Specific Polymerase Chain Reaction-Based Assay for the Identification of the Arbucular Mycorrhizal Fungus *Glamus intraradices*

1J.F. Gómez-Leyva, 2J.L. Lara-Reyna, 3L.V. Hernández-Cuevas and 4J.P. Martínez-Soriano
1Laboratorio de Biología Molecular, Instituto Tecnológico de Tlajomulco, Jalisco. Tlajomulco de Zúñiga, Jalisco, México
2Colegio de Postgraduados Campus Campeche, Campeche, Campeche México
3CICB, Universidad Autónoma de Tlaxcala, Tlaxcala, Tlaxcala, México
4CINVESTAV, Campus Guanajuato, Irapuato, Guanajuato, México

**Abstract:** Specific primers and Polymerase Chain Reaction (PCR) assays that identify Arbucular Mycorrhizal fungi (*Glamus intraradices*) were developed. Monoxenic cultures of fungi *G. intraradices* and *Gigaspora gigantea* in association with Ri T-DNA transformed carrot roots were established in order to obtain fungal DNA free of host and others contaminants. RAPD analysis using 10 AM fungi from genera *Glamus*, *Gigaspora* and *Acaulospora* allowed the determination of two amplified fragments that were specific to *G. intraradices*. The DNA fragments were cloned, sequenced and subsequently used to design SCAR species-specific primers. A set of primers, *GIN*cosF* and *GIN*cosR*, drove the amplification of a 630 bp fragment specific for *G. intraradices*, which was absent when DNA of other AM fungi or plants were used as templates. The assay allowed the detection of *G. intraradices* in colonized roots of carrot. The SCAR-based protocol described here may be a tool of great value in studies of Glomeromycota’s molecular systematic and ecology.

**Key words:** Arbucular mycorrhizal, molecular identification, monoxenic culture, RAPD, SCAR, *Agrobacterium rhizogenes*

**INTRODUCTION**

The Phylum Glomeromycota represents an interesting biological group because all its members fall in one of two types of symbiotic mutualistic associations. One of them is constituted for the monotypic family Geosiphonaceae whose only member *Geosiphon pyriformis* forms a peculiar symbiosis named endocytobiosis with cyanobacteria belonging to the genus *Nostoc* (Gehrig et al., 1996). The other symbiotic type is the very common arbucular mycorrhiza, which is formed by the rest of the fungal species. Recent data establish that AM is associated to the roots of 80% of plant species, which indicates the complexity in origin, evolution and diversification of this group (Wang and Qiu, 2006).

Traditionally, the taxonomic identification of AM fungi has been based on the morphological features of spores. Undoubtedly these structures contain important taxonomic information (Sieverding and Oehl, 2006) and the morphological approach to the identification of species although useful to estimate biodiversity it is also limited and controversial. The spores are persistence structures basically formed under unfavorable environmental conditions; their presence may be resultant of past events and not to show prevalent conditions during the sampled period (Lovelock and Ewel, 2005). Moreover, because most of the spores are produced outside the roots it could lead to an underestimation of the biodiversity of AM fungi on field samples. In order to solve these problems recent research has focused on the selection and implementation of tools to facilitate an accurate and reproducible identification of AM fungi (Reddy et al., 2005).

Molecular markers have been specifically developed for the detection and identification of pathogens with impressive accuracy. Furthermore, they have been successfully used in detecting fungi (Nazar et al., 1991; Simon et al., 1992; Berbee and Taylor, 1995). Nevertheless, the sequences used to design the assays have been targeted to ribosomal genes and consequently, they are universal and in some cases no species-specific (Schübler et al., 2001a), but genera or superior taxonomic ranks. Studies on *Glamus intraradices* have proven that
the ITS region of the ribosomal genes is variable; therefore this fact compromises the specificity of the test (Jansa et al., 2002; Reddy et al., 2005).

An interesting strategy to develop species-specific molecular markers is based on the isolation, further sequencing of DNA fragments amplified by RAPD PCR and the use of designed primers to specific target sequences that may be unique for a species. These molecular markers are known as Sequence Characterized Amplified Region or SCARs (Paran and Michelmore, 1993). This approach has been applied to the identification of different species of fungi, including some ectomycorrhizal (Gardebouaf et al., 1997). In contrast, this is not an easy task for MA fungi due to the mutualistic symbiosis with the plant, which makes difficult the isolation of uncontaminated DNA. The use of in vitro root-organ cultures (Bécard and Piché, 1992) may overcome this obstacle; however, there have been very few species on which this system has been successful obtained. The objective of this research was to establish monoxenic cultures of the fungi *G. intraradices* and *Gigaspora gigantea* to obtain pure DNA to further develop a specific test designed to identify the species by the use of a SCAR marker. This method would help to determine accurately the identity of these species in natural ecosystems; moreover, it may contribute to the estimation of the relative abundance of the fungus into the roots.

**MATERIALS AND METHODS**

**Fungal material and strains:** The AM fungi *Glomus intraradices* Schenck and Smith strain 0046TLX03, *G. intraradices* strain BEG 144, *G. caledonium* (Nicol. and Gerd.) Trappe and Gerd. strain BEG 20, *G. clarodendrum* Schenck and Smith strain 0003TLX01, *G. etunicatum* Becker and Gerdemann strain 0004MOR01, *G. fasciculatum* (Thaxter) Gerd. and Trappe emend. Walker and Koske, *G. mossea* (Nicol. and Gerd.), *Gigaspora gigantea* (Nicol. and Gerd.) Gerd and Trappe strain 0033TLX06, *G. margarita* Becker and Hall strain 0036TLX06, *Scutellospora dipurpurascens* Morton and Koske strain 0020TLX06, *S. pellucida* (Nicol. and Schenck) Walker and Sanders strain 0018TLX06, *Acaulospora laevis* Morton strain BEG 78, *A. laevis* Gerdemann and Trappe strain BEG 8, *A. longula* Spain and Schenck strain BEG 8 and *A. spinosa* Walker and Trappe strain 0039TLX01 were obtained in pure pot culture from AM fungi collections of the CICB from the Universidad Autónoma de Tlaxcala (TLX and MOR codes) and the European Bank of Glomales (BEG code) and kept in soil at 4°C.

**Production of hairy roots and monoxenic cultures:** The strains of *Agrobacterium rhizogenes* LBA 9402 and agropine type AR12 were used for DNA transformation of carrot (*Daucus carota* L.) to which the binary vector pBI121 was inserted. Both strains harbored the wild type Ri plasmid. Bacterial cells were grown at 28°C in YEB (10 g L⁻¹ yeast extract, 5 g L⁻¹ beef extract, 5 g L⁻¹ peptone, 5 g L⁻¹ saccharose, 0.49 g L⁻¹ MgSO₄·7H₂O supplemented with 50 mg L⁻¹ of rifampycin plus 100 mg L⁻¹ of kanamycin) to an OD 600 of 0.4. The infection process was performed on transversal disks of carrot co-cultivated with 0.5 mL of the bacterial suspension (OD 600 = 1) for 24 h in the darkness. Tissues were transferred to medium MS (Murashige and Skoog, 1962) supplemented with 500 mg L⁻¹ of ceptonaxime and incubated at 25°C. The bacteria-free hairy roots were removed and transferred to minimum media (M) (Bécard and Piché, 1992) and maintained as clones. Spores of *G. intraradices* and *G. gigantea* were extracted by wet sieving and surface sterilized by treatment with 2% Chloramine T plus 0.1% Tween 20 for 15 min at 4°C, vacuum was applied and the procedure was repeated twice. The spores were transferred to an antibiotic solution (100 mg L⁻¹ of gentamycin sulphate, 2000 mg L⁻¹ streptomycin sulphate) for 24 h at 4°C. After this treatment, the spores were rinsed with double distilled water and gently distributed over a plate containing M medium and incubated at 25°C for 3–4 days. Germinated spores were then transferred to the proximity of the transformed Ri T-DNA carrot roots having active growth to stimulate the colonization. In order to obtain a massive production of spores, the dual system developed by St-Arnaud et al. (1996) was used.

**PCR analysis of transformed roots:** Genomic DNA of the different root clones was extracted according to Doyle and Doyle (1990). Twenty nanogram of genomic DNA was used in a reaction volume of 25 µL containing 15 mM Tris-HCl (pH 8), 0.1% Triton X-100, 50 mM KCl, 1.5 mM MgCl₂, 100 µM of each dNTP, 2.5 U de Taq DNA polymerase (Promega, Madison, WI) and 10 pmol of each of the primers ROLB1 (5' AGT GAT CCT AAA TGG CTA TCC CTT CCA GCA A), ROLB2 (5' TTA GGC TTC TTT CTT CAG GTT TAC TGC AGC), VIRD1 (5' AGT TCG CAA GGA CTT AAG CCCA) and VIRD2 (5' GGA GTC TTT CAG CAT GGA GCA A) (Hamill et al., 1991). These primers drove the amplification of 780 bp and 450 bp fragments from the genes rol B and vir D1 of Agrobacterium, respectively. The amplification was performed using an MJ Research PT-100 thermocycler (MJ Research, Watertown, MA) using the following profile: an initial denaturation step of 94°C/3 min, followed
by a 25 cycles of 94°C/1 min, 55°C/1 min, 72°C/1.5 min and a final extension step of 72°C/7 min then held at 4°C. The amplified products were fractionated in 1.2% agarose gels.

**Fungal DNA extraction and RAPD analysis:** Spores of *G. intraradices* and *G. gigantea* were obtained from the distal part of the monoxenic culture by dissolving Phytagel in 10 mM sodium citrate and crushed in 40 μL of TE buffer (10 mM Tris-HCl pH 8.1, 1 mM EDTA) and heated to 95°C for 20 min in 40 μL of 30% w/v Chelex-100 resin (BioRad). Genomic DNA was separated from cellular debris by centrifugation at 14,000 rpm for 1 min; the resulting extract was diluted and used immediately for use in the PCR assays. Spores of the other AM fungi species were isolated from the soil of propagation pots using the wet sieving and decantation process (Gerdemann and Nicolson, 1963) and were later surface sterilized by treatment with 2% Chloramine T and antibiotics solutions. DNA extraction of these fungal species was conducted according to Lee and Taylor (1991). RAPD reactions were performed in a total volume of 50 μL containing 10 mM Tris-HCl (pH 8.8), 50 mM KCl, 1.5 mM MgCl₂, 0.1% Triton X-100, 100 μM each of dNTP (Invitrogen, Carlsbad, CA), 0.5 μM each random 10-mer (Bio-Synthesis Co., Lewisville, TX), 20 ng of genomic DNA and 2 units of Taq DNA polymerase (Promega, Madison, WI). Thermal Cycler MJ Research PTC-100 was programmed for an initial denaturation at 94°C/3 min, followed by 35 cycles of 94°C/1 min, 36°C/1 min, 72°C/2 min and a final extension at 72°C/7 min. Amplification products were separated in 1.4% agarose gels and stained with ethidium bromide; the DNA bands were visualized under UV light and photographed. Every experiment was performed by duplicate.

**Cloning and sequencing of RAPD markers:** Polymorphic RAPD fragments amplified from *G. intraradices* and *G. gigantea* were purified using the Wizard PCR Prep kit (Promega, Madison, WI) and cloned into the pGEM-T Easy vector (Promega, Madison, WI) as recommended by the manufacturer. The recombinant vectors were used to transform competent *Escherichia coli* cells DH5α. The selection of recombinants was performed by PCR using white colonies directly as source of template DNA that was amplified utilizing the RAPD primers employed for the first amplification. Plasmid DNA from recombinant colonies was purified using the High Pure Plasmid Isolation kit (Boehringer Ingelheim, GmbH, Germany). Insert size was verified by EcoRI digestions, followed by 1.5% agarose gel fractionation. The complete sequence of each cloned fragment was obtained by the use of an automated sequencing robot ABI PRISM 377 (Applied Biosystems, Foster City, CA). DNA sequences were compared by alignment by the DNAsis V 2.0 program for the Macintosh system. DNA sequences for *G. intraradices* were deposited to the GenBank databases.

**Design of SCAR primers and PCR conditions:** The search for DNA similarities was performed using the BLAST and BLASTX programs from the NCBI network service (Altschul et al., 1997). For each RAPD fragment several SCAR primers were designed using the complete DNA sequence for the non-occurrence of secondary structures. The absence of cross hybridization was checked using the PrimerSelect 3.11 software For Windows (DNASTar, Lasergene, Madison, WI). Primers were designed with GC content of 50-60% and synthesized by Invitrogen. The PCR specific assays were carried out in a total volume of 25 μL containing approximately 5 ng DNA, 1X reaction buffer (Promega), 200 μM of each dNTP (Invitrogen), 1.5 mM MgCl₂, 20 pmol SCAR primers, 2.5 U Taq polymerase (Promega). Amplifications with SCAR primers were performed in the MJ-Research PTC-100 as follows: initial denaturation at 94°C/2 min, followed by 30 cycles at 94°C/1 min, 53-58°C/1 min, 72°C/1 min and then a final extension at 72°C/7 min. As proper controls, DNA of every species was amplified by using the universal ITS primers ITS1 and ITS4 as described by White et al. (1990). Amplification products were fractionated by gel electrophoresis in 1.4% agarose gels in 1X TAE buffer (40 mM Tris-acetate, 1 mM EDTA, pH 8), stained with ethidium bromide and visualized under UV light.

**RESULTS AND DISCUSSION**

**Carrot roots transformation and monoxenic culture:** Spores are the only differentiated biologically structures of AM fungi that could be studied outside the host. Even though spores need to be disinfected to remove contaminants, the presence of multiple nuclei makes the interpretation of the RAPD analysis complex. In order to avoid that, we established monoxenic cultures of *G. intraradices* and *G. gigantea*. The culture system requires the maintenance of roots in an autonomous and undefined stage of continuous growth under controlled conditions. In this research, we used two types of *A. rhizogenes* strains such as LBA9402 and AR12 for the production of transformed roots (Fig 1A). Both strains transfer their T-DNA and induced hairy roots free of bacterial cells. Consequently, roots obtained with strain LBA9402 agropine-manopine type resulted in better vigor and branching growth compared with roots produced with
strain AR12. This may be due to a higher accumulation of auxins in roots (Nin et al., 1997). On the other hand, the disinfection method of spores reduced contamination to a 10% with a germination percentage of 70% of both species in agar-water. There are reports about several factors influence the in vitro germination rates of the spores, such as radical exudates, flavonoids, pHe conditions, presence of CO₂, low temperature storage and physiological status of the spore (Bécard et al., 1992; Chabot et al., 1992; Poulin et al., 1993; Juge et al., 2002). In our study germination was successful probably due to the optimal physiological situation of the spores or the disinfection process itself without need to add external components. Four weeks later of the in vitro infection initiation, a massive development of mycelium was observed on the medium accompanied by formation of Branched Absorbing Structures (BAS) as described by Bago et al. (1998).

The spore formation under the dual system (Fig. 1B) as described by St-Arnaud et al. (1996) started four months posterior to the inoculation of *G. intraradices* with an average of 800 spores per dish (Fig. 1C, D), whereas *G. gigantea* only produced 40 spores per dish during the same period.

**RAPD analysis and PCR detection:** Only three of the twenty five primers produced consistent amplification patterns. Primer 5'-TGCAGCACC (Bio-synthesis 70-09), which has a 70% content of G+C, drove the amplification of two specific fragments of 950 and 650 bp for *G. intraradices* (Fig. 2). These fragments were cloned and sequenced (GenBank accession numbers AY244447 and AY244448). The terminal ends of the sequenced fragments perfectly matched the sequence of the 10-mer used for the PCR amplification. From all stock of AM fungi species used in this study, only *G. claroideum*, *G. fasciculatum*, *G. gigantea* and *G. margarita* could be amplified, all other produce no satisfactory results, which reinforce that AM fungi DNA extraction and amplification is a difficult task for almost all species, specially if these species are not been propagated monoxenically.

Based on the disclosed DNA sequences, several set primers were designed, synthesized and used to optimize PCR assay and the amplification of a unique fragment to accurately identify *G. intraradices*. Primers GIN₉₀F (5'-TGC AGC ACC GCC TCC ACC) and GIN₉₀R (5'-TGC AGC ACC GTC GCT TGT TA) drove the amplification of an expected 930 bp fragment. This method encouraged the detection of *G. intraradices* strain 0046TLX03 in fragments of infected in vitro roots of *D. carota* and in vivo roots of *Sorghum* sp., colonized with *G. intraradices* strain BEG144 (data not shown).

In addition, consistent results were obtained with primer set GIN₉₀F (5'- GCA CCG CAA GTT AAG TAC
Fig. 2: Random amplified polymorphic DNA (RAPD) fingerprints using genomic DNA of mycorrhizal arbuscular fungi. Lanes 1-2 *Glomus claroideum*, 3-4 *G. fasciculatum*, 5-6 *G. intraradices* 0046TLX03, 7-8 *Gigaspora gigantea*, 9-10 *G. margarita*. M 100 bp DNA ladder. Arrows indicate species-specific and reproducible RAPD bands of 950 and 650 bp that were converted into SCAR markers.

Fig. 3: (A) Agarose gel electrophoresis of PCR products obtained using the *Glomus intraradices*-specific primers GIN<sub>10</sub>F/R designed in this study. (B) Agarose gel electrophoresis of ITS 1/4 primers used as internal controls. M 100 bp DNA ladder. Lanes (1) genomic DNA from *Glomus intraradices* 0046TLX03, (2) carrot root colonized with *G. intraradices* BEG144, (3) *G. fasciculatum*, (4) *G. claroideum*, (5) *G. mosseae*, (6) *Acaulospora laevis*, (7) *Daucus carota*.

CCA AC) and GIN<sub>10</sub>R (5'-CCG TGA TCA TGA TGT CTC AGG TT). Annealing temperatures of 54°C were used to produce the expected fragment of 630 bp. Lower temperatures allowed the amplification of an unspecific fragment of 800 bp. The test proved to be highly specific to *G. intraradices* and no amplified DNA was observed when genomic DNA from others AM fungi was used as template (Fig. 3A). Controls DNA of every species were successfully amplified by using the universal ITS primers (Fig. 3B).

There are reports of PCR-based tests that have been presented as specific for several fungi of the order Glomerales (cited as Glomales by Simon, 1996). Many of them have been developed by using ribosomal DNA as template such as the VANSI primer set proposed by Simon *et al.* (1993). Recently, it has been demonstrated that these primers are not specific and that primer homologous sequences are absent in at least 88 of the MA fungi analyzed (Lloyd-MacGilp *et al.*, 1996; Schübler *et al.*, 2001b; Sanders, 2003). Moreover, for
G. mosseae alone, at least 23 sequences with homology varying from 66 to 98% have been reported by Antonioli et al. (2000).

The nucleotide sequences corresponding to SCAR fragments reported here may be of vital importance for the development of quantitative PCR assays for the study of G. intraradices. The distribution of the AM fungus in the host roots could be now elucidated and consequently a better understanding of spore abundance in the soil and infection process.

This research has successfully demonstrated the use of monoxenic cultivation of two AM fungi and its further use to analyze RAPD patterns in order to develop specific PCR tests based on the selection of unique SCAR sequences for different species. Furthermore, the use of this approach contributed to the accurate detection and identification of G. intraradices on in vitro and in vivo tests. The test offers promising use for further studies on the ecology of the mycorrhiza-plant interaction in nature.

ACKNOWLEDGMENTS

This study was supported by funds from Consejo Nacional de Ciencia y Tecnologia (CONACyT) grant number 26357B. We thank Juan Carlos Ochoa-Sanchez, Magali Hernandez-Valencia and Juan Enrique Cortes-Valle for his technical support.

REFERENCES


