Supplementing Pochonia chlamydosporia with botanicals for management of Meloidogyne incognita infesting chickpea

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Supplementing *Pochonia chlamydosporia* with botanicals for management of *Meloidogyne incognita* infesting chickpea

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**ABSTRACT**

Root-knot nematodes pose a severe threat to worldwide agricultural development. Due to the high toxicity of chemical nematicides, eco-friendly control strategies against root-knot nematodes need to be established. A pot and *in vitro* experiment were performed to estimate nematicidal potential of *Pochonia chlamydosporia*. *P. chlamydosporia* was used alone or in combination with two botanicals for controlling *Meloidogyne incognita* in chickpea. The laboratory assessment was performed with four prepared concentrations (S, S/2, S/10, S/25) of fungal inoculum of *P. chlamydosporia* against egg hatching and second-stage juvenile’s mortality of *M. incognita*. All four concentrations reduced egg hatching and increased mortality of J2s. In pot experiment, *P. chlamydosporia* was used with chopped leaves of two botanicals *viz.*, *Ageratum conyzoides* and *Eichhornia crassipes* against *M. incognita* in chickpea. All the treatments found significantly suppressed root infestation caused by *M. incognita* and improved growth and physiological attributes of chickpea. The combined application of *P. chlamydosporia* + *A. conyzoides* was found highly effective, and *E. crassipes* alone was least. Therefore, using *P. chlamydosporia* with botanicals is a promising sustainable strategy in agriculture against *M. incognita* infected chickpea.

**Results**

The combined application of *P. chlamydosporia* + *A. conyzoides* was found highly effective, and *E. crassipes* alone was least. Therefore, using *P. chlamydosporia* with botanicals is a promising sustainable strategy in agriculture against *M. incognita* infected chickpea.

**Introduction**

Pulses seem to be the essential foods for vegetarians in the entire world. Pulses are continuously gaining prominence as an affordable protein source and a vital component for sustainable agriculture. Chickpea (*Cicer arietinum* L., Family-Fabaceae) is the world’s second-largest legume crop grown globally across an area of 14.6 M hac (FAOSTAT 2017). Numerous factors significantly reduce chickpea production, including insect pests, fungi, bacteria, and nematodes (Zwart et al. 2019). *Meloidogyne incognita* is a root-knot nematode (RKN) and a sedentary endo-parasite (SEP) that affects thousands of plant species, including pulses and vegetables, and is among the most dangerous agricultural ailments. In India, *Meloidogyne* species such as *M. incognita*, *M. javanica* have been estimated to show losses of 19–40% and 24–61% to chickpea production, respectively (Ali and Naimuddin 2010). RKNs reduce the growth of host crops and intervene with nodulation and nitrogen fixation (Rehman et al. 2012). Indeed, considering the high economic effect of RKNs, several management techniques in agricultural production have been introduced, *viz.*, use of chemical nematicides, biocontrol agents from botanicals, and resistant varieties. The most successful way of handling RKNs, primarily through the use of chemical nematicides, but, on the other hand, most of them are forbidden due to their toxic impact on flora and fauna, which has also disrupted soil structure and the climate. Applying biocontrol agents is suggested as a much better option and highly feasible for the RKN management programme. Fungal biocontrol agents may suppress the RKNs by various modes, producing toxins, antibiotics, and enzymes or interference with the detection of the nematode-plant host. Colonisation of root by *Pochonia chlamydosporia* provided good efficacy and advantages for host plant about growth parameters, defense mechanism against many pathogens *viz.*, nematodes, bacteria, and fungi (Siddiqi and Akhtar 2008). Several biocontrol agents *viz.* *P. chlamydosporia*, *Paecilomyces lilacinus*, and Trichoderma harzianum have...
The inoculum of root-knot nematode, *M. incognita* was maintained in a glasshouse on eggplant. Identification of *M. incognita* was performed based on perineal pattern (Sasser and Carter 1982). Eggs of *M. incognita* separated from infected eggplant roots by dipping roots in 0.05% NaOCl for 3-4 min (Hussey and Barker 1973). Then eggs were collected and washed with DDW on 25 μm pore size sieves. The egg masses were left for hatching in a BOD incubator, and J2s that hatched from the eggs on a sieve were collected after five days. After that, fresh egg masses and J2s were used for the hatching, mortality, and pots bioassay.

**Preparation of cultural filtrate of *P. chlamydosporia***

To obtain adequate inoculum, *P. chlamydosporia* inoculated by a sterile inoculation needle in Richard’s liquid medium (Riker and Riker 1936). The fungal mycelia mat on filter paper was washed in DDW, washed away excess water and nutrients with blotting paper. The fungal inoculum was prepared by mixing 10 g of mycelium mat in 100 ml of double distilled water and blending it (10,000 rpm) in a waring blender for 30 s. Ten millilitres of the fungal inoculum of *P. chlamydosporia* were used for inoculation in pots. The fungal inoculum obtained was labelled as ‘S’ (Standard) and being used as a stock suspension. Successive concentrations viz., S/2, S/10, S/25 were prepared by adding the required amount of DDW for hatching and mortality experiments (Mukhtar et al. 2013).

**Effect of different concentrations of fungal inoculum of *P. chlamydosporia* on the mortality of J2s of *M. incognita***

Four concentrations (S, S/2, S/10, S/25) of fungal inoculum of *P. chlamydosporia* were prepared to determine the effect on mortality of J2s. For the mortality test, 1 ml of DDW containing 150 freshly hatched J2s poured into petri dishes having 9 ml of different concentrations of fungal inoculum mentioned above. Petri dishes having only DDW were used as control. Each concentration had five replicates. The petri dishes were sealed with a lid and wrapped in parafilm. Petri dishes were placed at 28°C, and the number of live and dead J2s counted after 6, 12, 24, and 48 hrs of incubation using a stereoscopic microscope. The mortality of J2s was recorded according to the mean percentage of dead J2s. The J2s displayed flexibility or looked as winding shapes declared alive (El-Rokiek and El-Nagdi 2011), and if J2s did not show any motion and their body outline looks straight, they were found dead. Living and dead J2s were calculated under the binocular microscope, and the percentage of mortality was

**Materials and methodology**

**Plant, fungi, and root-knot nematode**

Chickpea cv. *Avrodhi* was used as a host crop. Seeds were surface sterilised in a 1.0% NaOCl for 15 min, washed 3–4 times in Double Distilled Water (DDW) for 10 s each. The pure culture of a fungal biocontrol agent, *P. chlamydosporia* (ITCC No. 6898), was purchased from the Indian Agricultural Research Institute (IARI), New Delhi, India. Fungal culture was preserved on the medium of Potato Dextrose Agar (PDA). The identification of pure fungal culture was based on microscopic (conidial, colourful, conidium, vesicle shaped) and macroscopic features (diameter, colour, and colony texture). For the mass production of *P. chlamydosporia*, Richards Medium was utilised. The RKN, *M. incognita*, was used as a pathogen.

**Maintenance of the inoculum of root-knot nematode, *M. incognita***

To obtain adequate inoculum, *P. chlamydosporia* inoculated by a sterile inoculation needle in Richard’s liquid medium (Riker and Riker 1936). The fungal mycelia mat on filter paper was washed in DDW, washed away excess water and nutrients with blotting paper. The fungal inoculum was prepared by mixing 10 g of mycelium mat in 100 ml of double distilled water and blending it (10,000 rpm) in a waring blender for 30 s. Ten millilitres of the fungal inoculum of *P. chlamydosporia* were used for inoculation in pots. The fungal inoculum obtained was labelled as ‘S’ (Standard) and being used as a stock suspension. Successive concentrations viz., S/2, S/10, S/25 were prepared by adding the required amount of DDW for hatching and mortality experiments (Mukhtar et al. 2013).

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determined using the formula (Sun et al. 2006).
Per cent mortality of J2s = 100 × Dead J2s/TotalJ2s

Effect of different concentrations of fungal inoculum of P. chlamydosporia on egg hatching of M. incognita

The egg inhibitory activity of various concentrations (S, S/2, S/10, S/25) of fungal inoculum of P. chlamydosporia was tested by egg mass dipping method. Six healthy egg masses of M. incognita were collected from the infected root of eggplant using forceps and poured into petri dishes with 10 ml of the different concentrations of fungal inoculum referred to above. Petri dishes were coated with parafilm to avoid evaporation and then placed at 28°C. Egg masses were dipped in DDW served as control. Each concentration had been repeated five times, excluding control. The hatching value was recorded by counting the number of hatched J2s in each replicate after 3, 5, and 7 days of incubation using a binocular microscope and calculating the per cent inhibition in egg hatching of each replicate using the following formula (Khan et al. 2019).

Per cent inhibition in egg hatching = (C0 - Tα) × 100/C0

Where,

C0 = Number of J2s that emerged from egg masses in control (DDW)
Tα = Number of J2s emerged from egg masses in each concentration of fungal inoculum of P. chlamydosporia

In planta bioassay (pot study)

A pot experiment was laid out in Plant Pathology and Plant Nematology glasshouse, Department of Botany, Aligarh Muslim University, Aligarh. Seeds of susceptible chickpea cv. Avrodhi was purchase from the local market of Aligarh. The earthen pots (15 cm wide) loaded with 1 kg of autoclaved soil in proportion to 3:1 (sandy loam: farmyard manure). Pots were amended with 15 g of freshly chopped leaves of A. conyzoides and E. crassipes. The pots were arranged according to the treatment set-up and put in a completely randomised glasshouse (Temperature-18-25°C; Relative humidity-95-100%). Pots were regularly watered for the complete decomposition of chopped leaves. After 10 days of adjustment, 4-5 sterilised chickpea seeds sown 1-2 in. deep in each pot. When the seedlings grew two sets of leaves, the plants were thinned. Each of the healthy and stable seedlings was selected per pot, and the remaining ones removed from the ground level, including control. Aqueous suspension with 2500 freshly hatched J2 of M. incognita was inoculated evenly around each seedling root. Two days after the inoculation, a sterile pipette added an S concentration level (10 mL) of P. chlamydosporia around chickpea roots. In control, pots were inoculated with 10 ml of DDW in place of fungal inoculum of P. chlamydosporia. Plants have been irrigated sufficiently throughout the experiment and carefully to eliminate errors during the examination. After 60 days, the plants were uprooted and washed with flowing tap water to remove adhered soil particles, and assessments were performed. The growth, physiological and pathological parameters of chickpea plants are presented in tables and figures.

Photosynthetic pigments analysis

Chlorophyll content estimation bioassay

The chlorophyll content of fresh leaves of each treatment was estimated using the Mackinney (1941) process. Take 1 g of leaves from each treatment, ground in a blender with 20 ml of 80% acetone, and centrifuge at 5000 rpm for 5 min. The supernatant was collected, and absorption determined at 645 and 663 nm against (blank) 80% acetone on the spectrophotometer (UV 1700 Shimadzu, Japan). The chlorophyll content of each treatment (mg g⁻¹ tissue) was evaluated using the following formula.

Total Chlorophyll g⁻¹ tissue = 20.2(A645) + 8.02(A663)

V × 1000 X W

where,

A = Absorbance of leaf sample read at 645 and 663 nm
V = Final volume,
W = Fresh weight of leaves,
D = Length of the path of light

Carotenoid content estimation bioassay

The carotenoid content in leaves samples was observed using the method given by MacLachlan and Zalik (1963). To determine carotenoid content, the protocol for preparation of extract of leaves of each treated pot is the same as of chlorophyll bioassay. The absorbance of the extracts was observed at wavelength 480 and 510 nm against blank (80%) acetone on the spectrophotometer. The carotenoid value was calculated using the following
formula.

\[
\text{Carotenoid Content} = 7.6(\text{O.D.} 480) - 1.49(\text{O.D.} 510)\times \frac{V}{\text{DXWX} 1000}
\]

\(\text{O.D.}\) = Optical density of extract of leaf sample read at 480 and 510 nm  
\(V\) = Final volume  
\(W\) = Fresh weight  
\(D\) = Length of the path of light

**Pathological analysis**

**Nematode population in soil**

At termination, soil samples were taken from each treated pot using a garden trowel then mixed thoroughly. Weigh 250 g of well-mixed soil from each treatment to estimate the population of J2s. Estimation of the nematode population (J2s) was done by Cobb’s sieving and decanting technique (Cobb 1918), followed by modified Baermann’s funnel method (Southey 1986). Counting of J2s was done under the stereoscope microscope (Motic SMZ 168 series).

**Statistical analysis**

Statistical analysis was performed using the Duncan Multiple Range Test (DMRT) of complete randomised block design using R (version 2.14.1 software). The standard error of the mean (±SE) and Least Significant Difference (LSD) was calculated at a 5% significance level. The Principal Component Analysis (PCA) was calculated using Origin software [version 2019b (9.65)]. Microsoft Excel assessed the coefficient of correlation.

**Results**

**Nematicidal effect of P. chlamydosporia on egg hatching of M. incognita in vitro**

All four selected concentrations (S, S/2, S/10, S/25) of fungal inoculum were significantly inhibited egg hatching of *M. incognita* at \((p < 0.05)\). There was an increased inhibition in egg hatching when the concentration of *P. chlamydosporia* was increased from S/25 level to S level. Maximum inhibition in egg hatching was observed at S concentration, while minimum inhibition was reported at S/25. S/10 and S/25 concentrations of *P. chlamydosporia* also showed a decrease in efficacy as compared to S and S/2. The individual inhibitory effect of the used concentration of *P. chlamydosporia* is given in Table 1.

**Nematicidal effect of P. chlamydosporia on second-stage juvenile’s mortality of M. incognita in vitro**

The findings showed an apparent effect of four concentrations (S, S/2, S/10, S/25) of *P. chlamydosporia* on the mortality of J2s of *M. incognita* after 6, 12, 24, and 48 hrs of the incubation period \((p < 0.05)\). The *P. chlamydosporia* were toxic to J2s of *M. incognita* compared to distilled water (control). The standard concentration (S) showed maximum mortality compared to other concentrations (S/2, S/10, S/25). All concentrations showed the highest mortality of J2s at 48 hrs of incubation period compared to 6, 12, 24 hrs. The S concentration was found highly toxic to J2s at 48 hrs of incubation, and this toxicity was found significant compared to the other concentrations. Similarly, the incubation period also significantly influenced the mortality of J2s and reached the maximum after 48 hrs of incubation compared to 6, 12, and 24 hrs. As the concentration of fungal inoculum increase from S/25 to S, there is an increase in the mortality of J2s. Individual toxic effect of four concentrations of *P. chlamydosporia* on J2s of *M. incognita* shown in Table 2.

**Pot experiment**

**The nematicidal potential of P. chlamydosporia alone or in combination with chopped leaves of botanicals on the growth attributes of chickpea**

The inoculum of *P. chlamydosporia* (10 mL) alone or combined with chopped leaves (15 g) of *A. conyzoides* and *E. crassipes* showed improvement in infected chickpea’s growth attributes. All treatments substantially \((p < 0.05)\) enhanced the chickpea growth attributes (Figure 1). Application of combined treatment of *P. chlamydosporia* + *A. conyzoides* gave the highest improvement in plant length \((56 ± 2.17\) cm), plant fresh weight \((137.4 ± 4.12\) g), number of pods and nodules \((34.80 ± 1.74\) and \(34.60 ± 1.83\)), and followed by *P. chlamydosporia* + *E. crassipes* (plant length \(-49.6 ± 1.33\) cm, plant fresh weight \(-111.20 ± 3.64\) g, number of pods \(-28.60 ± 2.60\), number of nodules \(-30.20 ± 1.74\)), *P. chlamydosporia* alone (plant length \(-43 ± 1.76\) cm, plant fresh weight \(-83 ± 2.65\) g, number of pods \(-23.40 ± 2.02\), number of nodules \(-25.60 ± 2.82\)), *A. conyzoides* alone (plant length \(-37.6 ± 1.01\) cm, plant fresh weight \(-60.80 ± 2.45\) g, number of pods \(-20.60 ± 1.30\), number of nodules \(-22.40 ± 1.77\)), and least was observed in *E. crassipes* alone (plant length \(-33.4 ± 1.53\) cm, plant fresh weight \(-52.80 ± 1.64\) g, number of pods \(-16.20 ± 1.74\), number of nodules \(-20 ± 1.15\)) as compared to those plant
treated with nematode only (plant length $-23.8 \pm 1.24$ cm, plant fresh weight $-44 \pm 2.16$ g, number of pods $-10.60 \pm 0.87$, number of nodules $-14.20 \pm 1.47$) (Figure 1). The findings revealed that there exists a strong linear relationship between galls/plant and plant length ($R^2 = 0.78$); plant fresh weight ($R^2 = 0.54$); number of pods ($R^2 = 0.73$); the number of nodules ($R^2 = 0.72$); chlorophyll content ($R^2 = 0.80$); carotenoid content ($R^2 = 0.67$). As the relation is positive, the increase in the number of galls increased the per cent reduction of growth attributes of chickpea (Figure 2).

**Nematicidal potential of *P. chlamydosporia* alone or in combination with chopped leaves of botanicals on the physiological attributes of chickpea**

The application of *P. chlamydosporia* (10 ml) alone or in combination with chopped leaves (15 g) of *A. conyzoides* and *E. crassipes* enhanced the physiological attributes (chlorophyll and carotenoid content) of chickpea. Significantly, maximum chlorophyll content ($2.51 \pm 0.05$ mg/g) and carotenoid content ($0.454 \pm 0.007$ mg/g) were achieved when combined treatment *P. chlamydosporia* + *A. conyzoides* was applied. Other treatments also enhanced physiological attributes (chlorophyll and carotenoid content) viz., *P. chlamydosporia* + *E. crassipes* ($2.34 \pm 0.03$ and $0.411 \pm 0.13$ mg/g), *P. chlamydosporia* alone ($1.98 \pm 0.04$ and $0.356 \pm 0.11$ mg/g), *A. conyzoides* alone ($1.87 \pm 0.04$ and $0.332 \pm 0.11$ mg/g), and *E. crassipes* alone ($1.65 \pm 0.07$ and $0.302 \pm 0.10$ mg/g) as compared to those plant treated with nematode only ($1.24 \pm 0.04$ and $0.264 \pm 0.08$ mg/g) (Figure 3).

**The nematicidal potential of *P. chlamydosporia* alone or in combination with chopped leaves of botanicals on the nematode population and galls/plant**

The treatments significantly ($p < 0.05$) reduced the pathological parameters such as nematode population, and galls/plant (Figure 4). Treatment with combined application of *P. chlamydosporia* + *A. conyzoides* achieved the highest reduction in galls ($8.40 \pm 0.41$) followed *P. chlamydosporia* + *E. crassipes* ($14.20 \pm 1.41$), *P. chlamydosporia* alone ($25.20 \pm 1.64$), *A. conyzoides* alone ($33.30 \pm 1.81$), *E. crassipes* alone ($38.60 \pm 2.05$) compared to the pot treated with nematode only ($112 \pm 7$). When counting the nematode population in 250 g soil, a significant reduction was seen compared with the control. In the same manner, *P. chlamydosporia* + *A. conyzoides* was most prominent in suppressing the nematode population ($455 \pm 16.07$). The least suppressive effect was found in *E. crassipes* alone ($1002 \pm 24.84$) as compared to pot treated with an only nematode ($3024 \pm 28.21$) (Figure 4). The outcome of the principal component analysis showed that the population of *M. incognita* in soil and galls per plant was strongly correlated with other parameters of chickpea. Scatter biplot showed that *P. chlamydosporia* combined with chopped leaves of *A. conyzoides* was found highly effective, reduced the infestation caused by *M. incognita* considerably, and improved chickpea’s growth attributes (Figure 5).

**Discussion**

In *in vitro* experiment, the tested concentrations viz., S, S/2, S/10, S/25 of fungal inoculum were effectively inhibited egg hatching of egg masses. We also found J2s mortality of *M. incognita* in the same fungal concentration (Tables 1 and 2). Standard concentration ‘S’ was found highly effective in reducing egg hatching of *M. incognita*, followed by S/2, S/10, S/25 (Table 1). However, contrary findings were also reported by Uddin et al. (2019) and Singh and Mathur (2010). They found in their studies that treatment with *P. chlamydosporia* gave no significant result in egg hatching inhibition. The maximum mortality was 85% observed at 48 hrs of incubation in standard ‘S’ concentration of *P. chlamydosporia* followed by S/2, S/10, S/25 (Table 2). In contrast, *chlamydosporia* showed 11.3-

**Table 1. Effect of different concentrations of *P. chlamydosporia* on egg hatching of *M. incognita* at 3, 5 and 7 days of incubation period.**

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Time (Days)</th>
<th>S</th>
<th>S/2</th>
<th>S/10</th>
<th>S/25</th>
<th>DDW</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>P. chlamydosporia</em></td>
<td>3</td>
<td>57.60±2.60</td>
<td>76.60±2.60</td>
<td>90±3.78</td>
<td>116±3.46</td>
<td>340±6.42</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>88.80±2.31</td>
<td>110.20±3.81</td>
<td>134.40±4.63</td>
<td>161.20±4.04</td>
<td>340±6.42</td>
</tr>
<tr>
<td></td>
<td>7</td>
<td>122.20±4.34</td>
<td>138.20±4.11</td>
<td>158.60±5.78</td>
<td>186.40±4.06</td>
<td>340±6.42</td>
</tr>
<tr>
<td><em>E. crassipes</em></td>
<td>3</td>
<td>64.05%</td>
<td>59.55%</td>
<td>53.55%</td>
<td>45.17%</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>73.88%</td>
<td>67.586%</td>
<td>60.47%</td>
<td>52.58%</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>7</td>
<td>110.20±3.81</td>
<td>112.20±4.34</td>
<td>138.20±4.11</td>
<td>158.60±5.78</td>
<td>340±6.42</td>
</tr>
</tbody>
</table>

Each value is an average of five replicate; DDW = Double Distilled Water (Control); SE = Standard Error; J2s = Second-stage juveniles; Values are given in parentheses represent percent inhibition in egg hatching over control; Values are given without parentheses represent number of the hatched J2s of *M. incognita* in different concentrations.
76.3% mortality of J2s of *M. incognita* after 72 hrs of exposure, according to a published report by Uddin et al. (2019). The nematicidal effect of fungal inoculum was increased when exposure time extended. Meyer et al. (2004) and Elbadri et al. (2008) reported in their study that the impact of fungal inoculum varied from concentration to concentration, thus confirming these findings. In our study, the concentration and incubation period were important factors. The nematicidal action of *P. chlamydosporia* may be due to direct parasitism of fungus mycelium on nematode, or perhaps secondary metabolites released in surroundings, which caused mortality of J2s after treatment. Fungal activity as the nematicidal impact on adult nematodes, suppression

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Time (hours)</th>
<th>Number of dead J2s (Mean ± SE) in different concentrations</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>S</td>
<td>S/2</td>
</tr>
<tr>
<td><em>P. chlamydosporia</em></td>
<td>6</td>
<td>56.20±2.69 (37.46%)</td>
</tr>
<tr>
<td></td>
<td>12</td>
<td>77.20±2.61 (51.46%)</td>
</tr>
<tr>
<td></td>
<td>24</td>
<td>98±2.30 (65.33%)</td>
</tr>
<tr>
<td></td>
<td>48</td>
<td>128±3.21 (85.33%)</td>
</tr>
</tbody>
</table>

LSD (*p* = 0.05) 8.89 8.75 5.98 6.93 0

Each value is an average of five replicates; DDW = Double Distilled Water (control); SE = Standard Error; J2s = Second-stage juveniles; Values are given in parentheses represent per cent J2s mortality over control; Values are given without parentheses represent number of the dead J2s of *M. incognita* in different concentrations.

Figure 1. Nematicidal effect of *P. chlamydosporia* alone or in combination with chopped leaves of *A. conyzoides* and *E. crassipes* on the growth attributes of chickpea infested with *M. incognita* (Pc + Ac = *P. chlamydosporia* + *A. conyzoides*; Pc + Ec = *P. chlamydosporia* + *E. crassipes*; Pc alone = *P. chlamydosporia* alone; Ac alone = *A. conyzoides* alone; Ec alone = *E. crassipes* alone).
of egg hatching and juvenile growth have also been tested (Meyer et al. 2004). Benedict and Brady (1972) found that antagonistic compounds were released, responsible for reducing the J2s population.

Similarly, in pot study, P. chlamydosporia alone or in combination with chopped leaves of A. conyzoides and E. crassipes significantly enhanced growth attributes of chickpea and decreased disease severity caused by M. incognita to varying degrees. The combined application of P. chlamydosporia with chopped leaves of A. conyzoides caused the highest reduction in root galling and nematode population (Figure 4). It improved the growth and physiological attributes of chickpea (Figures 1 and 3). However, contrary findings were also reported by Podestá et al. (2013), who found that fungus P. chlamydosporia had little effect on destroying the nematode eggs, reducing the number of galls by only 2.68%. Analysis of correlation coefficient exhibited that number of galls had a positive correlation with plant length, plant fresh weight, chlorophyll and carotenoid content and number of pods and nodules (Figure 2). Scattered points which are existing in graphs show whether two variables have a relationship or not. Maximum scattered points with minimum correlation were found between the number of galls and plant fresh weight ($R^2 = 0.544$) with positive correlation and maximum correlation with highly condensed points observed number of galls index and chlorophyll

![Figure 2. Relationship between the number of galls and per cent reduction in various growth attributes of chickpea (Pc + Ac = P. chlamydosporia + A. conyzoides; Pc + Ec = P. chlamydosporia + E. crassipes; Pc alone = P. chlamydosporia alone; Ac alone = A. conyzoides alone; Ec alone = E. crassipes alone).](image-url)
content ($R^2 = 0.800$) (Figure 3). Our finding confirmed with Rich et al. (1984), reported that significant positive correlations were observed between nematode numbers and plant yield of tobacco.

As we observed in our pot study that *P. chlamydosporia* and botanicals have a significant impact against nematodes. Our finding confirmed with Naz et al. (2021), reported that applying biocontrol agents in combination may be an attractive measure because the combined use of different organisms may produce synergistic effects that facilitate the management of RKNs. That might be due to the release of bioactive compounds after decomposing plant leaves, altering the current nematode pattern and enhancing chickpea’s growth. Adding botanicals to soil improves soil quality, acts as a nutrient reservoir and provides the perfect habitat for plant growth. Our findings are similar to Aktar and Malik (2000), they reported that the addition of organic matter increases the nutrient content and improves the soil texture, promoting the development of antagonistic microorganisms, while its breakdown produces toxic compounds against RKNs. Biocontrol agents colonised roots better in organic amendments soil, resulting in better protection of plants from various pathogens and thus improved plant growth. Once organic matter is applied in soil may release compounds with nematicidal effect and support the increase in native antagonist population in

**Figure 3.** Nematicidal effect of *P. chlamydosporia* alone or in combination with chopped leaves of *A. conyzoides* and *E. crassipes* on the physiological attributes of chickpea infested with *M. incognita* (*Pc + Ac = P. chlamydosporia + A. conyzoides; Pc + Ec = P. chlamydosporia + E. crassipes; Pc alone = P. chlamydosporia alone; Ac alone = A. conyzoides alone; Ec alone = E. crassipes alone).*

**Figure 4.** Nematicidal effect of *P. chlamydosporia* alone or in combination with chopped leaves of *A. conyzoides* and *E. crassipes* on the pathological attributes of chickpea (*Pc + Ac = P. chlamydosporia + A. conyzoides; Pc + Ec = P. chlamydosporia + E. crassipes; Pc alone = P. chlamydosporia alone; Ac alone = A. conyzoides alone; Ec alone = E. crassipes alone).*
the soil or serve as the substrate for the development and the establishment of antagonists in soil (Cannayane and Rajendran 2001). Egg parasitic fungi, P. chlamydosporia and P. lilacinus are attractive biocontrol agents for RKNs management programmes in a wide variety of crops (Anastasiadis et al. 2008; Ebrahim et al. 2008). The fungus P. chlamydosporia colonises and infects RKNs eggs and exposed females, reducing the number of infective second stage juveniles (Manzannilla-López et al. 2013). After decomposition of organic additives/matters, release some nutrients responsible for the improvement in plant nutrition (Reiner 2015) and the development of plant resistance against nematode infection (Reiner 2015). A synergistic impact observed on the management of M. incognita on chickpea was due to P. chlamydosporia with botanicals. A significant decrease in nematode infection could be due to release of secondary metabolites by Pochonia spp. and botanicals that have nematicidal behaviour. Several metabolites have been isolated from various plants with a wide variety of structures and their nematicidal activities are assessed against multiple species of nematodes (Faria et al. 2016; Khan et al. 2017). The assumption derived from the analysis is that biocontrol agents and botanicals enhanced growth attributes and yield of chickpea and reduced the infestation caused by M. incognita. Applying these commodities, which are readily available domestically, would be far more beneficial from the economic perspective of farmers if adequately used, especially for small-scale farmers who could not buy expensive chemical nematicides. Therefore, our approach to applying P. chlamydosporia alone or in combination with chopped leaves of selected botanicals in agricultural practices could benefit the soil’s physicochemical attributes and provide an environmentally friendly option for the sustainable management of root-knot nematode. Our findings supported the benefits of biocontrol agents and botanicals by keeping down the root-knot disease in chickpea plants. Therefore, using fungal inoculum (P. chlamydosporia) with selected botanicals may be a suitable environmentally friendly option to keep the nematode population below the threshold level.

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Declaration of interest statement
The authors declare that there is no conflict of interest.

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