Arbuscular mycorrhizal fungi and *Urochloa brizantha*: symbiosis and spore multiplication¹

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INTRODUCTION

Arbuscular mycorrhizal fungi (AMF) establish a mutualistic symbiotic association with the roots of most plant species, characterized by a functional integration of the plants with the AMF, resulting in bidirectional and simultaneous exchange of metabolites and nutrients between the microsymbiont and the host plant (Parniske 2008, Smith & Read 2008, Oehl et al. 2009).

ABSTRACT

Arbuscular mycorrhizal fungi (AMF) may exhibit distinct behaviors when associated with the same species of host plant, being necessary to understand their ecology, in order to optimize their management and maintenance in germplasm bank. This study aimed to evaluate different AMF associated with *Urochloa brizantha* (A. Rich.) Stapf by analyzing the symbiosis establishment time, spore multiplication and production of glomalin-related soil protein. The experiment was carried out in a completely randomized design, in a 6 x 5 arrangement (five AMF species, non-inoculated control treatment and five evaluation times). The following aspects were analyzed: plant growth, spore multiplication, mycorrhizal colonization and glomalin production. The highest number of spores occurred for *Acaulospora longula* and *A. colombiana*, exhibiting the highest mycorrhizal colonization at 76 days. The inoculation favored the root growth of *U. brizantha* at 15 days of cultivation, plant height and root dry matter at 60 days and shoot dry matter at 90 days, especially for the *Acaulospora* species. The inoculation with *A. colombiana, A. longula* and *Paraglomus occultum* resulted in increased glomalin at 120 days. *Gigaspora margarita* and *P. occultum* did not reach the maximum colonization and spore multiplication, indicating that a period of time longer than 120 days of cultivation is necessary.

KEYWORDS: Germplasm bank, glomalin,-glomeromycota.

INTRODUCTION

These fungi occupy a vital ecological niche and play key roles for the environmental balance in native or cultivated lands (Bücking et al. 2016). Unlike other microorganism groups, AMF are obligatory biotrophics and require a metabolically-active root system to establish the symbiosis and complete their life cycle, renewing their infective propagules (Smith & Read 2008, Siqueira et al. 2010). Due to the obligatory biotrophism, the multiplication of AMF is achieved in axenic in vitro conditions. The present study aimed to evaluate different AMF associated with *U. brizantha* by analyzing the symbiosis establishment, spore multiplication and production of glomalin-related soil protein.
cultivation using transformed roots, or in its classical form, in trap pots cultivated with plants (Selvakumar et al. 2018). *In vitro* cultivation enables the multiplication of viable, pure, decontaminated propagules in a shorter period of time, being effective for multiplication of fast-growing AMF, which establish symbiosis in 3 to 4 days (Dalpe & Seguin 2010). However, it presents limitations in the AMF development, such as reduced growth and reduced capacity to colonize roots. On the other hand, the AMF multiplication method that uses cultivation trap pots with substrate/soil is widely used, especially for being less artificial and more cost-effective and capable of producing large amounts of highly-efficient inoculants in the mycorrhizal colonization of plant roots (Schlemper & Stürmer 2014, Selvakumar et al. 2018).

Studies have shown that there is compatibility between AMF and host plants, resulting in a variation in spore multiplication, mycorrhizal colonization, extra-radicular hyphae growth and production of glomalin-related soil protein (Klironomos et al. 2005, Parniske 2008, Smith & Read 2008, Siqueira et al. 2010). Glomalin is a glycoprotein found in the soil, produced on the hyphae surface and by degradation of the AMF hyphae and spores (Lehmann et al. 2017).

The use of plants of the *Urochloa* genus for multiplication of FMA in germplasm banks is already widespread, but little is known about the host plant interaction with different species of AMF regarding the symbiosis establishment time, multiplication of infective propagules, mycorrhizal colonization and production of glomalin-related soil protein, therefore, producing different effects on the plant growth and on soil. This information is important for optimizing spore multiplication, management of soilborne AMF, planning of studies on inoculation conducted in the field or in controlled environments, and for keeping these fungi in germplasm banks. Therefore, the hypothesis of this study is that different AMF species may have different behaviors when colonizing the same host plant.

Thus, this study aimed to determine if there is a distinct behavior in the interaction of different AMF species associated with *Urochloa brizantha* (A. Rich.) Stapf and analyze the time of symbiosis establishment, spore multiplication and glomalin-related soil protein production.

**MATERIAL AND METHODS**

The study was carried out in a greenhouse, from September 2015 to February 2016, in a completely randomized design with split plots and five replications, considering the inoculation with five AMF species (Table 1) and one non-inoculated treatment, and five evaluation times (15, 30, 60, 90 and 120 days after the germination). The AMF were used as available in the AMF collection at the Universidade Federal de Lavras, in Lavras, Minas Gerais state, Brazil.

The substrate consisted of a mixture of 2:1 (v/v) of washed sand and Oxisol (USA 1999). The substrate chemical characterization was as it follows: pH value (water) = 5.4; H + Al = 2.9 cmol dm⁻³; Ca = 1.70 cmol c dm⁻³; Mg = 0.10 cmol c dm⁻³; K = 18 mg dm⁻³; P = 1.13 mg dm⁻³; OM = 2.11 dag kg⁻¹. The substrate was autoclaved at 121 °C for one hour, procedure conducted for two consecutive days, and then transferred to 1-kg capacity pots.

The inoculation was performed by transferring 150 AMF spores to each pot containing inoculum soil, which, in addition to the spores, also contained colonized hyphae and roots, which also act as AMF infective propagules. The inoculum used in this study was obtained by multiplication of AMF in cultivation pots (autoclaved soil) for 180 days. Then, the experiment inoculation was carried out.

Sowing was conducted using *Urochloa brizantha* (A. Rich.) Stapf seeds, which were

Table 1. Arbuscular mycorrhizal fungi (AMF) species used in this study, identified by a deposit code in the AMF collection of the Universidade Federal de Lavras, in Lavras, Minas Gerais state, Brazil.

<table>
<thead>
<tr>
<th>AMF species</th>
<th>Code</th>
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<tbody>
<tr>
<td><em>Acaulospora colombiana</em></td>
<td>(Spain &amp; Schenck) Kaonongbua, Morton &amp; Bever 537 UFLA</td>
</tr>
<tr>
<td><em>Acaulospora longula</em></td>
<td>Spain &amp; Schenck 242 UFLA</td>
</tr>
<tr>
<td><em>Acaulospora morrowiae</em></td>
<td>Spain &amp; Schenck 467 UFLA</td>
</tr>
<tr>
<td><em>Gigaspora margarita</em></td>
<td>Becker &amp; Hall 252 UFLA</td>
</tr>
<tr>
<td><em>Paraglomus occultum</em></td>
<td>(Walker) Morton &amp; Redecker 438 UFLA</td>
</tr>
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</table>

* The AFM species were firstly isolated from the Brazilian Savannah biome and propagated in a greenhouse using *Urochloa decumbens*.
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Disinfested in a solution of sodium hypochlorite at 0.5 % (v/v) during 5 min and washed with distilled water. Each pot contained 20 seeds and, at 15 days after the germination, the plants were thinned out, remaining 10 plants per pot.

During the conduction of the experiment, 20 mL of nutrient solution composed of 210 mg L⁻¹ of N, 234 mg L⁻¹ of K and 15.05 mg L⁻¹ of P were applied, which corresponded to 50 % of the recommended P concentration. The pots were randomized, and the substrate was maintained with 60 % of the moisture content of the total pores volume.

The study was carried out during 120 days, with periodic assessments at five times (15, 30, 60, 90 and 120 days after the germination). At each of these times, the plants were harvested and the following measurements were made: plant height, root length, shoot dry mass, root dry matter, number of spores, mycorrhizal colonization and easily extractable glomalin-related soil protein.

The number of spores was quantified in 50 mL of the substrate of each treatment, using the wet sieving method (Gerdemann & Nicolson 1963) and water centrifugation in water and sucrose solution (Jenkins 1964). Counting was made directly from corrugated plates under stereoscopic microscope.

Mycorrhizal colonization was quantified in 1-g samples of thin roots. The roots were washed, diaphanized with KOH (10 %) and H₂O₂, acidified with HCl (1 %) and stained with Trypan blue in lactoglycerol (0.05 %) (Phillips & Hayman 1970). The quantification of colonized roots was achieved by the interaction technique in reticulate plates (Phillips & Hayman 1970, Giovannetti & Mosse 1980).

The concentration of easily extractable glomalin-related soil protein was determined using the method proposed by Wright & Upadhyaya (1998). For this purpose, 1-g soil samples with 8 mL of sodium citrate solution (20 mM at pH 7.2) were autoclaved for 30 min at 121 °C, then centrifuged (3,200 rpm/20 min), and the glomalin-related soil protein extract was obtained from the soil samples and aggregates (Bradford 1976, Wright & Upadhyaya 1998). Afterwards, the supernatants were quantified by spectrophotometer following the Bradford (1976) method, and the glomalin-related soil protein concentrations (mg g⁻¹ of soil) were estimated using the standard protein curve equation (y = 0.0082x + 0.481; R² = 0.99**), which was obtained by Bovine Serum Albumin (BSA) as a purified standard protein (Purin 2005).

Data were subjected to the normality test and, for the mycorrhizal colonization and number of spores, they were transformed into Log (x + 1). Then, the analysis of variance (Anova) was carried out, and, when significant, the means were compared by the Tukey test at 5 % of probability, using the Sisvar software (Ferreira 2011). Regression analysis was conducted as a function of the cultivation time.

RESULTS AND DISCUSSION

All AMF species exhibited an increased number of spores during the cultivation time (Figure 1), with variation occurring according to the AMF species and the evaluation time, with emphasis on species of the *Acaulospora* genus. The regression equations, coefficients of determination (r²) and significance of the regression coefficients are also shown in Figure 1. Such response of different AMF associated with the same host plant species is considered common, considering that the sporulation capacity is a characteristic of each fungus species, being influenced by the host plant when relating to the compatibility and efficiency of the plant/AMF symbiotic system (Klironomos et al. 2005).

The greatest potential for spore multiplication in *U. brizantha* was observed for *A. longula* followed by *A. colombiana* and *A. morrowiae*, producing 859, 755 and 228 spores (50 mL of substrate), respectively (Figure 1). The regression equation for *A. longula* is: y = 2.627e⁻⁰·⁰⁴⁸⁵x, R² = 0.9821*; for *A. colombiana* is: y = 2.6528e⁻⁰·⁰²⁴⁸x, R² = 0.883*; for *A. morrowiae* is: y = 0.8454e⁻⁰·⁰⁴⁸₇x, R² = 0.954*; and for *G. margarita* is: y = 9.6432e⁻⁰·⁰₁x, R² = 0.9525*.

![Figure 1. Number of spores (NS) of AMF species associated with Urochloa brizantha, as a function of the cultivation time (15, 30, 60, 90 and 120 days of cultivation). * p < 0.05.](image-url)
respectively, at 120 days of cultivation, according to the regression equation (Figure 1). The high multiplication potential of A. longula in Urochloa plants is a typical characteristic of this kind of fungus, corroborating results found by Coelho et al. (2014). On the other hand, the lowest spore multiplication was observed for G. margarita and P. occultum, which exhibited, at 120 days of cultivation, 73 and 34 spores, respectively, in 50 mL of substrate (Figure 1). These results suggest that the use of U. brizantha as a multiplier of G. margarita and P. occultum would require a longer cultivation time (more than 120 days), as previously reported for G. margarita and G. gigantea (Oehl et al. 2009, Jiménez-Martínez et al. 2014).

The number of spores observed in this study shows the efficiency and infective potential of AMF of the Acaulospora genus, whose most species usually exhibit a high capacity of spore multiplication, a fact that confers a wide adaptation of the genus to the most different environments (Oehl et al. 2006). Furthermore, Acaulospora species are considered as having a more “aggressive” establishment, symbiosis formation and multiplication of infective propagules (Stürmer et al. 2006). This genus is very common in Brazilian soils, and the majority of the species are largely found in soils of tropical regions, distributed in different biomes in native and cultivated areas (Stürmer & Siqueira 2011, Ferreira et al. 2012, Assis et al. 2014).

The spore multiplication varied according to the cultivation time and AMF, showing that there is a difference in the interaction of the fungus species when associated with the same host plant. These results suggest that, for the studied AMF, it is necessary a different cultivation period to achieve the maximum multiplication of AMF spores associated with U. brizantha. This behavior has also been observed when assessing the interaction of the same AMF associated with different host plants and substrates (Kadian et al. 2018), showing that, as well as the AMF physiology, the host plant genotype also influences the mycorrhizal symbiosis.

The spore density was proportional to the AMF mycorrhizal colonization, where A. colombiana achieved the greatest mycorrhizal colonization (30 %) in a shorter period of cultivation (at 76 days), followed by A. longula (around 35 % of colonization at 96 days of cultivation), A. morrowiae (32 % at 120 days), P. occultum (around 27 % at 112 days) and G. margarita (16 % of mycorrhizal colonization at 120 days of cultivation), according to the regression equations (Figure 2). These results indicate that there are differences between AMF species, with respect to the establishment of mycorrhizal symbiosis with U. brizantha, with emphasis for A. colombiana, which showed the greatest colonization in a shorter cultivation time. On the other hand, the lowest mycorrhizal colonization was observed for G. margarita at 120 days (Figure 2).

For the inoculation with P. occultum, it was found a sporulation of 73 spores (50 mL of substrate), with 27 % of mycorrhizal colonization at 112 days of cultivation, and, for G. margarita, 34 spores (50 mL of substrate) with 16 % of colonization. The regression equations, coefficients of determination ($R^2$) and significance of the regression coefficients are also shown in Figures 1 and 2. The small number of spores for the species of the Gigaspora genus has also been reported in other studies, as by Jiménez-Martinez et al. (2014), who recorded 20 spores g$^{-1}$ of substrate for G. gigantea.

Studies show that, for the Gigaspora species, a natural germination of spores occurs more slowly, sometimes requiring a longer period of “dormancy” of the spores in the soil (Gazey et al. 1993, Oehl et al. 2009). However, in the present study, such “dormancy” period for G. margarita was not observed, since the soil-inoculum was used immediately after the multiplication of the infective propagules used for the inoculation of the U. brizantha plants.

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![Figure 2](image-url)
Arbuscular mycorrhizal fungi and *Urochloa brizantha*: symbiosis and spore multiplication

AMF species present different colonization strategies, according to the host plant species, cultivation conditions and characteristics of each AMF species (Siqueira & Klauber-Filho 2002, Parniske 2008, Stürmer & Siqueira 2013). This behavior was found in the present study for *A. colombiana*, *A. longula*, *A. morrowiae*, *P. occultum* and *G. margarita* associated with *U. brizantha*. Although it is a plant species commonly used for AMF multiplication and cultivation in culture collections, it was found a variation in the symbiosis establishment according to the cultivation time and AMF species.

For the evaluations of *U. brizantha* growth, in general, it was found a higher inoculation effect with *A. colombiana* and *A. longula* (p < 0.05) (Figures 3 and 4). The *U. brizantha* plants exhibited an increase of up to 91.1 % for shoot dry matter when inoculated with *A. longula*, and 47.08 % for *A. colombiana* at 120 days of cultivation, when compared with the non-inoculated plants.

Most of the inoculated AMF species, except for *G. margarita*, exhibited increased shoot dry matter volumes at 120 days of cultivation (p < 0.05) (Figures 3 and 4). In general, the lowest inoculation effect on the *U. brizantha* plants development was found for *G. margarita* (Figures 3 and 4). This behavior for *G. margarita* is related with the lower infective potential observed in the study, through the low sporulation rate, low mycorrhizal colonization and, consequently, a smaller effect on plant growth.

On the other hand, the inoculation with *A. longula* and *A. colombiana* resulted in more increased amounts of root dry matter of *U. brizantha* at 120 days of cultivation. Root dry matter is a key variable, contributing to the soil organic carbon apportionment, especially in tropical soils, where erosion and organic matter decomposition rates are higher (Lal & Logan 1995).

Carneiro et al. (1999) also found an increased production of root dry matter of *U. decumbens* in degraded areas, which raised from 3.5 Mg ha⁻¹ to 26.7 Mg ha⁻¹ of roots, when inoculated with AMF. The AMF capacity to enhance the growth of the *Urochloa* genus is another key aspect, due to the increased foliar biomass, which results in a greater photosynthetic activity (increased fixation of atmospheric CO₂), favoring the accumulation of organic carbon by roots and, accordingly, increasing the nutrients flow and cycling in the soil/plant system (Wang et al. 2016). Furthermore, increases for shoot dry matter are a desirable condition, considering the importance of the *Urochloa* genus in the formation of pasture/forage crops for animal feed.

As observed for the multiplication of spores and root mycorrhizal colonization, different AMF species also showed distinct behaviors for increased glomalin concentration during the cultivation time, according to the regression equations (Figure 5), especially for inoculation with the *Acaulospora* species, with an increase of up to 40 % for *A. longula*, *A. colombiana* and *P. occultum*, and 25 % and 19 % for *A. morrowiae* and *G. margarita*, when compared...
with the soil not inoculated with AMF, whose initial concentration was 2.67 mg g⁻¹ of glomalin in the soil. The glomalin-related soil protein content at 120 days of cultivation ranged from 3.74 mg g⁻¹ to 3.30 mg g⁻¹ of soil, respectively for the A. longula and G. margarita species, which exhibited a higher and lower glomalin-related soil protein concentration, respectively (Figure 5).

The reduced glomalin-related protein concentration in the soil samples not inoculated with AMF supports the fact that although glomalin is considered recalcitrant, degradation may occur over time by diverse factors present in the soil, especially by the soil microfauna activity (Figure 5).

The increased glomalin-related soil protein concentrations in the AMF-inoculated treatments represent an increase that ranged from 1,000 kg ha⁻¹ for G. margarita and 2,140 kg ha⁻¹ for A. longula. The accumulation of this glycoprotein in the soil is important due to its great capacity of retaining organic carbon, which contributes to the “reduction of the CO₂” emission, contributing to a greater apportionment of C and N in the soil organic fraction (Rillig et al. 2001, Cornis 2002, Wright & Nichols 2002, Lehmann et al. 2017). Therefore, from a practical point of view, glomalin-related soil protein represents a major input of organic C into the soil system, which favors the increase of diversity and biological activity and contributes to an improved soil structure and quality.

In general, inoculation with A. longula showed the best response over time for the number of spores, a high root dry matter increase of up to 89% and shoot dry matter with 91%, and, for glomalin-related soil protein, a 40% increase was recorded, representing a storage of up to 2,140 kg ha⁻¹ of glomalin.

The use of species of the Urochloa genus as an AMF-multiplier plant is already well-known, but still lacking information on the interactions of different AMF for mycorrhizal colonization and spore multiplication to be used in AMF collections. Thus, this study indicates that more information is needed for a better understanding of the symbiotic relationships between different AMF species associated with the same host plant, taking into account the large diversity of AMF species that are known.

These results provide information that may help to support a better management of native AMF, either in pasture or crop lands, for the production of inoculants or “trap cultivation” and for maintenance of AMF collections in germplasm banks.

**CONCLUSIONS**

1. *Acaulospora colombiana* exhibits the greatest mycorrhizal colonization in a shorter cultivation time, at 76 days;
2. The *A. longula* and *A. colombiana* species are more effective in producing spores after 120 days of cultivation with *Urochloa brizantha*;
3. *Paraglomus occultum* and *Gigaspora margarita* require more time of cultivation (over 120 days) for spore multiplication and mycorrhizal colonization of *U. brizantha*;
4. The highest concentration of glomalin is found in plants inoculated with *A. longula*, *A. colombiana* and *P. occultum*.

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**REFERENCES**

ASSIS, P. C. R. et al. Fungos micorrízicos arbusculares em campos de murundus após a conversão para sistemas...
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