Production and Testing of Biopesticide for Control of Postharvest Mold Infections on Fresh Fruits of Apple and Pear

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Abstract

The present research aimed at producing a biopesticide with proper formulation of invert emulsion (water-in-oil type) and effective strain of *Trichoderma harzianum* then, testing it against pathogens of postharvest mold infections (*Botrytis cinerea* & *Penicillium expansum*) on fresh fruits of apple and pear. The proper formulation of invert emulsion (IE#3) used for biopesticide production has the following ingredients (100% w/w): soybean oil (28.50%), coconut oil (19.50%), oil-soluble emulsifier: Tween 20 (2.0%), glycerine (4.25%), water-soluble emulsifier: dehymuls k (0.75%), sterile distilled water (22.5%) and conidial suspension of the effective strain of *T. harzianum* “TrichoPAL1” in water (22.5%, concentration 1 x 10⁷ conidia/ml). Testing efficacy of the produced biopesticide has indicated a significant reduction in the disease lesion diameter of mold infections on wounded apple and pear fruits stored at 20°C ± 1°C compared to the untreated fruits or control (reduction from up to 38.75 to about 7.50 mm, respectively, according to the type of mold infections and fruit type). Also, the treatment with the produced biopesticide has resulted in a long protection period from mold infections on wounded and un-wounded fresh fruits of apple and pear stored under controlled and semi-commercial conditions (up to 2.5 months according to the type of mold infections and fruit type). In conclusion, the overall results have demonstrated the effectiveness of produced biopesticide on stored fruits under controlled and semi-commercial conditions therefore, it is recommended to test this effectiveness on marketed fruits stored under variable conditions before applying it at a large scale.

Keywords

Apple Fruit, Pear Fruit, *Trichoderma harzianum*, *Botrytis cinerea*, *Penicillium expansum*, Biopesticides, Postharvest Mold Infections

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1. Introduction

The control of postharvest mold infections on fresh fruits is usually done by the conventional use of synthetic fungicides [1] [2]. Alternative approaches to chemical use are highly appreciated due to the increasing importance of integrated management strategies relying on biocontrol possibly combined with selective application of fungicides [3] [4]. The above strategies must achieve two demands: effectiveness and environmental safety. Also, integrated control could be a convenient strategy for newer attempts in the control of postharvest mold infections [5] [6].

Regarding the group of interesting biocontrol agents that can be involved in the integrated management strategies, the antagonistic fungi are considered the most promising agents because they are currently being developed as a novel approach for the control of many plant diseases [7]. However, these fungi suffer from the following disadvantages: i) their initial or acute biocontrol potential is much slower in comparison with chemical fungicides; ii) they need high humidity for conidial germination and subsequent development and sporulation; iii) they are susceptible to UV irradiation; and iv) some targeted plant diseases may develop defense mechanisms against the fungal attack. Some of these constraints could be overcome by developing, for example, formulations which improve the stability of these fungi, UV resistance, distribution on the surface of disease host-plant, and contact to the targets or can synergistically enhance efficacy by improving the virulence of these fungi [8] [9].

The use of antagonistic fungi as biocontrol agents of plant diseases, for example, postharvest mold infections, has been previously reported by some investigators [10]-[13], but without attempting to formulate these biocontrol agents during their application. Therefore, formulation of these fungi is crucial for increasing their efficacy during application against targeted plant diseases. Invert Emulsions (water-in-oil type) are one of the promising formulations that can be used for antagonistic fungi because these emulsions contain the necessary water for germination of fungal conidia and their development during or after application. The water introduced into emulsions is usually homogenized with oil at a high speed (20,000 rpm for 1.5 minutes using homogeniser), thus the water droplets in the final emulsions are completely surrounded by the oil droplets (water-in-oil emulsion). Therefore, oil droplets prevent the water droplets from quick evaporation. Under hot dry conditions, water used for preparation of the conidial suspension of these fungi will evaporate and dry up quickly but if this water is incorporated into invert emulsions as described above, it will evaporate very slowly following application due to the above reason [14].

Recently, research studies conducted in our laboratory under controlled environmental conditions have shown that certain strains of *Trichoderma harzianum* applied in invert emulsion are proved to be effective against fungal plant pathogens infecting the fresh produce of various crops including fruits postharvest [15]-[17]. Therefore, in the present research, biopesticides based on using new combinations (formulations) of invert emulsions with effective strain(s) of the antagonistic fungus—*T. harzianum* will be produced then tested against postharvest mold infections on apple and pear fruits stored under variable environmental conditions and in semi-commercial conditions. Therefore, the main objective of the present research was to produce an effective biopesticide against postharvest mold infections on apples and pears and then test its effectiveness. To arrive this objective, the following aims should be fulfilled: i) screening for selection of the most effective strain of *Trichoderma harzianum* against postharvest mold infections; ii) formulation of the selected strain of *T. harzianum* using new combinations of invert emulsion formulation; iii) testing the efficacy of the formulated strain of *T. harzianum* in invert emulsions against mold infections especially those caused by *Botrytis cinerea* and *Penicillium expansum* on apple and pear fruits stored in incubator or under semi-commercial conditions.

2. Materials and Methods

2.1. Isolation of Pathogens Causing Mold Infections on Fresh Fruits of Apple and Pear

The following postharvest mold infections of fruits were studied in the present research: Gray mold caused by *Botrytis cinerea* and Blue mold caused by *Penicillium expansum*. Local strains of *B. cinerea* (designated BCPAL) and *P. expansum* (designated PEPAL) were isolated on Potato Dextrose Agar medium (PDA) amended with Chloramphenicol antibiotic (250 mg/L W/V, added to the medium after autoclaving). The two strains were isolated from naturally infected fruits of apple (source: a local wholesale market). Pure cultures of these isolated strains were used after sub-culturing of them on the same culture medium (PDA). These cultures were kept in a cold room at 4°C ± 1°C for being used during bioassays’ conduction.
2.2. Isolation of Strains of the Antagonistic Fungus: *T. harzianum*

Local strains of *T. harzianum* were used in the present bioassays since this fungus has an antagonistic effect to many causal agents of fungal plant diseases including postharvest mold fungi of fresh fruits. Four local strains of *T. harzianum* were isolated from dead tomato roots previously infected with root rot fungi and then buried in the soil cultivated with various crops in Tulkarm district, Palestine. The isolated strains were designated “TrichoPAL1”, “TrichoPAL2”, “TrichoPAL3” and “TrichoPAL4”. They were first isolated on Potato Dextrose Agar medium (PDA) amended with Chloramphenicol antibiotic (250 mg/L W/V, added to the medium after autoclaving). Identification of these strains as isolates of *T. harzianum* was made according to the typical morphological characteristics of this fungus described by Grondona et al. 1997. After sub-culturing of these strains on oatmeal agar medium (OMA), pure cultures of these strains were kept in a cold room (4°C ± 1°C) for being used during bioassays’ conduction.

2.3. Screening Test for the Efficacy of Isolated *T. harzianum* Strains

Efficacy of the isolated strains of *T. harzianum* was tested against postharvest mold infections for screening the most effective strain. For this purpose, healthy apple and pear fruits were infected with *B. cinerea* (strain BCPAL) and *P. expansum* (strain PEPAL) then treated with the conidial suspension of *T. harzianum* isolates at a concentration of $1 \times 10^7$ conidia/ml each. Comparison of the efficacy of these strains was done according to the suppression level of the disease lesion diameter resulted from *B. cinerea* and *P. expansum* infection on fruits used in bioassays. The infection with *B. cinerea* and *P. expansum* on apple and pear fruits and then treating them with the antagonistic fungus (*T. harzianum*) was done by depositing 25 µl-droplet of conidial suspension of *B. cinerea* or *P. expansum* (concentration $1 \times 10^6$ conidia/ml) per fruit after inducing a small superficial wound on each fruit surface. A micropipette (100 µl capacity) is used for depositing of each droplet on each fruit. The control treatment is consisting of superficially wounded fruits treated with 25 µl-droplet of sterile distilled water. Inoculated and control fruits were then treated with conidial suspension of the following *T. harzianum* strains: “TrichoPAL1”, “TrichoPAL2”, “TrichoPAL3” and “TrichoPAL4”. This treatment was done by depositing 25 µl-droplet of conidial suspension (concentration $1 \times 10^7$ conidia/ml each) of each strain per fruit at the same site of disease inoculation using a 100 µl-micropipette. Four fruits representing 4 replicates per inoculated fruit type per pathogen per *T. harzianum* strain or control treatment were used. Each fruit was kept in a closed plastic pot of 9 cm diameter and 15 cm depth. A completely randomized design was used. The disease lesion diameter obtained 7 days after the treatment with the antagonistic fungus and incubation at 20°C ± 1°C was measured on each fruit then the mean lesion diameter of each disease on each fruit type per disease per *T. harzianum* strain was calculated. Finally, the Average mean of disease lesion diameter per strain of *T. harzianum* was calculated and used for comparison of the efficacy of tested strains to select the most effective one.

2.4. Preparation of Invert Emulsion Formulations and Selection of the Most Proper

New combinations of invert emulsion formulations (water-in-oil type) were prepared in this research. Each combination was mainly consisted of two phases: i) aqueous or water phase comprising of sterile distilled water, glycerine and water-soluble emulsifier, and ii) oil phase comprising of plant-origin oil(s) and oil-soluble emulsifier. The proportion of each phase in the final invert emulsion was 50% (w/w) and the percentage of ingredients in each phase was changed during preparation so that the resulting combination was accepted as a suitable formulation if it had a high stability and low viscosity. Mixing of the two phases during preparation of the emulsion was done at a high speed (20,000 rpm for 1.5 minutes using a homogenizer). This is done to ensure the homogeneity of the emulsion. Five combinations were prepared using the above technique in order to select the most proper one characterized by the highest stability and lowest viscosity. The stability of the prepared emulsion was judged by the non-separation of the two phases composing the emulsion along the time following preparation. The viscosity of the prepared emulsion was measured by a viscometer. The emulsion is considered “not viscous” if its viscosity <25 cps (centipoises), “relatively viscous” if its viscosity 25 - 35 cps and viscous if its viscosity >35 cps.

2.5. Introduction of the Most Effective Strain of *T. harzianum* into the Selected Formulation of Invert Emulsion

The most effective strain of *T. harzianum* obtained by screening in the above section was introduced into the se-
selected formulation of invert emulsion characterized by the highest stability and lowest viscosity. Introduction was done, at first, into the aqueous or water phase which finally become consisted of sterile distilled water, conidial suspension of the effective strain of *T. harzianum*, glycerine and water-soluble emulsifier. The ingredients of oil phase did not change and remained as indicated above. The ratio of these two phases in the final invert emulsion was the same as indicated above. Conidial suspension of the most effective strain of *T. harzianum* in the formulation was $1 \times 10^7$ conidia/ml. Mixing of the two phases after introduction of the conidial suspension of the antagonistic fungus was done as indicated in the previous section to ensure the homogeneity of the emulsion.

2.6. Testing Efficacy of Formulated Strain of *T. harzianum* against Postharvest Mold Infections on Fruits

The efficacy test was practiced against mold infections on wounded and unwounded fruits of apple and pear stored at 20°C ± 1°C in the incubator or at 20°C ± 5°C under semi-commercial conditions as follows.

2.6.1. Efficacy on Wounded Fruits of Apple and Pear Stored in Incubator at 20°C ± 1°C

To test efficacy of the most effective strain of *T. harzianum* formulated in invert emulsion against pathogens of postharvest mold infections, healthy fruits of apple and pear were, first, infected with *B. cinerea* and *P. expansum* then, treated with the formulated *T. harzianum* according to the protocol indicated in the previous section. Comparison of the efficacy after the treatment was done according to the suppression level of the disease lesion diameter resulted from *B. cinerea* or *P. expansum* infection on treated fruits. The technique of inoculation with the mold pathogens on apple and pear fruits and then treatment with the formulated *T. harzianum* was done by depositing 25 µl-droplet of conidial suspension of each one of mold pathogens (concentration $1 \times 10^6$ conidia/ml) per fruit after inducing a small circular superficial wound (2 mm diameter) on each fruit surface. A micropipette (100 µl capacity) was used for depositing the droplet on the wounded fruit. The control treatment is consisting of superficially wounded fruits treated with 25 µl-droplet of sterile distilled water. Inoculated and control fruits were then treated with 25 µl-droplet of formulated strain of *T. harzianum* (concentration $1 \times 10^7$ conidia/ml each) deposited at the same site of disease inoculation. In each bioassay, four types of treatments were used per pathogen per fruit type. The treatments were: formulated *T. harzianum* in invert emulsion, unformulated *T. harzianum*, blank formulation of invert emulsion and control (untreated with *T. harzianum*). Four fruits representing 4 replicates per treatment type were used. After that, each treated fruit was placed in a closed plastic pot (9 cm diameter and 15 cm depth) then stored in an incubator at 20°C ± 1°C. A completely randomized design was used. For each pathogen on each fruit type, the disease lesion diameter obtained 7 days after the treatment was measured then the mean lesion diameter of each treatment was calculated then used for comparison of the efficacy.

2.6.2. Efficacy on Un-Wounded Fruits of Apple and Pear Stored in Incubator at 20°C ± 1°C

To test efficacy of the most effective strain of *T. harzianum* formulated in invert emulsion against the fungal pathogens of postharvest mold infections, healthy un-wounded fruits of apple and pear were, first, infected with *B. cinerea* and *P. expansum* then treated with the formulated *T. harzianum* strain. Evaluation of the efficacy was done according to the time required for the appearance of disease symptoms (B. cinerea or P. expansum) on untreated and treated fruits with the formulated *T. harzianum*. The technique of inoculation with the fungal pathogens of mold infections on apple and pear fruits was done by spraying 200 µl spray volume of conidial suspension of these fungal pathogens (concentration $1 \times 10^6$ conidia/ml) per healthy un-wounded fruit then the inoculated fruits were placed in rectangular plastic boxes (20 × 10 × 10 cm; long, wide and high, respectively). A transparent plastic film was used to cover each box. A small calibrated hand sprayer (1 L capacity) was used for spraying the intended spray volume on the un-wounded fruits. The control treatment is consisting of un-wounded fruits sprayed with the same quantity of spray volume of sterile distilled water after infection with mold pathogens. Inoculated and control fruits were then sprayed with 200 µl spray volume of formulated *T. harzianum* (concentration $1 \times 10^7$ conidia/ml each) per fruit in the same manner as indicated above. In each bioassay, four types of treatments were used per pathogen per fruit type. The treatments were: formulated *T. harzianum* in invert emulsion, unformulated *T. harzianum*, blank formulation of invert emulsion and control (untreated with *T. harzianum*). Five fruits per box representing 5 replicates per treatment type were used. A completely randomized design was used. The time needed for the appearance of typical symptoms of each disease (gray or blue mold) on fruits of each treatment was determined and then used for evaluation of the efficacy. The plastic
boxes containing fruits used for bioassays were stored in an incubator at 20°C ± 1°C until the appearance of the disease lesion on the incubated fruits.

2.6.3. Efficacy on Un-Wounded Fruits of Apple and Pear Stored at 20°C ± 5°C (Under Semi-Commercial Conditions)

To test efficacy of the effective strain of T. harzianum formulated in invert emulsion against the fungal pathogens of postharvest mold infections on apple and pear fruits stored under semi-commercial conditions (20°C ± 5°C), rectangular plastic boxes (20 × 10 × 10 cm; long, wide and high, respectively) were used. A transparent plastic film was used to cover the fruits after treatment. These boxes are similar to those used for marketing of these fruits. The technique of inoculation with the fungal pathogens of mold infections on apple and pear fruits was done by spraying 1.0 ml of conidial suspension of these fungal pathogens (concentration 1 × 10⁶ conidia/ml) on healthy un-wounded fruits kept in each plastic box. A small calibrated hand sprayer (1 L capacity) was used for spraying the intended spray volume per box containing 5 healthy un-wounded fruits. The control treatment is consisting of 5 healthy un-wounded fruits kept in one box and sprayed with the same spray volume of sterile distilled water (1.0 ml). The inoculated and control fruits were then sprayed with 1.0 ml spray volume of formulated T. harzianum (concentration 1 × 10⁷ conidia/ml each) using the technique and equipment as indicated above. For each bioassay, two types of treatments were used per pathogen per fruit type. The treatments were: formulated T. harzianum in invert emulsion and control (untreated with T. harzianum). Five fruits per box representing 5 replicates per treatment type were used. A completely randomized design was used. The time needed for the appearance of typical symptoms of each disease (gray or blue mold) on fruits after the treatment was determined and then used for evaluation of the efficacy. The protection period represents the time extending from inoculation to appearance of the first symptoms of infection with the disease on treated fruits. During bioassays, the plastic boxes that contain fruits were stored in a store-room at 20°C ± 5°C until the appearance of the disease symptoms on incubated fruits. These conditions are similar to those used for storing fruits of apple and pear during marketing of the product.

2.7. Statistical Analyses

For the targeted diseases of mold infections, the mean % of disease lesion diameter on fruits treated with the formulated and unformulated T. harzianum was statistically analyzed using analysis of variance (ANOVA). Significant differences (at P = 0.05) between the treatment means were determined by F-test. Also, the mean separation was performed using Duncan’s Multiple Range test.

3. Results

3.1. Efficacy of the Isolated Strains of T. harzianum

As a result of the screening test, the strain of T. harzianum “TrichoPAL1” has shown the highest efficacy against B. cinerea (strain BCPAL) and P. expansum (strain PEPAL) in comparison with other strains of T. harzianum because this strain has significantly reduced (at P = 0.05) the average mean of disease lesion diameter of both pathogens on wounded apple and pear fruits from 21.25 mm diameter (in the untreated control) to 12.31 mm diameter (Table 1). Therefore, it has been selected as the most effective strain of T. harzianum for being introduced into the selected formulation of invert emulsion. The formulated strain of T. harzianum was then used for further bioassays as biopesticide. However, no significant differences were obtained between the average means of disease lesion diameter of both pathogens on wounded apple and pear fruits for the other strains of T. harzianum “TrichoPAL2”, “TrichoPAL3” and “TrichoPAL4” and the untreated control (Table 1), so they are not selected for further bioassays.

3.2. Selection of Invert Emulsion Formulation as a Carrier of T. harzianum

Among the different combinations (formulations) of invert emulsion prepared, the combination “IE # 3” was selected as the most proper because it has shown a high stability for long time with no viscosity (Table 2), so that it has been selected as a carrier of T. harzianum “strain TrichoPAL1” then tested as biopesticide against mold infections on fresh fruits. The selected formulation (IE # 3) has the following ingredients: i) a mixture of two oils of plant-origin (soybean oil: 28.50% w/w and coconut oil: 19.50% w/w); ii) oil-soluble emulsifier (Tween 20: 2.0% w/w); iii) sterile distilled water (22.5% w/w); iv) conidial suspension of the selected strain of T. harzianum in...
Table 1. Disease lesions of *Botrytis cinerea* (strain BCPAL) and *Penicillium expansum* (strain PEPAL) formed on wounded apple and pear fruits (varieties: Red Delicious-Starking and Bartlett, respectively) 7 days after inoculation and treatment with four strains of *Trichoderma harzianum* (incubation of treated fruits was done in an incubator at 20°C ± 1°C).

<table>
<thead>
<tr>
<th>Strains of <em>T. harzianum</em></th>
<th>Mean disease lesion diameter of <em>B. cinerea</em> and <em>P. expansum</em> (in mm) observed on wounded apple and pear fruits when treated with <em>T. harzianum</em></th>
<th>Average mean of disease lesion diameter (±SEM) per strain of <em>T. harzianum</em>**</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td><em>B. cinerea</em> on Apple fruit</td>
<td><em>B. cinerea</em> on Pear fruit</td>
</tr>
<tr>
<td>TrichoPAL1</td>
<td>14.25</td>
<td>10.50</td>
</tr>
<tr>
<td>TrichoPAL2</td>
<td>20.75</td>
<td>17.50</td>
</tr>
<tr>
<td>TrichoPAL3</td>
<td>19.50</td>
<td>21.25</td>
</tr>
<tr>
<td>TrichoPAL4</td>
<td>20.25</td>
<td>18.50</td>
</tr>
<tr>
<td>Untreated Control</td>
<td>20.00</td>
<td>21.25</td>
</tr>
</tbody>
</table>

*Four fruits of each fruit type representing 4 replicates per pathogen treatment per strain were used. Therefore, 4 replicates were performed for each test (20 tests were performed for the 4 strains + control). SEM: standard error of the mean or standard deviation of the mean which is = Standard Deviation of original distribution (SD)/Square root of Sample Size (n); **Means followed by the same letter are not significantly different at *P* = 0.05 using ANOVA and Duncan’s Multiple Range test.

Table 2. Combinations of invert emulsion formulation (IE) based on variable percentage of ingredients composing oil phase and water phase of the emulsion.

<table>
<thead>
<tr>
<th>Combination number of invert emulsion (IE)</th>
<th>% of ingredients (w/w) of water phase and oil phase composing the formulation of invert emulsion (IE)</th>
<th>Stability of prepared emulsion</th>
<th>Viscosity of prepared emulsion</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Soybean oil</td>
<td>Coconut oil</td>
<td>Tween 20</td>
</tr>
<tr>
<td>IE # 1</td>
<td>29.00</td>
<td>18.00</td>
<td>3.00</td>
</tr>
<tr>
<td>IE # 2</td>
<td>28.50</td>
<td>19.00</td>
<td>2.50</td>
</tr>
<tr>
<td>IE # 3</td>
<td>28.50</td>
<td>19.50</td>
<td>2.00</td>
</tr>
<tr>
<td>IE # 4</td>
<td>28.50</td>
<td>19.00</td>
<td>2.50</td>
</tr>
<tr>
<td>IE # 5</td>
<td>29.00</td>
<td>18.00</td>
<td>3.00</td>
</tr>
</tbody>
</table>

The total % of these ingredients (w/w) is 100 which is equally divided between the water phase and the oil phase (50% w/w each).

3.3. Testing the Efficacy of Formulated *T. harzianum* in Invert Emulsion against Mold Infections on Fresh Fruits

3.3.1. On Wounded Fruits of Apple and Pear Stored in an Incubator at 20°C ± 1°C

The mean of disease lesion diameter of *B. cinerea* and *P. expansum* on wounded apple fruits has significantly decreased (at *P* = 0.05) when treated with the formulated strain “TrichoPAL1” of *T. harzianum* in invert emulsion (combination IE # 3). This decrease was ranged from 35.75 and 18.75 mm to 6.25 and 3.75 mm for *B. cinerea* and *P. expansum*, respectively, in comparison with the control (Table 3). The same trend was obtained in the results on wounded pear fruits (significant reduction in the mean of disease lesion diameter of both pathogens due to treatment with the formulated *T. harzianum*) (Table 3). However, no significant differences are present between means of the disease lesion diameter of both diseases on both types of fruits regarding the two types of control treatments (untreated fruit control and Blank formulation of invert emulsion) (Table 3). This may indicate that no harmful effects may be caused by the ingredients of the selected formulation.

Water (TrichoPAL1: 22.5% w/w), v) glycerine (4.25% w/w), and water-soluble emulsifier (Dehymuls k: 0.75% w/w) (Table 2). The total % of these ingredients (w/w) is 100 which is equally divided between the water phase and the oil phase (50% w/w each).
Table 3. Disease lesions of Botrytis cinerea (strain BCPAL) and Penicillium expansum (strain PEPAL) formed on wounded apple and pear fruits (variety: Red Delicious-Starking and Bartlett, respectively) after inoculation and treatment with unformulated and formulated Trichoderma harzianum (strain TrichoPAL1) in invert emulsion (incubation of the treated and control fruits was done in incubator at 20°C ± 1°C).

<table>
<thead>
<tr>
<th>Treatments with T. harzianum and control</th>
<th>Mean lesion diameter (in mm) (±SEM) of B. cinerea appeared after 5 days of inoculation and treatment</th>
<th>Mean lesion diameter (in mm) (±SEM) of P. expansum appeared after 7 days of inoculation and treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Formulated TrichoPAL1 of T. harzianum</td>
<td>6.25 (±0.42) A**</td>
<td>3.75 (±0.45) A**</td>
</tr>
<tr>
<td>Un.formulated TrichoPAL1 of T. harzianum</td>
<td>14.50 (±0.78) B</td>
<td>7.50 (±0.81) B</td>
</tr>
<tr>
<td>Blank formulation of invert emulsion (untreated fruits)</td>
<td>31.25 (±0.69) C</td>
<td>16.25 (±0.85) C</td>
</tr>
<tr>
<td>Control (untreated fruits)</td>
<td>35.75 (±0.82) C</td>
<td>18.75 (±0.91) C</td>
</tr>
</tbody>
</table>

*Four fruits of each fruit type representing 4 replicates per pathogen treatment per strain were used. Therefore, 4 replicates were performed for each test (16 tests were performed for formulated strain, unformulated strain, blank formulation and control). SEM: standard error of the mean or standard deviation of the mean which is = Standard Deviation of original distribution (SD)/Square root of Sample Size (n); **Within each column, means followed by the same letter are not significantly different at P = 0.05 using ANOVA and Duncan’s Multiple Range test.

3.3.2. On Un-Wounded Fruits of Apple and Pear Stored in an Incubator at 20°C ± 1°C

The time required for the appearance of the first symptoms of infection with B. cinerea and P. expansum on un-wounded fresh apples stored in an incubator under controlled conditions (20°C ± 1°C) was the longest on apple fruits treated with the formulated T. harzianum (72 and 58 days for the two diseases, respectively) and the shortest on apple fruits of the control treatment (12 and 18 days for the two diseases, respectively) (Table 4). Intermediate time periods were required for the appearance of the first symptoms of infection with the above-mentioned diseases on apples treated with the unformulated T. harzianum or with the blank formulation (Table 4). These time periods represent the protection period on apple fruits needed to prevent infection with these diseases when treated with the formulated T. harzianum compared to the control. Lower values of protection periods against the same mold infections (B. cinerea and P. expansum) on un-wounded fresh pear fruits stored at 20°C ± 1°C were obtained when treated with the formulated T. harzianum compared to the control (65 and 50 days versus 11 and 19 days for the two diseases, respectively) (Table 4).

3.3.3. On Un-Wounded Fruits of Apple and Pear Stored in Semi-Commercial Conditions at 20°C ± 5°C

The time required for the appearance of the first symptoms of infection with B. cinerea and P. expansum on un-wounded fresh fruits of apple stored in semi-commercial conditions (20°C ± 5°C) was the longest on apple fruits treated with the formulated T. harzianum (61 and 44 days for the two diseases, respectively) and the shortest on apple fruits of the control treatment (10 and 15 days for the two diseases, respectively) (Table 5). These time periods represent the protection period on apple fruits needed to prevent infection with the diseases when treated with the formulated T. harzianum compared to the control. Lower values of protection periods against the same mold infections (B. cinerea and P. expansum) on un-wounded fresh pear fruits stored at 20°C ± 5°C were obtained when treated with the formulated T. harzianum compared to the control (50 and 40 days versus 9 and 12 day for the two diseases, respectively) (Table 5).

4. Discussion

Results obtained in the present research demonstrate the ability of the produced biopesticide to reduce significantly the mold infections on fresh fruits of apple and pear stored under controlled and partially controlled conditions. This ability is attributed to the capacity of the formulated strain of T. harzianum (TrichoPAL1) to suppress the growth of pathogens causing mold infections on treated fruits (B. cinerea and P. expansum) hence control these postharvest diseases. In addition to this ability, the formulation of invert emulsion (water-in-oil type) used for production of this biopesticide has significantly increased its efficacy against the targeted post
Table 4. Time period needed for the first appearance of disease lesion of *Botrytis cinerea* (strain BCPAL) and *Penicillium expansum* (strain PEPAL) on un-wounded apple and pear fruits (variety: Red Delicious-Starking and Bartlett, respectively) treated with unformulated and formulated *Trichoderma harzianum* (strain TrichoPAL1) (incubation of the treated and control fruits was done in incubator at 20°C ± 1°C).

<table>
<thead>
<tr>
<th>Treatments with <em>T. harzianum</em> and control</th>
<th>Time needed for the first appearance of lesions of <em>B. cinerea</em> (in days)*</th>
<th>Time needed for the first appearance of lesions of <em>P. expansum</em> (in days)*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>on un-wounded apple fruit</td>
<td>on un-wounded pear fruit</td>
</tr>
<tr>
<td>Formulated TrichoPAL1 of <em>T. harzianum</em></td>
<td>72</td>
<td>65</td>
</tr>
<tr>
<td>Un-formulated TrichoPAL1 of <em>T. harzianum</em></td>
<td>39</td>
<td>40</td>
</tr>
<tr>
<td>Blank formulation of invert emulsion</td>
<td>14</td>
<td>12</td>
</tr>
<tr>
<td>Control (untreated fruits)</td>
<td>12</td>
<td>11</td>
</tr>
</tbody>
</table>

*Four fruits of each fruit type representing 4 replicates per pathogen treatment per strain were used. Therefore, 4 replicates were performed for each test (16 tests were performed for formulated strain, unformulated strain, blank formulation and control).

Table 5. Time period needed for the appearance of the first disease lesion of *Botrytis cinerea* (strain BCPAL) and *Penicillium expansum* (strain PEPAL) on un-wounded apple and pear fruits (variety: Red Delicious-Starking and Bartlett, respectively) treated with formulated *Trichoderma harzianum* (strain TrichoPAL1) (incubation of the treated and control fruits was done in semi-commercial conditions (20°C ± 5°C)).

<table>
<thead>
<tr>
<th>Treatments with <em>T. harzianum</em> and control</th>
<th>Time needed for the first appearance of lesions of <em>B. cinerea</em> (in days)*</th>
<th>Time needed for the first appearance of lesions of <em>P. expansum</em> (in days)*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>on un-wounded apple fruit</td>
<td>on un-wounded pear fruit</td>
</tr>
<tr>
<td>Formulated TrichoPAL1 of <em>T. harzianum</em></td>
<td>61</td>
<td>50</td>
</tr>
<tr>
<td>Control (untreated fruits)</td>
<td>10</td>
<td>9</td>
</tr>
</tbody>
</table>

*Four fruits of each fruit type representing 4 replicates per pathogen treatment per strain were used. Therefore, 4 replicates were performed for each test (8 tests were performed for formulated strain and control).

It is well-known that the various strains of *T. harzianum* can establish a mycoparasitic relationship with their host pathogens during parasitism. This relationship could be done by interfering with the development and growth of pathogens after conidial germination through coiling of their mycelium over the pathogen mycelium or by disrupting the host fungus cell wall and consequent death of the pathogen [18] [19]. Antibiosis is another antagonistic relationship made by certain strains of *T. harzianum*. This antibiosis could be done by producing antifungal antibiotics in form of extracellular products (for example, hydrolytic enzymes like chitinases, glucanases...
and proteases) [18] [20]-[24]. Also, these strains could be propagated after successful host infection and distribution within the contaminated environment (horizontal transmission is most probable). The above characteristics make these antagonistic fungi excellent candidates as biocontrol agents in the integrated management strategies of many plant diseases including postharvest diseases [21] [25].

From a practical point of view, formulation of *T. harzianum* strains is very necessary for protecting the introduced fungus and enhancing its efficacy. Until present, many researchers have attempted to formulate strains and isolates of *T. harzianum* using appropriate formulations. For example, formulation of T39 isolate of *T. harzianum* in a powder form (commercially sold as Trichodex™) then application of it against pathogens of many foliar diseases such as *Botrytis cinerea*, *Pseudoperonospora cubensis*, *Sclerotinia sclerotiorum* and *Sphaerotheca fusca* (syn. *S. fuliginea*) in cucumber under commercial greenhouse conditions [26]. Also, formulation of the strain 1295-22 (T-22) of *T. harzianum* in a liquid form (commercialized as T-22 Planter Box™) containing the conidial suspension of the fungus used for the seed treatment against damping-off diseases [7] [27]. The same strain was also formulated in a liquid form but this form contains the entire thallus of the fungus colonized on clay particles (commercialized as RootShield™) for control of soil borne diseases [7] [27]. There is also another *T. harzianum* product available in the Czech Republic and Denmark for the greenhouse use called commercially (Supresvit™) in a dispersible powder containing conidia of the strain PV5736-89 and used for soil or potting mixes to control damping-off and root rots of ornamentals and forest tree-seedlings [7]. In comparison with the above formulations of *T. harzianum*, our formulation of invert emulsion (water-in-oil type) contains the suspended conidia of the fungus strain “TrichoPAL1” in the emulsion. These conidia were incorporated in the emulsion during its preparation so that the viability of these conidia could be preserved for long time realizing a significant reduction in the mean of disease lesion diameter of pathogens challenged. In a previous research, one of the prepared formulations of invert emulsion (water-in-oil type) that contained the conidia of an effective strain of the fungus “Ci306” did not significantly lose its viability after 6 weeks of their introduction into emulsion. Also, it achieved a high biological efficacy against *B. cinerea* on strawberry by decreasing the disease lesion diameter of the pathogen by 44.0% to 55.0% compared to the control [15]. This proves again the benefits of our formulation and its efficacy compared to other formulations.

5. Conclusion

An effective strain of *T. harzianum* was successfully formulated as biopesticide using a stable and non-viscous formulation of invert emulsion (water-in-oil type) then tested against target pathogens of postharvest mold infections on fruits of apple and pear. Treatments with the tested biopesticide have reduced significantly the mean of disease lesion diameter on previously infected fruits before treatment. Also, they have protected fruits for a long time from later infections (protection period up to 2.5 months). Due to above demonstration of biopesticide effectiveness, it is recommended to test it at a large scale for the control of mold infections on marketed fruits stored under variable conditions.

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References


