (JGI) international Genomic Encyclopedia of Bacteria and Archaea-Root Nodule Bacteria (GEBA-RNB) project that aimed to systematically sequence the genomes of diverse rhizobial strains (Reeve et al. 2014). The remaining strains were well-studied bacteria whose genomes have previously been sequenced. Together, these 116 selected strains represented the ten major genera of RNB and tumorigenic bacteria: *Agrobacterium*, *Azorhizobium*, *Bradyrhizobium*, *Burkholderia*, *Cupriavidus*, *Ensifer* (syn. *Sinorhizobium*), *Mesorhizobium*, *Methylobacterium*, *Microvirga*, and *Rhizobium*, isolated from 72 legume species and over 30 countries. This collection of strains included 27 fully described species, of which *R. leguminosarum* and its three biovars *phaseoli*, *trifolii*, and *viciae* comprised the largest number of strains (*n*=27).

To build up the database of MALDI mass spectra, each strain was grown at 28 °C on YM or TY solid media for 1, 2, 3, 5, and 7 days before being fingerprinted in quadruplicate using suspensions in 25 % formic acid. This procedure was found to be the most reliable for obtaining reproducible MALDI mass spectra for both free-living plate-cultured Rhizobia and bacteria analyzed directly from nodules, independently of the age of colonies or plant nodules (Ziegler et al. 2012). For each of the 116 reference strains, a reference SuperSpectrum[™] (SSp) was computed using at least 15 individual spectra of the highest quality and, depending on the range of biomarker masses targeted, the alpha-cyano-4hydroxycinnamic acid (CHCA; 3-12 kDa) or sinapinic acid (SA; 4-30 kDa) matrices (Pineda et al. 2003). The 232 SSp generated in this way together formed the "rhizobia-specific module" of the SARAMIS™ database.

Using deduced ribosomal proteins as markers for MALDI-TOF MS identification of rhizobia

Given the increasing number of sequenced bacterial genomes and the difficulty of obtaining numerous yet wellcharacterized and curated reference strains for further developing the rhizobia-specific SARAMISTM module, we searched for ways to replace the empirically derived MALDI-TOF mass spectra with databases consisting of deduced masses of biomarker ribosomal proteins, which had been calculated from genomic sequence data.

Genomes of the 116 selected RNB and tumorigenic bacteria that were accessed online using the JGI or National Center for Biotechnology Information (NCBI) homepages and the genome online database identifiers (GOLD ID) are listed in Table 1. The search revealed that each of the genomes included genes of 54 RPs that potentially could be used as biomarkers. Masses of the 54 RPs were deduced from genomic sequences and compiled into strain-specific lists of biomarkers. Where genes coding for RPs were either missing or found to be misannotated, ribosomal protein nucleotide sequences of closely related strains were used to identify the correct open-reading frame. Table S1 in the Supplementary Material shows the 163 RPs for which these genome

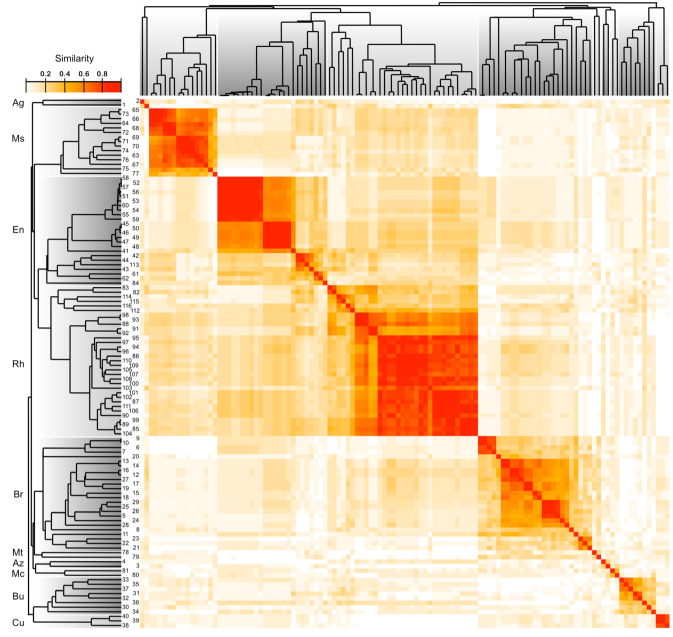


Fig. 1 Hierarchical cluster analysis showing similarities in the number of identical masses calculated for all of the ribosomal proteins (RPs) and in pairwise combinations of strains. Strains are numbered as in Table 1, and alternating shades of *grey* delimit the boundaries (from *left* to *right* and *top* to *bottom*) for the *Agrobacterium* (*Ag*), *Mesorhizobium* (*Ms*), *Ensifer* (*En*), *Rhizobium* (*Rh*), *Bradyrhizobium* (*Br*), *Methylobacterium* (*Mt*),

Azorhizobium (*Az*), *Microvirga* (*Mc*), *Burkholderia* (*Bu*), and *Cupriavidus* (*Cu*) genera. The degree of homogeneity or heterogeneity of clustering groups is illustrated in the heatmap. Levels of similarity between the theoretical spectra composed of 54 RPs are color-coded, from *white* (no similarity) to *dark red* (full identity)

annotation corrections needed to be made. In order to calculate masses of ribosomal proteins as accurately as possible, the most frequent posttranslational modifications were taken into account. For example, 131 Da corresponding to a cleaved methionine was subtracted from the predicted mass of the ribosomal protein when the amino acid in position 2 was either Gly, Ala, Pro, Ser, Thr, Val, or Cys. In addition, ribosomal protein L33 was always considered to be methylated (Arnold and Reilly 1999), except for *Burkholderia* strains, where our

experimental data showed no evidence for such methylation. Masses for RPs calculated in this way ranged from 4.3 kDa (L36 of *Cupriavidus taiwanensis* LMG 19424^T) to 63.2 kDa (S1 of *Bradyrhizobium* sp. ARR65), of which 71 % had a mass below the 30 kDa upper limit for MALDI-TOF spectra acquired using sinapinic acid as matrix. To introduce this dataset into the SARAMIS database, a text file listing all of the 54 RPs masses with arbitrary intensity values was compiled for each of our 116 bacterial strains. This complete dataset of deduced ribosomal protein masses was labeled as DRPM.

Figure 1 shows two complementary representations of a hierarchical cluster analysis using the Dice algorithm for similarities in the number of identical masses among the 54 deduced RPs in pairwise combinations of strains. In the cluster trees, genera boundaries are delimited using labeled and shaded domains with strains being numbered as in Table 1. In the central diagram, color codes ranging from white (no similarity) to dark red (full identity) show corresponding levels of similarity. Similar and more detailed analyses for strains belonging to each of the genera *Bradyrhizobium*, *Ensifer*, *Mesorhizobium*, and *Rhizobium*, respectively, are presented in Figs. S1–S4.

Except for Rhizobium giardinii by. giardinii type strain H152^T (strain #84), strains clustered into distinct and taxonomically correct genera with Rhizobia belonging to the Beta subdivision of Proteobacteria (Burkholderia and *Cupriavidus*) forming a distinct branch (shown at the very bottom or the very right in Fig. 1). Interestingly, the theoretical RPs spectrum of *R. giardinii* by. *giardinii* H152^T appeared to be more similar to those of Ensifer than Rhizobium strains, a finding consistent with the unclear phylogenetic position of H152^T reported previously (Laguerre et al. 2001; Mousavi et al. 2014; Ormeño-Orrillo et al. 2015). Closely related strains, such as the Ensifer medicae (#45 to 50) and Ensifer meliloti (#51 to 60) isolates, formed two compact clusters clearly separated from other Ensifer members by Ensifer arboris (#41) (see also Supplementary Fig. S2). A similar topology was reported for a 16S rRNA-based phylogenetic tree of Ensifer strains when describing the genome sequence of *E. arboris* LMG 14919^T (Reeve et al. 2013). The theoretical RPs spectrum also confirmed differences between Rhizobium etli CFN 42^T (#82) and strain CIAT 652 (#83) and supported the classification of CIAT 652 as belonging to Rhizobium phaseoli (López-Guerrero et al. 2012). The separate clusters formed by the Mesorhizobium loti strains (#67 to 73) indicated that these strains, previously identified on the basis of their ability to nodulate Lotus spp., were likely to belong to several different Mesorhizobium species. In contrast to the Ensifer, Mesorhizobium and Rhizobium genera, spectra of RPs calculated for bradyrhizobia showed significantly lower levels of similarity. This was not surprising given the heterogeneity of bradyrhizobial strains (Sánchez-Juanes et al. 2013). Figure 1 also showed that in spite of only two isolates of distinct species selected to represent each of the Agrobacterium, Azorhizobium, Methylobacterium, and Microvirga genera, the clustering of theoretical RPs spectra was unambiguous. This suggested that the limited number of representative strains currently included in the rhizobiaspecific module for these four genera did not prevent accurate strain identification. These results indicated that together, masses of ribosomal proteins were potentially sufficient markers for generating strain-specific MALDI-TOF MS fingerprints.

Ascribing ribosomal proteins to masses found in empirical MALDI-TOF MS spectra

In routine procedures for strain identification, factors such as temperature, growth media, and conditions can influence the quality of MALDI-TOF MS spectra, and the number and intensity of peaks that are recorded (Wieme et al. 2014). Thus, to maximize the chances of recovering as many ribosomal proteins as possible, cell lysate samples were used to obtain at least four high-quality MALDI-TOF MS spectra for each of the 116 bacterial reference strains. All mass peaks between 4 and 30 kDa found in the empirical spectrum of a given strain were matched against the ribosomal protein masses that were deduced from the corresponding genome sequence. To account for possible experimental variations in measurements, an error interval of 800 ppm was tolerated, and only mass peaks successfully ascribed to a ribosomal protein and found in at least half of the empirical spectra produced for each strain were further considered. Depending on the genus to which individual strains belonged, the numbers of reproducible masses successfully ascribed to ribosomal proteins were as follows: Agrobacterium 25, Azorhizobium 22, Bradyrhizobium 17, Burkholderia 22, Cupriavidus 21, Ensifer 22, Mesorhizobium 15, Methylobacterium 25, Microvirga 25, and Rhizobium 22. Of these assigned RP masses, 13 were consistently identified in all spectra, regardless of the strain or genus considered (Fig. 2). This subset of reproducible masses corresponded to ribosomal subunits L20, L24, L28, L29, L31, L32, L33, L36, S10, S15, S16, S17, and S20 (see Table S2 in the Supplementary Material). As purified cell lysates were used to generate the empirical spectra used for ascribing ribosomal proteins to detectable masses, rather than isolated ribosomes, this experimental dataset was recorded in the PARPM database.

To assess whether this PARPM database was able to provide accurate identification of strains using standard empirical spectra as queries, four to five of the sinapinic acid-prepared spectra that had initially been collected from each the 116 reference strains were analyzed again for the presence or absence of mass peaks that corresponded to the selected subset of 13 RPs. In 539 of the 571 spectra (94 %) examined in this way, up to 13 RP markers were successfully assigned, with eight markers found to be the minimal threshold for obtaining an accurate identification at the species level.

Congruent topologies for 16S-, MLSA-, rMLSA, or PARPM-based phylogenetic trees

Ribosomal proteins have previously been shown to be reliable mass-spectra markers for phylogenetic classification of various bacteria (Suarez et al. 2013; Teramoto et al. 2007b). To

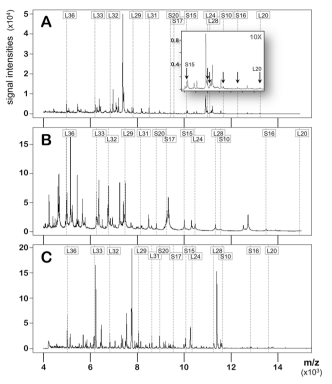


Fig. 2 Empirical spectra of strains *Bradyrhizobium* sp. WSM 2793 (panel **a**), *Ensifer fredii* NGR234 (**b**), and *Mircrovirga lotononidis* WSM3557^T (**c**) onto which the respective positions of the 13 ribosomal subunits (L36–L20 and 4–15 kDa in size) that were selected to construct the PARPM database are shown as annotated *dashed lines*. Note that although the intensities of peaks for several RPs appeared weak, signals were nonetheless recorded as protein hits in the automatic analysis mode. For example, low-intensity peaks matching ribosomal subunits are marked by *arrows* in the superimposed section of panel **a** in which signal intensities were magnified ten times ($10\times$)

test whether the subset of 13 RPs that comprised the PARPM database also provided a sufficient basis for phylogenetic analyses of the 116 selected strains of RNB and tumorigenic bacteria, we compared the topologies of trees reconstructed using the 16S rDNA sequences (16S rRNA); concatenated recA, gyrB, rpoB, dnaJ, and atpD sequences (MLSA); concatenated sequences coding for the ribosomal proteins rplT (L20), rplX (L24), rpmB (L28), rpmC (L29), rpmE (L31), rpmF (L32), rpmG (L33), rpmJ (L36), rpsJ (S10), rpsO (S15), rpsP (S16), rpsQ (S17), and rpsT (S20) (rMLSA); and the PARPM data (PARPM, see Fig. 3). As the 16S rRNA genes were missing from the draft genome sequences of R. leguminosarum bv. viciae strains 128c53, Ps8, and Vh3, as well as R. sullae WSM1592, the 16S rDNA sequences were amplified and deposited in GenBank (accession numbers KF678361-KF678364) before reconstructing the corresponding tree.

As shown in Fig. 3 and except for strain H152^T, the remaining 115 strains included in this analysis grouped into genera that were consistent with current rhizobial taxonomy. Depending on the data used for reconstructing the phylogenetic tree,

strain *R. giardinii* by giardinii strain H152^T clustered with either agrobacteria (16S rRNA) or members of the Ensifer genus (MLSA, rMLSA, and PARPM). Data presented in Fig. 3 illustrates the limited resolution obtained at the species level for trees reconstructed using DNA sequences. In contrast, the PARPM dataset offered an in-depth resolution including at the subspecies level and, in particular, for the agronomically important Bradyrhizobium, Ensifer, and Rhizobium genera. However, while the PARPM dataset consistently grouped the strains within genera and accurately reflected diversity and differences at the species and often subspecies level, it was not able to resolve rhizobial evolutionary relationships at higher taxonomic levels. Similarly, as genes that determine host range are part of the accessory genome, related strains of R. leguminosarum assigned to distinct symbiovars phaseoli, viciae, and trifolii were not resolved using PARPM (see positions of strains #85, 89, 99, and 104 in Fig. 3d).

Testing the rhizobia-specific module of the SARAMISTM database

The reliability and resolving power of the rhizobia-specific module was initially assessed on plate cultures of more than 100 unknown bacterial isolates obtained from root nodules of field-grown pigeon pea collected from Ivory Coast sites. According to MALDI-TOF MS results, most of the pigeon pea isolates clustered into two major subgroups, identified as either Bradyrhizobium elkanii or Bradyrhizobium spp. (data not shown). The identities of representative strains of both subgroups were later confirmed by 16S rRNA sequencing (see Table 2). Additionally, the symbiotic phenotypes of these strains were assayed on either the original host C. cajan or the more promiscuous legume V. unguiculata (Lewin et al. 1987). A similar procedure was carried out for a few isolates, such as strain CI-34D1, which were not recognized by MALDI-TOF MS as a potential RNB. In fact, strain CI-34D1 was identified by MALDI-TOF MS as a Brevibacillus sp. and 16S rDNA sequencing showed it was related to Brevibacillus reuszeri. As shown in Table 2, only the C. cajan isolates identified by MALDI-TOF MS as members of known rhizobial genera were found to nodulate and fix nitrogen with both pigeon pea and cowpea. As expected, the non-RNB strain CI-34D1 failed to nodulate either of the tested host plants. In further tests, endosymbiotic bacteria from nodules collected from roots of clover (Trifolium spp.) or French seradella (Ornithopus sativus) growing in Australian fields were analyzed directly by MALDI-TOF MS using the protocol established by Ziegler et al. (2012), which does not require isolation or cultivation of the RNB. Nodules were found to harbor bacteria identified by MALDI-TOF MS as either R. leguminosarum (clover symbionts) or Bradyrhizobium canariense (French seradella symbionts). To verify the accuracy of MALDI-TOF MS-based results, nodule bacteria were

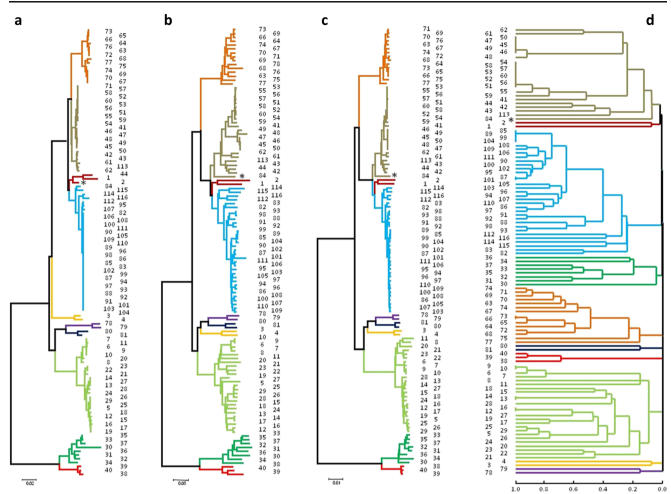


Fig. 3 Topology of trees obtained by cluster analysis of the **a** partial 16S rRNA sequences (1,006 nucleotides), **b** concatenated sequences of *recA*, *gyrB*, *rpoB*, *dnaJ*, and *atpD* genes (in total 9,531 bases), **c** concatenated sequences of the protein-coding *rplT*, *rplX*, *rpmB*, *rpmC*, *rpmE*, *rpmF*, *rpmG*, *rpmJ*, *rpsJ*, *rpsO*, *rpsP*, *rpsQ*, and *rpsT* genes (total of 2,991 nucleotides), or **d** binary matrix derived from PARPM database. Strains were numbered as in Table 1 and were listed next to each of the tree

isolated and the 16S rDNA sequences obtained for five (A-BRK13, A-G150, A-G169, A-Q1.7, and A-WB) out of the 41 isolates identified as *R. leguminosarum* and one (A-GQ1.6) of the four *B. canariense* strains (see Table 2). As all of these results corroborated the initial MALDI-TOF MS identifications, the rhizobia-specific module of SARAMISTM was found to correctly identify symbiotic bacteria whether grown on solid media or found inside nitrogen-fixing nodules.

Spectra that were generated for identifying bacteria isolated from Ivory Coast pigeon pea nodules were also matched against PARPM. Ultimately, all of the spectra of *C. cajan* isolates previously identified as members of the *B. elkanii* genus were found to best match the PARPM profiles of *B. elkanii* strains USDA76^T, USDA94, USDA3259, and USDA3254. In addition, the SA-prepared spectra of Rhizobia isolated from *Trifolium* sp. nodules were found to share 9 to 12 of the 13 RPs markers included in the PARPM accessions for

branches. To facilitate comparison of tree topologies, bacterial genera were color-coded as follows: *Agrobacterium (wine red), Azorhizobium* (orange), Bradyrhizobium (light green), Burkholderia (green), *Cupriavidus (red), Ensifer (olive), Mesorhizobium (brown), Methylobacterium (violet), Microvirga (dark blue),* and *Rhizobium* (light blue). Poorly resolved taxonomic position of *R. giardinii* bv. giardinii type strain H152^T marked by as asterisk

the reference *R. leguminosarum* bv. *trifolii* strains SRDI943 and WSM1325. These results indicated that identification of rhizobial isolates through a PARPM-based analysis was both feasible and reliable.

Discussion

This work highlights the robustness and accuracy of mass spectrometry for high-throughput identification of RNB and tumorigenic bacteria. In comparison to classical identification procedures, such as biochemical tests, restriction or amplified fragment length polymorphism (RFLP or AFLP), or sequencing of marker genes (e.g., 16S rRNA), MALDI-TOF MS is considerably faster, more cost-effective, and less labor-intensive. These advantages, along with the proven accuracy of MALDI-TOF MS-based identification of bacteria, have

Strain	Identification result as given b	Isolated from	Symbiotic phenotype on		
	MALDI-TOF MS	16S sequencing (closest related reference)		Pigeon pea	Cowpea
CI-36B	Bradyrhizobium elkanii	<i>B. elkanii</i> (USDA76 ^T /100 %)	Cajanus cajan	Fix+	Fix+
CI-1B	Bradyrhizobium sp.	B. elkanii (USDA76 ^T /99 %)	C. cajan	Fix+	Fix+
CI-34D1	Brevibacillus sp.	Brevibacillus reuszeri (DSM9887 ^T)	C. cajan	Nod-	Nod-
A-WB	Rhizobium leguminosarum	R. leguminosarum bv. trifolii (WSM1325)	Trifolium spumosum	nt	Nt
A-G150	R. leguminosarum	R. leguminosarum bv. trifolii (WSM1325)	T. subterraneum	nt	Nt
A-G169	R. leguminosarum	R. leguminosarum bv. trifolii (WU95)	T. subterraneum	nt	Nt
A-BRK13	R. leguminosarum	R. leguminosarum bv. trifolii (WSM1325)	T. spumosum	nt	Nt
A-Q1.7	R. leguminosarum	R. leguminosarum bv. trifolii (WSM1325)	Trifolium sp.	nt	Nt
A-Q1.6	Bradyrhizobium canariense	<i>B. canariense</i> (WSM471)	Ornithopus sativus	nt	Nt

 Table 2
 Characteristics of strains isolated from field nodules

Identity of selected nodule isolates was established using the Rhizobia-specific module of the upgraded SARAMIS[™] database (MALDI-TOF MS) or by matching the DNA sequence of the amplified 16S rRNA gene against the GenBank database (16S sequencing). Nodulation and nitrogen fixation properties of the isolated strains from Ivory Coast was examined using Magenta jars as described in the "Determining symbiotic properties of nodule isolates". Symbiotic proficiency of bacteria isolated from nodules of *Trifolium* spp. and *O. sativus* collected in Australia was not tested (nt) as no seeds of these particular legumes could be obtained

revolutionized the routine identification of microorganisms to species and subspecies level in clinical diagnostic and food microbiological laboratories (Croxatto et al. 2012; Mazzeo et al. 2006). MALDI-TOF MS has also been shown to be an excellent tool for identification of diverse rhizobial species of *Bradyrhizobium*, *Ensifer*, *Shinella*, and *Rhizobium* applicable to large populations of isolates in ecological and taxonomic studies (Ferreira et al. 2011; Sánchez-Juanes et al. 2013).

However, accurate identification of bacteria via MALDI-TOF MS depends upon the establishment of comprehensive spectral databases for a broad range of type or reference strains, and the currently available rhizobial databases are not representative of the phylogenetic diversity of RNB. We, therefore, developed a rhizobia-specific module for the SARA MISTM database that consisted of 114 fully sequenced and well-characterized strains belonging to nine of the most common genera of RNB, plus two agrobacterial accessions. It included 27 strains of R. leguminosarum bv. phaseoli, trifolii, or viciae and a number of strains of E. medicae and E. meliloti. Providing multiple strains for these species allowed us to analyze interstrain variability and thereby assess the effectiveness of MALDI-TOF MS for typing rhizobial strains, as has previously been shown for Neisseria meningitides isolates (Suarez et al. 2013). Our results showed that MALDI-TOF MS analysis using the rhizobia-specific database grouped RNB strains into genera and species congruent with the phylogenies obtained using marker gene sequences (16S rRNA, MLSA) and consistent with classifications previously obtained for cluster analysis of MALDI-TOF spectra of Ensifer and Rhizobium spp. (Ferreira et al. 2011). The database also provided an accurate and highly resolved analysis of strain diversity at the species and often subspecies level. This highlights the value of MALDI-TOF MS as a tool not only for identification, but also for microbial systematics (Welker and Moore 2011).

Using either pure plate cultures of bacterial isolates from Ivory Coast pigeon pea nodules or bacteria obtained directly from nodules of Trifolium spp. or French seradella, we then confirmed that the rhizobia-specific module could be used to accurately identify symbiotic rhizobia in field studies. When non-RNB were analyzed, the larger SARAMIS[™] bacterial database provided an identification that was in agreement with subsequent 16S rRNA analyses. This makes MALDI-TOF MS and the rhizobia-specific module an ideal replacement for the demanding and expensive sequencing of partial 16S rRNA or AFLP/RFLP analyses that have been performed to identify and determine the diversity of microsymbionts of wild legumes in recent large-scale studies (Aserse et al. 2013; Safronova et al. 2014; Yang et al. 2013; Zhao et al. 2014). In the economically important field of symbiotic nitrogen fixation in agricultural systems, knowledge of the biodiversity of rhizobia and of local populations is important for the design of successful inoculation strategies (Lindström et al. 2010). Identification of RNB to strain level offers the possibility of using MALDI-TOF MS in diagnostic tests, for example, in ecological studies to test the ability of inoculant strains to compete with naturalized soil populations of rhizobia (Denton et al. 2002).

As new legume symbionts are isolated, named and described, the rhizobia-specific module will need to be updated with additional reference strains, in particular, for genera that are poorly represented in this study (e.g., *Methylobacterium* and *Microvirga*) or not yet included (e.g., *Phyllobacterium*). Obtaining a pure copy of a reference strain for generating the corresponding MALDI-TOF mass spectra and superspectra is becoming ever more difficult due to increasing legal constraints that limit the transfer of microbiological material between countries. We, therefore, explored the possibility of using whole or partial rhizobial genome sequences as a basis for calculating protein masses that could be used as strainspecific biomarkers in MALDI-TOF mass spectra. These proteins needed to (i) be representative of the microorganism's core genome, (ii) fall within the mass range detected by available MALDI-TOF mass spectrometers, and (iii) constitute the most frequent mass peaks in empirical spectra. As proteins encoded by MLSA marker genes, such as *dnaJ*, *atpD*, *gyrB*, recA, and rpoB, had masses larger than 38 kDa (the molecular weight of RecA) and thus were too large for standard mass spectral analyses, we decided instead to focus on the RPs. A recent study found that a subset of biomarkers composed of 15 RPs and named as S10-spc-alpha operon gene-encoded ribosomal protein mass spectrum (SP10 GERMS) are highly conserved in eubacterial genomes and can discriminate bacteria at species and strain levels (Tamura et al. 2013).

By using the genome sequences of the 116 reference strains to deduce the masses of their RPs, then using hierarchical cluster analysis to search for similarities in the number of identical RP masses in pairwise combinations of strains (Fig. 1), we showed that RPs could be used as molecular markers to assign RNB to the genera or species to which they belonged. This confirms RPs as accurate markers for bacterial discrimination (Sato et al. 2012; Teramoto et al. 2007b) and, in a major finding, shows that they can be used to construct in silico reference MS spectra for strains whose genomes have been fully sequenced. When mass peaks found in empirical spectra of RNB were matched against genome-deduced masses of RPs, we identified subunits L20, L24, L28, L29, L31, L32, L33, L36, S10, S15, S16, S17, and S20 as the 13 most frequently ascribed ribosomal protein masses that together formed fingerprints that were specific for 89 of the 116 reference strains. This data was translated into a SARA MISTM-compatible format and stored into a specific database called PARPM. Querying the PARPM database with empirical mass spectra obtained using RNB grown on plates, or directly from nodules, resulted in correct identification of unknown isolates to the species level.

The rapid and constant evolution and revision of rhizobial taxonomy presents a challenge to the determination of microbial identity and phylogeny. Advances in genomic sequencing technologies and analyses now allow a much greater precision in the taxonomic placement of strains (Jolley et al. 2012; Kim et al. 2014; Klenk and Göker 2010). The increasing affordability of next-generation sequencing technologies is resulting in a vast increase in the number of sequenced bacterial genomes. However, it is of crucial importance that the sequenced strains are selected with care to reflect bacterial diversity. To this end, sequencing projects, such as the Genomic Encyclopedia of Bacteria and Archaea-type strains, ensure reproducibility of the results as well as provide a strong link between

genome data and biologically relevant features of each of the selected strains (Kyrpides et al. 2014b). The integration of sequencing technologies with MS databases has the potential to revolutionize rhizobial, or indeed microbial, diagnostics and systematics. Already more than 130 additional and complementing whole genomes of RNB have been accessed and stored as DRPM in the SARAMISTM database (data not shown). That type strains of novel species should have their genomes sequenced at the time they are deposited into culture collections as was recently proposed (Kyrpides et al. 2014a), will facilitate the development of this database.

Over the years, genome data has become essential for understanding the molecular basis of plant symbioses (Mora et al. 2014), and some of the mechanisms involved in the evolution and speciation of rhizobia (Tian et al. 2012). Our results now illustrate how such genomic datasets can become crucial elements for establishing new technological developments able to facilitate large-scale studies of rhizobial populations in fields and natural environments.

Acknowledgments This work was funded by the Swiss Commission for Technology and Innovation (CTI project 11225.1 PFLS_LS), the Swiss National Science Foundation (Grant Nos. 31003A-116591 and 31003A_146548), Mabritec, and Meat & Livestock Australia/ Australian Wool Innovation Grant B.PSP.0013. We are grateful to Susana Brom, Sofie Goormachtig, Eric Giraud, Alessio Mengoni, Philip Poole, José-Enrique Ruiz-Sainz, and Maged Saad for providing a number of reference strains, Damiana Ravasi and Paola Decristophoris for assistance with DNA sequencing of a number of marker genes, and Antoine Huyghe for his help in generating Fig. 1.

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