

SOP 001

Date: 11.09.2023

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# STANDARD OPERATING PROCEDURE

# METHOD FOR ISOLATION AND OBSERVATION OF AMF SPORES FROM SOIL

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## **METHOD DOCUMENT CONTROL LOG**

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#### Changes in this version compared to previous version:

- 1. Inclusion of workflow
- 2. Inclusion of quality control document

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## **ABBREVIATIONS**

• μm - micrometre

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- cm centimetre
- g/l gram per litre
- GLP Good Laboratory Practices
- M molar
- Min minute
- MI millilitre
- Rpm rotation per minute
- Sec second
- v/v volume per volume
- AMF- Arbuscular Mycorrhiza Fungi

#### **DEFINITIONS**

- a) **Contaminated by microorganisms**: Every plate, tube, pipette, or other instruments (glassware, pestles, ependorff tube...) which has been in contact with microorganisms and cannot be sterilized by the flame of a Bunsen burner is considered as contaminated.
- b) **Contaminated by toxic chemicals**: Every tube, flask, pipette or other instruments (Weighing boats, glassware...) which has been in contact with toxic chemicals is considered as contaminated.
- c) **Good Laboratory Practices (GLP):** The Principles of Good Laboratory Practices (GLP) have been developed to promote the quality and validity of results and of the analysis conducted in a laboratory. It is a managerial concept covering the organization and the conditions under which laboratory studies are planned, performed, monitored, recorded and reported. Its principles also include the protection of man and the environment.

#### **1. INTRODUCTION**

The most successful arbuscular mycorrhizae are formed with the fungal phylum Glomeromycota. Arbuscular mycorrhizal fungi penetrate the cortical cells of the roots to form unique structures

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such as internal hyphae, arbuscules and coils, vesicles. The fungi exist in the soil as spores, auxiliary cells or hyphal net.

The fungal structures: spores, arbuscules, vesicles, hyphae and auxiliary cells are all used to diagnose families, genera and species and the status of the symbiosis and its functions in plants and soils. Methods for evaluating functions range from morphological, biochemical to molecular. This procedure describes the detailed protocol to isolate arbuscular mycorrhizal fungal (AMF) spores from either field soil of artificial substrate mixtures used for the pot cultures of the AMF.

#### 2. SCOPE AND APPLICATION

Arbuscular mycorrhiza fungi (AMF) are soil microorganisms that can form mutualistic relationships with most terrestrial plants. AMF spores germinate, infect the root system, and form arbuscules structures inside the cells of the plants. Arbuscules serve as sites for nutrient exchange between the plant and the fungi. Another distinguishing feature of this symbiosis is the ability to form dense mycorrhizal network surrounding the root system aids in enhanced water and nutrient uptake by plants roots.

## **3. PRINCIPLE**

Mycorrhizal has been in use as a biofertilizer especially in agricultural lands for maize, barley, wheat and oats. The product boosts the performance and vitality of plants. Application of mycorrhiza in agricultural fields is also sustainable compared to continuous application of fertilizers. An example of Mycorrhiza product is MycoApply

## 4. PROCEDURES

## **5.1 MATERIALS, EQUIPMENT AND REAGENTS**

#### i. Materials

- a) Centrifuge with a swinging bucket rotor
- b) Waring blender
- c) Microscope

#### ii. Equipment

- a) Coverslip (preferably circular, variable diameter depending on spore size)
- b) Sieves (710  $\mu m$  and 45  $\mu m$  or 500 and 53  $\mu m)$
- c) Petri dishes, 9 cm in diameter
- d) Pipette and Pro pipette 5 ml



- e) Beakers (Small and big)
- f) Spatula

#### iii. Reagents

- a) Melzer's reagent (Chloral Hydrate, 484.3 g/l; Iodine, 7.3 g/l; Potassium Iodide, 24.2 g/l)
- b) Polyvinyl-Lacto-glycerol (PVLG) (Polyvinyl alcohol, 73.3 g/l; Lactic acid, 44% (v/v); Glycerol, 4.4% (v/v))
- c) Sugar/sucrose solution (2.5 M or 48%)

#### **5.2 ANALYSIS DESCRIPTION**

[For a detailed description of our methodology please visit or contact us]

## **5.2.1 Mounting slides and observations**

- Voucher specimens of permanent slides (slides sealed with colorless nail varnish at 5-10 days after mounting) are prepared as reference material.
- Prepare three slides: On the first one, place two drops of water at opposite ends of a microscope slide. On the second, place one drop of PVLG on each end and on the third one, put PVLG + Melzers (ratio 1 :1) on both ends.
- Pick carefully spores with a fine forceps or micro-pipette and place in each drop.
- Lower a coverslip, preferably circular with diameter of 13 mm onto each drop of mounting reagent + spore.
- For each slide, observe the spore as whole for diagnosis of appearance on one end and on the other end, crush the spore gently and severely. A needle or pin may be used to crush the spores. Caution must be taken not to break the coverslip.
- In case of bubbles, gently place ethanol though the space between slide and coverslip to clear bubble, absorb ethanol with tissue and then gently place the mounting reagent again.

## **5. MAINTENANCE**

Servicing of the Microscope, Luminar flow, Centrifuge and Blender to be done twice annually.

## 6. OCCUPATIONAL HEALTH AND SAFETY

• All activities performed under this SOP comply with the recommendations of ICRAF Health and Safety policy

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- Laboratory dress code: clean laboratory coats and low-heeled closed shoes worn at all times while performing the procedures.
- Hand gloves are worn when handling chemicals and the microorganisms
- Laboratory working areas are cleaned daily and benches regularly disinfected and sterilized using 70% alcohol during various procedures.

## 7. REFERENCES

 Brundrett, M., Abbott, L.K. and Jasper, D.A. (1999). Glomalean mycorrhizal fungi from tropical Australia. I. Comparison of the effectiveness and specificity of different isolation procedures. Mycorrhiza 8, 305-314.

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- Wagner, S.C., Skipper, H.D., Walley, F. and Bridges, W.B. (2001). Long-term survival of Glomus claroideum propagules from soil pot cultures under simulated conditions. Mycologia 93, 815-820.

## **6.** ANNEX



• For description of spores refer to <u>INVAM WEBSITE</u> and Schenck and Perez, 1990.