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TITLE PAGE

Combining ecophysiological and microbial ecological approaches to study the relationship between *Medicago truncatula* genotypes and their associated rhizosphere bacterial communities

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SUMMARY

Background and Aims: To assess how plant genotype and rhizosphere bacterial communities may interact, the genetic structure and diversity of bacterial communities in the rhizosphere soil of different *Medicago truncatula* genotypes were studied in relation to the plant carbon and nitrogen nutrition at the whole plant level.

Methods: The genetic structure and diversity of plant-associated rhizosphere bacterial communities was analysed by Automated Ribosomal Intergenic Spacer Analysis and 454-pyrosequencing. In parallel, the carbon and nitrogen nutrition of the plants was estimated by a phenotypic description at both structural level (growth) and functional level (using carbon and nitrogen isotope labeling and an ecophysiological framework).

Results: An early effect of the plant genotype was observed on the rhizosphere bacterial communities, while few significant differences were detected at the plant structural phenotypic level. However, at a functional level, the

different *Medicago truncatula* genotypes could be distinguished by their different nutritional strategies. Moreover, a comparison analysis showed that ecophysiological profiles of the different *Medicago truncatula* genotypes were correlated to the genetic structure and the diversity of the rhizosphere bacterial communities.

Conclusions: The exploration of the genetic structure and diversity of rhizosphere bacterial communities combined with an ecophysiological approach is an innovative way to progress in our knowledge of plant-microbe interactions in the rhizosphere.

Key words: bacterial communities; diversity; genetic structure; *Medicago truncatula*; nutrition; rhizosphere

INTRODUCTION

Rhizosphere microbial communities are able to interact with plants and, for some kind of interactions, play beneficial roles in plant health and growth (Buee et al. 2009; Franche et al. 2009; Harrison 1999; Raaijmakers et al. 2009; Richardson et al. 2009). Interactions between plants and microbes have been well documented (for reviews, see Buee et al. (2009); Harrison (1999); Hartmann et al. (2009); Jones et al. (2009a); Lambers et al. (2009)). As an example it has been demonstrated that plant genotype is one of the most important factors in the evolution of microbial communities (Buee et al. 2009). The influence of plant genotype is likely to occur through the release of rhizodeposits; significantly stronger rhizosphere effect towards bacterial communities was observed during the plant vegetative period than at later growth stages and can be ascribed to differential root exudation along the plant growth cycle (Mougel et al. 2006).

However, studies to date have avoided considering the bidirectional links between plants and microbial communities and focused either on the microbial or on the plant “side”. Consequently, a major objective is now to study the effect of the plant genotype on the shaping of the soil microbial communities, which could in return improve plant nutrient uptake. This challenge can now be attained by using novel and multidisciplinary approaches (Hartmann et al. 2009).

Recently developed microbiology techniques permit the characterization of microbial communities in complex soil environments (Ranjard et al. 2000; Singh et al. 2004). ARISA (for Automated Ribosomal Intergenic Spacer Analysis) was developed (Fisher and Triplett 1999), rapidly optimized and then validated as a sensitive and robust analysis method for studying soil microbial communities in both soil (Ranjard et al. 2003; Ranjard et al. 2001) and

the plant rhizosphere (Mougel et al. 2006). In addition, pyrosequencing, a high-throughput technique allowing the analysis of microbial diversity, was used to examine bacterial communities in soil samples (Roesch et al. 2007). However, as stated above, microbiological studies generally did not assess nutritional variations of the associated plants.

While most plants acquire soil mineral nitrogen (N), predominantly nitrate and to lesser extent ammonium, legumes also possess the capacity to acquire atmospheric dinitrogen (N₂), thanks to a symbiosis with bacteria belonging to the *Rhizobiaceae* family. Legumes contribute to diversifying cropping systems and decrease the need for external inputs (Jensen and Hauggaard-Nielsen 2003; Munier-Jolain and Carrouée 2003; Nemecek et al. 2008). Despite these advantages, legumes are still under exploited in European cropping systems mostly because of fluctuating yields resulting from abiotic and biotic constraints often affecting plant N nutrition. Plant N nutrition is known to interact closely with plant C nutrition (Marschner 1995; Minchin et al. 1981). Ecophysiological tools have been developed to understand the physiological bases involved in legume plant C and N nutrition (Moreau 2007; Moreau et al. 2012). However in these studies plants were grown in inert substrate inoculated with *Rhizobium* strain, the contribution of the entire rhizosphere microbial communities to plant nutrition was also not assessed.

The aim of our work was to assess plant genotype influence on the genetic structure of the rhizosphere bacterial communities in relation to the plant C and N nutritional strategies. The objective is to propose a global and multidisciplinary approach combining ecophysiology and microbial ecology to study plant-microbe interactions. A greenhouse experiment was conducted on seven genotypes of the legume plant model *Medicago truncatula*, which were contrasted in their genetic diversity (Ronfort et al. 2006). This study was confined to the vegetative period of *Medicago truncatula* during which roots and nodules are established and the rhizosphere effect towards bacterial communities is suspected to be stronger. On the one hand, microbiological tools were used to study the influence of the *Medicago truncatula* genotype on the genetic structure and diversity of the rhizosphere bacterial communities. On the other hand, the different nutritional strategies depending on the plant genotype were studied using an ecophysiological approach with three complementary approaches: a phenotypic growth description, C and N flux measurements using isotope labeling and an ecophysiological framework. Finally, the abundance of the rhizosphere bacterial communities was compared with the plant functional traits to bring some clues in the biological link between some phyla abundances and the plant nutritional strategies.

MATERIALS AND METHODS

Plant growth conditions and experimental design

The seven contrasted *Medicago truncatula* genotypes (DZA 045-6, DZA 315-16, DZA 315-26, F 83005-5, Jemalong A17, Jemalong J6 and SA 028064) used in this study have been chosen for their different geographic origins and genetic diversity (Ronfort et al. 2006). Seeds of *Medicago truncatula* genotypes were scarified, surface-sterilized (Mougel et al. 2006), vernalized at 4°C during 48h and germinated on 0.7% (w/v) water agar plates at 25°C in the dark. One germinated seed of each *Medicago truncatula* genotype was sown in pot containing 900 g of a silt-clay loam soil (Mas d'Imbert, France) which is poor in mineral N and rich in organic matter and hosts symbiotic microbes (Offre et al. 2007). This soil is characterized by low mineral nitrogen (N) content: 0.018 g.kg⁻¹ of NO₃-N and 0.002 g.kg⁻¹ of NH₄-N (NF ISO 14256-2). Physicochemical characteristics of this soil were determined as: 11% of sand, 51% of silt, 38% of clay, pH=8, 1.45% of organic carbon (NF ISO 10694), 0.1% of N (NF ISO 13878) and 0.1% of phosphorus (NF ISO 11885; inductively coupled plasma atomic emission spectroscopy (ICP-AES)).

Plants were cultivated up to the end of vegetative period at a density of 45 plants. m⁻² in a greenhouse, under a photoperiod of 15 h, a temperature of 26°/21°C (day/night). Supplementary artificial light was supplied with sodium lamps (MACS 400 W; Mazda, Dijon, France) to complement photosynthetically active radiation. In order to characterize the interactions between *Medicago truncatula* and the soil microbial communities in conditions where the plant can interact with N₂-fixing bacteria, plants were watered with a N-free solution (Moreau et al. 2008). An automatic watering scheme considering the weight of two control pots was used to maintain soil moisture at 45% of its maximum soil water-holding capacity and a manual control watering was also done each weeks.

At three sampling dates (340, 634 and 934 degree-days after sowing), representing three vegetative developmental stages (4 leaves, first ramification and second ramification) (Moreau et al. 2006; Mougel et al. 2006), four plants per genotype were harvested and the leaf area was measured (LI-3100 Area Meter, Li-Cor Inc., Lincoln, NE, USA); shoot and root biomass was determined after drying at 80°C during 48h. Data obtained for these three plant harvests were used to calculate integrative parameters. Only the third harvest data point (934 degree-days sowing) was used for both plant phenotypic and bacterial description, which with 33 ± 10 leaves per plant was considered as an early stage of plant development. Thus, at 934 degree-days after sowing, six plants per genotype were used for estimating C and N fluxes in plant. Among these 6 plants, three were used for ¹³C and ¹⁵N labeling while three other plants were taken as controls for non-labeled plants. Additionally, four of these six plants were randomly

selected for plant growth description (leaf area and dry shoot and below-ground biomass), calculation of integrative ecophysiological framework parameters and microbial description (Automated Ribosomal Intergenic Spacer Analysis and pyrosequencing analysis).

¹³C-labelling

Photosynthetic carbon (C) uptake was quantified with ¹³CO₂ (Jeudy et al. 2010; Voisin et al. 2003) at 934 degree-days after sowing. The labeling experiment was conducted in a gas-proof chamber as in Jeudy et al. (2010). The environmental parameters of this chamber were: a 14/10h day/night photoperiod, a temperature of 22°C, 60% relative-humidity, a photosynthetically active radiation of 650 μmol photons.m⁻².s⁻¹ (108 DULUX L 55 W lamps, APPRO5-21850, Saint Appolinaire, France). The atmospheric CO₂ concentration within the labeling chamber was measured and maintained at 380 μL L⁻¹ CO₂ (Jeudy et al. 2010). Plants were exposed for 10h to a ¹³C-enriched atmosphere (63.8% ¹³C/¹²C). Isotopic compositions of gases in the labeling chamber were controlled by a modular gas analyser (SIC MAHIAC S710, Hamburg, Germany) using non dispersive infrared.

After two days of chase, plants were separated into shoots and nodulated roots whose dry biomass was determined after oven drying at 80°C for 48h. Plant ¹³C content was analysed using a continuous-flow isotope ratio mass spectrometer (Sercon, Crewe, UK) coupled to a C-N elemental analyser (Thermo Electron NC2500, Courtaboeuf, France). The quantities of carbon (QC) in plant shoots and nodulated roots acquired during the labeling period were calculated as follows:

$$QC = DW \times \frac{\%C}{100} \times \frac{EC_{plant} - EC_{control}}{EC_{source} - EC_{control}} \quad (1)$$

where DW is the dry biomass, %C the percentage of carbon in dry biomass (w/w), EC_{plant} and EC_{control} are respectively the ¹³C abundance of the labeled plant or the non-labeled control plant and EC_{source} is the ¹³C enrichment of the atmosphere.

The rhizodeposition was quantified in the same way by analysing the quantity of ¹³C content in the rhizosphere soil. The labeling measurement was performed on three plants per genotype.

¹⁵N-labelling

Nitrogen (N) entering plants through symbiotic N₂ fixation was estimated using ¹⁵NO₃⁻ labeling and isotope dilution method (Salon et al. 2010). At sowing, 0.23 mg of K¹⁵NO₃ (10% ¹⁵N atom excess enrichment) was added in three pots without plant and in three pots for each sown genotypes. As control, 0.23 mg of K¹⁴NO₃ was supplied to three control pots without plant and for each genotype. In order to estimate mineralization, the decrease of ¹⁵N

of $^{15}\text{NO}_3^-$ was measured at 934 days after sowing using a diffusion method (Brooks et al. 1989; Mathieu 2005).

The symbiotic N_2 fixation (%Ndfa) was then estimated as follows:

$$\%Ndfa = 100 \times \frac{(E_{ref*} - E_{ref}) - (DW_{leg*} \times E_{leg*} - DW_{leg} \times E_{leg})}{(E_{ref*} - E_{ref}) - \varepsilon_{fix}} \quad (2)$$

where E_{ref*} and E_{ref} are the ^{15}N abundance of respectively labeled soil and nonlabeled control soil; DW_{leg*} and DW_{leg} are the dry biomass of respectively labeled plant and nonlabeled control plant; E_{leg*} and E_{leg} are the ^{15}N abundance of labeled plant and nonlabeled control plants respectively; $\varepsilon_{fix} = -1\delta\%$

Respiration

Respiration of root and microbial communities was measured at 934 degree-days in a growth chamber with similar environmental conditions as those used for the labeling experiment. The atmospheres of the root systems of each genotype (the three non-labeled plants per genotype), were individually separated from the shoot compartment using an air-tight plexiglass container. The respiration was then estimated by measuring the rate at which the CO_2 concentration within the container increased using an infrared gas analyzer (IRGA; Qubitac; Qubit Systems Inc., Kingston, Canada).

C and N nutritional strategies via an ecophysiological framework

The various plant genotypes could employ contrasted carbon (C) and nitrogen (N) nutritional strategies to reach similar growth. An ecophysiological framework (Moreau 2007; Moreau et al. 2012) was used to study these different C and N nutritional strategies. This framework linked four integrative variables (leaf area, total dry biomass, below-ground dry biomass and total amount of N in the plant) with four intermediate parameters:

i) RUE, which represents the Radiation Use Efficiency for biomass production. *RUE* was calculated as the slope of the linear function between total dry biomass (TDW) and the sum of intercepted PAR (Photosynthetically Active Radiation) ($\Sigma PAR_{intercepted}$)

$$TDW = \alpha + RUE \times \Sigma PAR_{intercepted} \quad (3) \quad \text{where} \quad \Sigma PAR_{intercepted} = \sum_{i=1}^D LA_i \times PAR_i \quad (4)$$

where *TDW* is the total dry biomass (g), α is the y-intercept, $\Sigma PAR_{intercepted}$ is the sum of PAR intercepted ($\text{MJ} \cdot \text{plant}^{-1}$), *D* is the day (day), LA_i is the leaf area at the day *i* ($\text{cm}^2 \cdot \text{plant}^{-1}$) and was calculated as reported by Chenu (2004), PAR_i is the incident PAR at the day *i* ($\text{MJ} \cdot \text{m}^{-2} \cdot \text{day}^{-1}$).

ii) RTR, which represents the Root:Total biomass Ratio. *RTR* was calculated as the slope of the linear function between below-ground dry biomass (RDW) and total dry biomass (TDW)

$$RDW = \alpha + RTR \times TDW \quad (5)$$

where *RDW* is the below-ground dry biomass (g) and *TDW* is the total dry biomass (g).

iii) SNU, which represents plant-Specific N Uptake. *SNU* was calculated as the slope of the linear function between total amount of N (N_{total}) and below-ground dry biomass (*RDW*)

$$N_{total} = \alpha + SNU \times RDW \quad (6)$$

where N_{total} is the total amount of N ($\text{g}\cdot\text{plant}^{-1}$) and *RDW* is the below-ground dry biomass (g).

iv) *NLA*, which represents the conversion factor of N to Leaf Area. *NLA* was calculated as the slope of the linear function between leaf area (*LA*) and total amount of N (N_{total}).

$$LA = \alpha + NLA \times N_{total} \quad (7)$$

where *LA* is the leaf area ($\text{cm}^2\cdot\text{plant}^{-1}$) and N_{total} is the total amount of N ($\text{g}\cdot\text{plant}^{-1}$).

Extraction and purification of total DNA from soil sample

DNA was extracted from two different compartments: rhizosphere soil and root tissue as previously described by Mougel et al. (2006). This sampling design enables to distinguish bacterial communities as a function of their proximity to plant influence (rhizosphere effect). The rhizosphere soil was obtained after manually separating the root system with adhering soil from the container and by washing the root system under agitation (vortex at 30 hertz during 1 minute) in 50 ml of sterile 0.025 M K_2SO_4 solution. The root system was discarded and the soil was collected after centrifugation at 9000 g for 10 min. The root tissue compartment included both bacterial communities in roots and on the root surface (rhizoplane). For the rhizosphere soil compartment and the root tissue compartment four replicates per genotype were used. Each sample was weighed, frozen in liquid nitrogen and stored at -80°C for further use in DNA extraction. The DNA extraction procedure was based on chemical and mechanical extraction as described previously by Ranjard et al. (2003) and by Mougel et al. (2006). DNA preparations were quantified as described previously (Mougel et al. 2006).

Automated RISAs fingerprinting

The bacterial ribosomal IGS were amplified with primers S-D-Bact-1522-b-S-20 (3' end of 16S genes) and L-D-Bact-132-a-A-18 (5' end of 23S genes) (Normand et al. 1996) for bacterial automated ribosomal intergenic spacer analysis (B-ARISA). PCR conditions, PCR template preparation for DNA sequencer loading and electrophoresis conditions were described by Ranjard et al. (2003).

454 sequencing and bioinformatics analysis

Rhizosphere bacterial diversity of the four *Medicago truncatula* genotypes was analysed using 454 sequencing. Genotype selection was based on results obtained on genetic structure of rhizosphere bacterial communities, where

three different genotype groups were identified. One representative genotype of each group (or two representative genotypes for the biggest group) was chosen (DZA 045-6, DZA 315-16, F 83005-5 and Jemalong A17).

A 16S rRNA gene fragment of the appropriate size and sequence variability for 454 pyrosequencing was amplified using the primers 530F (5'- ACTCCTACGGGAGGCAGCAG) (Acosta-Martinez et al. 2008) and 803R (5'- CTACCNGGGTATCTAAT-3'). Ten base pair tags ("Multiplex Identifier", MID) at 5' position were added to the primers to specifically identify each sample as recommended by the manufacturer (Beckman Coulter Genomics). For each rhizosphere soil analyzed, the DNA extracted from the four replicates was pooled to perform the PCR. DNA aliquots (2 ng) from each pool were used for a 50 µl PCR reaction under these conditions: 95°C for 3 min, 30 cycles of 1 min at 95°C (denaturation), 55°C for 30 s (annealing) and 72°C for 30 s (extension), followed by 5 min at 72°C. Adapter sequences were added to 300 ng per sample as recommended by the manufacturer and pyrosequencing was carried out on a Genome Sequencer FLX 454 (Beckman Coulter Genomics).

The bioinformatics analysis followed those developed by Terrat et al. (2011) corresponding to (i) the orientation for each kept read was checked, reverse-complemented where necessary (average read length: 400) and sorted according to the MID sequences, (ii) the reads were then filtered and eliminated if the exact primer sequences were not found, or if the sequences contained any ambiguity (Ns) or if its length was below 300 bases, (iii) a Python program was applied to obtain strict dereplication (i.e. clustering of strictly identical sequences). This led to a 30 to 40% reduction in the number of sequences. The dereplicated reads were then aligned and clustered into Operational Taxonomic Units (OTUs) defined by a 3% distance level using available tools from the Ribosomal Database Project (RDP) pyrosequencing pipeline (Cole et al. 2009). Then, another homemade filtering step was applied to potentially eliminate sources of errors (PCR chimeras and sequencing errors) based on OTUs definition. Briefly, for each sample, single-singletons (reads detected only once and not clustered) were deleted in order to eliminate these potential errors and to avoid an overestimated number of OTUs (deleted single-singletons represented only 7% of all reads but more than 30% of detected OTUs) (Behnke et al. 2011; Pawlowski et al. 2011). In fact, a majority of the single-singletons might be artifacts (PCR chimeras and large sequencing errors) resulting from noise in high-throughput pyrosequencing of the most abundant OTUs (Kunin et al. 2010; Reeder and Knight 2009). Subsequently, all high-quality sequence sets were another time aligned and clustered (at a 3% genetic distance) using tools from the RDP pyrosequencing pipeline (Cole et al. 2009). Rarefaction curves and diversity indices were finally determined as described previously by Terrat et al. (2011).

For taxonomy-based analysis, the Naïve Bayesian rRNA classifier of the RDP project was used (Cole et al. 2009) (the bootstrap value was set to 80%).

The sequences collected in this study have been deposited under accession number ERP000919 in the EMBL nucleotide sequence database.

Statistical analysis

All statistical analyses were performed using the statistical software package R 2.9.2 (R Development Core Team, 2009, Vienna, Austria). Only differences significant at $P < 0.05$ were considered.

Biomass, leaf area, photosynthesis rates, rhizodeposition, respiration, total amount of nitrogen (N) and symbiotic N_2 fixation were analysed using ANOVA tests. The means were compared using a Tukey-HSD post-hoc test.

Radiation use efficiency, root:total biomass ratio, specific N uptake and N to leaf area ratio were analysed using ANCOVA test, after a Bonferroni correction.

Two principal component analyses (PCA) were performed. The first one, which described the genetic structure of bacterial communities of the different *Medicago truncatula* genotypes (using ARISA data), was performed as reported by Ranjard et al. (2003) The second one described phenotypic characterization of the different genotypes of *Medicago truncatula*. A co-inertia analysis was used to correlate the genetic structure of the rhizosphere bacterial communities with the plant phenotypic characteristics. Co-inertia analysis is a standard multivariate analysis technique that describes the relationship between two data tables and serves as a general method to perform simultaneous analysis of two tables for studying species-environment relationship when the number of variables is higher than that of samples (Doledec and Chessel 1994; Dray et al. 2003). Monte-Carlo tests were used to check the significance of the PCA and the co-inertia. The two PCA, the co-inertia analysis and the Monte Carlo test were performed using the Ade4TkGUI software (Thioulouse and Dray 2007).

Co-inertia analysis allowed identifying the major ecophysiological variables and microbial markers that discriminate *Medicago truncatula* genotypes.

Pearson correlation coefficients were also calculated between the rhizosphere bacterial diversity (454 pyrosequencing data) of the four *Medicago truncatula* genotypes (DZA 045-6, DZA 315-16, F 83005-5 and Jemalong A17) and the ecophysiological variables.

Two dendrograms were drawn using the “ward” method of R statistical software package after calculating Euclidean distances. The first dendrogram of simple sequence repeat marker diversity was based on microsatellite data published by Ronfort et al. (2006). The second dendrogram, representing the genetic structure of rhizosphere bacterial communities of the different *Medicago truncatula* genotypes, was done by means of ARISA band intensity of the four replicates per genotype. A Mantel test was performed to test correlation between the two dendrograms obtained.

RESULTS

At an early stage of plant development, the *Medicago truncatula* genotype modifies the genetic structure of the bacterial communities in the rhizosphere soil but not in the root tissue

The genetic structure of bacterial communities in the rhizosphere soil compartment differed according to *Medicago truncatula* genotype (Fig. 1a). The first two components accounted for approximately 70% of the total variance. DZA 315-16 and DZA 315-26 were separated from the other genotypes on the first axis, which explained 43% of the total variance (Fig. 1a). DZA 045-6 was separated from the others on the second axis, which explained 27% of the total variance (Fig. 1a). In contrast, the genetic structure of the bacterial communities in the root tissue compartment (Fig. 1b) was not affected by the plant genotype.

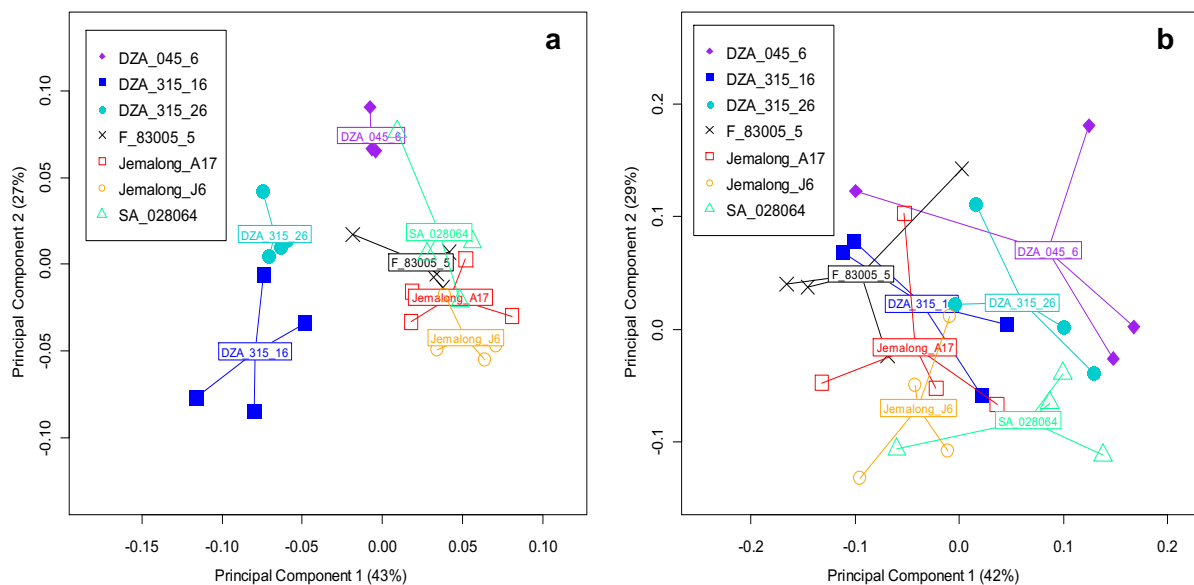


Fig. 1 Comparison of the genetic structure of bacterial communities by Principal Component Analysis (PCA) of bacterial automated ribosomal intergenic spacer analysis (B-ARISA) profiles from different compartments: **a.** rhizosphere soil and **b.** root tissue of different genotypes of *Medicago truncatula* at 934 degree-days. For the rhizosphere soil compartment ($P = 0.002$), **a.** represented the first two components combined account for approximately 70% of the total variance; For the root tissue compartment ($P = 0.06$), **b.** represented the first two components combined account for approximately 71% of the total variance

At an early stage of plant development, rhizosphere bacterial diversity is contrasted among *Medicago truncatula* genotypes

To precise bacterial community description, the rhizosphere bacterial diversity of four of the most contrasted *Medicago truncatula* genotypes (DZA 045-6, DZA 315-16, F 83005-5 and Jemalong A17) based on their rhizosphere bacterial genetic structures were analysed using 454 sequencing (Fig. 2). Diversity of the bacterial

communities in the root tissue compartment was not analysed because no significant effect of the *Medicago truncatula* was observed. Between 93 to 94 % of OTU were assigned to major bacterial phyla using our bioinformatic pipeline.

The rhizosphere bacterial diversity was contrasted among *Medicago truncatula* genotypes specifically between the proportion of *Proteobacteria*, *Acidobacteria* and *Bacteroidetes* phylum which represent between 81 to 83 % of the total OTUs (Fig. 2). The proportion of *Proteobacteria* varied among genotypes from 22% of the total sequenced bacteria for DZA 315-16 to 51% for Jemalong A17 (Fig. 2). Inside *Proteobacteria*, α -*Proteobacteria* that include the *Medicago truncatula* nitrogen fixing symbiots which are preferentially reproducing in association with *Medicago truncatula* host represented 55% of sequences among *Proteobacteria* (data not shown). The proportion of *Acidobacteria* varied among genotypes from 19% of the total sequenced bacteria for Jemalong A17 to 56% for DZA 315-16 (Fig. 2). The proportion of *Bacteroidetes* varied among genotypes from 3% of the total sequenced bacteria for DZA 315-16 to 11% for Jemalong A17 (Fig. 2).

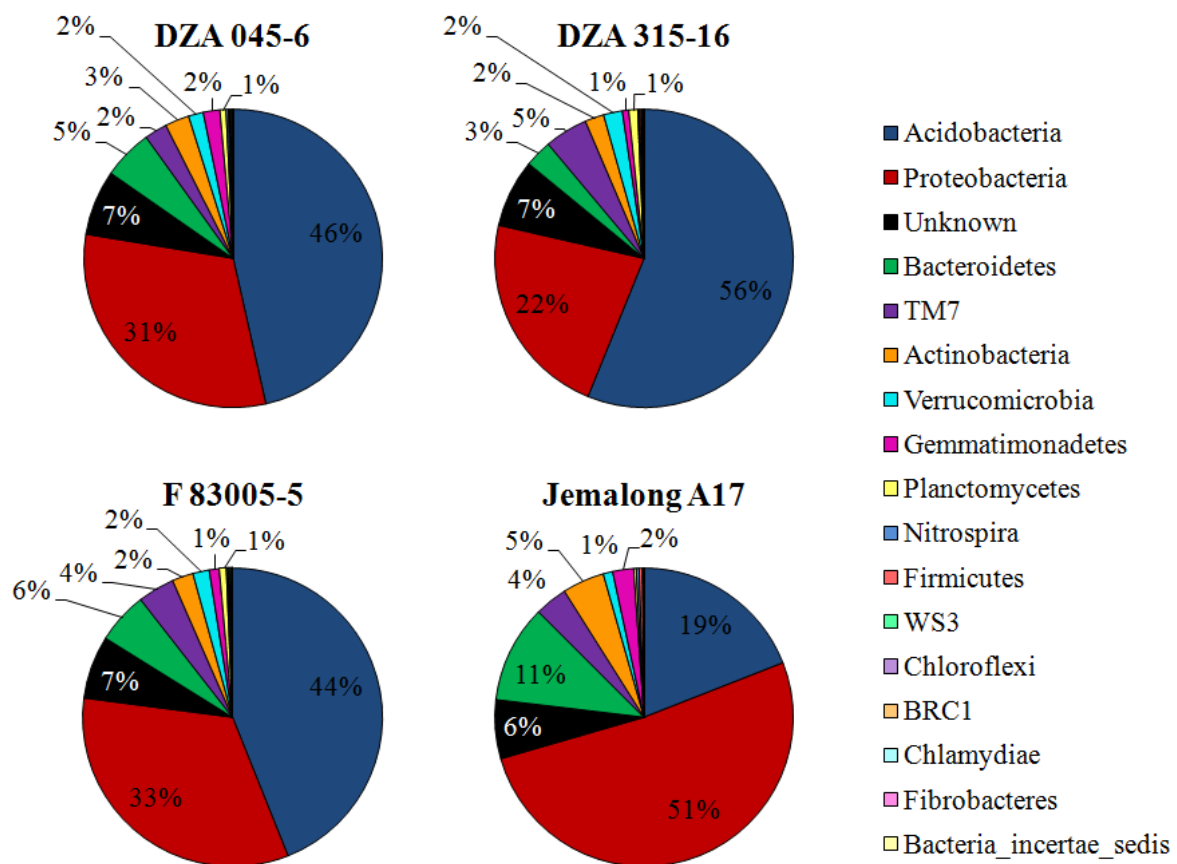


Fig. 2 Rhizosphere bacterial phyla of four *Medicago truncatula* genotypes (DZA 045-6, DZA 315-16, F 83005-5 and Jemalong A17) identified from 454 pyrosequencing

Therefore, the modifications of the genetic structure of the rhizosphere bacterial communities among the different genotypes of *Medicago truncatula* were representative of a contrasted rhizosphere bacterial diversity. As such, the *Medicago truncatula* genotype significantly affected the genetic structure and diversity of rhizosphere bacterial communities at an early stage of its development.

At an early stage of plant development, functional descriptors better discriminate *Medicago truncatula* genotypes than phenotypic structural descriptors

Plant growth was characterised with standard phenotypic structural descriptors such as total dry biomass (DW) and leaf area (Table 1). Responses ranged between 0.63 and 1.31 g for total DW and between 44 and 112 cm² for leaf area (Table 1). At 934 degree-days, the only difference in plant growth was observed for DZA 045-6, which was significantly smaller than DZA 315-16, F 83005-5 and Jemalong A17 (Table 1). These growth descriptors did not permit a distinction among the six other genotypes.

To go further than in previous microbiological studies, some plant functions involved in plant carbon (C) and nitrogen (N) nutrition were assessed using both an isotope labeling experiment and an ecophysiological framework (Moreau 2007). The isotope labeling enabled quantification of photosynthesis, rhizodeposition and symbiotic N₂ fixation (Table 1). The ecophysiological framework allowed the calculation of four intermediate parameters: the radiation use efficiency (RUE); the root:total biomass ratio (RTR); the specific nitrogen uptake (SNU) and the conversion factor of nitrogen to leaf area (NLA). The different genotypes were contrasted in their C and N nutrition. Jemalong A17 had higher RUE than DZA 315-16, DZA 315-26 and F 83005-5 (Table 1). DZA 045-6 had a higher photosynthesis than SA 028064 (Table 1). Rhizodeposition did not significantly vary with *Medicago truncatula* genotype, but DZA 315-16 and its microbial communities had a higher respiration than Jemalong A17 and Jemalong J6 (Table 1). The SNU and the root:total biomass ratio of DZA 315-16 were respectively higher and lower than for Jemalong J6 (Table 1). Symbiotic N₂ fixation and NLA did not differ among plant genotypes (Table 1).

Thus, phenotypic structural descriptors (dry biomass and leaf area) significantly differentiated DZA 045-6 from DZA 315-16, F 83005-5 and Jemalong A17. A detailed level of discrimination was achieved with functional descriptors (RUE, SNU and respiration) that significantly differentiated DZA 315-16 from Jemalong A17 and Jemalong J6.

Table 1 Developmental and physiological characterizations of the different genotypes of *Medicago truncatula*

Means were classified using a Tukey test ($P < 0.05$) and different letters indicate statistically different means among genotypes

RUE represents the radiation use efficiency and RUE is the correlation coefficient of the function between total biomass (g plant^{-1}) and the sum of intercepted PAR (MJ plant^{-1}); Root:total biomass ratio is the correlation coefficient of the function between below-ground biomass (g plant^{-1}) and total biomass (g plant^{-1}); SNU represents the specific nitrogen uptake and SNU is the correlation coefficient of the function between the total amount of nitrogen (g plant^{-1}) and below-ground biomass (g plant^{-1}); NLA represents the conversion factor of nitrogen to leaf area and NLA is the correlation coefficient of the function between leaf area ($\text{cm}^2 \text{ plant}^{-1}$) and the total amount of nitrogen (g plant^{-1})

RUE, root:total ratio, SNU and NLA were calculated from three dates (340, 634 and 934 degree-days)

Total biomass, leaf area, photosynthesis, rhizodeposition, respiration, total amount of nitrogen and symbiotic fixation were measured at 934 degree-days

Genotype	Total biomass (g)	Root:total biomass ratio	Leaf area (cm^2)	RUE (g of total biomass MJ^{-1} of intercepted PAR)	Photosynthesis ($\mu\text{g C}\cdot\text{s}^{-1}\cdot\text{cm}^{-2}$ of leaf area)	Rhizodeposition ($\mu\text{g C}\cdot\text{s}^{-1}\cdot\text{g}^{-1}$ of root)	Respiration ($\mu\text{g of CO}_2\cdot\text{s}^{-1}\cdot\text{g}^{-1}$ of root)	Total amount of N ($\text{mg}\cdot\text{plant}^{-1}$)	%Ndfa	SNU (mg of N g^{-1} of below-ground biomass)	NLA (cm^2 of leaves g^{-1} of N)
DZA 045-6	0.63 b	0.32 ab	44 b	3.84 ab	0.014 a	0.018 a	1.59 ab	18 a	76.8 a	86.1 ab	2216 a
DZA 315-16	1.31 a	0.28 b	112 a	3.52 b	0.010 ab	0.022 a	3.69 a	33 a	85.0 a	85.7 a	3186 a
DZA 315-26	0.92 ab	0.33 ab	84 a	3.14 b	0.010 ab	0.024 a	1.67 ab	24 a	70.7 a	78.4 ab	3137 a
F 83005-5	1.13 a	0.30 ab	95 a	3.29 b	0.010 ab	0.018 a	2.42 ab	30 a	74.0 a	79.0 ab	2990 a
Jemalong A17	1.28 a	0.34 ab	89 a	4.65 a	0.012 ab	0.020 a	0.97 b	31 a	64.6 a	69.5 ab	2699 a
Jemalong J6	1.08 ab	0.35 a	78 ab	3.92 ab	0.013 ab	0.021 a	1.10 b	24 a	77.3 a	60.8 b	2972 a
SA 028064	1.07 ab	0.28 ab	76 ab	3.98 ab	0.008 b	0.011 a	1.25 ab	25 a	76.2 a	78.0 ab	3004 a

Microbiological and ecophysiological data are correlated

In order to correlate microbiological and ecophysiological data, a co-inertia analysis was carried out by comparing the PCA of the genetic structure of the rhizosphere microbial communities (ARISA data) with a PCA analysis of the ecophysiological data (Online Resource 1). The co-inertia indicated that the differences observed between plant genotypes in both genetic structure of associated bacterial communities and ecophysiological profiles were correlated ($P = 0.026$) (Fig. 3). The co-inertia represented the first two components which accounted for approximately 71% of the total variance. The co-inertia analyses allowed to distinguish three different groups of genotypes: i) DZA 045-6; ii) DZA 315-16, and iii) Jemalong A17 and J6 (Fig. 3a, b).

Microbial molecular markers that mostly discriminated the three different groups of genotypes were ranged from 286 base pairs (bp) to 712 bp (Fig. 3): X692, X640, X670, X398, X712 and X362 on the first principal component; and X286, X640, X692, X670, X562 X392 and X394 on the second principal component. Ecophysiological variables that mostly discriminated the three different groups of genotypes were respiration, N fixation, leaf area, RUE and NLA on the first principal component and total biomass, leaf area, N total content and RUE on the second principal component (Fig. 3).

DZA 315-16 was separated from DZA 045-6, Jemalong A17 and J6 on the first axis, which explained 38% of the total variability (Fig. 3a, b). This difference could be explained, at the bacterial level, by two molecular markers of 692-693 bp and 712-713 bp, which were present for DZA 315-16 but not for DZA 045-6, Jemalong A17 and Jemalong J6 and by two molecular markers of 362-363 bp, 398-399 bp, which had a lower intensity for DZA 315-16 than for DZA 045-6, Jemalong A17 and J6 (Fig. 3c). At the plant ecophysiological level, DZA 315-16 was related to a high leaf area, a high respiration and a high NLA contrary to DZA 045-6, Jemalong A17 and J6 (Fig. 3d).

DZA 045-6 was separated from DZA 315-16, Jemalong A17 and J6 on the second axis, which explained 33% of the total variability (Fig. 3a, b). This difference could be explained, at the bacterial level, by both two molecular markers of 286-287 bp and 392-395 bp, which had a higher intensity for DZA 045-6 than for DZA 315-16, Jemalong A17 and Jemalong J6, and one molecular marker of 562-563 bp, which had a lower intensity for DZA 045-6 than for DZA 315-16, Jemalong A17 and Jemalong J6 (Fig. 3c). At the plant physiological level, DZA 045-6 was related to a low total biomass, a low leaf area and a low total amount of N, contrary to DZA 315-16, Jemalong A17 and Jemalong J6 (Fig. 3d).

DZA 045-6 and DZA 315-16 were separated from Jemalong A17 and J6 on both the first and second axis (Fig. 3a, b). This difference could be explained, at the bacterial level, by two other molecular markers of 640-641 bp and

670-671 bp, which had a higher intensity for Jemalong A17 and Jemalong J6 than for DZA 045-6 and DZA 315-16 (Fig. 3c). At the plant ecophysiological level, DZA 045-6 and DZA 315-16 were related to a higher SNU and N fixation and a lower RUE than for Jemalong A17 and J6 (Fig. 3d).

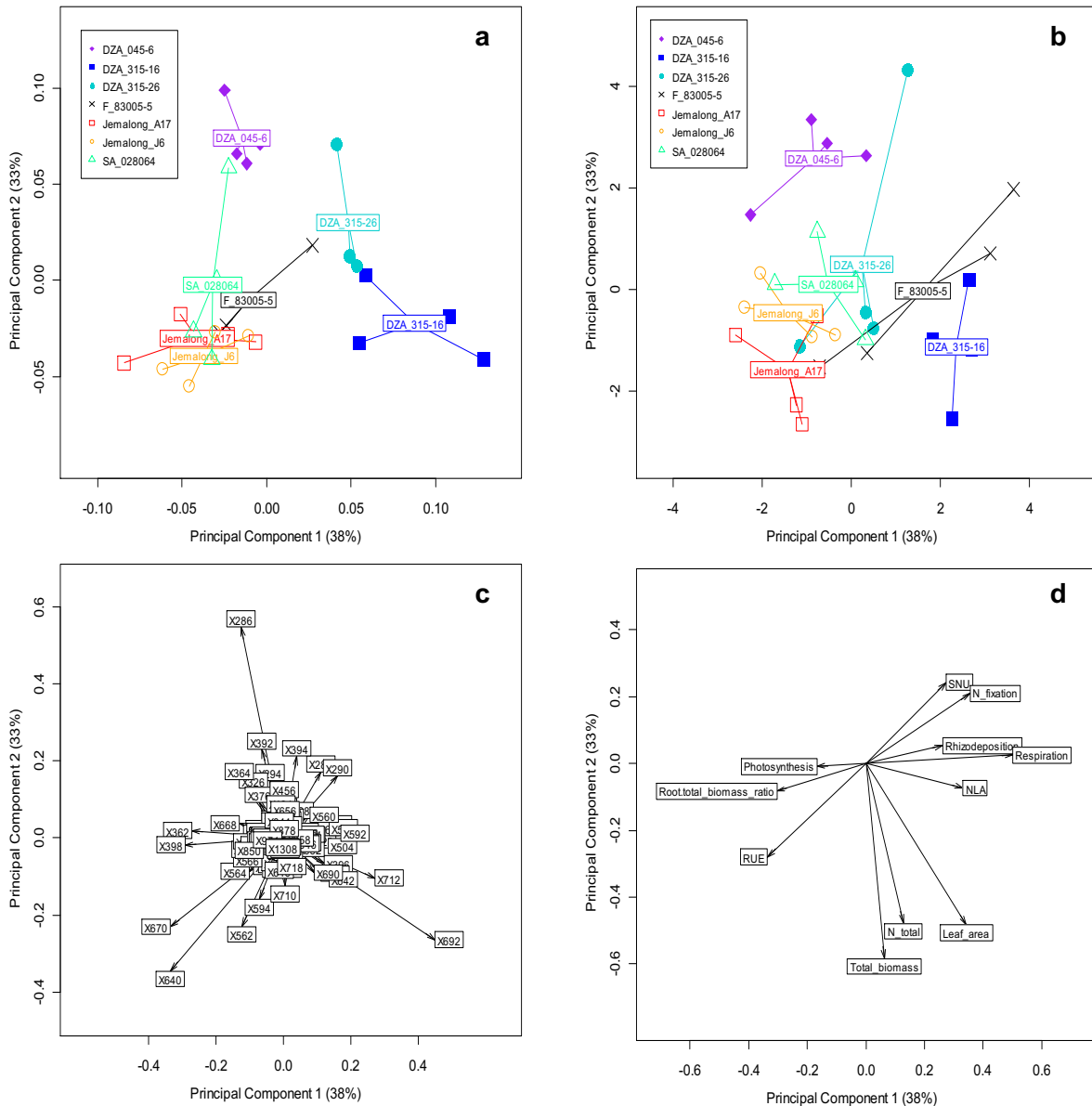


Fig. 3 Comparison of the ecophysiological data with the genetic structure of rhizosphere bacterial communities (B-ARISA) by co-inertia for different genotypes of *Medicago truncatula* at 934 degree-days ($P = 0.026$). **a.** Factorial plane of the genotypes discrimination in the co-inertia as a function of their B-ARISA profile ; **b.** Factorial plane of the genotypes discrimination in the co-inertia as a function of their ecophysiological profile ; **c.** Factorial plane of B-ARISA bands contribution in the co-inertia ; each number represented size of B-ARISA band (or DNA fragment) (e.g. X286 represent 286-287 base pair DNA fragments) ; **d.** Factorial plane of ecophysiological characteristics contribution in the co-inertia. The first two components of PCA represented account for approximately 71% of the total variance

The Pearson correlation coefficients between rhizosphere bacterial diversity data (454 pyrosequencing) and ecophysiological variable for the four *Medicago truncatula* genotypes (DZA 045-6, DZA315-16, F 83005-5 and

Jemalong A17) were calculated (Table 2). While *Acidobacteria*, *Verrucomicrobia*, *Planctomycetes* and *BCR1* abundances were positively correlated to N fixation *Proteobacteria*, *Bacteroidetes*, and *Actinobacteria* abundances negatively correlated to N fixation (Table 2). *Verrucomicrobia* abundance was negatively correlated to RTR while *Gemmatimonadetes* and *Nitrospira* abundances were positively correlated to RTR (Table 2). *Actinobacteria* and *WS3* abundances were respectively positively and negatively correlated to RUE; *WS3* abundance was positively correlated to SNU; *TM7* abundance was positively correlated to NLA; *Gemmatimonadetes* and *Fibrobacteres* abundances were negatively correlated to respiration (Table 2). *TM7* abundance was positively correlated to leaf area and *Bacteria incertae sedis* and *OP10* abundances were negatively correlated to total biomass and N total (Table 2).

Microbiological and genetic data are correlated

A comparison between two dendrograms, one based on bacterial ARISA data and one based on 13 *M. truncatula* microsatellites showed a highly significant correlation between the genetic structure of the rhizosphere bacterial communities and the genetic diversity of *Medicago truncatula* genotypes ($P < 0.001$) (Fig. 4).

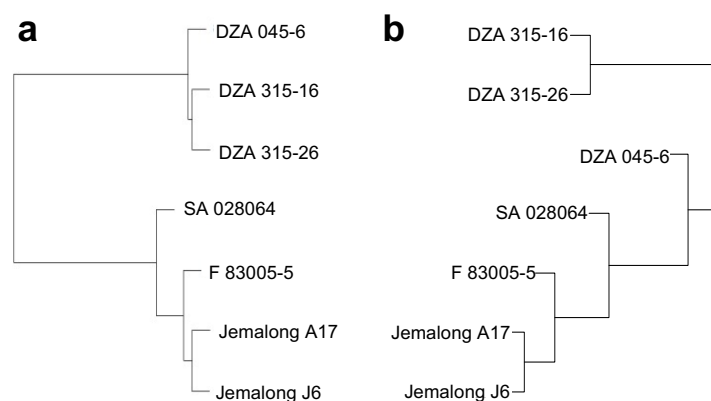


Fig. 4 Dendrogram comparison between the *Medicago truncatula* phylogenetic structure (a) and the genetic structure of their associated rhizosphere bacterial communities (b). Phylogenetic dendrogram of the seven *Medicago truncatula* genotypes based on simple sequence repeat marker diversity (modified from Ronfort *et al.* 2006). Dendrogram of the genetic structure of the *Medicago truncatula* rhizosphere bacterial communities based on the bacterial ribosomal intergenic spacer data.

A Mantel test showed a significant correlation ($P < 0.001$) between these two dendrograms

Table 2 Pearson correlation coefficients between rhizosphere bacterial diversity (454 pyrosequencing data) and ecophysiological variables

Only significant correlations at $P < 0.05$ ranged from -1 to -0.95 and from 0.95 to 1 were presented; ns: not significant

* The minimum – maximum interval is given based on the phylum abundance of the four different *Medicago truncatula* genotypes

Leaf area ($\text{cm}^2 \text{ plant}^{-1}$), total biomass (g plant^{-1}), total amount of nitrogen (N total) (mg plant^{-1}), symbiotic N fixation (%Ndfa), respiration ($\mu\text{g of CO}_2 \text{ s}^{-1} \text{ g}^{-1}$ of root), rhizodeposition ($\mu\text{g C s}^{-1} \text{ g}^{-1}$ of root) and photosynthesis ($\mu\text{g C s}^{-1} \text{ cm}^{-2}$ of leaf area) were measured at 934 degree-days. RUE represents the radiation use efficiency (g of total biomass MJ^{-1} of intercepted PAR), SNU represents the specific nitrogen uptake (mg of N g^{-1} of below-ground biomass), NLA represents the conversion factor of nitrogen to leaf area (cm^2 of leaves g^{-1} of N) and RTR represents the root:total biomass ratio.

	Phylum relative abundance (min – max)*	Leaf area	Total biomass	N total	RUE	N fixation	SNU	Respiration	NLA	RTR	Rhizodeposition	Photosynthesis
<i>Acidobacteria</i>	19.0 – 56.1 %	ns	ns	ns	ns	0.98	ns	ns	ns	ns	ns	ns
<i>Proteobacteria</i>	22.4 – 51.5 %	ns	ns	ns	ns	-0.97	ns	ns	ns	ns	ns	ns
<i>Bacteroidetes</i>	2.9 – 10.6 %	ns	ns	ns	ns	-0.97	ns	ns	ns	ns	ns	ns
<i>TM7</i>	2.5 – 4.7 %	0.98	ns	ns	ns	ns	ns	ns	0.96	ns	ns	ns
<i>Actinobacteria</i>	2.1 - 4.6 %	ns	ns	ns	0.98	-1.00	ns	ns	ns	ns	ns	ns
<i>Verrucomicrobia</i>	1.0 2.0 %	ns	ns	ns	ns	0.97	ns	ns	ns	-0.96	ns	ns
<i>Gemmatimonadetes</i>	0.7 – 2.3 %	ns	ns	ns	ns	ns	ns	-0.98	ns	0.96	ns	ns
<i>Planctomycetes</i>	0.3 – 1.0 %	ns	ns	ns	ns	0.97	ns	ns	ns	ns	ns	ns
<i>Nitrospira</i>	0.2 – 0.3 %	ns	ns	ns	ns	ns	ns	ns	ns	0.99	ns	ns
<i>Firmicutes</i>	0.1 – 0.4 %	ns	ns	ns	ns	ns	ns	ns	ns	ns	ns	ns
<i>WS3</i>	0.1 – 0.2 %	ns	ns	ns	-0.95	ns	1.00	ns	ns	ns	ns	ns
<i>Chloroflexi</i>	< 0.14 %	ns	ns	ns	ns	ns	ns	ns	ns	ns	ns	ns
<i>BRC1</i>	< 0.11 %	ns	ns	ns	ns	0.95	ns	ns	ns	ns	ns	ns
<i>Chlamydiae</i>	< 0.05 %	ns	ns	ns	ns	ns	ns	ns	ns	ns	ns	ns
<i>Fibrobacteres</i>	< 0.03 %	ns	ns	ns	ns	ns	ns	-0.99	ns	ns	ns	ns
<i>Bacteria incertae sedis</i>	< 0.02 %	ns	-0.97	-0.97	ns	ns	ns	ns	ns	ns	ns	ns
<i>OP10</i>	< 0.01 %	ns	-0.97	-0.97	ns	ns	ns	ns	ns	ns	ns	ns
<i>Unknown</i>	6.4 – 7.4 %	ns	ns	ns	ns	ns	ns	ns	ns	ns	ns	ns

DISCUSSION

The interactions between plants and soil microbial communities are bidirectional: on the one side, plant directs the selection of microbes via rhizodeposition (Hartmann et al. 2009) and on the other side, microbes can influence plant health, development and growth, for example, by enhancing nutrient availability and uptake (Richardson et al. 2009). Nevertheless, plant-microbe interactions are usually studied focusing either on the plant or on the microbe side. To our knowledge, our study is the first one assessing the plant genotype's effect on the genetic structure of the entire rhizospheric and endophytic bacterial community in relation with plant growth and carbon (C) and nitrogen (N) nutrition at the whole plant level.

The plant genotype has an early effect on the rhizosphere bacterial communities

The rhizosphere harbours multiple bacterial communities. Nevertheless, in the literature, plant genotype effects were particularly studied on specific rhizobacterial communities such as plant growth-promoting bacteria (Dalmastrì et al. 1999; Mazzola et al. 2004; Paffetti et al. 1996). Our data confirmed the importance of the plant genotype in shaping the genetic structure and diversity of the entire bacterial rhizosphere for the model legume *Medicago truncatula*. Although *Medicago truncatula* is able to fix atmospheric N₂, this effect was not limited to only rhizobial strains. Besides, no significant difference was observed in nodulation intensity among *Medicago truncatula* genotypes (Online resource 2). To date, the influence of plant genotype on the entire rhizosphere bacterial communities has only been demonstrated for three genotypes of *Solanum tuberosum* L. (Andreote et al. 2009; van Overbeek and van Elsas 2008) and for eight accessions of *Arabidopsis thaliana* (Micallef et al. 2009). Furthermore, the effect of the *Medicago truncatula* genotype on bacterial communities is seen early during the growth cycle. This is in agreement with two previous studies, one on canola at the rosette stage (Dunfield and Germida 2003), the other on three different potato genotypes at the first growth stage (Andreote et al. 2009). Altogether, these studies support the idea of the high sensitivity of bacterial communities to intra-species plant genotype variation. Studies on plant-bacterial community interactions should focus on early stages of plant development.

At an early stage of development, functional descriptors are essential to discriminate plant genotypes

Few microbiological studies have assessed plant phenotypic variations in relation to the modifications of rhizosphere microbial communities. The observed variations were generally based on structural descriptors (Mougel et al. 2006; Parra-Colmenares and Kahn 2005). In our study, structural phenotypic descriptors such as biomass and leaf area only discriminated DZA045-6 from the other genotypes.

To thoroughly characterize the different *Medicago truncatula* genotypes in relation with their nutritional strategies, an ecophysiological framework with integrative variables (Moreau 2007) was used. Our study enabled us to identify three contrasted nutritional strategies. The first one was illustrated by DZA 045-6, which had a lower biomass than the other genotypes. This can be explained by the low conversion of N to leaf area of DZA 045-6, as leaf area is instrumental to acquire carbon resources and thus biomass. The second nutritional strategy was that of DZA 315-16. This genotype allocated more biomass to its shoots which displayed the least C uptake efficiency among the studied genotypes. The third strategy, illustrated by Jemalong A17 and Jemalong J6, was to allocate more biomass to their below-ground parts, which displayed the least specific N uptake efficiency among the studied genotypes. As such, these differences in biomass allocation by the various genotypes seem to reflect a compensation process, termed the “functional equilibrium” theory (Brouwer 1983; Farrar and Jones 2000). The dry biomass distribution between root and shoot can be regulated by equilibrium between root activity (water and nutrient absorption) and shoot activity (photosynthesis). The PCA of *Medicago truncatula* ecophysiological profiles (Online Resource 1) illustrate this theory well, as RUE is positively correlated with root:total biomass ratio and is negatively correlated with SNU. Such contrasted nutritional strategies have been identified in several species during their vegetative growth (Lemaire et al. 2008; Moreau 2007) but, to our knowledge, they have not been studied in relation to the rhizosphere microbial communities.

Thus, at an early stage of the plant development, while plant structural descriptors did not reveal effects of plant-microbe interactions on the plant, functional descriptors could discriminate plant genotypes.

Rhizosphere bacterial diversity could be controlled by labile carbon availability and rhizosphere CO₂ concentration

By traditional culture methods, 95-99% of the microbial communities present in the soil are not accessible. Thanks to the development of culture-independent approaches, the most dominant bacterial phyla present in the rhizosphere soil could be identified: *Proteobacteria*, *Actinobacteria*, *Acidobacteria*, *Verrucomicrobia*, *Planctomycetes*, *Bacteroidetes* and *Firmicutes* (da Rocha et al. 2009). In the rhizosphere of *Medicago truncatula*, the two most dominant bacterial phyla were *Proteobacteria* and *Acidobacteria* (Fig. 2). This result is in agreement with a meta-analysis performed by Fierer et al. (2007) showing the high abundance of *Proteobacteria* (50-60%) and *Acidobacteria* (10-15%) in the rhizosphere. More interestingly, in our study, we found a high *Acidobacteria* abundance (between 19 to 56%) and a modified *Acidobacteria:Proteobacteria* ratio between DZA315-16, DZA045-6, F83005-5 and Jemalong A17 (Fig. 2). If *Proteobacteria* phylum was successful rhizosphere colonizers, the high abundance of *Acidobacteria* is more surprising. Two major hypotheses could be used to

explain these results in relation to the ecological life strategies of *Proteobacteria* and *Acidobacteria* and to the plant ecophysiological parameters.

Acidobacteria is considered as an oligotrophic phylum (K-strategists) whereas *Proteobacteria* is considered as a copiotrophic phylum (r-strategists). Ecological attributes of copiotrophs were their greater ability to consume labile organic C pools and their high growth rates in non limiting resource conditions. In contrast, oligotrophs exhibit slower growth rate and are suppose to outcompete copiotrophs in low nutrient availability conditions due to a higher substrate affinities (Fierer et al. 2007). *Acidobacteria* are composed mostly of non-cultivable bacteria, which may have become adapted to elevated concentrations of CO₂ and concentrations of O₂ lower than the atmospheric O₂ concentrations (Stevenson et al. 2004) and their abundance and diversity were strongly positively correlated with pH (Jones et al. 2009b). The rhizosphere environment has generally a lower pH compare to soil, lower oxygen and higher carbon dioxide concentrations and is a rich nutrient environment with a great quantity of labile organic C.

Among ecophysiological variables, radiation use efficiency (RUE) was significantly higher in Jemalong A17. The RUE, which corresponds to the efficiency of conversion of atmospheric CO₂ in plant biomass could be directly correlate to the quantity of rhizodeposits. In our study the rhizodeposition was not different among *Medicago truncatula* genotypes but we did not measured the quantity of labile carbon which could affect *Acidobacteria:Proteobacteria* ratio in the soil. Indeed, according to Fierer et al. (2007), contrary to *Proteobacteria*, *Acidobacteria* abundance is negatively correlated to carbon mineralisation rates. This point could explain the high abundance of the r-strategists *Proteobacteria* phylum compare to *Acidobacteria* in the rhizosphere of Jemalong A17. Rhizosphere respiration, corresponding to root and microbial respiration, was four-fold higher in the rhizosphere of DZA 315-16 but was not directly correlated to *Acidobacteria* phylum (Table 2). The N fixation was positively correlated to *Acidobacteria* phylum abundance (Table 2). A stronger rhizosphere respiration and nodule activity could explain a more important CO₂ concentration in the rhizosphere of DZA 315-16 which would positively select *Acidobacteria*. Altogether, the abundance and modification of the ratio between these two major phyla could be the mirror of modifications of labile carbon availability, CO₂ concentration and pH in the rhizosphere.

Quantitative rhizodeposition analyse is not sufficient to understand the basis of plant-microbe interactions

The total amount of plant C deposited in the rhizosphere is known to greatly vary with the plant ecophysiology, as determined by environment, genetics and physiology (Nguyen 2003). Micallef et al. (2009) explained the different bacterial communities present in the plant rhizosphere by quantitative and qualitative differences in the exudation

pattern of the different *Arabidopsis thaliana* accessions. In our study, no significant difference could be observed for the total amount of C lost by plants via rhizodeposition. Although rhizodeposition was correlated with a microbial marker (X562), the bacterial community diversity represented by this marker is unknown. The dissection of taxonomic structures and the metabolic/ecological functions of the microbial communities could be assessed using metagenomics approaches. However, this result does not exclude quantitative difference in root exudation or qualitative differences in the signalling pathways among plant genotypes as it was shown for the legume–Rhizobia molecular dialogue (Bais et al. 2006; Oldroyd and Downie 2008). To understand further the basis of plant-microbe interactions, qualitative rhizodeposition analysis should be promoted instead of quantitative analysis.

A global and multidisciplinary approach including ecophysiology, microbial ecology and quantitative genetics is a key tool in deciphering plant-microbe interactions

In our study, ecophysiology of the different *Medicago truncatula* genotypes was assessed using two complementary levels of discrimination: the first based on phenotypic structural descriptors; the second based on functional descriptors. Ecophysiological profiles of the different *Medicago truncatula* genotypes were correlated to the genetic structure of the rhizosphere bacterial communities. Nevertheless, microbial approach used (ARISA) did not permit to assess diversity and functions of the microbial communities. A direct causal link between the assembly of the rhizosphere bacterial and plant physiological functions could not be ascertained in the present study. Establishing this connection will be a challenge for the future.

Our study highlighted that *Medicago truncatula* plant genotypes-associated microbial communities are diverse and potentially under selection by their host plant. In our study few bacterial phyla (*Proteobacteria* and *Acidobacteria*) show their abundance modified in relation with plant functional traits. At this stage we need to generalize this result in order to define diversity profile of bacteria associated with root in relation with plant and microbial traits. High-throughput sequencing technology will facilitate surveys of taxonomic loci as well as loci implicated in impacting plant phenotypes (e.g., nitrogen fixation, root architecture, leaf area,...). Our study demonstrated the importance today to have a multidisciplinary approach in plant-microbe interactions. In addition, the relationship between plant functional traits and microbial diversity and fitness needs to be quantified simultaneously. Moreover, the highly significant correlation observed between the genetic structure of the rhizosphere bacterial communities and the genetic diversity of *Medicago truncatula* genotypes (Fig. 4) should be confirmed with a *Medicago truncatula* core collection whose genetic diversity is established and that presents a greater number of genotypes. Furthermore, in our study, the environmental effect on *Medicago truncatula* associated rhizosphere

bacterial communities was not evaluated *ie* one soil-type and one cultural condition. Therefore, in our experiment, the genetic effect is probably overestimated. In fact, according to van Overbeek and van Elsas (2008), potato-associated bacterial communities were affected by both plant growth stage and experimental factors (soil and year) rather than by plant genotype. It should be of great interest to observe the ecophysiological behaviour of a *Medicago truncatula* core collection in variable environmental conditions to distinguish genetic and environmental effects on rhizosphere bacterial communities. If relevant, a quantitative genetic approach could be implemented to identify the genetic basis of the plant-microbial community interactions. In summary, it will be necessary to combine ecophysiology, microbial ecology and quantitative genetics to increase our understanding of plant-microbe interactions. In the near future, this could be useful for plant breeders, who will be able to select their plant genotype as a function of the associated soil microbial communities so as to obtain better crop yields (biofertilization) and pathogen resistance (biocontrol).

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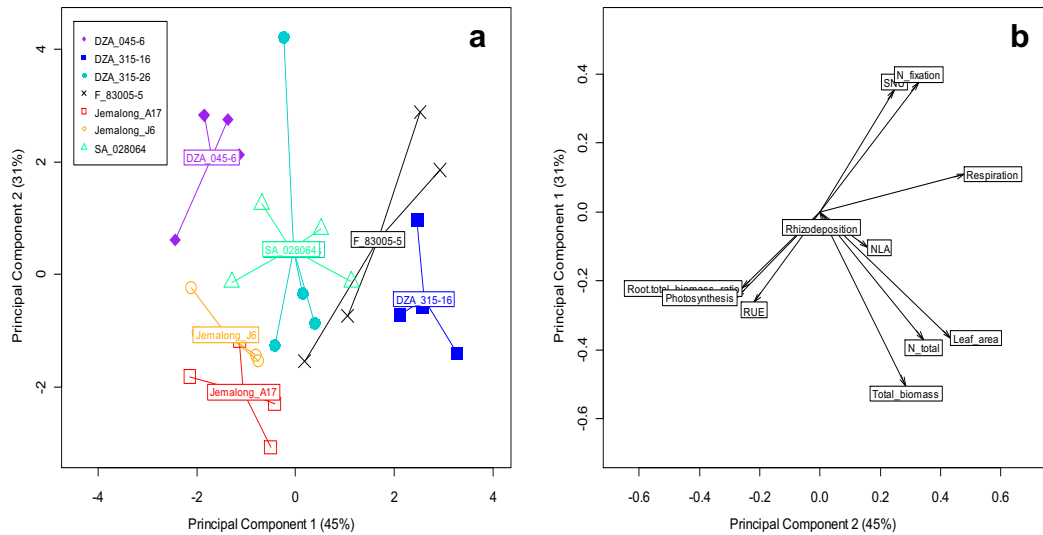
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Online Resource 1 Comparison by principal component analysis of ecophysiological profiles of different *Medicago truncatula* genotypes at 934 degree-days. (a) Factorial plane of the genotypes discrimination in the PCA. (b) Factorial plane of ecophysiological characteristics contribution in the PCA.

Online resource 2 Level of symbiotic association at 934 degree-days for different genotypes of *Medicago truncatula*.

Means were classified using a Tukey test ($P < 0.05$) and different letters indicate statistically different means between genotypes

^a Nodulation was visually assessed, using a qualitative scale (Moreau et al. 2008). The scale takes into account the intensity of nodulation, the size of nodules, and their colour. Nodule colour reflects their functionality (a pink colour indicates the production of leghaemoglobin that is present only in nitrogen fixing nodules). The scale includes five scores: 0 (absence of nodules), 1 (only some white nodules of small size), 2 (both white and pink nodules of small size), 3 (mainly pink nodules of larger size), and 4 (many pink nodules of large size)

^b Mycorrhization parameters represented frequency (F%) and intensity of colonization of the root cortex (M%) and arbuscular abundance of the mycorrhized root cortex (A%) were determined on five root systems per genotype. A nonvital staining procedure was used to allow visualization of total AM fungi (Tisserant et al. 1993). The intensity of root cortex colonization by AM fungi was determined as previously described (Trouvelot et al. 1986) using MYCOCALC software (<http://www.dijon.inra.fr/mychintec/MycoCalc-prg/download.html>).

Genotype	Nodulation intensity ^a	Mycorrhization parameters ^b		
		F (%)	M (%)	A (%)
DZA 045-6	3.8 a	90.8 a	31.1 a	27.0 a
DZA 315-16	4.0 a	92.7 a	35.2 a	29.8 a
DZA 315-26	4.0 a	89.3 a	32.7 a	29.5 a
F 83005-5	4.0 a	95.3 a	44.8 a	38.9 a
Jemalong A17	3.5 a	86.7 a	36.2 a	29.5 a
Jemalong J6	3.8 a	90.7 a	43.3 a	36.6 a
SA 028064	3.3 a	90.8 a	42.6 a	39.7 a