Original Research Article

- 2 Development of Bioformulations for the Management of Blackgram Dry Root Rot
- 3 Caused by *Rhizoctonia bataticola* (Taub Butler)
- 4 Abstract

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- 5 An attempt was made to control dry root rot using consortia of bioinoculants. A total of 10
- 6 fungal (Trichoderma) and 30 bacterial (Pseudomonas and Bacillus) isolates were collected
- 7 and screened for their antagonistic activity against mycelial growth of Rhizoctonia
- 8 bataticola under in vitro condition. Among these, Trichoderma (TL1), Pseudomonas
- 9 fluorescens (PfUL(A)) and Bacillus subtilis (BsOP2) isolates exhibited maximum inhibition.
- As results of the compatibility of the biocontrol agents revealed that *P. fluorescens* strains
- were compatible with B. subtilis and Trichoderma but B. subtilis strains were not compatible
- with *Trichoderma* strains. The biocontrol consortia consisting of *P. fluorescens* (PfUL(A))
- and B. subtilis (BsOP2) + FYM + Neem cake was found to be promising in reducing dry
- 14 root rot incidence under field conditions. The biocontrol consortia also induced high level of
- 15 defence related enzymes v phenylalanine ammonia lyase, catalase, peroxidise and
- 16 polyphenol oxidase activity.
- 17 Key words: Black gram, Dry root rot, Bacillus, Pseudomonas, Biocontrol consortia

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Introduction

- 20 The productivity of blackgram or urdbean (Vigna mungo (L.) Hepper) was reduced due to
- various diseases with an estimated yield loss of 20 to 30 percent. Fungicides are widely used
- as seed or soil treatment to combat various root diseases. However, use of fungicides causes
- 23 environmental hazards and development of resistance in pathogen. In recent years, more
- 24 emphasis has been given to the use of bioagents and organic amendments. Several
- 25 antagonistic organisms have been successfully used as biocontrol agents for controlling soil
- borne pathogens (Droby, 2001; Karthikeyan et al., 2005; Meyer and Roberts, 2002). At
- 27 present most of the biocontrol agents are applied singly to combat the growth of the
- 28 pathogens. Although the potential benefits of a single biocontrol agent application has been
- 29 demonstrated in many studies, it may also partially account for the inconsistent performance
- 30 because a single biocontrol agent is not likely to be active in all kinds of soil environment and
- 31 all agricultural ecosystems (Raupach and Kloepper, 1998). One of the strategies for

overcoming such inconsistent performance is to combine two or more beneficial microbes in a biocontrol formulations. Combinations of biocontrol agents have the potential for more extensive colonization of the rhizosphere, more consistent expression of beneficial traits under a wider range of soil conditions and antagonistic to a larger number of plant pathogens than biocontrol strains applied individually. Thus, more emphasis was laid on the combined use of two or more strains of biocontrol agents, which turned out to be more successful than either of them alone, as reported by several workers (Nandakumar *et al.*, 2001; Bharathi *et al.*, 2004; Thilgavathi *et al.*, 2007; Senthilraja *et al.*, 2010). Therefore, the present study was undertaken to evaluate the efficacy of biocontrol consortia consisting organic amendments *viz.*, neem cake and FYM against root rot disease of blackgram.

Materials and Methods

Isolation of pathogen and biocontrol agents

The dry root rot pathogen *R. bataticola* was isolated from infected black gram plants using potato dextrose agar (PDA) medium. The biocontrol agents *Trichoderma*, *Pseudomonas* and *Bacillus* were isolated from rhizosphere soils of black gram using Trichoderma selective medium (TSM) (Elad and Chet, 1983), King's B medium (KB) (King *et al.*, 1954) and Nutrient Agar (NA) medium (Rangaswami, 1972), respectively. The individual colonies of *Trichoderma* were identified based on the morphological characters (Webster and Lomas, 1964). Similarly, the bacterial isolates were characterized based on standard biochemical tests (Hildebrand *et al.*, 1992). Antagonism of *T. viride* against *R. bataticola* was assayed with the dual-culture (Dennis and Webster, 1971).

Compatibility of biocontrol agents

The bacterial strains were tested for their compatibility with each other following the method of Fukui *et al.* (1994). The compatibility of the fungal biocontrol agent with the bacterial strains was tested by their mycelial overgrowth on the bacterial strains without any inhibition zone, using the dual culture technique (Dennis and Webster, 1971).

Seed treatment

Blackgram seeds (ADT5) were surface sterilized with 2% sodium hypochlorite for 30 sec. then rinsed in sterile distilled water and dried overnight. Ten ml of the bacterial antagonist containing 3×10^8 cfu/ml was taken in Petri plate. To this 100 mg of carboxy methyl

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cellulose (CMC) was added as an adhesive material. One gram of seeds was soaked in 10 ml of bacterial suspension (containing 3x108cfu/ml) for 2 h and dried overnight in a sterile Petri plate. *Trichoderma* isolates multiplied in *Trichoderma* special broth were harvested for mycelial mats along with spores, then the contents were mixed with sterile distilled water and (20x108cfu/ml) was checked through dilution plate technique and subjected to seed treatment as above.

Assessment of plant growth promotion

- 71 Plant growth-promoting activity of the best isolates of *Pseudomonas* sp. (PfUL(A),
- 72 PfAL1 and PfCBE9), Bacillus sp. (BSOP2, BCBE1 and BKK3) and Trichoderma sp. (TL1,
- 73 TCBE3 and TOKK1) were assessed based on the seedling vigour index by the standard roll
- towel method (ISTA, 1993).

Preparation of biocontrol consortia

- Talc based formulation of *Pseudomonas, Bacillus* and *Trichoderma* were prepared as
- the methods discussed by (Vidhyasekaran and Muthamilan, 1995; Ramakrishnan et al. 1994),
- 78 The biocontrol consortia of fungal and bacteria was prepared by mixing equal quantity of talc
- 79 based formulation both the biocontrol agents w/w.

Effect of talc-based bioformulations with organic amendments

81 Potting medium (red soil:sand:cow dung at 1:1:1, w/w/w) was autoclaved for 1 hour 82 for two consecutive days and filled in pots and incorporated with sand:maize inoculum 83 (50g/pot) of R. bataticola. Ten gram of talc based bioformulation was mixed with ten gram of 84 FYM and neem cake then applied per pot as soil application (capacity of pot 10% of potting 85 medium filled). Seeds of black gram cv. ADT-5 were surface sterilized with 2% sodium hypochlorite, seed treated with sown 20 seeds per pot. Carbendazim at the rate of 2 g kg⁻¹ 86 87 of seed was applied as a chemical check. Ten seedlings were maintained per pot up to 20 88 days. Further, five seedlings were maintained until harvest. The pathogen inoculated and 89 uninoculated served as control. Soil drenching of 0.1 per cent carbendazim was included as 90 chemical check. Three replications (three pots per replication) were maintained and the pots 91 were arranged in a randomized manner. The treatments of experiment were, T1 -92 **Psudomonas** sp (PfUl(A))Neem cake Farm Yard Manure, 93 T2 - Bacilllus sp (BsOP2) + Neem Cake + FYM, T3- Trichoderma (TL1) + Neem cake +

FYM, T4 - PfUl(A)+TL1+ Neem cake + FYM, T5 - BsOP2+ TL1+Neem cake + FYM,

T6 –PfUl(A)+BsOP2+ + TL1+Neem cake + FYM, T7 – Carbendazim (0.1%), T8 –Inoculated control and T9–Healthy control. The incidence of root rot was recorded and expressed as percentage of disease incidence.

Assay of defence enzymes

Samples of blackgram were collected from individual treatments at 2 days interval starting from zero, 1, 3,5,7 and 9th days after challenge inoculated with the pathogen to study the induction of defence related enzymes in response to treatment and inoculation of pathogen in black gram plants under glass house conditions. One g of root sample was homogenized with 2 ml of 0.1 M sodium phosphate buffer (pH 7.0) at 4°C. The homogenate was centrifuged for 20 min at 10,000 rpm and the supernatant was used to determine Phenylalanine ammonia lyase (PAL), Catalase (CL), Peroxidase (PO) and Polyphenol oxidase (PPO). Peroxidase activity was assayed as described by Hammerschmidt *et al.* (1982) and was expressed as changes in absorbance at 470 nm min⁻¹ g⁻¹ of fresh tissue. PPO activity was determined following the procedure given by Mayer *et al.* (1965) and was expressed as changes in absorbance at 470 nm min⁻¹ g⁻¹ of fresh tissue. PAL activity was assayed following the method of Ross and Sederoff (1992) and was expressed as nmoles of cinnamic acid min⁻¹ g⁻¹ of fresh tissue. Catalase activity was determined following the procedure given by Kato and Shimizu (1987) and was expressed as changes in absorbance min⁻¹ g⁻¹ of fresh tissue.

Field studies

Two field experiments were conducted to evaluate the efficacy of bioformulation on dry root rot disease incidence with organic amendments. The experiments were laid out in a randomized block design (RBD) with three replications. The treatments consisted of, T1 – Psudomonas sp (PfUl(A)) (10g ST + 2.5kg SA) + Neem cake (150 kg/ha)+ Farm Yard Manure (2 ton/ha), T2 – Bacillus sp (BsOP2) (10g ST + 2.5kg SA) + Neem Cake + FYM, T3 - Trichoderma (TL1) (4g ST + 2.5kg SA) + Neem cake + FYM, T4 - PfUl(A) (5g ST + 1.25kg SA) +TL1 (5g ST + 1.25kg SA) + Neem cake + FYM, T5 - BsOP2 (5g ST + 1.25kg SA) + TL1 (5g ST + 1.25kg SA) + Neem cake + FYM, T6 – PfUl(A) (5g ST + 0.85kg SA) + BsOP2 (5g ST + 0.85kg SA) + TL1 (2g ST + 0.85kg SA) + Neem cake + FYM, T7 -Carbendazim (2g ST + 0.1% Soil drenching), T8 – control. All the treatments were given as seed treatment and soil application. Seeds were soaked in double the volume of sterile distilled water containing the talc-based formulation (10g kg⁻¹ of seed) (Vidhyasekaran et al.,

- 127 1997). In the field, the biocontrol consortia applied 2.5kg/ha along with 2 tons of FYM and 128 150 kg neemcake. Seed treatment 2g/kg of seed and soil drenching 0.1 per cent 129 carbendazim was used as the chemical check for comparison. The observations were 130 recorded on dry root rot incidence, plant height, number of pods per plant and number of 131 seeds per plant.
- The data were statistically analyzed using the IRRISTAT version92 developed by the International Rice Research Institute Biometrics unit, the Philippines (Gomez and Gomez, 1984).

Results and Discussion

In vitro screening of biocontrol agents

A total of 10 *Trichoderma*, 20 *Pseudomonas* sp and 10 *Bacillus sp*, isolates were screened for their antagonistic activities against mycelial growth of *R. bataticola*. All the ten *Trichoderma* isolates are inhibited the mycelial growth of *R. bataticola*. Among them, TL1 recorded the least mycelial growth (4 cm) with 55.6 % inhibition over control. This was followed by TCBE3 and TOKK1 isolates with 4.4 cm and 4.6 cm mycelial growth and 51.10 and 48.9 % inhibition over control, respectively (Table 1). Out of twenty *P. fluorescens* isolates tested, PfUL(A) recorded maximum inhibition of 4.7 cm mycelial growth (41.1%). This was followed by isolates *viz.*, PfAL1 and PfCBE9 isolates recorded 41.10% reduction over control (Table 2). In the case of *Bacillus* isolate (BSOP2) recorded maximum inhibition of 5 cm mycelia growth (44.4%). This was followed by isolates *viz.*, BCBE1 andBKK3 which recorded 33.33% and 31.10% reduction, respectively (Table 3). Our results are in conformity with the findings of many workers (Wong and Baker, 1984; Cook, 1993; Rajkumar *et al.*, 2005).

Plant growth promotion

The biocontrol strains of *Pseudomonas* (PfUL(A)), *Bacillus* (BsOP2) and *Trichoderma* (TL1) produced black gram seedlings with a significantly higher vigour index, 3943.3, 3825.6 and 3706.1, respectively, than the control. Interestingly, *Pseudomonas* (PfUL(A)), *Bacillus* (BsOP2) and *Trichoderma* (TL1) also produced higher germination percentage 98.6%, 98.6%98.6% and seedling length 41.2cm, 39.5cm, 40cm, respectively. The untreated control seedlings had the lowest vigour index (2409.4) (Table 4). Bharathi *et al.* (2004) evaluated the efficacy of 13 plant growth promoting rhizobacterial strains against chilli fruit rot and dieback incited by *Colletotrichum capsici*. The results reveal

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in this study corroborate earlier studies and indicate a future possibility that plant growth promoting rhizobacteria bioformulations can be used to promote growth and health of economic crops (Karthikeyan *et al.*, 2005; Loganathan *et al.*, 2010).

Compatibility of biocontrol agents

Strains of *Pseudomonas* (PfUL(A)), *Bacillus* (BsOP2) and *Trichoderma* (TL1) were tested in vitro for compatibility. Strains that overgrew each other were compatible with each other, whereas strains that were separated by an inhibition zone were incompatible. No inhibition zone formed between PfUL(A) + TL1, PfUL(A) + BsOP2 indicating that these strains were compatible. Inhibition in growth was found between BsOP2+TL1 indicating that these strains were incompatible. Several authors have suggested that combinations of introduced biocontrol agents have to be compatible with each other for better and more consistent disease suppression (Raaijmakerset al., 1995). Several authors have suggested that combinations of introduced biocontrol agents have to be compatible with each other for better and more consistent disease suppression (Raaijmakers *et* al., 1995). In the current study, the isolates of T. viride (TVL1), P. fluorescens (PfUL(A)) and B. subtilis (BSOP2), organic amendments, such as neem cake and FYM showed greater antagonistic activity against M. phaseolina in vitro. The results are consistent with the findings of several research workers who demonstrated the use of antagonistic microorganisms (T. viride, P. fluorescens and B. subtilis), organic amendments against various soil borne fungal pathogens (Thilgavathi et al., 2007; Karthiba et al., 2011).

Glasshouse study

Those treatments that had been most effective in inhibiting the mycelial growth of *M. phaseolina* were selected for pot culture studies. Of these treatments, a combination of fungal and bacterial strains reduced the incidence of root rot more strongly than did the individual strains. The result from the pot culture experiment revealed that among different treatments received individual and combinations of biocontrol agents along with FYM+Neem cake, the treatment combination of PfUL(A)+ BSOP2 performed better in 257 reducing root rot incidence of blackgram. The recorded disease incidence was 20 per cent, this was followed by 26.7% incidence was observed

191 in the combination of PfUL(A)+TVL1. The untreated check was recorded 66.75 per 192 cent incidence. The highest germination was also recorded in seeds treated 193 with the mixture of PfUL(A)+ BSOP2 + neem cake + FYM (95%). The treatment differed significantly from all other treatments, as well as from the 194 195 untreated control which was only 71.7 per cent (Table 5). Several authors have 196 suggested that combinations of introduced biocontrol agents have to be 197 compatible with each other for better and more consistent disease suppression (Raaijmakers et al., 1995). Similarly, the incorporation of biocontrol agents 198 199 with organic amendments and the efficacy of neem and FYM in fungal disease 200 management have been reported by many workers (Sundaravadana, 2002; Gopal, 201 Thilgavathi *et al.*, 2007). An important prerequisite for the effectiveness of strains appears to be the compatibility of the co312 202 inoculated microorganisms (Raaijmakers et al., 1995; Georgakopoulos et al., 203 2002). In the present study, the isolates of T. viride (TVL1), P. fluorescens 204 (PfUL(A)) and B. subtilis (BSOP2) were compatible with each other and with neem 205 206 cake.

Induction of defence related enzymes

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The peroxide activity increased significantly up to seven days in all the treatments (treated with biocontrol agents, organic amendments and inoculated with the pathogen) and thereafter it declined. Among the various bioformulations, application of treatment of PfUl(A)+BsOP2+ Neem cake + FYM followed by challenge inoculation and showed higher induction of peroxidase (1.678 changes in absorbance min⁻¹ g⁻¹ of fresh tissue). The induction reached a maximum level on 7 days after challenge inoculation. The activity of the enzyme thereafter declined with a decreasing rate than the inoculated control. Plants treated with bioformulation PfUl(A)+BsOP2+ Neem cake + FYM also recorded a higher level of PO activity throughout study period than the other treatments. The inoculated control showed reduction of PO activity starting from 7th day and then decreased to lower level than uninoculated control (Fig1). The same trend was observed in the PPO, PAL and Catalase (Fig 2, 3 and 4). These three enzyme activity was increased significantly up to seventh day in all the treatments and thereafter declined. Combined application of PfUl(A)+BsOP2+ Neem cake + FYM and challenge inoculated with the pathogen recorded higher PPO (1.38 changes

in absorbance min⁻¹ g⁻¹ of fresh tissue), PAL (1.235 changes in absorbance min⁻¹ g⁻¹ of fresh tissue) and Catalase (0.912 changes in absorbance min⁻¹ g⁻¹ of fresh tissue) activity respectively than individual applications. The next highest activity was observed in the plants treated with the combinations of BsOP2+ TL1+Neem cake + FYM. Peroxidase considered as an important PR proteins (Van Loon, 1997) and a key enzyme in the biosynthesis of lignin and other oxidative phenols. Increase in peroxidase expression in combined biocontrol agent treated test plants was significant, compared to untreated (absolute) and negative control (pathogen infested) plants. Some workers have reported the role of peroxidase in cell wall-building processes by oxidation of hydroxyl cinnamyl alcohols into free radical intermediates, phenol oxidation, polysaccharide cross linking, cross linking of extension monomers, lignification and suberization. These defense related genes are sleeping genes and it is needed to activate them by appropriate stimuli. P. fluorescens has been used in induced systemic resistance by some earlier workers (Van Loon et al., 1998). Zdor and Anderson (1992) noticed that rhizosphere colonization of various bacteria induced PO activity in bean. The higher PO activity was observed in cucumber roots treated with Pseudomonas corrugate challenged with Pseudomonas aphanidermatum (Chen et al., 2000) and seedlings treated with Pseudomonas spp. challenged with similar pathogen in chilli (Kavitha et al., 2005).

Field study

The greatest reduction in dry root rot incidence was observed in plots treated with the mixture of PfUL(A)+ BsOP2 + neem cake + FYM (25.65 PDI) followed by PfUL(A)+ TL1 + neem cake + FYM (29.02PDI) as compared with the untreated control (55.85PDI). The biocontrol agents not only reduced disease incidence and also enhances the plant growth. Mixture application of PfUL(A)+ BsOP2 + FYM + neem cake recorded the maximum plant height of 103 cm with yield (785kg/ha) and compared to control (64cm, 580kg/ha). This was followed by PfUL(A) + TL1 + FYM + neem cake recorded 98 cm plant height and 710 kg/ha yield (Table 6).Soil application of biocontrol agents *viz.*, *T. viride*, *T. harzianum*, *P. fluorescens and B. subtilis* effectively reduced root rot caused by soil borne pathogens in several crops (Saravanakumar *et al.*, 2007; Thilgavathi *et al.*, 2007; Loganathan *et al.*, 2010).The *P.fluorescens* strains reduced the root rot infection through several mechanisms including production of lytic enzymes (Velazhahan *et al.*, 1999), siderophores (Scher and Baker, 1982), salicylic acid (Klessig and Malamy, 1994) and hydrogen cyanide (Bakker and Schippers, 1987). *B. subtilis* strains known to inhibit several soil borne diseases such as *Fusarium* wilt of red gram (Podile and Dube, 1985) and *R. solani* (damping off of

- peppermint) (Kamalakannan et al., 2003). Organic amendments are recommended as
- biological means to reduce the incidence of several soil borne diseases. Roy (1989) reported
- 257 that the activity of R. solani in organic amended soil was temporarily checked which was due
- 258 to increase in CO₂ and decrease in N content of soil. Soil amendment with FYM led to
- 259 increased disease control efficacy of fungal antagonist *Trichoderma* spp. against *Fusarium*
- wilt of cumin (Gopal, 2002). Seed treatment with *P. fluorescens* along with soil amendment
- 261 like mustard cake, vermicompost and FYM provided a better protection against
- 262 *Macrophomina* root rot of chickpea (Khan and Gangopadhyay, 2008).

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Table 1. Efficacy of different *Trichoderma* isolates against the growth of *M. phaseolina in vitro*.

Trichoderma isolates	Growth of <i>M. phaseolina</i> Per cent inhibition over	
		control
TCBE3	4.4 ^b	51.1 ^b (45.6)
TTV1	8.2 ^t	8.9 ^e (16.9)
TMDU1	5.5°	38.9 ^{bc} (38.6)
TSOP4	5.8 ^{cd}	35.6° (36.6)
TVL1	4.0 ^a	55.6 ^a (48.2)
TVL2	6.8 ^e	24.4 ^d (29.6)
TOR1	6.2 ^d	31.3 ^{cd} (33.9)
TKM2	5.4°	40.0 ^{bc} (39.2)
TOKK1	4.6 ^b	48.9 ^{ab} (44.4)
TTNL1	6.3 ^{de}	30.0 ^{cd} (33.2)
Control	9.0 ^g	0.00

Values are mean of three replications. In a column, means followed by a common letter(s) are not significantly different (P = 0.05) by DMRT.

Table 2. Efficacy of different *Pseudomonas* isolates against the growth of *M. phaseolina in vitro*.

Pseudomonas isolates	Growth of <i>M. phaseolina</i>	Per cent inhibition over		
		control		
PfCBE3	8.5 ^{gh}	5.6 ^h (12.4)		
PfPPN3	8.1 ^{tg}	10.0 ^{tgh} (18.1)		
PfPPN4	6.8 ^d	24.4 ^{cde} (29.5)		
PfUL(A)	4.7 ^a	44.4 ^a (41.8)		
PfKK1	7.7 ^{ef}	14.4 ^{efg} (22.1)		
PfKK2	5.6 ^{bc}	37.8 ^{\]7b} (37.9)		
PfAL1	5.3 ^b	41.1 ^b (39.9)		
PfOKK2	7.4 ^e	17.8 ^{def} (24.8)		
PfCBE2	8.5 ^{gh}	5.6 ^h (12.4)		
PfCBE9	5.3 ^b	41.1 ^b (39.9)		
PfV1	6.6 ^d	26.7 ^{bcd} (31.1)		
PfKB1	8.6 ^{gh}	4.8 ^h (11.4)		
PfKB3	8.3 ^g	7.8g ^h (15.6)		
PfVT1	7.4 ^e	17.8 ^{def} (24.8)		
PfSOP2	6.8 ^d	24.4 ^{cde} (29.5)		
PfTVL1	$6.0^{\rm c}$	33.3 ^{bc} (35.2)		
PfTVL4	5.6 ^{bc}	37.8 ^b (37.9)		
PfMDU1	8.1 ^{fg}	10.0 ^{fgh} (18.1)		
PfMDU2	7.5 ^e	16.7 ^{def} (29.0)		
PfPPOZ1	8.2 ^{fg}	8.9 ^{fgh} (16.9)		
Control	9.0 ^h	0.00		

Values are mean of three replications. In a column, means followed by a common letter(s) are not significantly different (P = 0.05) by DMRT.

Table 3. Efficacy of different *Bacillus* isolates against the growth of *M. phaseolina in vitro*.

Bacillus isolates	Growth of M. phaseolina	Per cent inhibition over		
		control		
BCBE1	$6.0^{\rm b}$	33.33 ^{ab} (35.2)		
BCBE2	8.2 ^{de}	8.9 ^d (16.9)		
BPPN5	7.3°	18.9° (25.6)		
BKK3	6.2 ^b	31.1 ^{ab} (33.9)		
BOKK3	8.3 ^e	7.8 ^d (15.6)		
BV2	7.4 ^c	24.4 ^{bc} (29.6)		
BKB3	8.2 ^{de}	8.9 ^d (16.9)		
BSOP2	5.0^{a}	36.7 ^a (37.2)		
BV3	7.6°	15.6 ^{cd} (23.1)		
BMDU2	7.7 ^{cd}	14.4 ^{cd} (22.1)		
Control	9.0 ^f	0.00		

Values are mean of three replications. In a column, means followed by a common letter(s) are not significantly different (P = 0.05) by DMRT.

Table 4. Growth promotion activities of biocontrol agents on blackgram seedlings

Biocontrol	Seedling length	Germination %	Vigour index
agents			
TL1	40.0 ^{ab}	98.6 ^a (83.28)	39440 ^b
TCBE3	38.6 ^{bcd}	98.6 ^a (83.28)	38260 ^d
TOKK1	38.6 ^{bcd}	94.6 ^d (76.57)	36710 ^g
PfUL(A)	41.2ª	98.6° (83.28)	40820 ^a
PfAL1	38.1 ^{cde}	97.3 ^b (80.57)	37270 ^f
PfCBE9	36.8 ^{ef}	96.0° (78.48)	35520 ⁱ
BSOP2	39.5 ^{bc}	98.6 ^a (83.28)	39140 ^c
BCBE1	38.2 ^{cde}	97.3 ^b (80.57)	37360 ^e
BKK3	36.2 ^f	97.3 ^b (83.28)	35420 ^j
Carbendazim (2g/kg)	37.5 ^{def}	97.3 ^b (80.57)	36680 ^h
Control	34.6 ^g	93.3 ^d (75.0)	32420 ^k

Values are mean of three replications. In a column, means followed by a common letter(s) are not significantly different (P = 0.05) by DMRT. Values in parentheses are arcsine transformed.

Table 5. Effect of biocontrol consortia and organic amendments on germination and dry root rot incidence in blackgram

Treatments	Germination %	% increase over control	Per cent disease incidence	% decrease over control
PfUL(A)	83.3 ^d (65.9)	16.3	33.3 ^{cd} (35.2)	50
BSOP2	86.7° (68.6)	20.9	30.0 ^{bc} (33.2)	55
TL1	81.7 ^e (64.7)	14.0	36.7 ^d (37.3)	45
PfUL(A)+ BSOP2	95.0 ^a (77.1)	32.6	20.0° (26.5)	70
PfUL(A) + TL 1	91.7 ^b (73.3)	28.0	26.7 ^b (31.0)	60
PfUL(A) + BSOP2 + TL1	83.3 ^d (65.9)	16.3	43.3 ^e (41.1)	35
Carbendazim (0.1%)	81.7 ^e (64.7)	14.0	20.0° (26.5)	70
Control	71.7 ^f (57.9)	-	66.7 ^g (54.8)	-

Values are mean of three replications. In a column, means followed by a common letter(s) are not significantly different (P = 0.05) by DMRT. Values in parentheses are arcsine transformed.

Table 6. Effect of biocontrol consortia and organic amendments on the incidence of dry root rot growth and yield parameters in blackgram

Treatment details	Shoot	Root	Plant	No. of	No. of	Percent	Yield
	Length	Length	Height	pods/	Seeds/pod	disease	Kg/ha
	(Cm)	(Cm)	(Cm)	plant		incidence	
(PfUl(A)) + Neem	61 ^c	35 ^b	96°	100 ^d	8 ^a	31.52 ^d	650 ^e
cake + Farm Yard							
Manure							
(BsOP2) + Neem	62 ^b	35 ^b	96 ^c	107 ^b	8 ^a	30.40^{c}	700°
Cake + FYM							
(TL1) + Neem	58 ^d	32 ^c	90 ^d	75 ^e	7 ^b	32.46 ^e	605 ^f
cake + FYM							
PfUl(A)+BsOP2+	66 ^a	37 ^a	103 ^a	110 ^a	8 ^a	25.65 ^a	785 ^a
Neem cake + FYM							
PfUl(A)+	65 ^a	36 ^a	98 ^b	106 ^b	7 ^b	29.02 ^b	710 ^b
TL1+Neem cake +							
FYM							
PfUl(A) + BsOP2	63 ^b	35 ^b	98 ^b	104 ^c	7 ^b	31.24 ^d	680 ^d
+TL1+Neem							
cake+ FYM							
Carbendazim	53 ^e	25 ^d	78 ^e	65 ^t	8 ^a	24.85 ^a	700 ^b
Control	45 ^f	19 ^e	64 ^f	35 ^g	6°	55.85 ^f	580 ^g

Values are mean of three replications. In a column, means followed by a common letter(s) are not significantly different (P = 0.05) by DMRT

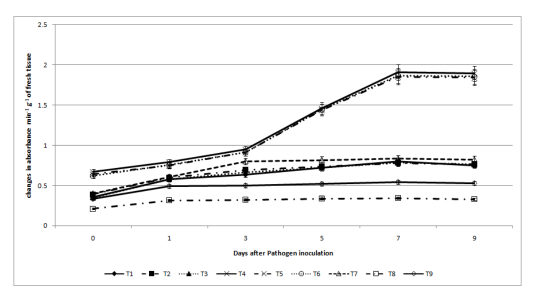


Fig 1. Induction of peroxidise activity in blackgram plants treated with biocontrol consortia and organic amendments. T1–TL1, T2–PfUL(A), T3–BSOP2, T4–PfUL(A) + TL1, T5–PfUL(A)+BSOP2, T6–PfUL(A)+BSOP2+TL1, T7–Carbendazim (0.1%), T8–Inoculated control, T9–Healthy control

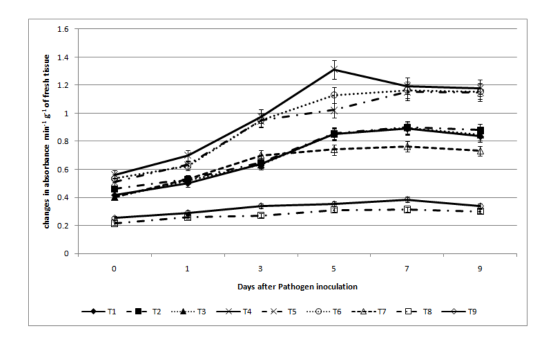


Fig 2. Induction of polyphenol oxidase activity in blackgram plants treated with biocontrol consortia and organic amendments. T1–TL1, T2–PfUL(A), T3–BSOP2, T4–PfUL(A) + TL1, T5–PfUL(A)+BSOP2, T6–PfUL(A)+BSOP2+TL1, T7–Carbendazim (0.1%), T8–Inoculated control, T9–Healthy control

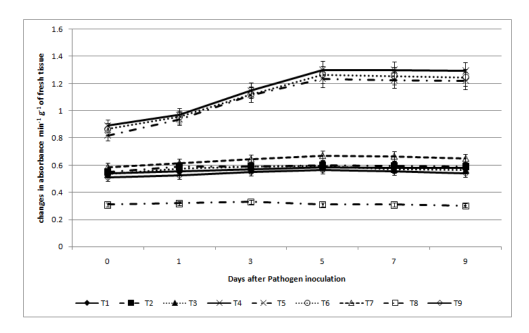


Fig. 3. Induction of phenylalanine ammonia lyase activity in blackgram plants treated with biocontrol consortia and organic amendments.T1–TL1, T2–PfUL(A), T3–BSOP2, T4–PfUL(A) + TL1, T5–PfUL(A)+ BSOP2, T6–PfUL(A)+BSOP2+TL1, T7–Carbendazim (0.1%), T8–Inoculated control, T9–Healthy control

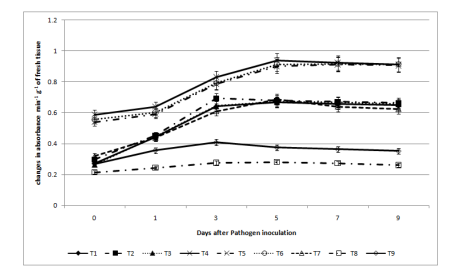


Fig. 4. Induction of catalase activity in blackgram plants treated with biocontrol consortia and organic amendments. T1–TL1, T2–PfUL(A), T3–BSOP2, T4–PfUL(A) + TL1, T5–PfUL(A)+BSOP2, T6–PfUL(A)+BSOP2+TL1, T7–Carbendazim (0.1%), T8–Inoculated control, T9–Healthy control