Occurrence and Distribution of Entomopathogenic Fungi in Agricultural Soil of Durg District of Chhattisgarh, India

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Abstract

Entomopathogenic fungi are important natural enemies of arthropods and can be used for biological control of ticks. They are widely distributed in a wide range of habitats including aquatic, forest, agricultural soil and pasture habitats. Sixty crop roots soil samples were collected from nearby field and grazing areas of animals in Durg district of Chhattisgarh. A total of seven fungal isolates were recovered from the organic environment of Durg, Chhattisgarh and they were belonging to Genus; Aspergillus, Penicillium, Cladosporium, Fusarium, Rhizopus, Metarhizium and Trichoderma. The fungal colonies were isolated from soil samples and surface sterilized ticks were treated with aqueous fungal suspension. The mortality of ticks was observed on third to seventh day when hyphae covered the whole body surface. Out of the seven isolates explored from soil samples, four isolates were found to infect ticks namely, Fusarium sp., Rhizopus sp., Metarhizium sp. and Trichoderma sp.

Introduction

Tick and tick borne diseases are one of the biggest public health and veterinary problems in the world. These ectoparasites have an impact on the production and health of the animals through sucking blood or by transmitting the infectious agents such as viruses, bacteria, rickettsiae and protozoa (Eskezia and Desta, 2016).

Use of Chemical acaricides for control of ticks was considered as one of the best methods, but ticks have developed resistance against a range of currently-used acaricides such as organophosphates, carbamates, synthetic pyrethroids and amidines (Martins et al., 1995). The use of insecticides also produces environmental pollution with residues in milk, meat, vegetables and producing detrimental effects to human health. Biological control of ticks using
Entomopathogenic fungus is proved to be most economical and safest method to overcome the risk of environmental pollution and acaricidal resistance (Lacey et al., 2001). The bio-control potentials of entomopathogenic fungi (EPF) against animal ticks have shown promising results (Rao and Narladkar, 2017). The ability of EPF to penetrate the cuticle of arthropods, to kill several stages of the same pest and the relatively specific virulence of a single strain to one or a small group of pests make them good candidates as biocontrol agents (Samish et al., 2004). The present work was aimed to isolate and characterize the EPF from agricultural soil of Durg District in Chhattisgarh.

Materials and Methods

Isolation and morphological studies

Soil samples were collected during the period from September, 2018 to October, 2019 from ten villages namely Anjora, Rasmada, Thanaud, Kharpi, Sriloda, Nagpura, Damoda, Dhaba, Dhanora and Mohlai. From each village, six fields which were under cultivation of either paddy or maize or seasonal vegetable crops were selected for collection of soil samples. The distance between two villages was about 3 to 5 kilometers and the distance between two fields was about half to 1 kilometer. A total of Sixty soil samples of 25 gram each were collected in zipped polythene bags from sampling point nearby crop plant root i.e. from the area with 5 cm in diameter and 5 cm deep around the crops using a trowel with a total area of 16 m² from each field as per the method described by Amy et al., (2009). The soil samples were sieved through 2 mm mesh size to remove course material. The isolation of fungi from soil samples was enumerated by using serial soil dilution and soil plate method (Waksman, 1922) on Potato Dextrose Agar medium. Four serial dilution of soil samples were undertaken in 15 ml test tubes. Dilutions of 10², 10³ and 10⁴ were made to avoid overcrowding of fungal colonies. Potato Dextrose Agar Medium was prepared at a final concentration of 2.5% in conical flasks and autoclaved. Antibiotic solution using tetracycline and neomycin (w/v) was added to the medium at the rate of 0.02% after autoclaving to suppress bacterial growth. The molten medium was poured in radiation sterilized Petri plates (90 mm) in aseptic condition and allowed to solidify. One ml of the suspension of soil sample of each concentration was added to sterile Petri plates, in triplicates of each dilution, containing sterile Potato Dextrose Agar medium using micropipette. The plates were gently rotated to disperse the sample uniformly on agar plate. The plates were then incubated at 29°C and 75% relative humidity for 7 days. Plates were regularly monitored for fungal surface colonies. One isolate of each fungal growth from each soil sample was selected at random and further sub cultured. The subcultures were maintained on Potato Dextrose Agar Slants.

The ticks collected from the body of animals were rinsed with distilled water and then treated with 1% Potassium hypochlorite solution to prevent bacterial contamination. The surface sterile ticks were then placed on each fungal isolate and observed for fungal growth. The fungal isolates infecting ticks were separated and assumed as EPF. Isolation of entomopathogenic fungi was also done by placing the sterile ticks on petri plates containing soil sample. The soil samples were uniformly spread on Petri plates and moisture was maintained by adding sterile distilled water. Then sterile ticks were placed on soil samples and regularly observed for growth of fungal colonies. Ticks infected with fungus were isolated and then placed on petri plates containing growth media. The fungal colonies
were then further subcultured to obtain the pure culture.

Results and Discussion

Out of the sixty soil samples collected from agricultural fields all the soil samples showed presence of fungal colonies. The fungal colonies were observed on third day whereas full grown fungal colonies were observed on fifth to seventh day at 10^4 serial dilution. Number of colonies in culture plates was counted on third day. The colonies were found diffused with each other on seventh day. From a single soil sample, different fungal colonies were isolated. Out of the sixty soil samples examined for fungal isolation, a total of 2286 fungal colonies were recovered. On an average, 38.1 colonies were isolated from each soil sample. All the sixty soil samples were recorded as 100% positive for presence of fungal isolates. Out of the 38.1 colonies isolated from soil samples some of the colonies were found morphologically similar to each other on the basis of colony morphology. Some of the fungal colonies were diffused with each other hence could not identified. A total of seven fungal isolates were recovered from organic environment of Durg Chhattisgarh belonging to Genus; Aspergillus, Penicillium, Cladosporium, Fusarium, Rhizopus, Metarhizium and Trichoderma. The Aspergillus was found most predominant soil dwelling fungal species followed by Penicillium, Cladosporium, Fusarium, Rhizopus, Metarhizium and Trichoderma. The percent recovery of fungal isolates of genus Aspergillus was 37.81% followed by Penicillium, 16.91%; Cladosporium, 10.94%; Fusarium, 6.96%; Rhizopus, 5.22%; Metarhizium, 5.22%; Trichoderma, 4.22% and unidentified was 12.68%. Out of the seven isolates explored from soil samples, four isolates were found to infect ticks and were observed as EPF namely, Fusarium sp, Rhizopus sp, Metarhizium sp. and Trichoderma sp. These fungal isolates were morphologically identified based on colony morphology, shape and colour of colony, mycelium and conidial structure.

The rare occurrence of Metarhizium sp. isolated from natural habitats was observed by Chandler et al., (1997), Bidochka et al., (1998), Meyling and Eilenberg (2005) and Thakur and Sandhu (2010), the findings are in accordance with the present work. In the present study the EPF species isolated from soil belonging to genera Fusarium, Rhizopus, Metarhizium and Trichoderma were recorded.

The present findings were in accordance with Gouli et al., (2013) who also isolated EPF such as Aspergillus, Fusarium and Penicillium. In present investigation, 5.22% of EPF Metarhizium sp. was observed. Khudhair et al., (2014) reported 18.1% of EPF belonging to species Metarhizium anisopliae in Iraqi province agro-ecosystems using Galleria mellonella bait trap technique. They reported highest entomopathogenic frequency rate with 55.3% followed by lowest rate with 17%.

Tkaczuk et al., (2015) reported Metarhizium anisopliae and B. bassiana formed more colony forming units in soils from organic fields. M. anisopliae was the most frequently isolated fungus detected in 92% of the soil either from organic or conventional fields. Rasheed et al., (2004), Noor Zaman et al., (2012) and Ratna Kumar et al., (2015) isolated fungal genera like Aspergillus, Alternaria, Curvularia, Fusarium, Penicillium and Rhizopus from different crop fields. These observations were similar with present findings. In our investigation among the fungal isolates obtained from the soil samples, the genera Aspergillus and Penicillium were most dominant fungal species. The most common isolates in present
investigation were *Aspergillus fumigatus*, *A. niger*, *A. ustus*, *A. versicolor*, *Penicillium chrysogenum*, *P. notatum*, *Cladosporium sphaerospermum*, *Rhizopus oryzae*, *Metarhizium majus*, *Fusarium oxysporum* and *Trichoderma harzianum*.

Chandini and Rajeshwari (2017), reported that the *Penicillium* and *Aspergillus* were the dominant fungi followed by *Chaetomium*, *Trichoderma* and *Fusarium* in Mattavara forest. Fungal frequency of *Fusarium oxysporum* was noted by them was 33.83% which was 6.96% in the present study. Raja et al., (2017) with the similar findings investigated the fungal namely *Aspergillus niger*, *A. clavatus*, *A. sydowii*, *As. variabilis*, *A. fumigatus*, *Penicillium chrysogenum*, *Colletotrichum gloeosporioides*, *Mucor* sp. *Rhizopus stolonifer*, *Rhizopus oryzae*, *Cunninghamella bertholletiae*, *Scopulariopsis brumptii*, *Cladophialophora sp*. Most of the investigated species are similar with species investigated in the present work. In the present study the rare occurrence of EPF in various cultivated organic agricultural field areas was observed. This may be due to regular use of chemical pesticides, tropical temperature zone and irrigation practices which may be responsible for rare abundance of EPF. The soil moisture has a direct effect on the population of fungi hence, at higher moisture the tolerance and
colonization is badly affected (Adams et al., 1999). Hummel et al., (2002), in a long-term field study found that the application of certain pesticides significantly reduces the occurrence of EPF in the soil.

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