OCCURRENCE AND STRUCTURE OF ARBUSCULAR MYCORRHIZAL FUNGAL COMMUNITIES IN CASSAVA AFTER CULTIVATION OF COVER CROPS AS OBSERVED BY THE “PCR-DGGE” TECHNIQUE

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ABSTRACT

Cassava (Manihot esculenta Crantz) is a highly mycotrophic crop, and prior soil cover may affect the density of arbuscular mycorrhizal fungi (AMFs), as well as the composition of the AMFs community in the soil. The aim of this study was to evaluate the occurrence and the structure of AMFs communities in cassava grown after different cover crops, and the effect of the cover crop on mineral nutrition and cassava yield under an organic farming system. The occurrence and structure of the AMFs community was evaluated through polymerase chain reaction (PCR) and denaturing gradient gel electrophoresis (DGGE). A randomized block

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Cassava (or manioc) (Manihot esculenta Crantz) is cultivated throughout Brazil because it can adapt to the variable soil and climate conditions of the country. This crop is closely associated with food security, as it is consumed in several different ways, and cassava products are obtained from the processing of its flour, or by consumption of its roots. In addition, all parts of the plant, such as leaves, stalks, and residue, can be used for animal feed, fertilizer, and pest control (Alves et al., 2009; IBGE, 2010).

Cassava is generally grown by small farmers, with low technological input in terms of nutrient intake. The constant use of cassava by these traditional communities may be influenced both by propagation and multiplication of the plant; propagation material (stem cuttings) is available for new planting after each crop, and the cassava plant is hardy and adapts to environmental variations (Alves, 2006).

The leaf structure of cassava is characterized by low biomass production, with limited soil surface coverage and low input of organic matter; these situations lead to soil loss (Souza and Souza, 2006). In addition, high plant spacing, commonly
adopted for the crop, and leaf shedding during the dormancy period result in higher exposure of the soil to erosive agents (Carvalho and Fukuda, 2006). An alternative to minimize these problems is to use cover crops prior to planting cassava, the main purpose of which is to cover the soil, thus reducing erosion and incorporating organic matter, as well as to increase the fertility and microbial activity in the soil (Bettiol et al., 2002).

In addition to protecting the soil, cover crops play an important role in propogating the populations of arbuscular mycorrhizal fungi (AMF's) (Karasawa and Takebe, 2012). These microorganisms are obligate biotrophic fungi, and they contribute to the nutrition of various crops by increasing the absorption of nutrients that have low soil mobility, as is the case of P (Moreira and Siqueira, 2006). The diversity and the structure of AMF's communities present in the soil are influenced by symbiotic plants (Gomide et al., 2009), which may possibly exert selection pressure on AMF, given the various biochemical mechanisms for recognition among the organisms involved, resulting in some degree of specificity for such symbiosis (Vandenkoonhuyse et al., 2003).

Cassava is highly mycotrophic and has high mycorrhizal dependency since cropped plants have a poorly developed root system in their early stages (Colozzi and Nogueira, 2007). Therefore, the multiplying property of cover crops is an alternative for increasing the inoculum potential of AMF's in the soil (Souza et al., 1999), thus bringing benefits to successive crops of cassava. Legumes and grasses are some of the plant species that contribute most to multiplication of AMF's in the soil, whereas other common cover crops, such as oilseed radish, are non-mycotrophic (Gomide et al., 2009). However, little is known about the level of interference of monocropping or intercropping of cover crops on AMF's communities; cover crops may differ in their ability to select fungi and to affect their multiplication, which can cause changes in the AMF's community (Carrenho et al., 2002).

Spores have been used for assessment of AMF's communities in different agro-ecosystems, (Souza et al., 2010), but further taxonomic knowledge is required to determine the occurrence of those fungi. Nevertheless, AMF's spore density is often unrelated to plant root formation since other propagules, such as hyphae and colonized root fragments, may lead to mycorrhization of plants. In this context, the use of molecular techniques is a tool for better evaluation of the effects of management systems on AMF populations present in the soil (Smith and Read, 2008). In addition, these techniques make it possible to characterize and provide an in-depth description of the structure and succession of microorganisms in the soil (Novais et al., 2010). One of the common molecular techniques, PCR-DGGE (polymerase chain reaction-denaturing gradient gel electrophoresis), is currently used to evaluate the profile of AMF communities from agricultural soils (Gollotte et al., 2004; Liang et al., 2008).

Since cassava is highly mycotrophic, cover crops must cause differences in AMF's community composition, besides affecting performance of the cash crop. The objective of this study was to evaluate an AMF's community, through PCR-DGGE, in cassava (Manihot esculenta Crantz) grown after cover crops, as well as determine the influence of such cultivation on mineral nutrition and yield of cassava grown in an organic system.

**MATERIAL AND METHODS**

The experiment was conducted in an area with an Argissolo Vermelho-Amarelo Distrófico (330 g kg⁻¹ clay) at the Experimental Station of the Empresa de Pesquisa Agropecuária e Extensão Rural de Santa Catarina - Epagri (Agricultural Research Corporation of Santa Catarina), located in the municipality of Urussanga, SC, Brazil, from 7/16/2012 to 8/16/2013. Over the previous seven years, the area grew various species under organic management. The organic fertilization, with respective amount of N-P-K applied, and the results of soil chemical analysis are in table 1.

A randomized block design was used with four replications, and six treatments with different cover crops: oats (O), vetch (V), oilseed radish (R), O + V intercropping, and O + V + R intercropping, as well as a control treatment (fallow), which was mowed every 15 days. The plots measured 19.45 m².

Plots with O, V, and R received 120, 140, and 20 g of seeds, respectively, and the intercropped treatments received O, V, and R at the rate of 30, 40, and 10 g of seeds, respectively. At 110 days after planting, the cover crops were mowed, and cassava stem cuttings (genotype Oriental) began to be planted. Stem cuttings of 0.15 m length and three to four buds were used. Spacing between rows and plants was 0.90 m, with five rows and six stem cuttings per row.

At 110 days after cover crop planting and prior to cassava planting, soil samples were collected from the 0.00-0.10 m layer. Four subsamples were collected from each plot, for a total of 24 composite samples. The samples were used for characterization of the species and total spore counts, evaluation of the Most Probable Number (MPN) of infective propagules, and structural analysis of AMF's communities (Bagyaraj and Stürmer, 2010).

For total spore count, spores were extracted from 50 mL of the soil by the wet sieving technique (Gerdemann and Nicolson, 1963), followed by water and sucrose-gradient centrifugation (Jenkins, 1964). Spore density in the soil was determined by counting...
the spores in a plate under a 40X magnification stereomicroscope (Carl Zeiss MicroImaging, Göttingen, Germany). A sample composed of 24 collected soil samples was sent to the Botany Laboratory of FURB for taxonomic identification of the AMFs species.

For evaluation of the MPN, 60 mL of soil samples were sieved through a 2 mm mesh, diluted in a 10¹⁰ to 10³ decimal series, with five replicates, and homogenized with 540 mL of autoclaved soil. A *Rhizophagus clarus* (UFSC 06) inoculum, containing 953 spores in 50 mL of soil, was used as a reference. The resulting mixtures were placed in 100 mL tubes, sown with sorghum (*Sorghum bicolor*) and kept in a greenhouse for 45 days (Bagyaraj and Stürmer, 2010). Afterwards, the roots were bleached (Koske and Gemma, 1989) and evaluated for the presence or absence of mycorrhizal colonization, and the MPN was determined as described by Alexander (1982).

For evaluations of root colonization and the structure of AMFs communities in the cassava roots, collections were made at 33 and 110 days after planting (DAP) of the crop. The roots were collected randomly, disregarding the plants present in the surroundings, packed in brown paper bags, and kept under refrigeration. The percentage of mycorrhizal colonization was determined by the grid-line intersect method proposed by Giovannetti and Mosse (1980), after whitening and staining the roots (Koske and Gemma, 1989).

Table 1. History of the experimental area, information about previous crops, fertilization and physical-chemical analysis of the *Argissolo Vermelho-Amarelo Distrófico*

<table>
<thead>
<tr>
<th>Year</th>
<th>Crop</th>
<th>Natural fertilization</th>
<th>N-P-K applied kg ha⁻¹</th>
<th>Chemical analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>2006</td>
<td>Tomato</td>
<td>Chicken manure</td>
<td>147 - 209 - 149</td>
<td>22 g kg⁻¹ OM; pH(H₂O) 5.8; SMP Index 6.5; 28.5 mg kg⁻¹ P and 160 mg kg⁻¹ K (both extractable by Mehlich-1); 0.0 cmol, kg⁻¹ Al³⁺, 7.7 cmol, kg⁻¹ Ca²⁺, and 1.3 cmol, kg⁻¹ Mg²⁺ (both extractable by 1 mol L⁻¹ KCl)</td>
</tr>
<tr>
<td></td>
<td>Brassica</td>
<td>Chicken manure + aviary litter</td>
<td>450 - 401 - 232</td>
<td></td>
</tr>
<tr>
<td>2007</td>
<td>Green fertilizers(1)</td>
<td>Chicken manure</td>
<td>450 - 401 - 232</td>
<td>NR(2)</td>
</tr>
<tr>
<td></td>
<td>Cassava</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2008</td>
<td>Tomato</td>
<td>ND(3)</td>
<td>ND</td>
<td>22 g kg⁻¹ OM; pH(H₂O) 5.9; SMP Index 6.2; 78.3 mg kg⁻¹ P and 314 mg kg⁻¹ K (both extractable by Mehlich-1); 0.0 cmol, kg⁻¹ Al³⁺, 5.4 cmol, kg⁻¹ Ca²⁺, and 2.5 cmol, kg⁻¹ Mg²⁺ (both extractable by 1 mol L⁻¹ KCl)</td>
</tr>
<tr>
<td></td>
<td>Maize</td>
<td>ND</td>
<td>ND</td>
<td></td>
</tr>
<tr>
<td>2009</td>
<td>Tomato</td>
<td>Aviary litter</td>
<td>398 - 457 - 238</td>
<td>NR</td>
</tr>
<tr>
<td></td>
<td>Potato</td>
<td></td>
<td>200 - 380 - 280</td>
<td></td>
</tr>
<tr>
<td>2010</td>
<td>Tomato</td>
<td>Chicken manure</td>
<td>396 - 1260 - 208</td>
<td>23 g kg⁻¹ OM; pH(H₂O) 5.5; SMP Index 6.1; 17.5 mg kg⁻¹ P and 222 mg kg⁻¹ K (both extractable by Mehlich-1); 0.0 cmol, kg⁻¹ Al³⁺, 6.6 cmol, kg⁻¹ Ca²⁺, and 0.0 cmol, kg⁻¹ Mg²⁺ (both extractable by 1 mol L⁻¹ KCl)</td>
</tr>
<tr>
<td></td>
<td>Brassica</td>
<td></td>
<td>264 - 840 - 138</td>
<td>NR</td>
</tr>
<tr>
<td>2011</td>
<td>Tomato</td>
<td>Chicken manure</td>
<td>264 - 840 - 138</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Potato</td>
<td></td>
<td>264 - 840 - 138</td>
<td></td>
</tr>
<tr>
<td>2012</td>
<td>Green fertilizers(1)</td>
<td>IE(4)</td>
<td>17 g kg⁻¹ OM; pH(H₂O) 6.1; SMP Index 6.5; 147.9 mg kg⁻¹ P and 241.1 mg kg⁻¹ K (both extractable by Mehlich-1); 0.0 cmol, kg⁻¹ Al³⁺, 3.2 cmol, kg⁻¹ Ca²⁺, and 1.5 cmol, kg⁻¹ Mg²⁺ (both extractable by 1 mol L⁻¹ KCl)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cassava</td>
<td>IE</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(1) Green fertilizers: intercropping - oats, ryegrass, and oilseed radish. (2) NR: Chemical analysis performed. (3) ND: not available. (4) IE: experiment set up.

Evaluation of the structure of AMFs communities, both for soil samples and cassava roots, was performed by the PCR-DGGE method. Total DNA extraction for the soil samples used the PowerSoil® DNA Isolation extraction kit (MOBIO Laboratories, Inc., Carlsbad, CA, USA), according to manufacturer’s instructions. Prior to total DNA extraction from the root samples, the root surface was disinfected by immersion for 1 min in 70 % ethanol, 3 min in sodium hypochlorite solution (2.5 % active chlorine) (v/v), 1 min in 70 % alcohol, and two rinses for 1 min in sterile distilled water (Coombs and Franco, 2003). Subsequently, approximately 0.5 g of disinfected roots was macerated in liquid nitrogen, and DNA was extracted by the CTAB 2 % method (Doyle and Doyle, 1990).

With the DNA extracted for both the soil samples and the roots, partial amplification of the 28S rDNA gene was performed with specific primers for fungi: LR1 (5′ GCA TAT CAA TAA GGG GAG GAA 3′) and LR2 (5′ GTC GTT TAA AGG CAT CAT GTC 5′) (van Tuinen et al., 1998; Trouvelot et al., 1999). The PCR product was used in the second amplification with AMF-specific primers, FLR3 (5′ TGG AAA GGG AAA CGA TTG AAG T 3′) and FLR4 (5′ TAC GTC AAC ATC CTT AAC GAA 3′) (Gollotte et al., 2004).
Amplification in both reactions was performed in a buffer for Taq DNA polymerase, containing 0.2 mmol L\(^{-1}\) dNTPs, 3 mmol L\(^{-1}\) MgCl\(_2\), 1 U Taq DNA polymerase (Invitrogen, São Paulo, Brazil), 10 mmol L\(^{-1}\) of each primer, 10 ng of extracted DNA or DNA from the product of the first PCR (Gollotte et al., 2004). PCR amplification conditions were 5 min at 94 °C, 35 cycles for 1 min at 93 °C, 51 min at 58 °C, and 1 min at 72 °C, and final extension for 10 min at 72 °C.

Equal quantities of amplicons of the second PCR reaction (300 ng) were analyzed by gel electrophoresis with 8 % (w/v) bisacrylamide acrylamide: (37.5:1.0, w:w) containing a gradient of 15 to 55 % formamide and urea. Electrophoresis was carried out at 200 V and constant 60 °C for 4.5 h using a “DCode” system (BioRad, Hercules, CA, USA) and 1X TAE buffer. Gels were stained with Sybr Green (Life Technologies, São Paulo, SP, Brazil) and gel images were made by Gel Logic 220 Pro Imaging System (Carestream Health, New York, USA).

Cassava plants were evaluated at 33 and 110 DAP for leaf P content. Four young leaves near the plant apex were collected, dried at 65 °C to constant weight, and crushed in a Willey mill with 0.5 mm sieves. Leaf P analysis was performed according to Tedesco et al. (1995).

At 33 DAP of cassava, initial plant stand was evaluated for four plants per plot, considering height and stem diameter. At the end of the experiment, at 304 DAP, cassava yield was estimated for four plants per plot. During this period, weeds present in the area were identified.

To evaluate the variables of number of spores, MPN, and mycorrhizal colonization, the data were subjected to analysis of variance, and mean values were compared by Tukey’s test at 5 % probability. Data on number of spores were transformed to log(x + 1), while MPN and arcsine (% colonization/100)\(^{0.5}\), respectively.

The structure of AMFs communities was assessed with the software Gel Compar II (BioSystematica, Wales, UK), using hierarchical cluster analysis, the Jaccard coefficient, and the UPGMA clustering method. The matrix for presence-absence of amplicons, generated by the Jaccard coefficient, was used for calculation of the Venn diagram (http://bioinformatics.psb.ugent.be/webtools/Venn/).

**RESULTS AND DISCUSSION**

**Occurrence of AMF**

Cover crops produced significant effects (p<0.01) on the quantity of infective propagules of AMF in the soil (Table 1). Spore density ranged from 3.8 to 9.8 mL spores mL\(^{-1}\); the highest amount was found in the oats treatment. Similar amounts of spores are commonly found in agricultural areas after oats cultivation (Lermen et al., 2012), but they are considered to be small when compared to soils of natural ecosystems (Stürmer and Siqueira, 2011). The low occurrence of spores may be due to the no-till cropping system, which maintains the balance of the physical-chemical properties of the soil and does not stimulate spore formation. This does not occur in the conventional system, where AMF’s species survival is related to the formation of resistant propagules (Carrenho et al., 2010).

The inoculum potential of AMFs as determined by the MPN method showed that intercropping with oats, vetch, and oilseed radish resulted in a six-fold increase in inoculum potential compared to the treatment with vetch only, whereas the control treatment showed intermediate inoculum potential. However, such MPN values are approximately 100 times lower than those of the *R. clarus* inoculum used in this study as a reference. Nevertheless, the introduction of monocropped or intercropped oats increased the number of AMFs propagules in the soil compared to the vetch treatment, showing that the monocropping or intercropping of oats increases the number of propagules of AMFs. This may be due to the fasciculate roots of this grass and also because it is recognized as a multiplier of AMFs (Gomide et al., 2009). Studies have shown that the production of spores and inoculum potential are not directly associated with plant developmental stages or any climatic or seasonal variable, and there may be situations with no relationship between sporulation and AMFs root colonization (Merryweather and Fitter, 1998).

During spore count, nine AMF species were identified, including *Acaulospora mellea*, *A. morrowiae*, *Dentiscutata heterogama*, *Funneliformis mossae*, *Glomus* sp1, *Glomus* sp2, *Glomus* sp3, *Paraglomus ocov re cultum*, and *Scutellospora* sp (Redecker et al., 2013). In a cassava crop area after 82 days of pre-cultivation with legumes and sorghum, Souza et al. (1999) reported the occurrence of 16 AMFs species, with the genus *Glomus* showing the highest frequency. In addition, the authors found that pre-cultivation of sorghum resulted in significant increases in the number of AMFs propagules in the soil, a similar behavior observed for oats (monocropped or intercropped) in the present study.

The AMFs species identified were widely found in the experimental area, without significant effects of the treatments on the number of identified morphotypes. For this reason, priority was given to assessing the effects of cover crops on AMFs communities by means of the PCR-DGGE technique.

Although there is low inoculum potential in the soil after cover crop cultivation, cassava showed
high mycorrhizal colonization values at 33 DAP (Balota et al., 1999) (Table 2), with the highest rate occurring in the O + V treatment. Even the mowed treatment showed mycorrhizal colonization of 70%, which is considered to be high (Sieverding, 1991).

This high mycorrhizal colonization may be due to the high degree of mycotrophism of cassava in its early growth stages, since cassava has a poorly developed root system (Colozzi and Nogueira, 2007). Indeed, field studies have reported percentages of cassava mycorrhizal colonization ranging from 31 to 85% (Balota et al., 1999; Burns et al., 2012), which confirms the results of this study. Pre-cultivation of the cover crop had little influence on cassava mycorrhizal colonization at 110 DAP, and this may be associated with lower mycorrhizal dependency in the advanced stages of the crop in terms of P nutrition (Ramos and Martins, 2010), as well as root thickening, which hinders the establishment of mycorrhizal symbiosis (Zangaro and Moreira, 2010).

The occurrence of AMFs, assessed by the presence of AMF propagules in the soil and also by cassava mycorrhizal colonization, is influenced not only by cover crop treatments but also by the history of previous crops with mycotrophic plants, such as tomatoes, green manure crops, corn, and potato (Table 1), as well as the presence of weeds commonly found in the area during fallow.

While the experiment was conducted, around 20 weed species were identified, including *Cynodon dactylon* sp. L. (Cynodon), *Digitaria ciliaris* (tropical crabgrass), *Amaranthus viridis* L. (Slender Amaranth), *Rumex obtusifolius* L. (bitter dock), *Commelina erecta* sp. L. (white mouth dayflower), *Veronica persica* Poir. (bird’s-eye), *Cyperus rotundus* sp. L. (nut grass), *Amaranthus deflexus* L. (perennial pigweed), *Stellaria media* (L.) Vill (chickweed), *Sida spinosa* L. (prickly fanpetals), *Silene gallica* L. (catchfly), *Lupinus albus* (white lupin), *Impomoe anil* (L.) Roth (Japanese morning glory), *Brachiaaria plantaginea* (Link) Hitchc (Alexander grass), *Plantago tomentosa* Lam. (plantain), *Phenax sonneratti* (Poir.) Wedd. (Breaks stone), *Oxalis corymbosa* DC. (violet wood-sorrel), *Bidens* sp. (beggar-ticks), *Ageratum conyzoides* L. (goatweed), and *Ipomoea batatas* Poir. (sweet potato), identified according to Lorenzi (2008). Most of these species belong to plant families with the ability to form mycorrhizal association (Purin et al., 2006). Therefore, they have the ability to form an AMF propagule bank that can affect crops grown in succession.

### Structure of AMF communities

There was no similarity between treatments regarding the structure of AMFs communities in the soil, showing the heterogeneous distribution of these fungi and the lack of effect of monocropping or intercropping (Figure 1). The experimental area has a history of use of an organic farming system with the cultivation of various crops (Table 1), and this has a marked effect on the spread of AMF propagules in the area, thus making the structure of these fungal communities heterogeneous. Maybe only one cover crop cycle was not enough to cause notable selectivity of the AMF in the soil. The organic farming system promotes soil cover, incorporation of crop residues, use of organic fertilizers, and crop rotation. Moreover, the practice of organic farming allows better soil fertility conditions, increases organic matter content, improves infiltration of water and aeration, and promotes thermal equilibrium and the formation and stabilization of aggregates (Purin et al., 2006). This scenario may explain the low inoculum potential of AMFs in the soil (Table 2), because the formed aggregates protect, to some extent, the propagules of these fungi, reducing their capacity to associate with the cover crop. However, these propagules are still present in the soil and are eventually identified by PCR-DGGE techniques (Berbara et al., 2006).

The structure of AMFs communities in cassava roots was analyzed at 33 and 110 DAP. The dendrogram shows that there was 100% similarity between the intercropping treatments O + V and O + V + R after 33 DAP (Figure 2a). At 110 DAP, oilseed radish with pre-cultivation was also similar to the intercropping treatments, while the control (mowing) showed low similarity to the other cover crops in both evaluation periods. These results indicate that the profiles of AMFs communities found in the cassava roots are influenced by the cover crops and by the plant developmental stage and the fungal communities are influenced by

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**Table 2. Number of spores and infective propagules of mycorrhizal fungi and mycorrhizal colonization of cassava after cultivation of cover crops**

<table>
<thead>
<tr>
<th>Treatment</th>
<th>No. of infective propagules</th>
<th>Mycorrhizal colonization</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>AMF spores</td>
<td>MPN</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mowing (Control)</td>
<td>4.2 ab</td>
<td>1.2 ab</td>
</tr>
<tr>
<td>Oats (O)</td>
<td>9.9 a</td>
<td>2.1 ab</td>
</tr>
<tr>
<td>Vetch (V)</td>
<td>4.9 ab</td>
<td>0.6 b</td>
</tr>
<tr>
<td>Oilseed radish (R)</td>
<td>5.2 ab</td>
<td>1.1 ab</td>
</tr>
<tr>
<td>O + V</td>
<td>6.6 ab</td>
<td>2.3 ab</td>
</tr>
<tr>
<td>O + V + R</td>
<td>3.8 b</td>
<td>3.9 a</td>
</tr>
<tr>
<td><em>Rhizophagus clarus</em> (reference)</td>
<td>19</td>
<td>183</td>
</tr>
</tbody>
</table>

CV (%) 7.55 2.50 4.88

(1) Days after planting. (2) Means followed by the same lowercase letter among treatments vertically, and (3) means followed by the same capital letter among sampling times horizontally do not differ by the Tukey test at 5%.

the plant developmental stage as well. Samples of roots in pasture areas were evaluated and Vandenkoornhuyse et al. (2003) found temporal changes in the AMFs communities. Therefore, long-term management changes, such as use of fertilizers (Lumini et al., 2010; Lin et al., 2012), may affect the permanence and, consequently, the diversity of AMFs in the soil.

The control treatment (mowing) had fewer amplicons compared to the treatment with single plants at 33 and 110 DAP (Figure 2b). Moreover, fragment distribution was modified along the cassava cycle, and the treatment with oilseed radish showed AMFs specific communities at 33 DAP. However, the AMFs communities were similar among the intercropped and the oilseed radish treatments, with common amplicons among them. Therefore, changes in the succession of fungal communities in cassava roots are evident, and this may be related to the mycorrhizal symbiosis dynamics throughout this crop cycle. As mentioned earlier, in the later stages of plant development, the dependence on AMFs may be reduced by restricting the continuity of certain groups of fungi in the roots. Glomeraceae species show intraradicular hyphae that differentiate into rich lipid globular structures called vesicles (Smith and Read, 2008), which could allow these fungi to survive in the roots, even when there is a limitation in the supply of assimilates by the symbiotic plant. The PCR-DGGE technique reveals its limitation in the impossibility of specific identification of AMF (Øvreås et al., 1997); however, the occurrence of various Glomus species in the soil may indicate that this fungal group has the potential to colonize cassava roots even in advanced crop stages. Therefore, future studies employing the sequencing technique (Lumini et al., 2010; Lin et al., 2012) could be used to identify AMF species in symbiotic succession with cassava roots throughout that crop cycle.

**Plant population, P leaf content and yield of cassava**

Cassava growth, P nutrition, and yield were not affected by cover crop treatments. At 33 DAP, mean values for plant height and diameter were 12 and 0.4 cm, respectively. Mean leaf P content ranged from 5.2 to 6.4 g kg⁻¹ during cassava cultivation, regardless of the cover crop treatment used. Leaf P contents ranging from 2 to 5 g kg⁻¹ are suitable for cassava crops (Howeler, 1987; CQFS-RS/SC, 2004) and, therefore, the values found in this study, regardless of the collection period, are higher than the values recommended in the literature. This can be associated with the high soil fertility (Table 1). At the end of the experiment at 304 DAP, cassava had an average estimated yield of 21 Mg ha⁻¹ and no significant effect from cover crops. This yield is higher than the average yield of the cultivar Oriental in municipalities of the State of Santa Catarina, which is 15.4 Mg ha⁻¹ (Epagri, 2014). Therefore, soil fertility associated with high mycorrhizal colonization of cassava in the early stages of development (Table 2) contributed to P nutrition and yield for this crop.
**Pre-cultivation of cover crops did not affect the structure of AMFs communities in the soil. However, in cassava roots grown in succession, the plant developmental stage interferes with the structural clusters of the AMFs community.**

Despite high availability of P in the soil, the mycorrhizal colonization of cassava was high, and may has influenced leaf phosphorus content and crop yield, regardless of the cover plants used.

One cover crop cycle is not enough to completely clarify the structure of the AMFs communities in cassava crops.

**REFERENCES**


Occurrence and Structure of Arbuscular Mycorrhizal Fungal Communities...


