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Isolation, Identification and Pathogenicity Assay of Fungal Species on Horse Purslane (*Trianthema portulacastrum* L.,)

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Authors' contributions

This work was carried out in collaboration among all authors. All authors read and approved the final manuscript.

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Original Research Article

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ABSTRACT

A field survey was carried out in Tamil Nadu agricultural university, Coimbatore, to locate Horse purslane (*Trianthema portulacastrum*) infected with fungal pathogens. Heavily infested areas were recognized and symptomatic leaves were collected. Fungal species account for leaf spot disease of Horse purslane were Isolated from infected leaves on PDA medium. Based on morphological characteristics two foliar pathogens of Horse purslane were identified as *Gibbago trianthemae* Simmons and *Curvularia tuberculata Sivan*. Pathogenicity of two isolates was tested by applying spore treatment to Horse purslane (test plants). Leaves of test plants showed naturally occurring symptoms after inoculation with pathogens. Among the two isolates, *Gibbago trianthemae* was shown to be more effective against Horse purslane than *Curvularia tuberculata*.

Keywords: Biological control; Curvularia tuberculata; Gibbago trianthemae; infestation.

1. INTRODUCTION

Horse purslane (*Trianthema portulacastrum*), an annual herb in the family Aizoaceae, is one of the main weeds in agriculture [1]. *Trianthema*

portulacastrum is also known as Desert horse purslane, Giant pigweed (English), and *Vishakhapara* (Hindi). It was discovered as a noxious weed plant in several agricultural crops in tropical and subtropical areas including India,

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where it offers competition and reduces crop yields by 30% to 60% [2]. It is an introduced weed in India, however competition for yields in various agricultural and vegetable crops like mustard, maize, pigeon pea, soybean, potato, and onion crops, it has turned into a noxious weed [3,4]. Due to the great seed production and brief dormancy of Horse purslane, several generations can occur in a single season. When all of these adverse traits are taken into account, effective control tactics are needed for managing Horse purslane.

Many farmers are switching to organic farming and looking at different weed control options due to concerns about ecological, environmental and health issues associated with the widespread use of herbicides [5,6]. An effective substitute for synthetic herbicides and a sustainable method of environmental protection would be biological control. Utilizing live organisms like insects, nematodes, bacteria, or fungi can help manage weed populations biologically. It has been revealed that many parasitic, aquatic, woody, climbing, and herbaceous weeds are susceptible to plant pathogens [7]. The biological management of weeds using plant pathogens has become increasingly relevant since it is effective. safe, practicable, safe. and environmentally acceptable [8,9].

Among plant pathogens, the most diverse group is the fungi, which may infect many different types of plants and cause a wide range of illnesses. In recent years, fungal pathogens have been known to cause stem blight and leaf spot diseases in Horse purslane, which ultimately causes complete defoliation and drying of the weed [10]. In the present study, with the aim of controlling Horse purslane biologically, isolation and identification of pathogens responsible for the infection of Horse purslane was done and tested for pathogenicity.

2. MATERIALS AND METHODS

2.1 Isolation of Mycoflora from *Trianthema portulacastrum*

Diseased leaves of *Trianthema portulacastrum* were collected from farms at the Tamil Nadu Agricultural University, Coimbatore. To get rid of dirt and other debris, symptomatic leaves were carefully cleaned under running water. The infected leaf tissue was divided into small fragments of 1 cm-diameter, surface sterilized

with ethyl alcohol for 1-2 minutes, and then washed with distilled water. Leaf fragments are soaked for one minute in mercuric chloride solution, and then cleaned with sterile water before being placed in Petri plates with potato dextrose media supplemented with streptomycin sulphate, which inhibits bacterial growth. With all necessary precautions taken prevent to contamination, leaf fragments were inoculated in Petri plates, under laminar air flow. Petri plates were placed in an incubator after inoculation to create ideal conditions for pathogen growth, specifically i.e. 25±2°C under 12hrs dark/light period. Sub culturing was done from seven days old culture, and allowed pathogen to grow completely.

2.2 Pathogen Identification

Microscopic observations were carried out from completely grown culture plate. The identification features of each isolates such as colony diameter, colour, texture, sporulation, shape and sizes of conidiophores and conidia were carefully studied. Fungal isolates were identified with the help of pertinent literature [11,12,13].

2.3 Development of Liquid Formulations

Each Petri plate of a nine-mm diameter containing potato dextrose agar inoculated with discs of *Gibbago trianthemae* and *Curvularia tuberculata* in the middle, which were cultured for 20 days at room temperature until confluent growth and sporulation was achieved. Three to four discs from each Petri plate were inoculated in conical flasks with potato dextrose broth and allowed to form a complete mat. Following the completion of the mat, it was removed, crushed in a mixer, filtered through filter paper, and diluted with water to various concentrations. A wetting agent was added while spraying to ensure that every leaf gets an even coating of moisture.

2.4 Pathogenicity Test

During the survey, Horse purslane seeds were collected from agricultural fields. The collected seeds were dried and preserved in healthy condition without any contamination. The seeds were planted in plastic pots filled with sterilized soil to grow the plants. A green house with a 12 hour light/dark photoperiod was used to sustain the pots containing weed seedlings. Prior to the

Treatments	Concentration of formulations(spores/ml)
T ₁	Gibbago trianthemae, Curvularia tuberculata each @2×10 ^⁵ spores/ml
T_2	Gibbago trianthemae, Curvularia tuberculata each @4×10 ⁶ spores/ml
T ₃	Gibbago trianthemae, Curvularia tuberculata each @6x10 ⁶ spores/ml
T_4	Gibbago trianthemae, Curvularia tuberculata each @8×10 ⁶ spores/ml
T ₅	Gibbago trianthemae, Curvularia tuberculata each @10x10 ⁶ spores/ml
T ₆	control

inoculation of the pathogen spores, test plants were handled carefully to prevent pre-infection by other contaminants.

2.4.1 Inoculation of test plants

The assay on the infection process was conducted on the healthy, greenish plants growing in the greenhouse chamber. 12 pots were used for the inoculation of two pathogens with five different concentrations, along with a control. To prevent the spores from drying out, inoculations were carried out by 6 and 7 pm after plants sunset. The were sprayed with formulations of the two pathogens, Gibbago trianthemae and curvularia tuberculata at concentrations of 2, 4, 6, 8 and 10 with 10⁶ spores/ml, as well as a control [8], when the test plants had 10 to 15 leaves. Spore inoculum along with a 0.02% wetting agent is applied to test plants. Control plants received the same treatment except that they were sprayed with sterile water + 0.02% wetting agent. At intervals of 10 days, leaves are checked for symptoms.

The leaves were rated for disease based on the spread of the disease on them, by using the ranking scheme [13].

- 0- no sign of illness, a healthy plant;
- 1- mild symptoms, a plant with few symptoms on 1 to 10% of the leaf area;
- 2- moderate symptoms, a plant showing larger diseased areas on 11 to 25% of the leaf area;
- less severe symptoms, includes enlarged lesions that cover 26 to 50% of the leaf area;
- severe, symptoms cover 50 to 75% of the leaf area; and
- 5- Very severe, more than 75% of leaf area covered with symptoms

Disease intensity / Percent Disease Index were calculated using formula given by [14].

Percer	nt	Disease
Indox-	sum of ratings	v100
muex-	no of leaves observed×highest rating	X 100

3. RESULTS AND DISCUSSION

Two pathogens were isolated from infested leaf portion of Horse purslane and by microscopic observations they were identified as *Gibbago trianthemae* and *Curvularia tuberculata*.

3.1 Morphological Characters of Gibbago trianthemae

Culture colour is a grevish-white with cottony growth [Fig 1]. The mature culture appears black in colour. On PDA, the subsurface mycelia development was black and dense and the fuzzy aerial mycelium appeared with moderate levels of sporulation. Mycelia was pale straw-colored and septate. Conidia produced in culture were characterized by means of secondary conidiophores.Conidiophores simple or 1-2 transeptate, pale proliferated. 1-4 strawcolored.Conidia solitary, almost completely ellipsoid, with 2 entire longitudinal septa intersecting at right angles in each conidium half, 1-4 whole or partial transverse septa, and a few shorter septa in transverse sectors.



Fig. 1	I. C	olony	of	Gibbago	trianthemae
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3.2 Morphological Characters of *Curvulariatuberculata*

Culture on PDA is brown [Fig 2], mycelium on a natural substrate is typically submerged, and the hyphae are branching, septate, and brown, smooth or verrucose. Stomata generally large, upright, black, cylindrical, sometimes branched.

Mycelial colour ranges from grey to black and has septa.Conidia are straight or curved, ovoid, obclavate or ellipsoidal, 3-5 septate, intermediate cells brown to dark brown, end cells sub hyaline to pale or dark brown, mature conidia tuberculate. Young conidia are sub hyaline and smooth. First septum in the conidium is usually median, second septum often delimiting the basal cell but variations in septal formation may occur.



Fig. 2. Colony of Curvularia tuberculata

3.3 Growth of Two Isolates

On PDA media *Curvularia tuberculata* growth was faster when compared to *Gibbago trianthemae*.

3.4 Assay on the Pathogenicity of Fungal Species

At 10, 20, 30, and 40 days after inoculation (spore treatment) on Horse purslane, the

pathogenicity of fungal species was recorded. From each pot of green house plants, a random sample of leaves was taken to figure out the infection rate. The leaves were assessed using a disease rating scale based on the increase in disease area, and a percent disease index was calculated for various treatments. Test plants inoculated with *Gibbago trianthemae* showed more symptoms [Fig 3] and high percent disease index values than *curvularia tuberculata* [Fig 4].

Table 1. Colony growth of Gibbago trianthemae

Days after sub culturing	Colony diameter(cm)
1	0.5
5	4
10	5
15	7

Table 2. Colony growth of Curvulariatuberculata

Days after sub culturing	Colony diameter(cm)
1	1
5	5
10	6
15	9

Table 3. Disease intensity on test plants inoculated with Gibbago trianthemae

S.No	Treatment Details Percent Disease Index		(%)		
		10DAT	20DAT	30DAT	40DAT
1	T₁- <i>Gibbago trianthemae</i> @2x 10 ⁶ cfu/g	0.00	9.77	19.54	28.40
		(0.00)	(18.20)	(26.22)	(32.19)
2	T₂- <i>Gibbago trianthemae</i> @4×10 ⁶ cfu/g	0.00	16.02	21.48	34.72
		(0.00)	(23.59)	(27.57)	(36.08)
3	T₃ - <i>Gibbago trianthema</i> e@6×10 ⁶ cfu/g	0.00	21.21	27.03	43.34
		(0.00)	(27.39)	(31.31)	(41.17)
4	T₄- <i>Gibbago trianthemae</i> @8×10 ⁶ cfu/g	7.95	25.96	31.84	50.78
		(16.37)	(30.63)	(34.35)	(45.44)
5	T₄- <i>Gibbago trianthemae</i> @8×10 ⁶ cfu/g	12.56	30.30	42.34	68.50
		(20.75)	(33.20)	(40.59)	(55.90)
6	T ₆ -control	0.00	0.00	0.00	0.00
		(0.00)	(0.00)	(0.00)	(0.00)
SEd		0.39	1.34	1.63	2.28
CD 5%		0.96	3.27	3.99	5.59
CD 1%		1.45	4.96	6.04	8.47

Datainparenthesesare arcsine √per cent angular transformed values DAT=Days after treatment

S.No	Treatment Details	P	Percent Disease Index (%)		
		10DAT	20DAT	30DAT	40DAT
1	T₁-Curvularia tuberculata @2×10 [°] cfu/g	0.00	0.30	12.39	16.24
		(0.00)	(3.09)	(20.59)	(23.74)
2	T₂ - <i>Curvularia tuberculata</i> @4×10 ⁶ cfu/g	0.00	9.06	16.86	21.92
		(0.00)	(17.49)	(24.23)	(27.87)
3	T₃ - <i>Curvularia tuberculata</i> @6×10 [°] cfu/g	0.00	12.17	20.22	26.57
		(0.00)	(20.42)	(26.71)	(31.02)
4	T₄ - <i>Curvularia tuberculata</i> @8×10 [°] cfu/g	0.00	13.41	23.50	32.33
		(0.00)	(21.46)	(28.96)	(34.64)
5	T₅ - <i>Curvularia tuberculata</i> @10×10 ⁶ cfu/g	0.00	17.29	26.67	41.48
		(0.00)	(24.56)	(31.09)	(40.09)
6	T ₆ -control	0.00	0.00	0.00	0.00
		(0.00)	(0.00)	(0.00)	(0.00)
SEd		-	1.00	1.49	1.70
CD 5%		-	2.46	3.66	4.17
CD 1%		-	3.72	5.54	6.32

Table 4. Disease intens	ty on test	plants inoculated	with curvularia	tuberculata
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Data in parentheses are arcsine \sqrt{per} cent angular transformed values DAT=Days after treatment

Table 5. Pathogenicity of isolates on Horse purslane after spore treatment

Fungal Species	Isolated part	Score	Disease intensity
Gibbago trianthemae	Leaf	4	Severe symptoms
Curvularia tuberculata	Leaf	2	Moderate symptoms



A. Leaf spot symptoms



B. Increased lesions

Fig. 3. Plants treated with Gibbago trianthemae



C. Completely dried plant



D. Leaf spot symptoms



E. Partially dried plant



F. Control plant

Fig. 4. Plants treated with Curvularia tuberculata

4. CONCLUSION

Two fungal species were isolated and identified from diseased leaves of Horse purslane *(Trianthema portulacastrum)* which was major weed in agricultural crops in research area. Pathogenicity test was done by spore treatment on test plants grown in green house conditions and it was revealed that the isolate *Curvularia tuberculata* displayed mild symptoms on the host plant, and has low percent disease index than *Gibbago trianthemae*, which is more pathogenic, quickly infected the host plants. Hence *Gibbago trianthemae* can be used as potent biocontrol agent against Horse purslane.

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COMPETING INTERESTS

Authors have declared that no competing interests exist.

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