STANDARDIZATION OF THE SPORE DENSITY OF AM FUNGAL INOCULUM FOR EFFECTIVE COLONIZATION

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Abstract- A pot culture experiment was conducted to develop quality standards for the Arbuscular Mycorrhiza (AM) inoculum in terms of spore density in order to ensure the efficacy of the AM product when used for crop production as a P mobilizing biofertilizer. Plant growth parameters like root volume and total plant dry weight of maize (PEHM5) which was used as trap crop skyrocketed to its maximum when inoculated with 5-6 spores g⁻¹ of AM inoculum. Effective root colonization of 100% was recorded in maize when 5-6 spores, was present. Total root length colonization of the maize variety PEHM5 was registered to be more than 90%, when treated with an AM inoculum containing 5-6 spores g⁻¹. Moreover, a significant increase of the spore number was observed, when sterilized vermiculite + 10% soil was used as substrate for the AM inoculum production with increased spore density of 5-6 spores g⁻¹. Thus, spore density of 5-6 spores was standardized to be one of the quality control parameter of AM inoculum which can be adopted to certify a high quality AM product for crop production.

Key words- Arbuscular mycorrhizal (AM) fungi, inoculum, spore density and colonization

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Introduction
Arbuscular mycorrhizal (AM) fungi have shown to have a pivotal role in agriculture and land reclamation (Klironomos et al., 2002). AM inoculation of plants in field offers the possibility of reducing fertilizer and pesticide application and it improves the resistance to diseases caused by soil borne pathogens. Therefore, AM fungi are gaining popularity as 'Biofertilizer', 'Bioprotectant' and 'Biocontrol agent' and the industry of mycorrhizal inoculum production is expanding around the world (Feldmann and Grolkan, 2002). The growing number of new small to medium sized companies (SME’s) around the world producing inocula of mycorrhizal fungi has increased in the last decade. Still, there are major problems in bringing high quality and fit for purpose AM fungal products to target markets. Because AM fungi are obligate symbionts, the production of AM fungal inocula of the best quality is still challenging.

Moreover, the quality control of AM inoculum producing companies is not yet under control of independent institutions. Therefore, more accurate methods are required to ensure a high quality and consistent AM inoculant. At present, regulation of the product varies between countries and companies for the AM inoculant and each company still has its specific demands and target markets (Feldmann, 2007). Thus, by formulating quality standards for AM inoculants especially in terms of spore density, since spores of AM are only instrumental to bring a better colonization in plants and bring in the P mobilization that aid in ultimate plant growth, it would be more feasible to manipulate these fungi for increased agricultural productivity and plant health assurance, so that the end users can be satisfied by the producers and both will be benefited. Hence, the present investigation was undertaken to develop quality standards for AM inoculant in terms of spore density.

Materials and methods
This experiment was undertaken to standardize the suitable spore density for arbuscular mycorrhizal fungal inoculum using vermiculite as substrate and maize as host plant under pot culture conditions.
Standardization of the Spore Density of AM Fungal Inoculum for Effective Colonization

Production of arbuscular mycorrhizal (AM) mother inoculum for the Study
The arbuscular mycorrhizal fungal strain of Glomus intraradices (Tamil Nadu Agricultural University, Coimbatore) was obtained and multiplied in the net house at the Department of Agricultural Microbiology, TNAU, Coimbatore. Maize (Zea Mays) was used as host plant to multiply the arbuscular mycorrhizal fungi. Fifty live spores of Glomus intraradices was placed under germinated seedlings of maize into pots containing 1:1 sterile sand: soil mixture. The pots were watered regularly along with a spray of Hogland's plant nutrient solution (once in a week) and infection was allowed to develop in the host plant for at least three months before the inoculum was harvested. The ‘mother’ inoculum (containing spores of Glomus intraradices, hyphae and colonized root bits) was obtained by cutting all the roots of host plants to 1cm size and mixing it with the pot mixture.

The ‘mother’ inoculum was used for a series of pot culture experiments conducted to develop the quality standards for arbuscular mycorrhizal inoculum.

Isolation of AM spores from the mother inoculum
AM spores were isolated using the method of wet sieving and decantation with slight modifications as described by Gerdemann and Nicolson (1963) from the ‘mother’ inoculum. About 100 g of the AM inoculum was taken in a one litre beaker and added with ten times its volume of water i.e., water added to the sample in 1:10 ratio. This suspension was stirred vigorously using a glass rod and allowed to settle. The suspension was then decanted or poured through a sieve arrangement from coarse sieve to fine sieve. The sizes of the sieves were arranged as 180 µ, 106 µ and 45 µ from top to bottom. This process was repeated several times, until the suspension becomes colourless. The sievings were transferred to a separate beaker and filtered through Whatman No. 1 filter paper. The filter paper was observed under Nikon SMZ - 10 stereo zoom microscope for the presence of spores. The young, viable spores of Glomus intraradices were collected using capillary tube into a sterilized vial containing Ringer’s solution (Sodium chloride - 5.0 g, Calcium chloride - 0.01 g, Potassium chloride - 0.01 g, Magnesium chloride - 0.01 g and distilled water - 100 ml) and were maintained at 4°C until use.

Treatment Details

- **T1** - Uninoculated control
- **T2** - 1 to 3 spores g⁻¹ of AM inoculum
- **T3** - 5 to 6 spores g⁻¹ of AM inoculum
- **T4** - 10 to 12 spores g⁻¹ of AM inoculum

**Levels**
- **S1** - Vermiculite alone
- **S2** - Vermiculite + 10 % sterile soil

The treatments were replicated five times and were arranged in completely randomized block design.

Substrate
Raw vermiculite (Grade IV) obtained from Tamil Nadu Minerals, Chennai was used in this experiment, with and without the amendment of 10% sterile field soil collected from Eastern Block, TNAU Coimbatore. The substrate was sterilized at 15 lbs pressure for 30 minutes and used for the study. The substrates as per the treatments were filled in black pots (15 x 15 cm), about 1 kg pot⁻¹.

Preparation of AM inoculum
About 5 g of sterile vermiculite was mixed with the respective quantity of AM spores of Glomus intraradices isolated from the ‘mother’ inoculum and was used as the inoculum for the conduct of this experiment. AM inoculum developed with different spore densities were applied approximately 2.5 cm below the pot mix. The rate of inoculum was 5 g pot⁻¹ filled with 1 kg of substrate.

AM multiplication and sampling
Seeds of maize (variety PEHM 5) obtained from the State Department of Agriculture, Thondamuthur was used for the study. The seeds were surface sterilized with 0.1 percent mercuric chloride for 3 minutes before sowing and were sown 3 cm below the surface of the substrate. About five plants were maintained in each pot. The pots were watered regularly. Once in 15 days each pot was given with 20 ml of 2x Hoagland solution minus phosphate (Hoagland and Arnon, 1950). Plant samples were taken from each treatment on 45 and 60 days after sowing.

Observations
Root volume and plant dry weight were recorded. Arbuscular mycorrhizal colonization in root, spore density and total microbial load in the substrates were also estimated. In addition, total root length colonization of maize by AM was calculated.

Plant dry weight
The uprooted plants were sun dried for one day and then dried in hot air oven at 60 to 70°C for three consecutive days till the constant dry weight was obtained. The dry weights were recorded and expressed in g plant⁻¹.

Root volume
Volume of the root samples were determined by volume displacement analysis (Burdette, 1979). Initially, a graduated cylinder that would be large enough to fit the root being measured was selected. After filling the graduated cylinder with water to a satisfactory level, the volume of water alone was recorded. After that, the root was carefully placed in the graduated cylinder and the volume of water and root was recorded. In order to calculate the volume of the irregularly shaped root, the volume of the water alone was subtracted from the volume of the water and root and the values were expressed in cm³ plant⁻¹.

Per cent AM colonization = Number of root bits colonized

AM fungal colonization in roots of the host plant
Roots of maize uprooted on 45th and 60th days after sowing were assessed for AM fungal colonization by following root clearing and staining technique developed by Phillips and Hayman (1970). The uprooted roots were thoroughly washed in tap water to remove any soil or vermiculite particles. The roots were then cut into small bits of about 1 cm long and immersed in 10 % potassium hydroxide solution. This was autoclaved with 5 lbs pressure for 10 minutes, and washed with water till the brown colour disappeared. Then the roots were immersed in 2 % hydrochloric acid for about 5 to 10 minutes, and the acid was decanted. Tryphan blue 0.05 %
prepared with lactoglycerol was added (45 ml of lactic acid + 45 ml of glycerol + 10 ml of distilled water was mixed and added with 0.05 g Tryphan blue) after fixation with 2 % HCl. After staining, the root bits were kept as such overnight. Observation of the stained root bits for fungal infection under Olympus CX 40 microscope was carried out after destaining the stained roots with lactoglycerol (1:1) or glycerol 50 % in order to remove the excess stain. Per cent of AM fungal colonization was calculated using the formula,

$$\text{Per cent AM colonization} = \frac{\text{Number of root bits colonized}}{\text{Total number of root bits examined}} \times 100$$

Total Root Length colonization (TRLC) in Maize
Total root length colonized by hyphae, arbuscules and vesicles was estimated by grid line intersection method (McGonigle et al., 1990). Maize roots were stained using tryphan blue stain. For each subsample two to four slides were used. However all the slides for a subsample were treated as a single unit, and not as subsamples. A graticule with a cross hair was prepared and attached in the eye piece. The roots were aligned parallel to the long axis of the slides and observed at magnification of 200x. The alignment of the roots parallel to the horizontal line of the cross hairs was adjusted to right angle to the long axis of the root by the rotation of the vertical crosshairs. Where the centre of the crosshairs entered a root through an end rather than a side, i.e. the point of exit from the root through its side was taken as the point of intersection. Roots too wide to fit into the field of view at 40x magnification were examined in two or more width portions. The plane of focus was completely moved through the root to examine each intersection and a note of whether the vertical crosshairs actually cut any arbuscules, coils, vesicles or hyphae was made. The number of intersections was counted in the following categories viz., arbuscules, vesicles, coils or hyphae. When the vertical crosshairs cut one or more arbuscules or vesicles, the appropriate category was incremented by one and a similar increment was given for intersections where hyphae only were crossed. The arbuscular colonization (AC) and other categories (hyphae, coils, vesicles) colonization were calculated by dividing the count for the arbuscules and categories respectively by the total number of intersections examined. Finally the total root length colonization (TRLC) was calculated using the formula as given below:

$$\text{TRLC} = \text{AC} + \text{VC} + \text{HC} + \text{CC}$$

where,

- AC - Arbuscular colonization
- VC - Vesicular colonization
- HC - Hyphal colonization
- CC - Coil colonization

Quantification of the AM spore density
AM fungal spores were extracted from the rhizosphere substrate of maize by using the Wet sieving and Decantation Technique. Total number of viable spores were counted by following the procedure as follows.

Sievings from each sieve (180 µ and 106 µ) collected separately in to small beakers was transferred again into the fine sieve (45 µ). The sievings from fine sieve was collected using a fine get of water into a gridded filter paper funnel so as to remove the excess water. The filter paper was gridded at one half of the sheet and it was folded in such a way that the marked portion was the receiving surface during filtration. After the filtration the filter paper was removed gently and spreaded on a bigger size Petriplate with spores & other debris and observed under stereo zoom microscope. Presence of viable spores were counted by moving the Petriplate and the total number of viable spores were calculated corresponding to the weight of soil taken for analysis and the result was expressed as number 100 g⁻¹ of the sample sieved.

Statistical analysis
The data obtained were analyzed by a three and two factorial analysis of variance (p = 0.05) wherever possible, with days after sowing, treatments and substrates as experimental factors using AGRES software, with mean separation by Least significant difference (LSD) as per the methods detailed by the Panse and Sukhatme (1978). The analysis for microbial population count was based on the log and arcsine transformed values.

Result
With the objective of standardizing the spore density of AM fungal inoculum for effective colonization, a pot culture study was conducted using maize (PEHM 5) as host plant and vermiculite, vermiculite + 10% soil as substrate materials.

Root volume and total dry weight
A critical examination of the data presented in Table 1 indicated that inoculation of AM spores exhibited significant increase in root volume and total dry weight of maize grown in either vermiculite alone or with 10 % soil amendment in vermiculite at 45 and 60 days after plant growth. Progressive increase in root volume was noticed with an increase in age of the crop (Plate 1). At each range of spore density, the increase in root volume was observed and it ranges from 2.5 to 3.0 cm³ when grown in vermiculite and 2.8 to 3.1 cm³ when grown in vermiculite + 10 % soil, recording 47.0 to 76.4 per cent and 64.7 to 82.3 per cent increase over control respectively at 60 days of plant growth. There were no significant difference between the superior treatments that received 5 to 6 spores g⁻¹ (T₃) and 10 to 12 spores g⁻¹ (T₄) of AM inoculum. The treatment which received 1 to 3 spores g⁻¹ (T₂) was significantly lower from T₁ and T₄ when grown in vermiculite and vermiculite + 10 % soil. The interaction effects were not significant. Additionally, AM inoculum containing a spore density of 10 to 12 spores g⁻¹ was superior, which influenced the total dry weight of maize at all stages of the crop growth. It recorded the maximum of 6.2 g in vermiculite, 6.8 g in vermiculite amended with 10 % field soil and 105.6 and 151.8 per cent increase respectively over control at 45 days after sowing. The similar trend was maintained on 60 days. The interaction effect between days and substrates was observed not significant. Additionally, AM inoculum containing a spore density of 10 to 12 spores g⁻¹ was superior, which influenced the total dry weight of maize at all stages of the crop growth. It recorded the maximum of 6.2 g in vermiculite, 6.8 g in vermiculite amended with 10 % field soil with 129.8 and 151.8 per cent increase respectively over control at 45 days after sowing. The similar trend was maintained on
60 days. The interaction effect between days and substrates was observed not significant.

AM fungal colonization in roots of Maize

As per the method of root clearing and staining root colonization was assessed. The results presented in Table 2 tend to indicate a sort of significant increase on root colonization of maize when inoculated with varying levels of AM spores at both the stages of plant growth. AM inoculum application recorded root colonization of 88 to 95 per cent at 45 days of plant growth in vermiculite. When 10 % field soil was amended with vermiculite the root colonization of maize by AM fungi was still enhanced upto 96 per cent on 45 days and 100 per cent on 60 days. Inoculation of AM spores at 10 to 12 spores g\(^{-1}\) recorded maximum root colonization (Plate 2). However, there were no significant difference between the superior treatment that received 10 to 12 spores g\(^{-1}\) and 5 to 6 spores g\(^{-1}\) of AM inoculum. The interaction effects between treatments and days, treatments and substrates were observed significant.

Plate 1- Effect of AM inoculum with different spore densities on the root volume of Maize

Plate 2- Effect of AM inoculum with 10 to 12 spores g\(^{-1}\) on root colonization of Maize

Assessment of total root length colonization of Maize

Total root length colonized by hyphae, arbuscules, coils and vesicles of AM fungi was estimated by grid line intersection method (Mc Gonigle et al., 1990) at 60 days after sowing. The results showed that among the four treatments analysed, T\(_4\) which received 10 to 12 spores g\(^{-1}\) of AM inoculum had significantly maximum AM fungal total root length colonization (90 per cent) followed by T\(_3\) that received 5 to 6 spores g\(^{-1}\) of AM inoculum (88 per cent) when vermiculite was used as the substrate material (Table 3). Similarly, still further significantly enhanced total root length colonization was recorded in the treatment T\(_4\) (98 per cent), followed by T\(_3\) (96 per cent) when vermiculite + 10% soil was used as the substrate material. Uninoculated control had no AM fungal colonization. The interaction effect between treatments and substrates was significant.

Plate 1- Effect of AM inoculum with varying levels of spore densities on the viable spore count of Glomus intraradices

Plate 2- Effect of AM inoculum with 10 to 12 spores g\(^{-1}\) on root colonization of Maize

Viable spore count of Glomus intraradices in the treated substrate

The spore density was found to vary in each treatment. Viable spore numbers of AM fungi in substrate was significantly influenced with the range of spores present in the AM inoculum (Figure 1). Among the substrates used, vermiculite + 10 % soil recorded maximum viable spore count compared to vermiculite especially at 60 days after sowing.

Among the substrates sampled at 60 days after plant growth, addition of AM inoculum containing 10 to 12 spores g\(^{-1}\) marked a highest viable spore density of 145 spores 100 g\(^{-1}\) vermiculite followed by the addition of 5 to 6 and 1 to 3 spores g\(^{-1}\) with 133 and 115 spores 100 g\(^{-1}\) vermiculite respectively. In the same way vermiculite + 10 % soil that received 10 to 12, 5 to 6 and 1 to 3 spores g\(^{-1}\) showed 148, 145 and 124 spores 100 g\(^{-1}\) respectively. There were no significant differences between days and substrates followed by treatments, days and substrates.

Discussion

Infected root segments, infective propagules and spores isolated from open pot cultures of AM fungi inoculated plants have been...
the usual source of AM inoculum for commercial purpose (Sylvia et al., 1993). Spores of AM are the important component of inoculum and are more suitable for large scale production as well as for biochemical and molecular investigations of the AM symbiosis (Thompson, 1987). Hence a preliminary attempt was undertaken to standardize the spore density of AM inoculum for effective colonization using vermiculite and vermiculite + 10 % soil as substrates and maize (cultivar PEHM 5) as host plant. Three different levels of spore densities were tested (1 to 3, 5 to 6 and 10 to 12 spores g⁻¹ of AM inoculum).

With respect to root volume, T₄ (10 to 12 spores g⁻¹ of AM inoculum) recorded the highest. The similar trend was observed in total dry weight of maize also. The results showed that both T₄ and T₃ (5 to 6 spores g⁻¹ of AM inoculum) significantly improved the growth of maize and were not statistically significant from each other in their ability to stimulate root volume and total plant dry weight. Similarly, maximum root colonization, total root length colonization and viable spore count was observed to be maximum in T₄ followed by T₃ at 60 days when vermiculite amended with 10 % soil was used as the substrate for plant growth.

In the present investigation, amendment of 10 % soil in sterile vermiculite was found more suitable for AM fungal production and colonization, rather than vermiculite alone. The lower pore space volume of vermiculite and finer size of soil is an optimal combination for AM multiplication. For AM production, vermiculite amended with 10 % soil is most preferred since soil act as a nutrient supplement for the host plant grown in vermiculite. Moreover, soil is available in nature abundantly as free and usage of vermiculite will be costlier. Therefore, 5 to 6 spores g⁻¹ of AM inoculum should be considered as one of the quality standard that could ensure a high quality AM inoculum.

Conclusion
Thus, the result of the present study throws light on the standardized level of spore density that has to be present in the AM inoculum for effective colonization. It is confirmed that when a spore density of 5 to 6 spores are present in g⁻¹ of AM inoculum, it is much more enough to enhance the growth as well as mycorrhizal responses of host plant so that it will ultimately result in the better responses at field level and enhance the crop productivity. Therefore, 5 to 6 spores g⁻¹ of AM inoculum should be considered as one of the quality standard that could ensure a high quality AM inoculum.

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Table 1 - Effect of AM inoculum with different spore densities on root biomass and total dry weight of Maize

| Treatments | 45 DAS | 60 DAS | Mean | 45 DAS | 60 DAS | Mean | Total dry weight (g plant⁻¹) Mean |
|------------|--------|--------|------|--------|--------|------|----------------------------------|
|            | S1     | S2     | S1   | S2     | S1     | S2   |                                  |
| T1 - Uninoculated control | 1.0 | 1.3 | 1.6 | 2.0 | 2.3 | 2.7 | 2.9 | 3.1 | 3.5 |
| T2 - 1 to 3 spores g⁻¹ of AM inoculum | 2.2 | 2.3 | 2.5 | 2.8 | 2.6 | 3.1 | 2.9 | 3.1 | 3.6 |
| T3 - 5 to 6 spores g⁻¹ of AM inoculum | 2.5 | 2.7 | 2.8 | 3.0 | 2.9 | 3.2 | 6.1 | 6.7 | 7.4 |
| T4 - 10 to 12 spores g⁻¹ of AM inoculum | 2.6 | 2.8 | 3.0 | 3.1 | 3.0 | 3.2 | 6.2 | 6.6 | 7.5 |

DAS - Days after sowing, S1 - Sterile vermiculite alone, S2 - Sterile vermiculite + 10 % soil, Values represent mean of five replications, Values in parenthesis indicate per cent increase over control, ** - significant at p<0.01, * - significant at p<0.05, NS - not significant.

Table 2 - Effect of AM inoculum with different spore densities on root colonization of Maize

| Treatments | 45 DAS | 60 DAS | Mean | 45 DAS | 60 DAS | Mean | AM root colonization (%) Mean |
|------------|--------|--------|------|--------|--------|------|-------------------------------|
|            | S1     | S2     | S1   | S2     | S1     | S2   |                                 |
| T1 - Uninoculated control | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| T2 - 1 to 3 spores g⁻¹ of AM inoculum | 88 | 92 | 90 | 95 | 97 | 96 | 100 |
| T3 - 5 to 6 spores g⁻¹ of AM inoculum | 90 | 95 | 93 | 96 | 100 | 100 | 100 |
| T4 - 10 to 12 spores g⁻¹ of AM inoculum | 77.7 | 80.1 | 78.9 | 89.2 | 89.2 | 89.2 | 89.2 |

DAS - Days after sowing, S1 - Sterile vermiculite alone, S2 - Sterile vermiculite + 10 % soil, Values represent mean of five replications, Values in parenthesis indicate the per cent transformed values, ** - significant at p<0.01, * - significant at p<0.05, NS - not significant.

Table 3 - Effect of AM inoculum with different spore densities on total root length colonization of Maize at 60 days after sowing

| Treatments | VC | AC | CC | HC | TRLC | Mean |
|------------|----|----|----|----|------|------|
|            | S1 | S2 | S1 | S2 | S1   | S2   | S1   | S2   | S1   | S2   |
| T1 - Uninoculated control | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| T2 - 1 to 3 spores g⁻¹ of AM inoculum | 14 | 15 | 17 | 18 | 12 | 12 | 42 | 46 | 85 | 91 |
| T3 - 5 to 6 spores g⁻¹ of AM inoculum | 15 | 15 | 21 | 22 | 8 | 13 | 44 | 46 | 88 | 96 |
| T4 - 10 to 12 spores g⁻¹ of AM inoculum | 15 | 17 | 22 | 24 | 8 | 9 | 45 | 48 | 90 | 96 |

S1 - Sterile vermiculite alone, S2 - Sterile vermiculite + 10 % soil, VC - Vesicular colonization, AC - Arbuscular colonization, CC - Coil colonization, HC - Hyphal colonization, TRLC = Total root length colonization, Values represent mean of five replications, Values in parenthesis indicate per cent transformed values, ** - significant at p<0.01.