GERMINATION AND GERM TUBE GROWTH OF THE ARBUSCULAR MYCORRHIZAL FUNGI 

**GIGASPOR A L B I D A** IN DIFFERENT SUBSTRATES

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**ABSTRACT**

The role of the substrate on germination of *Gigaspora albida* Schenck & Smith was investigated. Spores were desinfested with 0.5% sodium hypochlorite (20 min.) and placed on Petri dishes over a Millipore filter, with one of the following media: a- 1% water-agar; b- water-agar + aqueous extract of roots of *Panicum miliaceum* L.; c- salt medium of Murashige & Skoog (MS) or sterilized sand; and incubated in the dark at room temperature (28ºC ± 2). The experimental design was at random with four treatments and four replicates. Germination was evaluated every 7 days until the 28th day. The water-agar medium was the most feasible for spore germination at the 7th day, followed by the MS medium at the 14th day. Conversely, the sand and the root extract medium did not allow high germination. Spores maintained in water-agar also presented longer germ tubes than spores in the other treatments. Auxiliary cells were observed at the beginning of formation of hyphal branching in all treatments, however they were more numerous in the water-agar medium.

**Key words:** water-agar, MS medium, root extract, Glomales, *Panicum miliaceum*.

**INTRODUCTION**

Arbuscular mycorrhizal fungi (AMF) are associated with plant roots and enhance host growth through increased nutrient uptake. The life cycle of these fungi is initiated by germination of spores which, isolated or in sporocarps, are formed in the soil or sometimes inside the roots (36).

Spores are the most important propagules for most AMF and the impact these organisms produce on their hosts will depend on the ability for fast spore germination and colonization (38).

Spore germination may be affected by many factors, such as: the need for a dormancy period (32), root exudates and/or volatiles; soil moisture, temperature, pH, light and CO₂ (3, 10, 13, 35); flavonoids (3, 12); as well as by presence of bacteria (1), but is independent of the presence of a susceptible host. When Daniels and Graham (6) supplied a 0.75% water-agar medium with a soil extract (200 g soil/L), they observed an increase in the germination rate of *Glomus mosseae* (Nicol. & Gerd.) Gerd. & Trappe spores and mentioned that this resulted from the nutrients extracted from soil. Paula *et al.* (28) observed that the rates of germination and mycelial growth of AMF increased 90% and 200%, respectively, when the water-agar medium was supplemented with plant cells. The greatest increase was observed with the addition of cells of *Pueraria phaseoloides* Benth. to the medium. Improvement of germination of two *Gigaspora* species, *G. rosea* Nicolson & Schenck and *G. decipiens* Hall & Abbott was obtained when spores were placed in different concentrations of cellulase from *Trichoderma reesei* E. G. Simmons and other fungi (30). Aminoacids may inhibit, activate, or have no effect on mycelial growth of *Gigaspora gigantea* (Nicol. & Gerd.) Gerd. & Trappe pre-germinated in water-agar, depending on the concentration in which they are available in the medium (9). Root membrane permeability under low phosphorus nutrition, lead to net loss of metabolites at levels enough to sustain germination and growth of AMF during pre and post infection (14).

High level of germination in non sterilized soil was reported, although autoclaved, pasteurized or gamma-irradiated soils...
may inhibit the process (7). Germination is also affected by temperature: the ideal temperature for germination of Acaulospora laevis Gerd. & Trappe was 20ºC, whilst the optimal for hyphal growth was 15 to 25ºC (39). However, higher germination and hyphal elongation of an isolate of this species in temperatures that varied from 30 to 35ºC was also observed (24).

Another important factor for germination of AMF is the pH (15). Spores of A. laevis maintained a few weeks at 6ºC, were exposed to different pH levels, and germination was higher in low pH. In addition, only 3% of spores germinated in water-agar, while the rate of germination of spores incubated between filter membrane in soil was higher than 70%.

Many factors such as moisture, aeration, light conditions, microbial activity, and nutrient toxicity affect the abundance, viability, and infectivity of propagules, and therefore directly affect the storage potential of the spores (5, 22, 35). AMF isolated from temperate environments should be maintained in environments between 20 to 25ºC. Some germination of spores, whereas those from tropical regions temperatures from 4 to 10ºC to break dormancy and enhance viability, and infectivity of propagules, and therefore directly affect the storage potential of the spores (5, 22, 35). AMF isolated from temperate environments should be maintained in environments between 20 to 25ºC. Some species, such as Gigaspora albida Schenck & Smith may produce multiple germ tubes in vitro (20) which were related to strategies for fungal survival (18).

Considering that information regarding AMF spore germination is still incomplete and that this part of the life cycle is essential for species preservation, the objective of this research was to evaluate the effect of different substrates on germination of spores of G. albida.

**MATERIALS AND METHODS**

Spores of G. albida (INVAM 927), grown in pot culture with bahiagrass (Paspalum notatum Flugge) as host plant, were extracted from soil by wet sieving and sucrose centrifugation (11, 17). Healthy spores were washed with distilled water, immersed in 0.5% sodium hypochlorite (NaOCl) for 2 min., and then washed four times in sterilized distilled water (26).

The following substrates were used: A – 1% water agar (Bacto agar); B. MS salts - Sigma (23), supplemented with 0.2 ml/L of claforan (sodic cephotaxima) and 1% agar; C – aqueous root extract of Panicum miliaceum L. (50 g of macerated roots plus 100 ml of distilled water) with 1% agar and 0.2 ml/L of previously autoclaved claforan; D – sand, sterilized in a microwave oven (8) and placed in covered 46 x 15 x 6.7 cm Nalgene sterilizing trays (Fisher Scientific, Inc., Orlando, Florida). Media A and B were autoclaved at 121ºC, 1 atm., for 20 min.

Five spores were transferred to Millipore filters (Gelman Sciences Inc., Ann Arbor, Michigan), and placed on 9 cm Petri dishes containing media A, B, and C. Spores were also placed between two Millipore filters and transferred to trays with autoclaved sand (medium D). Each Petri dish and box was considered as a replicate, with a total of four replicates per media. Both, Petri dishes and trays with spores were incubated in the dark at room temperature (28ºC ± 2).

Evaluations were made every 7 days for 28 days. The hypha were stained with drops of 0.05% aqueous Trypan blue and germination was observed at 40x using a stereomicroscope. Spores which produced germ tubes longer than their diameter were considered to have germinated (2). Germinated spores were expressed as percentage of the total number of evaluated spores. Hyphal length was estimated using the gridline intersect method (25).

The experimental design was a complete factorial of 4 treatments x 4 evaluations periods, each with 4 replicates. Data expressed as percentage were arcsin transformed before analyses of variance. The Statistica 5.0 program (37) was used for statistical analysis.

**RESULTS AND DISCUSSION**

Germination of this G. albida isolate was affected by substrate. Spores on water agar and MS medium presented the highest germination percentages, which were significantly greater than those in sand and water-agar with root extract (Table 1).

Root exudates of Troyer cirtange (Poncirus trifoliata x Citrus sinensis) and Sudangrass positively influenced germination and germ tube length and branching of Glomus epigaeum Daniels and Trappe (13). Hyphal growth of G. margarita Becker & Hall was stimulated by cell exudates or extracts of Pueraria phaseoloides but was inhibited under high concentration of these substances (27). The results here obtained were different; only 33% of the spores germinated in the root extract medium. Extract of Asparagus roots did not affect germination of some AMF (7). Although these root extracts are from mycorrhizal susceptible hosts (21), other factors are involved on AMF germination and colonization.

| Table 1. Germination of spores and length of germ tubes of Gigaspora albida in: a - water-agar 1%, b - medium with MS salts, c - water-agar 1% plus Panicum miliaceum root extract, and d - sand (28 days of incubation). |
|-------------------|-------------------|-------------------|
| **Substrate**     | **Germination (%)** | **Length of germ tube (mm)** |
| Water-agar 1%     | 100 a             | 59.62 a           |
| MS                | 70 ab             | 43.32 ab          |
| Sand              | 40 b              | 39.00 b           |
| Root extract      | 22.5 b            | 51.56 ab          |

* Mean number of germinated spores. Media followed by the same letter do not differ by the Tukey test at 0.05.
Differences in spore germination of AMF due to agar quality were reported: among some types, the Bacto agar allowed higher (83%) germination (6). The authors concluded that germination increased according to the increase of agar purity. When using Bacto agar we obtained 88% of germination, what indicate that it might be favorable to different AMF species.

Germination of *G. albida* increased over time, depending on the substrate. In water agar, spores reached 100% of germination at the 7th day. It was reported that the highest germination percentage of *Glomus mosseae* in 0.75% water agar occurred on the 26th day (6). In soil and MS medium, the highest level of germination was registered at the 14th day, while spores in root extract had a progressive increase of germination over time but reached only 40% of germination up to the 28th day of incubation (Fig. 1). In this case, it is possible that the germination had been related to some inhibitory substance present in the root extract, which became inactive with time. Siqueira et al. (34) referred inhibition of germination when organic substances were added to the medium.

More than one germ tube may be produced during germination of spores of *G. albida* in water agar medium supplied with thiamine (20). However, the same was not observed here. Thus, multiple emission of germ tubes may result from addition of thiamine or other stimulatory substance to the medium. Siqueira et al. (34) mentioned that this substance is part of root exudates. It should also be considered that emission of many germ tubes may be stimulated by substrate conditions, genetic information of the spore, and by physical pressure (35). Emission of more than one germ tube in *Gigaspora* species may also be due, among other factors, to formation of a germ wall and to the high amount of nuclei near the wall (4).

The higher rate of germination in the MS medium was obtained after 14 days of incubation. Higher germination and micelial growth of *G. gigantea*, *G. margarita*, and *Scutellospora heterogama* was observed when the MS medium was 10x diluted (29). They concluded that high germination was due to decrease in nutrient concentration. Excess of mineral salts in the medium may inhibit germination (36). However, this was not observed here, even using the recommended amount of salts in the MS medium.

Addition of a small amount of phosphorus to the agar medium may enhance germination and growth of germ tube (35). The results here obtained are in agreement with this observation, once that the MS medium contained potassium phosphate (170 mg.L⁻¹). Concentration up to 500 mg.L⁻¹ of P (KH₂PO₄) did not inhibit germination of some AMF spores (18). Conversely, decrease in germination of spores of *Glomus mosseae* was observed when P was added to the medium (31). In a similar way, it was reported that in pasteurized soil, *G. etunicatum* spore germination decreased with increasing P levels up to 60 mg.L⁻¹ (40).

The highest rate of germination in sand was obtained at the 14th day, with almost 65% of the spores germinated. However, rate of germination higher than 70% in spores of *A. laevis* placed between filter membranes in soil was observed (15). Disinfestation of the sand through microwave may have impaired germination once that the soil microbiota was eliminated (7, 16) and some studies have shown that germination may depend upon a microbial stimulus in non sterile soil (6).

Auxiliary cells were formed in all substrates, mainly after initiation of the branching process in the first hyphae. These structures were more frequently formed in the water agar (Fig. 2). Formation of auxiliary cells are not dependent of the growth medium but are supposedly related with the intrinsic capacity of the spores (33).
Evaluation of the germ tube length over number of germinated spores showed that these parameters are not related; that is, higher rate of germination does not imply formation of longer germ tubes and the opposite is also true. This was observed in studies (2) with G. albida (GABD 185) and G. gigantea (GGGT 109). After germination, AMF have an energy-saving mechanism which allows long-term infectivity of mycelium even in absence of the host (19) and this would probably depend on spore size (2). Spores maintained in water agar and in the root extract medium presented greater germ tube lengths than those in MS or sand; however, only the treatment in water agar and MS differed significantly (Table 1).

Difficulties in following hyphal growth outside the Millipore filters impaired the evaluation of the effect of time over germ tube length. The lower hyphal length obtained in this work, when compared with other values (3), might be due to the amount of CO$_2$ and flavonols used by the other authors. These factors were not studied in this experiment. On the other hand, the values here obtained were higher than those registered in a previous investigation (2) where germ tubes formed by G. albida, in soil, were shorter than 50 mm, after 21 days of incubation. This indicates that several factors determine germ tube growth, even of a same AMF species.

Considering that the effect of the substrate on germination and germ tube growth was confirmed, it might be recommended the use of water agar to improve both processes when working with G. albida.

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REFERENCES


