Genetic and symbiotic characterization of rhizobia nodulating legumes in a mining area in southeast Brazil

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Materials and Methods

Origin of the strains

Eighteen strains used in this study were isolated from nodules taken from forest seedlings of Machaerium nyctitans (7), Platypodium elegans (8), and Ormosia arborea (8).
arborea [3] produced in nurseries in mining areas in the municipalities of Nova Lima (20°07'5" S, 43°57'1" W, altitude 742 m) and Sabará (19°51'44" S, 43°47'38" W, altitude 705 m) in the state of Minas Gerais. Isolation and strain characterization were performed in solid 79 culture medium [Fred and Waksman, 1928]. All the strains previously identified at the genus level by sequencing of the 16S rRNA gene were analyzed phylogenetically (Table 1). The BR4101 strain of Ormosia nitida was also tested.

**Authentication and symbiotic efficiency**

Three experiments were conducted in a greenhouse. The aim of each experiment was to authenticate the strains, i.e., prove they are able to nodulate the legume species from which they were isolated, and test the symbiotic efficiency of each strain in its species of origin. The seeds of M. nyctitans and P. elegans were surface disinfected with 2% hypochlorite for 3 h, without the need for breaking dormancy. For O. arborea, dormancy was broken through immersion in sulfuric acid P.A. for 15 min. After that, the seeds of the three species were washed in sterile distilled water for 15 more min to eliminate residues. The seeds were placed in trays containing sterile sand for germination and were moistened with sterile distilled water. After seedlings formed, they were transplanted in 250 cm³ seedling plug pots containing 1:1 [v/v] mixture of sterile sand and vermiculite. At the time of transplanting, the seedlings were inoculated with 1 mL of inoculant of each one of the 18 bacterial strains in the respective host species, and the BR4101 strain was inoculated into the three species. For preparation of the inoculants, each strain was grown in liquid 79 medium in an incubation chamber shaken at 150 rpm. The cultivation time of the inoculants was determined according to the period for growth of each genus to achieve the log phase: six days for the Bradyrhizobium strains [slow growth], and three days for the Rhizobium strains [fast growth]. The treatments of the three experiments included absolute control - without inoculation and low mineral nitrogen content [LN]; control without inoculation and with high mineral nitrogen content [HN]; control with a bacterial strain from Ormosia nitida [BR4101] [efficient for this species [Faria and Uchôa, 2007]] and identified by sequencing of 16SrRNA as belonging to Ochrobactrum [Moreira and Siqueira, 2006]; and treatments inoculated with the strains from each forest species (Table 1), all with low mineral N content. The strain BR4101 was tested on all the leguminous tree species because there are no strains approved as inoculants by the Brazilian Ministry of Agriculture [MAPA] for any of the species studied [MAPA. IN SDA 2011].

The seedling plug pots received fertigation (50 mL per plug pot twice a week) through the application of a nutrient solution [Hoagland and Arnon, 1950] modified according to nitrogen content: 52.5 mg N L⁻¹ for the HN treatment, and 5.25 mg N L⁻¹ for the LN treatment and for the treatments inoculated with the strains as described by Guimarães et al. [2012].

The experiments were evaluated at 60, 90, and 150 days after transplanting, considering the size of the seeds for M. nyctitans, P. elegans, and O. arborea (smaller to larger).
larger, respectively]. This is because the larger the seed, the greater the energy reserve and the longer the time for cotyledon senescence, which may affect the beginning of the symbiosis establishment, i.e., the appearance and functionality of the nodules. The Soil Plant Analysis Development (SPAD) index was determined 60 days after transplanting (SPAD1) for all the forest species studied. For *P. elegans* and *O. arborea*, a second measurement was taken at the end of the experiment (90 and 150 days, respectively) (SPAD2). After collection, the following parameters were evaluated: number of nodules (NN), nodule dry matter (NDM), shoot dry matter (SDM), root dry matter (RDM), total dry matter (TDM) by the sum of SDM and RDM, relative efficiency (RE) calculated by the expression $RE = \frac{[SDM \text{ of inoculated treatment/SDM of HN control}] \times 100}$, shoot nitrogen content in g N per plant (SNC), and total nitrogen content of the plant in g N per plant (TNC). SNC was evaluated by the semi-micro Kjeldahl method (Kirk, 1950) and used for calculation of the TNC by the expression $SNC \times SDM$.

**DNA extraction**

Genomic DNA of the strains was extracted using the Wizard Genomic DNA Purification kit, following the manufacturer’s recommendations. The quality and concentration of the extracted DNA was analyzed using a Colibri spectrometer.

**Amplification and sequencing of housekeeping genes (atpD and gyrB)**

The *gyrB* gene was amplified using the primers 343F (5’ TTCGACCAGAAATCTCTAAYAAGG 3’) and 1043R (5’ AGCTTGTCCTTSGTCTGCG 3’). The *atpD* gene was amplified using the primers 352F (5’ AGCTTGTCCTTSGTCTGCG 3’) and 871R (5’ AGCTTGTCCTTSGTCTGCG 3’). The DNA fragments obtained from the PCR products were verified in 1 % (w/v) agarose gel and visualized in a UV transilluminator. The PCR products were sequenced by Wenseq in a 3500 xLGenetic analyzer (Applied Biosystems) sequencer with 24-50 cm capillaries and Pop7 polymer.

**PCR amplification of the nifH, nodC, and nodD symbiotic genes**

The *nifH* gene was amplified using the primers nifHF (5’ AAAGYYGWWATCGGAYAARTCCACAC 3’) and nifHR (5’ TGTGTGGCSGCTACATGCATGCGATCAT 3’) with Promega GoTaq DNA Polymerase ([500 U-M3005]), and the thermal cycler was used under the conditions described in Sarita et al. (2005). The *nodD* gene was amplified using the primers nodDF (5’ GATYTGCAATGAAATCGKAGAG 3’) and nodDR (5’ TCATAGACANACATCCACAGAT 3’) with Promega GoTaq DNA Polymerase ([500 U-M3005]), and the thermal cycler was used under the conditions described in Sterner and Parker (1999). The *nifH*, *nodC* and *nodD* bands were visualized in 1 % (w/v) agarose gel through the UV transilluminator.

**Phylogenetic analysis**

The sequences quality and contigs assembly of the housekeeping (*atpD* and *gyrB*) and symbiotic (*nifH*) genes were verified using the BioNumerics 7.6 software system. The MUSCLE (Edgar, 2004) multiple alignment algorithm was used to align the sequences. Phylogenetic trees were constructed on the MEGA 7 software (Kumar et al., 2016) by the neighbor-joining (NJ) method (Saitou and Nei, 1987) using the Kimura 2 parameter (Kimura, 1980) with bootstrap of 1000 replications. The sequences used in phylogenetic analysis were those that have a base pair number of 205 for the *nifH* gene for both *Rhizobium* and *Bradyrhizobium*, as well as 210 and 207 for the *gyrB* and 336 and 352 for the *atpD* genes, for *Rhizobium* and *Bradyrhizobium*, respectively. Sequences of type strains of *Bradyrhizobium* and *Rhizobium* available in the Genbank (National Center for Biotechnology Information, NCBI) that had at least 95 % similarity with the sequences studied (calculated by BLAST) were included in the phylogenetic analysis. Each phylogenetic analysis was evaluated with mathematical language to obtain the percentage of dissimilarity between sequences and their similarity, calculated by the expression $1 - x \times 100$, where “$x$” is the value of dissimilarity obtained by comparisons between the base pairs of each sequence.

**Statistical analysis**

A completely randomized experimental design (CRD) was used, with five replications for the experiment with *P. elegans* and three replications for experiments with *M. nyctitans* and *O. arborea*. Analysis of variance (ANOVA) was performed for the variables RDM, SDM, TDM, RE, SPAD1, SPAD2, SNC, and TNC, and their means were compared with the HN and LN controls by the Dunnett test at 5 % probability using the R software program in the gylma [Pena and Slate, 2006], agricolae [Mendiburu, 2017], and asbio [Ken, 2020] packages. In addition, Pearson linear correlation analysis was applied to all the variables analyzed in the symbiosis experiments, using the biotools package in the R software program (Silva et al., 2017).
Results

Authentication and symbiotic efficiency

In the LN and HN treatments, there was no nodule formation, confirming the absence of contamination in the experiments. The strain recommended for inoculation in *O. nitida* (BR4101) exhibited positive nodulation in the experiments with *O. arborea* and *M. nyctitans*. However, nodulation of this strain in *P. elegans* was not detected.

In the experiment with *M. nyctitans*, of the seven strains tested, all nodulated in the species of origin, six of *Bradyrhizobium* and one of *Rhizobium* (Table 2). Among these strains, it was possible to amplify the *nifH* gene of *Bradyrhizobium* UFLA01-814, UFLA01-834, UFLA01-839, UFLA01-860, UFLA01-883 and UFLA01-1164, and *Rhizobium* UFLA01-1172; the *nodC* gene of *Bradyrhizobium* UFLA01-814 and UFLA01-883; and the *nodD* gene of *Bradyrhizobium* UFLA01-834, UFLA01-839, and UFLA01-883. In the experiment with *P. elegans*, of the eight strains tested, six nodulated in the species of origin, three from *Bradyrhizobium* and three from *Rhizobium* (Table 3). Among these strains, it was possible to amplify the *nifH* gene of *Rhizobium* UFLA01-1128 and UFLA01-1137, and *Bradyrhizobium* UFLA01-1136 and the *nodD* gene of *Bradyrhizobium* UFLA01-874 and UFLA01-1136, and *Rhizobium* UFLA01-1128 and UFLA01-1134. In the experiment with *O. arborea*, of the three strains tested, all nodulated in the species of *Ochrobactrum*.

Table 2 – Performance of strains of *Jacarandá bico-de-pato* (*Machaerium nyctitans*) evaluated as regards number of nodules (NN), nodule dry matter (NDM), shoot dry matter (SDM), root dry matter (RDM), total dry matter (TDM), relative efficiency (RE), SPAD index (SPAD), shoot nitrogen content (SNC), and total nitrogen content per plant (TNC) at 60 days after sowing in an experiment in a greenhouse under axenic conditions. Mean data of three replications.

| Inoculation treatment (Genus*) | NN | NDM | SDM | RDM | TDM | RE | SPAD 1 | SNC | TNC |
|-------------------------------|----|-----|-----|-----|-----|----|--------|-----|-----|
| UFLA01-814(B)                 | 25 | 23.3 | 0.23 Aa** | 0.13 Aa | 0.36 Aa | 67.76 Aa | 23.43 Bb | 16.70 Ba | 3.89 Bb |
| UFLA01-834(B)                 | 45 | 36.7 | 0.24 Aa | 0.12 Aa | 0.36 Aa | 70.18 Aa | 24.27 Bb | 17.37 Bb | 4.22 Bb |
| UFLA01-839(B)                 | 49 | 39.67 | 0.29 Aa | 0.07 Aa | 0.36 Aa | 83.83 Aa | 36.00 Aa | 28.17 Aa | 8.35 Ab |
| UFLA01-860(B)                 | 41 | 1.67 | 0.34 Aa | 0.11 Aa | 0.34 Aa | 66.07 Aa | 25.50 Bb | 17.50 Bb | 3.98 Bb |
| UFLA01-883(B)                 | 41 | 1.10 | 0.34 Aa | 0.10 Aa | 0.34 Aa | 67.38 Aa | 25.40 Bb | 21.47 Bb | 4.98 Bb |
| UFLA01-1164(B)               | 55 | 15.0 | 0.18 Ba | 0.10 Aa | 0.28 Aa | 51.79 Ba | 28.07 Bb | 23.03 Bb | 4.12 Bb |
| UFLA01-1172(R)               | 23 | 1.33 | 0.15 Ba | 0.07 Aa | 0.22 Ba | 43.37 Ba | 25.77 Bb | 20.23 Bb | 2.78 Bb |
| BR4101(O)                   | 10 | 2.67 | 0.24 Aa | 0.15 Aa | 0.40 Aa | 70.57 Aa | 26.13 Bb | 16.20 Bb | 3.93 Bb |
| HN (52.5 mg N L–1)           | 0  | 0     | 0.34 Aa | 0.12 Aa | 0.46 Aa | 100 Aa  | 38.47 Aa | 34.27 Aa | 11.72 Aa |
| LN (5.25 mg N L–1)           | 0  | 0     | 0.26 Aa | 0.15 Aa | 0.41 Aa | 74.25 Aa | 25.57 Bb | 17.77 Bb | 4.55 Bb |

*B = Bradyrhizobium, R = Rhizobium, O = Ochrobactrum; **Data followed by the same uppercase letter in the column do not differ from the HN treatment, and followed by the same lowercase letter in the column do not differ from the LN treatment; significant by the Dunnett test (p < 0.05). HN = Control treatment with mineral nitrogen fertilization. LN = Control treatment without inoculation and with low mineral nitrogen content.

Table 3 – Performance of strains of *Jacarandá do Campo* (*Platypodium elegans*) evaluated as regards number of nodules (NN), nodule dry matter (NDM), shoot dry matter (SDM), root dry matter (RDM), total dry matter (TDM), relative efficiency (RE), SPAD index (SPAD), shoot nitrogen content (SNC), and total nitrogen content per plant (TNC) in an experiment in a greenhouse under axenic conditions. Mean data of five replications.

| Inoculation treatment (Genus*) | NN | NDM | SDM | RDM | TDM | RE | SPAD 1 | SNC | TNC |
|-------------------------------|----|-----|-----|-----|-----|----|--------|-----|-----|
| UFLA01-1127(R)              | 1.6 | 1.4 | 0.17 Aa** | 0.28 Aa | 0.45 Aa | 115.73 Aa | 32.67 Ab | 26.18 Aa | 17.22 Aa |
| UFLA01-1128(R)              | 0.2 | 0.6 | 0.12 Aa | 0.20 Ba | 0.32 Ba | 96.74 Aa | 33.43 Aa | 26.03 Aa | 22.50 Aa |
| UFLA01-1129(R)              | 0.0 | 0.15 Aa | 0.28 Aa | 0.43 Aa | 110.11 Aa | 33.62 Aa | 29.10 Aa | 19.12 Aa |
| UFLA01-874(B)              | 3.8 | 2.6 | 0.14 Aa | 0.30 Aa | 0.37 Aa | 98.17 Aa | 34.60 Aa | 31.52 Aa | 18.26 Aa |
| UFLA01-1134(R)              | 0.0 | 0.14 Aa | 0.24 Aa | 0.37 Aa | 96.49 Aa | 34.02 Aa | 28.18 Aa | 18.12 Aa |
| UFLA01-1132(R)              | 0.4 | 0.12 Aa | 0.35 Aa | 0.48 Aa | 102.7 Aa | 38.70 Aa | 32.14 Aa | 20.34 Aa |
| UFLA01-1137(R)              | 0.4 | 1.0 | 0.32 Aa | 0.49 Aa | 116.71 Aa | 36.62 Aa | 32.40 Aa | 18.60 Aa |
| UFLA01-1136(B)              | 14.2 | 8.4 | 0.13 Aa | 0.27 Aa | 0.41 Aa | 88.05 Aa | 35.67 Aa | 25.90 Aa | 11.08 Aa |
| BR4101(O)                  | 0.0 | 0.14 Aa | 0.19 Ba | 0.34 Aa | 99.9 Aa | 32.95 Aa | 29.17 Aa | 17.10 Aa |
| HN (52.5 mg N L–1)         | 0  | 0     | 0.16 Aa | 0.41 Aa | 0.57 Aa | 94.9 Aa  | 34.33 Aa | 33.42 Aa | 26.64 Aa |
| LN (5.25 mg N L–1)         | 0  | 0     | 0.15 Aa | 0.35 Aa | 0.50 Aa | 94.28 Aa | 39.20 Aa | 34.70 Aa | 20.74 Aa |

*B = Bradyrhizobium, R = Rhizobium, O = Ochrobactrum; **Data followed by the same uppercase letter in the column do not differ from the HN treatment, and followed by the same lowercase letter in the column do not differ from the LN treatment; significant by the Dunnett test (p < 0.05). HN = Control treatment with mineral nitrogen fertilization. LN = Control treatment without inoculation and with low mineral nitrogen content.
origin, one of Bradyrhizobium [UFLA 01-1161] and two of Rhizobium [UFLA 01-1147 and UFLA 01-1156] (Table 4). Among these strains, it was possible to amplify the nifH gene of the strain UFLA01-1156.

On analysing the morpho-physiological variables of *M. nyctitans* (Table 2), the treatment inoculated with the *Bradyrhizobium* UFLA01-839 strain stood out in relation to the SPAD index [36.00] as it did not show any difference from the control with mineral nitrogen fertilization (HN) [38.47] and was superior to the control without inoculation and with low nitrogen (LN) content [25.57]. The SPAD index was reflected in the shoot N content (SNC) and in the total N content (TNC), with positive correlation of 0.8 \((p < 0.001)\) between these variables. In addition to the SPAD and TNC index, the plants inoculated with UFLA01-839 also exhibited greater nodular dry matter, which had positive correlation of 0.2 and 0.1 with SNC and TNC, respectively \((p < 0.05)\). The treatment inoculated with the *Rhizobium* UFLA01-1172 strain exhibited relative efficiency of 43 %, less than all the other treatments, including the control without inoculation and with low N contents, which exhibited relative efficiency above 50 % (Table 2). The treatments increased the RDM of the plants; however, inoculation with the strains *Bradyrhizobium* UFLA01-1164 and *Rhizobium* UFLA01-1172 did not supply the nitrogen requirements of the plants and were inferior to the HN control (Table 2).

Analysis of the morpho-physiological variables of the authentication experiment in *P. elegans* (Table 3) revealed no differences between the SPAD1 and SPAD2 values for the treatments and the controls, and there was no difference between the controls (HN - LN). For all the treatments, there was a reduction from SPAD1 to SPAD2 (Table 3) and both indices exhibited a positive correlation of 0.5 \((p < 0.001)\). The treatment inoculated with the *Bradyrhizobium* UFLA01-1136 strain exhibited the highest NN and NDM (Table 3). Nevertheless, this was not reflected in a relative efficiency higher than that of the other treatments. The HN and LN controls were not significantly different from each other for any of the variables evaluated.

Table 4 – Performance of strains of *Olho-de-cabra* (*Ormosia arborea*) evaluated as regards number of nodules (NN), nodule dry matter (NDM), shoot dry matter (SDM), root dry matter (RDM), total dry matter (TDM), relative efficiency (RE), SPAD index at 60 days after sowing (SPAD 1), shoot nitrogen content (SNC), and total nitrogen content per plant (TNC) in an experiment in a greenhouse under axenic conditions. Mean data of three replications.

| Inoculation treatment (Genus*) | NN | NDM | SDM | RDM | TDM | RE | SPAD1 | SPAD2 | SNC | TNC |
|-------------------------------|----|-----|-----|-----|-----|----|-------|-------|-----|-----|
| mg per plant                  | mg g per plant | % | mg N g per plant | mg N per plant |
| UFLA 01-1156(R)               | 2  | 1.33 | 32.32 | 15.47 | 9.05 |
| UFLA 01-1161(B)               | 0.3 | 24.67 | 39.26 | 28.93 | 11.23 |
| UFLA 01-1147 (R)              | 1  | 0.33 | 29.36 | 27.87 | 11.90 |
| BRM101 (O)                    | 4  | 45.67 | 55.26 | 25.97 | 16.70 |
| HN (52.5 mg N L⁻¹)            | 0  | 0.71 | 39.26 | 27.87 | 11.90 |
| LN (15.25 mg N L⁻¹)           | 0  | 0.71 | 39.26 | 27.87 | 11.90 |

*Data followed by the same uppercase letter in the columns do not differ from the HN treatment, and followed by the same lowercase letter in the columns do not differ from the LN treatment; significant by the Dunnett test \((p < 0.05)\). HN = Control treatment with mineral nitrogen fertilization. LN = Control treatment without inoculation and with low mineral nitrogen content.

The *Rhizobium* UFLA 01-1156 and UFLA 01-1147 strains exhibited positive nodulation in the experiment with *O. arborea* (Table 4). The UFLA 01-1156 strain had SPAD2 and SNC values statistically similar to the HN treatment (Table 4). The highest values of SDM, SPAD, SNC, and TNC were observed in the treatment with mineral nitrogen (HN), exceeding the values of the inoculated treatments, including the value from the strain isolated from the same genus (*O. nitida*). This strain [BR4101] nodulated *O. arborea* and favored the accumulation of SDM (1.22 g per plant), with relative efficiency (55 %) similar to the control with mineral nitrogen (HN) (Table 4). The plants inoculated with the UFLA01-1156 strain (*Rhizobium*) obtained SPAD2 and shoot N content (SNC) values similar to the HN treatment. For the treatment inoculated with the UFLA01-1156 strain, there was positive correlation of SDM with the SNC and TNC variables, at 0.2 and 0.9, respectively \((p < 0.001)\).

**Phylogenetic analysis**

The phylogenetic analyses of the strains by the *gyrB* and *atpD* genes corroborate identification in *Rhizobium* and *Bradyrhizobium* obtained by phylogenetic analysis of the 16S rRNA gene (Table 1).

It was possible to amplify the *gyrB* gene from 10 of the 18 strains, four of them belonging to *Rhizobium* and six to *Bradyrhizobium* (Figure 1A and B). Among the strains identified as *Rhizobium*, there was clustering of the strains UFLA 01-1134, UFLA 01-1128 and UFLA 01-1127 of *P. elegans* with 100 % similarity to each other and 100 % bootstrap, which were nearest to *R. tropici* CIAT899’, with similarity of 92 % and 62 % to bootstrap. The UFLA01-1156 strain of *O. arborea* had 96 % similarity to the nearest species, *R. leucaenea* CFN 299’, 58 % bootstrap. Among the strains identified as *Bradyrhizobium*, UFLA01-1164 and UFLA01-839 of *M. nyctitans* were positioned separately between themselves and from all the other type strains of *Bradyrhizobium*. The UFLA01-1164 strain had 97 % similarity to the
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nearest species, *B. daqingense* CCBAU 15774, and the UFLA01-839 strain showed 99% similarity to the nearest species, *B. viridiflatus* SEMIA 690. The UFLA01-814 and UFLA01-860 strains of *M. nyctitans* created a separate cluster, 65% bootstrap, which belong to the *Bradyrhizobium elkanii* superclade. This group had 96% similarity to the nearest species, *B. pachyrhizi* PAC 48. The UFLA01-1136 strain of *P. elegans* was at a phylogenetic position isolated from the other species, with greatest proximity to *B. uaiense* UFLA03-164, with 98% similarity. Finally, the UFLA01-1132 strain was positioned separately from all *Bradyrhizobium* species. Its closest species was *B. diaezoefficiens* USDA 110, with 98.2% similarity.

From the *atpD* gene sequencing six strains of *Rhizobium* and two of *Bradyrhizobium* were identified [Figure 2A and B]. Among the strains identified as *Rhizobium*, a similar response was observed for the strains UFLA 01-1128 and UFLA 01-1127 of *P. elegans* compared to the *gyrB* gene. These strains were nearer...
Figure 2 – Phylogenetic trees based on sequences of the atpD gene of *Bradyrhizobium* (352 bp) (A) and *Rhizobium* (336 bp) (B) strains authenticated in regard to nodulating capacity in *M. nyctitans* (Mn), *P. elegans* (Pe), and *O. arborea* (Oa) and 24 type strains of known species exhibiting at least 95 % similarity with the sequences studied (calculated by BLAST). Phylogeny determined by the neighbor-joining and bootstrap method of 1000 replications. The designation (T) indicates biobar reference strains (former species type strain).
to R. hainanense I66\textsuperscript{c}, with 57 % bootstrap and 99 % similarity. The UFLA01-1156 strain of O. arborea was similar to R. jaguaris CCGE525\textsuperscript{c}, with 59 % bootstrap and 98 % similarity; and the strain UFLA01-1147 of O. arborea was similar to R. lentis BRL27\textsuperscript{c} at 96 % similarity. Among the strains identified as Bradyrhizobium, there was separate positioning of the strains UFLA01-1164 and UFLA01-839 of M. nyctitans, just as in the phylogenetic tree of the gyrB gene. However, in the phylogenetic tree of the atpD gene (Figure 2A), the UFLA01-1164 strain had greater similarity to B. americanum CMVU44\textsuperscript{c}, with 90 % bootstrap and 99 % similarity, and the UFLA01-839 strain had 99 % similarity to B. tropiciagri SEMIA6148\textsuperscript{c}.

In the phylogenetic tree of the nifH gene for seven out of the 11 strains that had this gene amplified (Figure 3), the strains of Bradyrhizobium UFLA01-814 and UFLA01-860 of M. nyctitans were clustered with greatest proximity to B. mercantei SEMIA6399\textsuperscript{d}. The UFLA01-839 of M. nyctitans and UFLA01-1136 of P. elegans are clustered but are separated from the other type strains from the Bradyrhizobium and Rhizobium genera. UFLA01-874 of P. elegans and UFLA01-1172 and UFLA01-1164 of M. nyctitans were clustered with the greatest proximity to B. stylosanthis BR446\textsuperscript{c}. All the clusters formed except UFLA01-839 and UFLA01-1136 are supported by high bootstrap values. Bradyrhizobium strains UFLA01-874 and UFLA01-1164 and Rhizobium UFLA01-1172 strain grouped together in a separate clade with high statistical support (98 %). This result elicits interest as it indicated a possible horizontal gene transfer.

**Discussion**

Although there was no response in plant dry matter, the contribution of nitrogen [N\textsubscript{2}] fixed by the bacteria can be quantified by the SPAD index, which is indirectly related to the chlorophyll content in the plant shoots, depending on the intensity of green in the leaves. In the case of Machaerium plants, the SPAD index was able to show differences between the treatments before the increase in dry matter, proving to be a useful analysis for a short-term experiment. These data corroborate the data obtained by Jaramillo et al. [2013], who obtained a SPAD response before differentiation in dry matter in an experiment on inoculation efficiency in Vigna unguiculata (cowpea). In addition, there was a positive correlation of the SPAD index with SNC, which made

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**Figure 3** – Phylogenetic tree based on sequences of the nifH (205 bp) gene of 7 strains authenticated in regard to nodulating capacity in M. nyctitans (Mn) and P. elegans (Pe) and 28 type strains of known species exhibiting at least 95 % similarity with the sequences studied (calculated by BLAST). Phylogeny determined by the neighbor-joining and bootstrap method of 1000 replications. The designation (T) indicates biovar reference strains (former species type strain).
it possible to use the values of this index as a rapid response in relation to shoot N content. Nyoki and Ndakidemi (2014) also cite the benefits of inoculation with rhizobia and their effect on chlorophyll formation, corroborating the data obtained in this study.

Pastorini et al. (2016) found no nodules on plants of *Macharia*um *brasilense* Vogel without inoculation at 90 days after transplanting to a substrate composed of a mixture of commercial organic substrate and sand (1:2). Nodules were observed only 180 days after transplanting. Nevertheless, in this study, nodulation of the *Macharia*um *nyctitans* plants was found at 60 days after transplanting, showing that through the inoculation process, it is possible to advance rhizobia symbiosis in the seedlings. The two strains of *P. elegans* that did not nodulate this species may be invasive bacteria, endophytic bacteria of the nodule, nodule-forming bacteria that are not specific for this species, or nodule-forming bacteria that failed to encounter conditions favorable to their nodulation. One of the conditions necessary for nodulation may be related to the point in time of inoculation, i.e., if inoculation is performed at the time of sowing, the time at which the seedling will require nitrogen (at deficiency) should be ascertained because if the seedling still has seed reserves, it will not need to establish symbiosis and will not produce signaling flavonoids that will activate expression of the *nod* genes of rhizobia, which may render the nodulation process inviable (Ndakidemi and Dakora, 2003). This delay in establishing symbiosis may be reflected in late responses in the efficiency tests, taking into consideration that the experiments were conducted in 60, 90, and 150 days in a greenhouse. Thus, certain forest species require a longer time for conducting the experiment up to evaluation. If the difference between each treatment and the control without inoculation with low mineral-N content fails to produce significant differences in the vegetative growth parameters evaluated, then it is necessary for the plants to be subjected to a longer time of exposure to the treatments so that the resultant differences will be significant. In addition, low nodulation or absence of it should be carefully observed, since microorganisms present in identifiable systems can perform another type of process that promotes vegetative growth, such as solubilization of phosphate and production of phytohormones (Costa et al., 2016).

The two superclades of *Bradyrhizobium* (*B. elkanii* and *B. japonicum*) are well defined phylogenetically (Avontuur et al., 2019). Strains representative of the two superclades isolated from *M. nyctitans* and *P. elegans* were authenticated, whereas from *O. arborea*, the only strain of *Bradyrhizobium* authenticated belonged to the *B. elkanii* superclade. Identification of strains forming different clusters in these two clades, or only separating themselves in an isolated manner from the other type species within the phylogenetic trees of the housekeeping genes (*atpD* and *gyrB*), separated or concatenated, with similarity values below 95 %, indicates the possibility of new species not yet described. In a study describing a new species of *Bradyrhizobium*, Costa et al. (2018) found similarity values of 96-93 and 97-95 % between type species and the strains tested when they analyzed sequences of housekeeping genes (*atpD* and *gyrB*). These values were also found our study, thereby corroborating the identification of possible new species. The *Bradyrhizobium* strain UFLA01-839, which is among those with the potential for being a new species, stood out in the authentication and symbiotic efficiency experiment with *M. nyctitans*, and it can be considered for future tests of selection of inoculant strains for this forest species. Although, no final conclusions can be formed about specificity of these tree species regarding rhizobia genera, it is noteworthy that the highest affinity of *M. nyctitans* with *Bradyrhizobium* and of *P. elegans* was with *Rhizobium*.

The results obtained from phylogenetic analysis of the housekeeping genes performed in this study corroborate other studies, which show that sequencing of housekeeping genes improves discrimination of species of both *Bradyrhizobium* and *Rhizobium*, complementing the analysis of the 16S rRNA gene (Guimarães et al., 2015; Bourebaba et al., 2016; Rouhrazzi et al., 2016; Degefu et al., 2017). In addition, results show that the *gyrB* gene is useful in discriminating species, as shown by Guimarães et al. (2015). This type of analysis was efficient in separating strains with greater potential for being new taxons. However, new studies that involve more comprehensive analyses of the genome of these microorganisms have become increasingly necessary for description of species (Konstantinidis and Tiedje, 2005; López-Guerrero et al., 2012; Okazaki et al., 2015; Kishi et al., 2016; Costa et al., 2018).

The high diversity found among strains isolated from the different forest species show how necessary it is to select specific strains for each one of them. This can be seen in various studies that tested *Mimosa* spp. inoculated with *Burkholderia* (Araújo et al., 2017) or *Inga* sp. inoculated with *Bradyrhizobium* (Silva et al., 2014; Porto et al., 2017). They exhibit different plant/microorganism combinations, reaffirming the need for determining which combinations are most efficient.

In addition, another important factor in the description of new species is their phylogenetic relationship based on symbiotic genes such as *nifH*. Horizontal transfer of these genes represents an important tool in the evolutionary force, which is able to confer adaptive advantages to determined rhizobia species (Parker, 2012; Remigi et al., 2016). There were strains both with and without affinity between the phylogenies of the *nifH* gene and the housekeeping genes. Thus, we can advance the hypothesis that there may have been both vertical and horizontal transmission of the *nifH* gene between the strains studied (Michel et al., 2020).
Conclusions

Nodulation of the three forest species was confirmed in 16 of their strains of origin, belonging to both *Rhizobium* and *Bradyrhizobium*. Phylogenetic analysis of the *atpD* and *gyrB* housekeeping genes indicated high diversity in *Rhizobium* and *Bradyrhizobium* strains with the potential for representing new species, such as UFLA01-1127, UFLA01-1128, UFLA01-1129, UFLA01-1134, UFLA01-1156, UFLA01-1172 in *Rhizobium*, and UFLA01-814, UFLA01-839, UFLA01-860, UFLA01-1164 and UFLA01-1132 in *Bradyrhizobium*.

Inoculation of the UFLA01-839 (*Bradyrhizobium* sp.) strain into the *Machaerium nyctitans* seedlings resulted in more efficient symbiosis. It was not possible to ascertain the symbiotic efficiency of the strains in *Platypodium elegans* and *Ormosia arborea*, probably due to the greater amount of energy reserves of the seeds, which supplied the demand of the seedlings in the present study. Therefore, a longer evaluation time for these species is recommended in future studies.

The SPAD index was efficient in detecting efficiency in fixing N₂ before the determination of shoot dry matter.

Identification and characterization of the 18 strains of rhizobia will allow them to be studied in future projects to test their symbiotic efficiency under axenic conditions, as well as to identify them at species level by genome sequencing allowing for future research to test their symbiotic efficiency in fixing N₂ before the determination of shoot dry matter.

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