Record of Arbuscular Mycorrhizal Fungi (AMF) in Pippali Plant under the Agro Ecological Conditions of Jorhat District, Assam, India

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ABSTRACT

Arbuscular mycorrhizal status of Pippali a medicinal plant species in the Jorhat district of Assam, India were surveyed during 2018. Percent colonization, spore density and diversity of arbuscular mycorrhizal fungi associated with the rhizospheric soil and roots of pippali growing wild as well as under cultivated conditions were investigated. It was found that 100% of the surveyed sample was of mycorrhizal infestation. The spore colonization per cent of AMF ranged from 30.80 to 66.50% spores per 100 g of soil. The maximum spore population (185.80 spores/100 gm of soil) was obtained on soil samples collected from Gibbon wild life sanctuary. This was followed by soil sample collected from Katoni par Soil sample collected from Upper deori gaon, Nakachari gaon, and Nagajanka showed spore population of 168.80 to 154.65 spores/100 ml of soil. Result showed that root colonization by arbuscular mycorrhizal fungi both in roots and soil of Pippali. But, percentage of root and soil colonization varies according to location. Variation in the spore density and percent colonization among different sampling sites could be attributed to host specificity, adaphic and climatic conditions. After comparing with synoptic key the genus of the mychorrhiza was identified tentatively as Glomus spp.

Keywords
A.M fungi, Assam, Pippali, Root and soil colonization, Spore density

Article Info
Accepted: 10 December 2018
Available Online: 10 January 2019

Introduction

Agriculture is the dominant land use pattern in the state Assam. It account for about 54.11 percent of the total geographical area of the state. Including persons dependent on plantation, more than 80 percent of total populations of Assam is dependent on agriculture. Agriculture plays an important role in revenue earning in Assam economy. The Assam experiences a plenty of rainfall and possess a fertile land which is extremely advantageous for cropping. Among the different districts of Assam, by understanding the existing agro ecological conditions of the study area is seen suitable for a growing of Pippali plant for the study and record of arbuscular mycorrhizal fungi in this particular plant.

Jorhat district is located between 26°20’ N and 27°15’ N latitudes and between 94°00’ E longitudes. On the north, it is bounded by Lakhimpur, east by Sivasagar, west by Golaghat districts of Assam and on the south by Nagaland state. Regarding the soils, basically alluvium in nature and climate belongs to four main types, viz., humid
continental severe winter, moist in all seasons and short summer, sub tropical monsoons, mild and dry winter, warm and humid summer, sub tropical monsoon, mild and dry winter and sub tropical monsoon with very heavy rain.

A large number of microorganisms are associated with the roots of plant in their natural habitat. Among them mycorrhiza plays a highly evolved association with the root system. Mutualistic symbiosis is one of the most abundant symbiotic activities in ecosystems. Arbuscular mycorrhizae are symbiotic associations formed between fungi, which belong to the phylum Glomeromycota (Schubler et al., 2001) and the root system of many plant species. Arbuscular mycorrhizal fungi (AMF) are known to play an important role in improvement of plant nutrition and growth of several ornamentals and vegetable crops (Rouphael et al., 2010; Koltai 2010). Amongst the microorganisms, AMF are able to solubilize insoluble phosphates and improve plant P nutrition (Zarei et al., 2006).

Several studies have been performed on changing secondary compound patterns of medicinal plant symbiosis with mycorrhizal fungi such as terpenoids (Akiyama and Hayashi 2002), phenols (Zhu and Yao 2004), phenylpropanoids (Weiss et al., 1997), glucosinolates (Vierheilig et al., 2000), carotenoids (Maier et al., 1995) and flavonoids (Larose et al., 2002).

Pippali (Piper longum), sometimes called Indian long piper (Pipli) is a flowering vine in the family Piperaceae, cultivated for its fruit, which is usually dried and used as a spice and seasoning. The plant itself, is a native of India. The fruit contain the alkaloid piperine, which contributes to their pungency. Long piper is a climber, of South Asian origin (Deccan peninsula). The word piper itself is derived from the Sanskrit word for long piper, pippali. It is a slender, aromatic, climber with perennial woody roots, creeping and jointed stems, and fleshy fruits embedded in spikes. Leaves are numerous, 6.3 to 9.0 cm, broadly ovate or oblong oval, dark green and shining above, pale and dull beneath. The older leaves are dentate, dark in color and heart shaped. The younger leaf is ovate in shape and contains 5 veins on them. Flowers are monoecious and male and female flowers are borne on different plants. The plant flowers in rains and fruits in early winters.

Pippali is certainly one of the most widely used of all Ayurvedic herbs. It is one of the best herbs for enhancing digestion, assimilation and metabolism of the foods we eat. It is also highly prized for its ability to enhance assimilation and potency of herbs in a synergistic formula (this is called the Yogavahi effect).

The present study was based on record of arbuscular mycorrhizal fungi (AMF) in Pippali plant under the agro ecological conditions of Jorhat district, Assam.

Materials and Methods

Collection of Rhizosphere soil sample and root sample of Pippali

Field survey was carried out in order to collect the root and rhizospheric soil sample of Pippali Plant from different blocks of Jorhat district, Assam (Fig. 1). Rhizospheric soils at a depth of 0-30 cm from 10 different locations were collected in sterile polyethylene bags (Table 1). Approximately 100gm of rhizospheric soil was collected and were air – dried and stored at 4°C for processing. For the collection of root sample, feeder roots were collected immediately after digging the plant. For the maintenance and preservation of roots, collected sample were washed under running tap water and rootlets were selected and cut into small pieces and fixed in acetic acid solution.
Study the root colonization of arbuscular mycorhizal fungi

To study the root colonization of Arbuscular mycorrhizal fungi in the roots of Pippali the standard procedure of Philip and Haymann, (1970) was followed. Per location few roots were selected randomly and sections were made and placed in glass slides and observed 10 section from each slide for 3 slides under microscope for root colonization. Number of root colonized were counted and percentage of root colonization was calculated by following formula,

\[
\text{Root colonization percentage} = \frac{\text{no. of infected root section}}{\text{no. of total root section}} \times 100
\]

Isolation of AMF spores from soil sample

Isolation of AMF spores from soil sample was done by following the Wet Seiving and decanting methods of Gerdemann and Nicolson, (1963). Spore population was then expressed in terms of number of spores per 100 gm of dry soil.

Characterization of arbuscular mycorrhizal fungi

The AMF isolated from Pippali plant as loose clusters of spores or as sporocarps and chlamydospores were characterized on the basis of spore morphology (size, shape, colour) wall character and hyphae attachment up to generic or if possible to species level using synoptic key of Trappe (1982) and INVAM manual of Schenck and Perez (1988).

Results and Discussion

Study of root colonization of AMF

Colonization of Pippali plant by AMF was ascertained by the presence of fungal structure in root segments. There was distinct variation in per cent AMF colonization of root samples collected from Pippali growing pockets from different locations of Jorhat. The highest AMF colonization (66.50%) was found in Gibbon wild life sanctuary followed by Katonipar (64.6%). This was followed by Upper Deorigaon, Karanga, Experimental garden, Department of Horticulture (AAU) and Nagajanka with spore colonization of 6.42%, 55.5%, 54.5% and 52.7% respectively. Lowest root colonization percentage was found in Lahdoigarh (30.80%) (Table 2).

Isolation of AMF spores from soil sample

It was observed that AMF propagules in different soil samples ranged between 110.00 to 185.80 per 100 g of soil sample. The highest spore load of 185 AMF spores per 100g soil samples in Pippali rhizosphere was observed in soil sample collected from Gibbon Wild Life sanctuary followed by Katonipar (176.00 spores per 100 g of soil sample) (Table 3). Soil sample collected from Upper Deorigaon, Nakachari, and Nagajanka showed spore population of 168.80 to 154.65 spores/100 ml of soil. The soil of Lahdoigarh was found registered lowest (110.55/g soil).

Identification of arbuscular mycorrhizal fungi

AMF spores isolated from Pippali from 10 different locations of Jorhat belonging to the genera *Glomus* were identified up to generic level on the basis of morphological characters using standard taxonomic key (Trappe, 1982; Schenck and Perez,1988).

Microscopic study showed that spore borne singly, round to globose shaped with smooth double layered wall or fused, spore measuring 60-75 X 90-110 µm in size, spore colour was yellow brown to dark brown, spore wall cavity absent (Plate 2). After comparing with synoptic key the genus of the mycorrhiza
was identified tentatively as *Glomus* spp as per the available standard key (Schenck and Perez, 1990). Studies on the distribution of AMF fungi in different locations revealed the predominance occurrence of AMF spores belonging to genus *Glomus* sp. which were dominant in distribution and abundance.

The distribution pattern of AMF in 10 different locations of 4 blocks of Jorhat districts of Assam revealed prevalence of spores of genera of AMF viz., *Glomusspp.* *Glomusspp* were the common AMF fungi in the majority of Pippali soils. *Glomusspp* are known to be the most common AMF throughout the world (Gerdeman and Trappe, 1974., Blaszkowski, 1989; Talukdar, 1993). The frequent detection of spores of *Glomus* spp. in the soils of selected Pippali growing of Jorhat district, Assam in the present investigation therefore supports the earlier findings. The occurrence of the five genera of AMF fungi were also reported earlier from Assam in tea and other crops by various workers (Anonymous, 1982; Hazarika, 1992; Baruah, 1994; Chaudhury, 1998; Das and Barthakur, 1999).

| Sl no. | Location                                | GPS data            | Date of survey | Collection made |
|-------|-----------------------------------------|---------------------|----------------|-----------------|
| 1     | Experimental garden, Department of Horticulture (AAU) | 26.724°N and 94.195°E | 02/12/17       | Plant and soil sample |
| 2     | Gibbon Wild Life Sanctuary (GWS)        | 26.716°N and 94.383°E | 02/12/17       | -do-           |
| 3     | Nakachari (NAK)                         | 26.702°N and 94.450°E | 02/12/17       | -do-           |
| 4     | Nagajanka (NAG)                         | 26.718°N and 94.470°E | 02/12/17       | -do-           |
| 5     | Karanga (KAR)                           | 26.695°N and 94.238°E | 03/12/17       | -do-           |
| 6     | Upper DeoriGaon (UDG)                   | 26.829°N and 94.119°E | 05/12/17       | -do-           |
| 7     | NamdeoriGaon (NDG)                      | 26.827°N and 94.086°E | 05/12/17       | -do-           |
| 8     | Katoni Par (KP)                         | 26.820°N and 94.074°E | 10/01/18       | -do-           |
| 9     | PukhuriporiaGaon (PPG)                  | 26.800°N and 94.070°E | 10/01/18       | -do-           |
| 10    | Lahdoigarh (LAH)                        | 26.785°N and 94.324°E | 25/01/18       | -do-           |
Table 2 Root Colonization (%) of arbuscular mycorrhizal fungi in sample collected from different locations

| Location                        | Root colonization(%) |
|---------------------------------|----------------------|
| Karanga(KAR)                    | 55.50 (48.16 e*)     |
| Upper DeoriGaon (UDG)           | 62.40 (52.17 b**)    |
| NamdeoriGaon (NDG)              | 46.50 (42.99 de)     |
| Katoni par (KP)                 | 64.60 (53.48 ab)     |
| Lahdoigarh (LAH)                | 30.80 (33.70 i)      |
| PukhuriporiaGaon (PPG)          | 43.50 (41.26 e)      |
| Gibbon Wild Life Sanctuary (GWS)| 66.50 (54.63 a)      |
| Nakachari (NAK)                 | 48.80 (44.31 d)      |
| Nagajanka (NAG)                 | 52.70 (46.54 c)      |
| Experimental garden, Department of Horticulture (AAU) | 54.50 (47.58 c) |
| SED(±)                          | 1.73                 |
| CD (p=0.05)                     | 3.49                 |

* Data in parentheses are angular transformed value, ** Data followed by same alphabets are statistically at par

Table 3 Spore count of Arbuscular Mycorrhizal Fungi per 100 gm. of soil of different locations

| Location                        | Spores/ 100 gm of soil |
|---------------------------------|------------------------|
| Karanaga (KAR)                  | 152.55 d               |
| Upper DeoriGaon (UDG)           | 168.80 c               |
| NamdeoriGaon (NDG)              | 134.50 f               |
| Katoni Par (KP)                 | 176.00 b               |
| Lahdoigarh (LAH)                | 176.00 b               |
| PukhuriporiaGaon (PPG)          | 144.55 e               |
| Gibbon Wild Life Sanctuary (GWS)| 185.80 a               |
| Nakachari (NAK)                 | 155.55 d               |
| Nagajanka (NAG)                 | 154.65 d               |
| Orchard (AAU)                   | 148.00 e               |
| S Ed(±)                         | 2.40                   |
| CD (p=0.05)                     | 4.10                   |

*Data in parentheses are angular transformed value, ** Data followed by same alphabets are statistically at par
Plate 1. Survey and collection of sample (root, soil) of Pippali plant from different locations of Jorhat district
Extent of root colonization by AMF also varied in different locations of Pippali plant. Colonization was highest in Gibbon wildlife sanctuary followed by Katoniper whereas lowest in Lahdoigarh area. The variation in root colonization in different location might be related to the soil environmental condition, fertility status, application of chemicals or it may also be attributed to mixed AMF species present in different locations. The Phosphorous level of soil had direct relation with VAM association (Mikola, 1981). The highest AMF population in Gibbon wild life sanctuary might be due to undisturbed condition of soil, which might favor more colonization by AMF fungi for by increasing the fertility status of soil. Similarly, low colonization of Lahdoigarh area of Pippali rhizospheric soil related to high level of available soil phosphorous as the place is highly related to cultivation which made the plant less dependent on AMF fungi for phosphorous nutrition. Changes in soil fertility related to disturbed and undisturbed soil due to amendment with mineral fertilizers can markedly affect the activity of soil mycorrhiza by influencing the level of root infection and spore production. Morita and Konishi (1989) reported that the average percentage of tea root colonization of lightly fertilized soil was two times higher than heavily fertilized fields. The higher root colonization in Gibbon wildlife sanctuary might be due to presence of efficient strain of AMF fungi, which can infect the Pippali root at higher rate.

Acknowledgement

The author acknowledges the Director of Post Graduate Studies, Professor and Department of Plant Pathology, Director of Research (Agri), Assam Agricultural University, Jorhat, Assam for the constant support and guidance during period of investigation. Due acknowledgement also goes to ICAR-AICRP on MAP and Betelvine, ICAR, New Delhi.

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**How to cite this article:**

Deori, M. and Pranab Dutta. 2019. Record of Arbuscular Mycorrhizal Fungi(AMF) in Pippali Plant under the Agro Ecological Conditions of Jorhat District, Assam, India. *Int.J.Curr.Microbiol.App.Sci.* 8(01): 1011-1019. doi: [https://doi.org/10.20546/ijcemas.2019.801.110](https://doi.org/10.20546/ijcemas.2019.801.110)