

International Journal of Environment and Climate Change

12(11): 2662-2673, 2022; Article no.IJECC.92279 ISSN: 2581-8627 (Past name: British Journal of Environment & Climate Change, Past ISSN: 2231–4784)

Management of Postharvest Anthracnose Disease of Papaya (*Carica papaya*) Using Antagonistic Microorganisms

Reshma N. Yadravi^a, D. L. Rudresh^{b*#}, S. L. Jagadeesh^a, D. S. Ambika^c and Parvati Pujer^d

 ^a Department of Post-Harvest Technology, College of Horticulture, Bagalkot, University of Horticultural Sciences, Bagalkot-587104, Karnataka, India.
^b Department of Agricultural Microbiology, College of Horticulture, Bagalkot, University of Horticultural Sciences, Bagalkot-587104, Karnataka, India.
^c Department of Plant Pathology, College of Horticulture, Bagalkot, University of Horticultural Sciences, Bagalkot-587104, Karnataka, India.
^d Department of Biotechnology and Crop Improvement, College of Horticulture, Bagalkot, Viversity of Horticultural Sciences, Bagalkot-587104, Karnataka, India.

Authors' contributions

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

Article Information

DOI: 10.9734/IJECC/2022/v12i1131269

Open Peer Review History:

This journal follows the Advanced Open Peer Review policy. Identity of the Reviewers, Editor(s) and additional Reviewers, peer review comments, different versions of the manuscript, comments of the editors, etc are available here: https://www.sdiarticle5.com/review-history/92279

Original Research Article

Received 10 July 2022 Accepted 19 September 2022 Published 21 September 2022

ABSTRACT

The antagonistic potential of microbial agents isolated from different sources like compost, rhizosphere soil, healthy fruits surface were investigated against *Colletotrichum gloeosporiodes under In vitro and In vivo*. The results of *In vitro* evaluation of 22 isolates of antagonists against *Colletotrichum gloeosporiodes* by dual-culture test showed *Trichoderma harzianum* UHSF-15 (with 62.50% inhibition of radial growth), *Bacillus amyloliquifaciens* UHSB-BCA-15 isolated from banana rhizosphere soil (50.00% inhibition) and *Bacillus amyloliquifaciens*UHSB-BCA-16 isolated from kiwi fruit surface (47.85% inhibition) to be most effective. *In vivo* screening of five effective microbial agentsselected from *In vitro* studies revealed that, *Bacillus amyloliquifaciens* UHSB-BCA-15 be most effective with lowest per cent disease index of 13.89% followed by *Bacillus*

#Assistant Professor;

^{*}Corresponding author: E-mail: rudreshdl@hotmail.com;

*amyloliquifaciens*UHSB-BCA-16 with 16.67% and carbendazium at 0.1% with 20.83% at 9d after storage. The higher per cent disease index of 58.33% was recorded in untreated control fruits. In another experiment where *C. gloeosporoidess*pore suspension $(1 \times 10^{6} \text{ CFU mL}^{-1})$ was challenge inoculated on fruits, the fruits treated with biocontrol agent *Bacillus amyloliquifaciens* UHSB-BCA-15 at concentrations of $1 \times 10^{8} \text{ CFU mL}^{-1}$ for 5 minutes was found to be effective in reducing postharvest infection to a greater level with 33.33% disease index followed by *Bacillus amyloliquifaciens* UHSB-BCA-16 with 50.00% disease index and carbendazium at 0.1% with 52.77% disease index compare to control (91.66% disease index) at 9d after storage.

Keywords: Bacillus amyloliquifaciens; microbial agents; microorganisms; anthracnose disease.

1. INTRODUCTION

Papaya is an important climacteric fruit crop of the world, which ishighly nutritious with good medicinal value. The rapid ripening and softening of the fruit makes it highly susceptible to many postharvest diseasesduring transit, storage and market. Among the postharvest diseases, the fungal rots cause losses up to 40% of their market value [1]. Among the fungal diseases, anthracnose caused bv Colletotrichum *gleosporoides* alone accounts for about 10.05% of total postharvest loss [2]. The infections are usually initiated in the field at early stages of fruit development on the surface of the fruit but the pathogen remains quiescent until the fruit reaches climacteric phase [3]. The initial symptoms of infection are water soaked, sunken spots at center of these spots, later turning black and then pink when the fungus produces spores [4]. Currently the control of anthracnose, a major postharvest disease of papaya relies mainly on the use of synthetic fungicides [5,6], but the use of chemicals may pose several adverse effect on consumers as the fruit is consumed within short period of time and may leave residue on fruits. Non chemical control methodslike physical and biological methods are of great importance. The physical treatment like hot water treatment may leads to fruit injury. Hence, control of the disease through use of microbial agents holds a great potential [7,8].

The biological control is a biological-based, non-toxic and environmentally safe approach for post-harvest disease control during the storage of harvested commodities [9,10]. Biological control includes the use of wide range of and antagonistic bacteria, fungi other microorganisms on the surface of the fruits and vegetables and they have shown to protect fruits and vegetables against growth and postharvest infection of pathogens (Cota et al., 2008). Antagonists like Bacillus subtilis, B. cereus, Pseudomonas aeruginosa, P. *cepacia, P. fluorescens, P. putidaa, Streptomyces* spp., *Trichoderma harzianum*are being used as potential biocontrol agents against the post-harvest pathogens in many fruit crops [5,11].

In the present investigation the antagonistic microorganisms were isolated from different sources like compost, rhizosphere soil and different fruits surfaces collected from different places and were evaluated for their efficacy under in vitroconditionby dual culture test. Five effective antagonistic organisms from in vitro evaluation were evaluated in vivo by treating the papaya fruits with suspensions of microbial agents against natural anthracnose inoculation infection and challenge with Colletotrichum gloeosporiodes spores. The fruits were observed for the per cent disease index.

2. MATERIALS AND METHODS

2.1 Isolation and Identification of Pathogen Colletotrichum gloeosporiodes

Six pathogen (Colletotrichum gloeosporioides) isolates wereisolated from anthracnose infected papaya (Carica papaya) fruitscollected from different places by standard serial dilution plating technique using potato dextrose agar (PDA) media(Boiled Potato extract from 200 g potato per liter; dextrose, 20 gL⁻¹; agar 20 gL⁻¹), supplemented with chloramphenicol (100 mg L ¹). The C. gloeosporioides spores and mycelia from the infected portion of the fruits were scraped using sterile scalpel and the pathogen sample was transferred to 9 ml of sterile water blank and serially diluted up to 10⁵ dilution in a Laminar Air Flow chamber and subsequently 1 ml aliquot from 10^3 , 10^4 and 10^5 dilution were transferred on to sterile Petri dishesand2 0 to 25 ml of molten PDA media was poured and rotated in both clock-wise and anti-clockwise direction. The plates were then allowed to solidify for 30 minutes and incubated at 27 \pm 1°C for 5 d and observed periodically. After development of fungal colonies, pure colonies were transferred on to freshly prepared PDA plates. The pure culture of fungus was sub-cultured on PDA slants and preserved in refrigerator at 5°C until used. The pathogenisolated from infected papaya fruits was identified using morphological features and bv microscopic observations [12] as Colletotrichum gleosporioides.The pathogen isolates were tested for their pathogenicity on healthy papaya fruits and the isolate with higher virulence index was selected for isolation of antagonistic microbial agents against the pathogen.

2.2 Isolation of Biocontrol Agents against *C. gloeosporiodes*

2.2.1 Isolation of antagonistic microbes from compost and rhizosphere soil samples by employing poisoned food technique

For isolation of microbial agents, fivecompost samples and 35 rhizosphere soil samples from rhizosphere of healthy papaya plants in anthracnose affected field as well as from other fruit crops like. Mango, Banana and Pomegranate were collected. Ten grams of compost or rhizosphere soil was suspended in 100 ml of sterile 0.85% NaCl solution and kept in rotary shaker at 150 rpm for 30 min to dispense the spore chain or cells into water. The suspension was allowed to settle for 10 minutes and subsequently serially diluted up to 10⁻⁴. From the serial dilutions $(10^{-2}, 10^{-3} \text{ and } 10^{-4} \text{ dilutions})$ one ml of aliquot was transferred to sterile Petri dishes and approximately 20 ml of cooled molten potato dextrose agar pre-seeded with spore suspension of pathogen (*C. gloeosporioides*at the rate of one ml of 10^9 CFU mL⁻¹spore suspension per 50 ml of PDA media) was poured to these plates and allowed to solidify for one hour in Laminar Air Flow Chamber and later the plates were incubated at 27 ± 1°C for 5 d in BOD incubator (Make: Remi). After incubation period the plates were observed for fungal and bacterial colonies with a clear zone of inhibition around them. The colonies with inhibition zone were pure cultured on new media plates by quadrangle streaking method on PDA and Nutrient Agar media (NA) for fungal and bacterial pure respectively. The isolates cultured antagonist isolates were also maintained in PDA

and NA slants for further screening against *C. gloeosporioides*.

2.2.2 Isolation of antagonistic microbes from healthy fruit surfaceby employing poisoned food technique

A thin layer of matured healthy fruit skin was scraped using sterile peeler and subsequently transferred to 100 ml water blanks. The blanks with fruit scrapes were agitated at 150 rpm for 30 min and subsequently subjected for serial dilution up to 10⁻⁴. Same procedure was followed for 19 different fruits Papaya, like Banana, Pomegranate, Apple, Ber, Kiwi fruit and Citrus fruits like Lime, Sweet Orange and Mandarine collected from the farmers field as well as from the local fruit markets. The dilutions 10^{-2} , 10^{-3} and 10⁻⁴ were used for isolation of antagonistic microbial agents by employing poisoned food technique (2.2.1). The colonies with inhibition zone were pure cultured on Petri plate by multiple streaking methods on PDA, NA and Yeast Peptone Dextrose Agar (YPDA) media for fungal, bacterial and yeast isolates respectively. cultures of antagonists The pure were maintained in PDA, NA and YPDA slants for further screening against C. gloeosporioides.

2.2.3 Collection of biocontrol agents

The efficient and proven biocontrol agents against fungal plant pathogens *viz., Trichoderma harzianum*UHSF-15*, Bacillus subtilis* UHSB-01 and *Pseudomonas fluorescens* UHSB-6 were collected from Biofertilizer Laboratory, Department of Agricultural Microbiology, College of Horticulture, Bagalkot for screening against *Colletotrichum gloeosporiodes.*

2.3 *In vitro* Screening of Antagonistic Microbial Isolates against *C. gloeosporiodes*

The microbial agents showing inhibition zones against pathogen in poisoned food technique were screened for their efficacy by dual culture method [13] on PDA media. For dual culture test, approximately 25 ml of molten PDA was poured into Petri plates and allowed for solidification. Five millimetre diameter disc of pathogen was cut with the help of sterile cork borer and placed near the periphery of PDA plate. Similarly antagonistic fungal isolate was placed on the other periphery at an angle of 180°, in case of bacterial and yeast antagonists, the antagonists

were streaked at the centre of the plates. The plates inoculated with only pathogen was served as control. The plates were incubated at $27 \pm 1^{\circ}$ C for seven days. Each treatment was replicated thrice. The extent of antagonistic activity by the antagonists was recorded on seventh day by measuring growth of the pathogen in dual culture plate and control plate. The per cent inhibition of pathogen (I) was calculated by using the formula suggested by Vincent [14].

I = [100 X (Growth of the pathogen in control plate in millimetre - Growth of the pathogen in dual culture plate in millimetre)]/Growth of the pathogen in control plate in millimetre.

2.4 *In vivo* Screening of Microbial Agents against *C. gloeosporiodes*

2.4.1 Selection of antagonistic microbial agents for *In vivo* studies

The five effective isolates of microbial agents from *in vitro* studies were selected and evaluated under *in vivo* for their efficacy against *C. gloeosporioides*against natural infection on fruits and against challenge inoculation with *C. gloeosporioides*.

2.4.2 Preparation of liquid culture of microbial agents

The overnight grown cultures of bacterial antagonists, which were grown on NBand the fungal and yeast isolates grown on PDBfor 5 d were used to prepare the aqueous suspension of microbial agents. The cultures of bacterial isolates were prepared by inoculating 2 loops of each culture to 250 ml of sterile nutrient broth and subsequent incubation on a rotary shaker at 150 rpm for 48 h. Similarly Fungal and yeast cultures were grown for 5 d in an BOD incubator at 25 \pm 1°C in Potato Dextrose Broth (PDB) by inoculation of 5 mm disc of fungi and 2 loop full of yeastculture.

2.5 In vivo Screening of Antagonistic Microbial Agents against C. gloeosporiodes

2.5.1 Selection of antagonistic microbial agents for *In vivo* studies

The five effective isolates of microbial agents from *In vitro* studies were selected and evaluated under *In vivo* for their efficacy against *C*.

gloeosporioidesagainst natural infection on fruits and against challenge inoculation with *C. gloeosporioides.*

2.5.2 Preparation of liquid culture of microbial agents

The overnight grown cultures of bacterial antagonists, which were grown on NBand the fungal and yeast isolates grown on PDBfor 5 d were used to prepare the aqueous suspension of microbial agents. The cultures of bacterial isolates were prepared by inoculating 2 loops of each culture to 250 ml of sterile nutrient broth and subsequent incubation on a rotary shaker at 150 rpm for 48 h. Similarly Fungal and yeast cultures were grown for 5 d in an BOD incubator at 25 \pm 1°C in Potato Dextrose Broth (PDB) by inoculation of 5 mm disc of fungi and 2 loop full of yeastculture.

2.6 *In vivo* Screening of Efficient Microbial Agents against Natural Anthracnose Infection of Papaya

The uniform size papaya fruits (Caricapapaya) of Red Lady variety at optimum matured stage were collected from the papaya fruit orchard maintained at College of Horticulture, Bagalkot, Karnataka, India and the fruits were dipped in the aqueous suspension of effective microbial agents (prepared by addition of 10 ml broth cultures of isolates containing 1x10⁸CFUmL⁻¹ in one liter of water) for 5 minutes and allowed to dry for 10-15 minutes. The treated fruits were kept in clean crates in separate sets according to treatments under ambient condition. Each treatment was replicated thrice, and six fruits were maintained per replication with completely randomized design. The standard chemical, carbendazium, treatment at 0.1 per cent for 5 minutes was used as a chemical control and hot water dipping treatment at 52° C for 3 minutes was used as physical treatment. The untreated natural fruits served as untreated control. During the study the fruits were observed for the disease symptoms and the extent of infection was scored using 0-4 scale (0=No infection. 0.1-25.0% fruit surface infected=1, 25.1-50.0% fruit surface infected=2, 50.1-75.0% fruit surface infected=3 and 4=>75.0% fruit surface infected) [2]. Per cent Disease Index (PDI) of postharvest disease of fruits was calculated by the formula given by Wheeler [15]. PDI= (Sum of all disease rating/ Total number of ratings) x (100/ Maximum disease grade).

2.7 Screening of Antagonistic Microbial Agents against Challenge Inoculation of Post Harvest Anthracnose Disease Pathogen (*C. gloeosporioides*) on Papaya Fruits Under *In vivo*

2.7.1 Preparation of *C. gloeosporioides* inoculum

The Spore suspension of *C. gloeosporioides*was prepared by growing *C. gloeosporioides* on potato dextrose broth at $25\pm 1^{\circ}$ C for 7 d.The culture suspension was then filtered through two layers of sterile muslin cloth. The concentration of conidia in the filtered suspension was diluted to 10^5 conidia ml⁻¹ using sterile saline solution (0.9% NaCl) with the help of haemocytometer [16].

The mature papaya fruits brought from the farmers field maintained at Bagalkot, Karnataka were surface sterilized with 0.5 per cent of Sodium hypochlorite for 3 minutes and rinsed with distilled water for three times. Subsequently the fruits were inoculated with conidial suspension of C. gloeosporioides containing 10⁵CFU mL⁻¹for 5 minutes and later taken out and placed in a clean dry tray and allowed to dry. After three hours of pathogen inoculation, the fruits were treated by spraying with aqueous suspension of efficient microbial agents (10⁸CFU mL⁻¹) at the rate of 10 ml per liter of water and allowed to dry for 10-15 minutes. The treated fruits were kept in clean crates in separate sets according to treatments and incubated in clean area in Laboratory for observation. Each treatment was replicated thrice and six fruits were kept per replication. During the study the fruits were observed for the disease symptoms and the extent of infection was scored using 0-4 scale [2] as mentioned above (2.4.3). Per cent disease index (PDI) was calculated using formula mentioned in previous section (2.4.3) and per cent disease decrease over control were calculated with the formula, Percent disease decrease over control = (Per cent infection in control - Per cent infection in treatment)/ Per cent infection in control.

2.7.2 Identification of efficient antagonistic microbial agents

The most efficient biocontrol isolates identified in this study were identified by 16S rRNA sequencing [17].

2.8 Statistical Analysis

The data obtained in this experiment were subjected to statistical analysis by ANOVA for completely randomized design (CRD analysis). Statistical analysis was performed using Web Agri Stat Package (WASP) Version 2 [18]. The level of significance used in F and t test was p=0.01. Critical difference and SE.m \pm values were calculated whenever F- test was found significant.

3. RESULTS AND DISCUSSION

3.1 Identification of Colletotrichum gloeosporoides

The ten days old cultures of Colletotrichum gloeosporoidesproduced white mycelium on potato dextrose agar (PDA) media which later turned into grey, dark grey, black with regular margins. The mycelium of pathogen was hyaline, septate and branched. Sporulation was abundant with maximum fruiting bodies at center of the plate as against mycelium growth towards On reverse side of culture periphery. plates, pathogen colony depicted gravish to dark grey concentric rings with sector formation. The size of the conidia varied from 9-20 X 3-7.5 µm when observed under the microscope. Similarly, conidia shape was found oblong or cylindrical shape with rounded ends having hyaline, aseptate with one to two oil globules. The similar morphological and microscopic characters of C. gloeosporoides were observed by Archana [19] and Jayalaxmi [20].

3.2 Isolation and *In vitro* Evaluation of Microbial Agents against *Colletotrichum gloeosporioides*

A total of 19 microbial agentsshowing clear zone of inhibition (Fig. 1) against *C. gloeosporiodes*in poisoned food technique were isolated from 59 samples used. Among the 22 microbial agentsevaluated against *C. gloeosporoides* under *In vitro*, *Trichoderma harzianum* UHSF-15 was found to be the most effective with 62.50% inhibition of radial growth followed by isolateno. 10 with 50%, which was on par with isolateno.9 (47.85%), isolateno. 7 (47.14%) and isolateno.16 (46.43%). The minimum inhibition of radial growth of *C. gloeosporioides*was recorded in isolateno. 4 (12.4%) (Table 1, Fig. 2.).

The higher antagonistic potential of *T.harzianum* against *C. gloeosporioides* was reported by Patel and Joshi [21]; Ashoka et al. [22]; Prabhakar et al. [23]. The antagonistic activity of *Trichoderma*

spp. Against *C. gloeosporioides* might be attributed to competition for space and nutrition, in addition todirect parasitism action by coiling and drawing the nutrition from pathogen mycelia [24] and by production of volatile antibiotics like dermadin and trichodermin against pathogen [25].

Next to T. harzianumUHSF-15 the microbial agentsisolated from different sources like isolate no. 10 (50.00%) (isolated from the soil sample collected from Banana rhizosphere), isolate no. 9 (47.85%) (isolated from kiwi fruit collected from local fruit market), isolate no. 7 (47.14%)(isolated from compost sample) and isolateno. 16 (isolated from (46.95%)Ber fruits surface)werefound to reduce the radial growth of C. gloeosporioidesto a significantly greater level. Similar results were noticed by Shi et al. [26], putida where Pseudomonas isolated from pericarp of papaya fruit was found effective against anthracnose of papaya and reduced incidence up to 54%. The bacterial strain MJM5763, isolated from yam cultivation field has inhibited effectively the growth of C. gloeosporioides (65%), Pestalotiaspp. (57.5%), Fussarium oxysporum (48%) and Ralstonia solani (40%) (Palaniyandi et al., 2011). Senthil et al. [27] reported that Bacillus subtilis strains (EPC-16 and EPC-8), were effective in suppressing mycelia growth of grape pathogens Penicillium expansum and Asperaillus carbonarius under in vitro condition. Similarly, the two Bacillus strains TB09 and TB72 were reported to produce volatile organic compounds that effectively reduced the anthracnose incidence up to 94.28% and 87.06% respectively [28]. Calvo et al. [29] reported antifungal activity of bacteria strain BUZ-14 (Bacillus amyloliquefaciens) against many post-harvest pathogens of orange, apple, grape and stone fruits like Botrytis cinerea, Moniliniafruticola, Monilinialaxa, Pencillimdigitatum, Pencilliumexpansum and P. italicum, which produces many bioactive compounds viz, iturin, fengycin, bacilysin, bacillomycin and difficidin for their antifungal activity.

3.3 *In vivo* Evaluation of Microbial Agents against *Colletotrichum gloeosporioides*

3.3.1 Evaluation of microbial agents against natural incidence of anthracnose on papaya fruits under *In vivo*

The lowest per cent disease index was recorded in fruits treated with T_5 (BCA₅ -isolate no. 10)

with 13.89% followed by T_4 (BCA₄-isolate no. 9) with 16.67% and T_7 (Chemical treatment with carbendazium at 0.1%) with 20.83% during storage of papaya fruits (Table 2; Fig. 3). The per cent disease index in untreated control fruits was 58.33%. The decreased disease incidence in microbial agents treated treatments might be attributed to the antagonistic mechanism exhibited by the isolates of microbial agents as described by Sharma et al. [30] viz., competition for the nutrients and space, direct parasitism, production of antibiotics or production of volatile organic compounds against pathogen.

3.3.2 Evaluation of microbial agents against challenge inoculation of postharvest anthracnose disease pathogen (*C. gloeosporioides*) on papaya fruits under *In vivo*

Postharvest treatment of anthracnose pathogen (C. gleosporoides) inoculated fruits with biocontrol agentisolate no. 10 for 5 min was most effective in reducing postharvest infection with 33.33% disease index (PDI) followed by isolate no.9 with 50.00% and chemical treatment with carbendazium at 0.1 per cent with 52.77% disease index at 9 d after storage (Table 3: Fig. 4). The higher per cent disease index of 91.66% was recorded in pathogen inoculated control fruits, which might be due to pathogen infection at earlier stage during fruit ripening or at climacteric phase. Reduced phenolic compounds during fruit ripening and susceptibility of fruit skin to chitinase enzyme which was secreted by pathogen [31] might be responsible for the disease development in the papaya fruit during storage.

3.3.3 Identification of efficient microbial agents

The most efficient biocontrol isolates *viz.*, isolate no. 10 and isolate no. 9 observed in the present investigation were identified as *Bacillus amyloliquifaciens* (Both isolates) and were given isolate no. UHSB-BCA-15 and UHSB-BCA-16 respectively.

In the present study the identified isolates were anthracnose effective against pathogen C.gloeosporoides by exhibiting some antagonistic mechanisms. Similary Kim and Chuan (2012)reported that В. amyloliquefaciens MET0908 secreted an extracellular β -1,3-glucanase, against Colletotrichum lagenarium, an anthracnose causing pathogen of watermelon which is a key enzymein the decomposition of fungalhyphal walls. Hu et al. [32] reported that the antagonism of *Bacillus* is mainly due to the production of the antagonistic substances by its secondary metabolism pathways. According to the reports of European Food Safety Authority (2008), some stains of *B. amyloliguefaciens*do not possess the

genes encoding *Bacillus*enterotoxins or the key gene implicated in the synthesis of emetic toxins, or does not demonstrate phenotypic characteristic of toxin production. Hence these isolates can be used effectively to manage the postharvest anthracnose disease during storage with better fruit quality.

Microbial agents	Radial growth of C.	Inhibition over control
	gloeosporoides (cm)	(%)
Isolate no. 1	4.20	40.00 (6.33)
Isolate no. 2	5.55	20.71 (4.53)
Isolate no. 3	5.45	22.14 (4.70)
Isolate no. 4	6.15	12.14 (3.49)
Isolate no. 5	4.00	42.85 (6.55)
Isolate no. 6	5.60	18.90 (4.35)
Isolate no. 7	3.70	47.14 (6.85)
Isolate no. 8	3.95	42.07 (6.48)
Isolate no. 9	3.65	47.85 (6.92)
Isolate no. 10	3.50	50.00 (7.07)
Isolate no. 11	4.00	42.86 (6.49)
Isolate no. 12	4.00	42.86 (6.55)
Isolate no. 13	4.10	41.43 (6.45)
Isolate no. 14	4.50	35.71 (6.00)
Isolate no. 15	5.25	24.95 (5.00)
Isolate no. 16	3.75	46.95 (6.89)
Isolate no. 17	5.20	25.50 (5.05)
Isolate no. 18	4.35	37.50 (6.12)
Isolate no. 19	3.85	44.35 (6.77)
Pseudomonas fluorescensUHSB-06	4.20	39.95 (6.33)
Bacillus subtilis UHSB-01	4.85	30.71 (5.53)
Trichoderma harzianumUHSF-15	2.65	62.50 (7.90)
Mean	4.38	37.23
SE m ±	0.32	0.10
CD (5 %)	1.14	0.37

Table 1.	. In vitro evaluation	of microbial a	igents against C	C. aloeosporoides by	v dual culture test
		•••••••••••••••••••••••••••••••••••••••			

Note: The values in the parenthesis are square root transformed value and the statistical analysis was done for transformed value







Fig. 2. Effective isolates of biocontrol agents inhibiting C. gloeosporoides in dual-culture test

Table 2. Effect of microbial agents treatment on incidence of anthracno	ose disease
(C. gloeosporoides) in papaya at 9 d of storage under ambient cor	ndition

Treatments	Per cent disease index (PDI)	Disease decrease over control (%)
T ₁ - BCA ₁ (isolate no. 22)	25.00 (5.05)	57.14
T ₂ - BCA ₂ (isolate no. 16)	33.33 (5.73)	42.85
T_3 - BCA ₃ (isolate no. 7)	27.78 (5.30)	52.37
T ₄ - BCA ₄ (isolate no. 9)	16.67 (4.09)	71.42
T₅ - BCA ₅ (isolate no. 10)	13.89 (3.79)	76.19
T ₆ - Hot water dip	33.33 (5.73)	42.86
T7- Chemical treatment	20.83 (4.56)	65.14
T ₈ - Untreated control	58.33 (7.64)	-
Mean	28.65	
SEm ±	0.47	
CD (1%)	1.42	

Note: The values in the parenthesis are squareroot transformed values and statistical analysis was carried out for transformed values.

BCA: Biocontrol agent; Chemical: Carbendazium at 0.1 per cent; Hot water dip (52° C for 3 minutes)

Yadravi et al.; IJECC, 12(11): 2662-2673, 2022; Article no.IJECC.92279



Fig. 3. Effect of biocontrol agents treatment on incidence of anthracnose disease (*C. gloeosporoides*) in papaya at 9 d of storage under ambient condition

Table 3. Effect of microbial agents treatment on incidence of anthracnose diseasein papaya fruits challenge inoculated with pathogen (*C. gloeosporoides*) at 9 d after storage under ambient condition

Treatments	Per cent disease index (PDI)	Disease decreased over control (%)
T ₁ - BCA ₁ (isolate no. 22)	58.33 (7.62)	36.36
$\mathbf{T}_2 - BCA_2$ (isolate no. 16)	77.77 (8.80)	15.15
$\mathbf{T}_3 - BCA_3$ (isolate no. 7)	58.33 (7.62)	36.36
$T_4 - BCA_4$ (isolate no. 9)	50.00 (7.05)	45.45
T ₅ - BCA ₅ (isolate no. 10)	33.33 (5.77)	63.64
T ₆ - Hot water dip	63.88 (7.99)	30.31
T ₇ - Chemical treatment	52.77 (7.24)	42.42
T ₈ - Untreated control	91.66 (9.58)	-
Mean	60.76	
SEm ±	0.29	
CD (1%)	0.87	

Note: The values in the parenthesis are squareroot transformed values and statistical analysis was carried out for transformed values.

BCA: Biocontrol agent; Chemical: Carbendazium at 0.1 per cent; Hot water dip (52° C for 3 minutes)



Fig. 4. Effect of biocontrol agents treatment on incidence of anthracnose disease in pathogen (*C. gloeosporioides*) challenge inoculated papaya at 9 d after storage under ambient condition

4. CONCLUSION

The two isolates of antagonistic microorganisms *viz.*, Isolate No.10 (*Bacillus amyloliquifaciens* isolate no. UHSB-BCA-15) isolated from banana rhizosphere soil and Isolate No. 9 (*Bacillus amyloliquifaciens* Isolate no. UHSB-BCA-16)), isolated from kiwi fruit surface can be effectively used for postharvest treatment of papaya against Anthracnose caused by *Colletotrichum gleosporoides.*

COMPETING INTERESTS

Authors have declared that no competing interests exist.

REFERENCES

1. Abeywickrama K, Wijerathna C, Rajapaksha N, Sarananda K, Kannangara S. Disease control strategies for extending storage life of papaya (*Carica papaya*), cultivars "Red Lady" and "Rathna",. Ceylon J Sci (Biol. Sci.). 2012;41(1):27-34. DOI: 10.4038/cjsbs.v41i1.4535

- 2. Pramod G, Swami AP, Srinivas P. Postharvest diseases of papaya fruit in Coimbatore markets. Ann Plant Prot Sci. 2007;15(1):140-4.
- 3. Ademe A, Ayalew A, Woldetsadik K. Evaluation of antifungal activity of plant extracts against papaya anthracnose (*Colletotrichum gloeosporioides*). J Plant Pathol Microbiol. 2014;4(10):1-4.
- Rana SK. Fungal diseases of papaya. Diseases of fruit crops. Oxford and IBH Publishing Colo. Pvt Ltd New Delhi (India). 2001:81-91.
- 5. He D, Zheng XD, Yin YM, Sun P, Zhang HY. Yeast application for controllingapple postharvest diseases associated with

Penicillium expansum. Bot Bull Acad Sin. 2003;44:211-6.

- Sansone G, Rezza I, Calvente V, Benuzzi D, Tosetti MISd. Control of Botrytis cinerea strains resistant to iprodione in apple with rhodotorulic acid and yeasts. Postharvest Biol Technol. 2005;35(3):245-51. DOI: 10.1016/j.postharvbio.2004.09.005
- 7. AOAC. Official methods of analysis. 16th ed. Arlington, VA: AOAC. 1995;45.1.14.
- Kim PI, Chung KC. Production of an antifungal protein for control of ColletotrichumlagenariumbyBacillus amyloliquefaciens MET0908. FEMS Microbiol Lett. 2004;234(1):177-83. DOI: 10.1016/j.femsle.2004.03.032, PMID 15109737.
- 9. Annonymous. Indian Horticulture Database, National Horticulture Board; 2016.
- Luo S, Wan B, Feng S, Shao Y. Biocontrol of postharvest anhracnose of mango fruit with Deberomycesnepalensis and effects on storage quality and post-harvest physiology. J Food Sci. 2015;80(11): 2555-63.
- Parthasarathy S, Rajalaxmi J, Narayanan P, Arunkumar K, Prabhakar K. Biocontrolpotential of microbial antagonists against postharvest diseases of fruit crops: a review. Bot Sci. 2017;6(1):17-24.
- 12. Bartnett HL, Hunter BB. Illustrated genera of imperfect fungi. In: Burgess Publishing Co, editor, Minneapolis. 3rd ed. 1972; 241.
- Dennis C, Webster J. Antagonistic properties of species groups of Trichoderma II. Production of volatile antibiotics. Vol. 57. Trans: British Mycology Society. 1971;41-8.
- Vincent JM. Distortion of fungal hyphae in presence of certain inhibitors. Nature. 1927;159:845-50.
- 15. Wheeler BEJ. An introduction to plant diseases. New York: John Wiley & Sons Limited; 1969.
- 16. Sariah M. Potential of Bacillus spp. as a biocontrol agent for anthracnose fruit rot of chilli. Malays Appl Biol. 1994;23:53-60.
- 17. de Oliveira EJ, Rabinovitch L, Monnerat RG, Passos LKJ, zahner V. Molecular charecterisation of Brevibacillus laterosporus and its potential use in

biological control agents. Appl Environ Microbiol. 2004;70(11):6657-64.

- DOI: 10.1128/AEM.70.11.6657-6664.2004
- 18. Jangam AK, Thali P. ICAR research complex, goa, Ela, old goa. India; 2010.
- Archana S. Interactive proteomics and molecular characterization of PGPE mediated defense response in mango against colletotrichumgloeosporioidesan incitant of anthracnose disease, Ph. D. Agriculturists [thesis],Tamilnadu Agric. Univ. Coimbatore; 2014.
- Jayalakshmi K. Studies on anthracnose of pomegranate caused by Colletotrichum gloeosporioides(Penz.) Penz. &Sacc [M.Sc. thesis]Univ. of Agric. Sci. Dharwad, India; 2010.
- Patel KD, Joshi KR. Antogonistic effect of some bioagents *In vitro* against *Colletotrichum gloeosporioides* Penz. and Sacc. the causal agent of leaf spot of turmeric. J Mycol Plant Pathol. 2001;31:126.
- 22. Ashoka S. Studies on fungal pathogens of vanilla with special reference to Colletotrichum *gloeosporioides* (Penz.) Penz. and Sacc. M.Sc. (Agri.) [thesis], Uni. Agric. Sci. Dharwad, India; 2005.
- 23. Prabhakar K, Raguchander Τ. Saravanakumar D, Muthulakshmi Ρ. Parthiban VK, Prakasam V. Management postharvest disease of mango of anthracnose incited by Colletotrichum gloeosporioides.phytopatho. Plant Prot. 2008;41(5):333-9.
- 24. Raheja S, Thakore BBL. Effect of physical factor, plant extracts and bioagent on *Colletotrichum gloeosporioides* Penz. the causal organism of anthracnose of Yam. J Mycol Plant Pathol. 2002;32:293-4.
- 25. Godtfredsen WO, Vangedal S. Trichodermin, а new sesquiterpene antibiotic. Acta Chem Scand. 1965;19(5):1088-102. DOI: 10.3891/acta.chem.scand.19-1088, PMID 5850137.
- Shi J, Liu A, Li X, Feng S, Chen W. Identification of endophytic bacterial strain MGP1 selected from papaya and its biocontrol effects on pathogens infecting harvested papaya fruit. J Sci Food Agric. 2010;90(2):227-32. DOI: 10.1002/jsfa.3798, PMID 20355035.

- Senthil R, Prabhakar K, Rajendran N, Karthikeyan G. Efficacy of different biological control agents major postharvest pathogens of grapes under room temperature storage conditions, Phytopathology Mediterrian. 2011;50: 55-65.
- Zheng M, Shi J, Shi J, Wang Q, Li Y. Antimicrobial effects of volatiles produced by two antagonistic Bacillus strains on the anthracnose pathogen in postharvest mangos. Biol Control. 2013;65(2): 200-6.

DOI: 10.1016/j.biocontrol.2013.02.004

29. Calvo H, Marco P, Blanco D, Oria R, Venturini ME. Potential of a new strain of Bacillus amyloliquefaciesBUZ-14 as a biocontrol agent of postharvest fruit diseases. Food Microbiol. 2017;63:101-10. DOI: 10.1016/j.fm.2016.11.004, PMID 28040156.

30. Sharma RR, Singh D, Singh R. Biological control of postharvest diseases of fruits and vegetables by microbial antagonists: a review. Biol Control. 2009;50(3):205-21.

DOI: 10.1016/j.biocontrol.2009.05.001

- 31. Chau KF. A histological study of anthracnose on *Carica papaya*. Phytopathology. 1983;73(8):1113-6. DOI: 10.1094/Phyto-73-1113
- 32. Hu HQ, Li XS, He H. Characterization of an antimicrobial material from a newly isolated Bacillus amyloliquefaciens from mangrove for biocontrol of Capsicum bacterial wilt. Biol Control. 2010;54(3): 359-65.

DOI: 10.1016/j.biocontrol.2010.06.015

© 2022 Yadravi et al.; This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/4.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Peer-review history: The peer review history for this paper can be accessed here: https://www.sdiarticle5.com/review-history/92279