Interaction efficiency of *Trichoderma* spp. and some plant extracts against ear-cockle disease

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**ABSTRACT**

Ear-cockle disease is one of the important aerial diseases of wheat. It is caused by wheat seed gall nematode *Anguina tritici*. This study aimed to evaluate the interaction effect between *Trichoderma* spp. as bioagents and some plant extracts. The latter are used as alternative substances to control the seed gall nematode in both laboratory and field conditions. The bioassay of juveniles in the second stage (J2) of *A. tritici* was conducted using the biopesticide Biocont-T-WP (*Trichoderma harzianum*), isolate of *Trichoderma hamatum* T-113, and nematic and seaweed extract for their effect on the viability of *A. tritici* J2. The lab-bioassay was achieved on the daily accumulated J2 mortality percentage during 1, 3, 5, 7, 9, and 11 days of incubation, while the field-bioassay was evaluated for wheat growth, yield, and infection attributes. The lab-bioassay showed that Biocont-T has a higher mortality effect followed by the nematic extract with averages of 40.56% and 12.52%, respectively. The accumulative J2 mortality percentage increased gradually and reached a maximum on the 11th day of incubation. In the field-bioassay, mixing wheat seed Ibaa-99 with Biocont-T decreased infection percentage 77.7% and the number of galls spike–1 to 1.63 compared to the control 5.66 galls spike–1.

1. INTRODUCTION

Wheat (*Triticum aestivum*) is a major crop in Iraq. Wheat yield faced different types of pests that may affect it negatively. Wheat is cultivated all over Iraq with an approximate area that exceeds 1.7 million ha. The Iraqi people consume, on average, nearly 7 million t of wheat per year [1].

Ear-cockle disease is one of the aerial diseases affecting wheat, caused by wheat seed gall nematode *Anguina tritici* (Steinbuch), which is considered a major pest in Iraq in addition to all wheat production areas in the world. The juveniles (J1) emerge from the wheat galls in the soil, attack plants, and move into the germinated seedlings. Next, J1 will move up to the spikes hurting new grains, turning the seed into seed galls, then causing major loss in the grain yield quality and quantity [2]. During the harvesting process, galls might fall into the soil and be a source of new infection that would be generated in the following season [3].

In Iraq, the first record of *A. tritici* was reported by [4], which recorded it as plant parasite nematode [5]. Al-Beldawi et al. [6] revealed that most wheat cultivars in Iraq are susceptible to infection with ear-cockle disease except var. Mexipak which showed resistance to this nematode. The infection rate is affected by several factors such as the number of inoculums (1 gall cm–2) and time and method of infection [7]. Stephan et al. [8] revealed that the infection in wheat fields recorded in most of Iraq’s provinces ranged between 22.9% and 75%, while Ami et al. [9] also reported that the infection rate in wheat in northern Iraq reached 50%.

The ear-cockle disease reduces not only human consumption but also the marketing price of wheat. Protein and gluten contents of the flour of infected wheat with seed galls will diminish as well [10].

Chemical control is an effective method to control the ear-cockle disease. A wide range of nematicides is used to control this disease. However, nematicides’ high toxicity, high hazard, and long persistence in the environment besides appearing resistant...
strains for this kind of pesticides make it a real challenge to control this pest [11]. Thus, many researchers have attempted to test new compounds to control this pest. These attempts include the possibility of using some insecticide and fungicide (either chemical or organic) in addition to plant extracts to control this pest and overcome appearing resistant strains besides long-term residue [12,13].

Based on few studies conducted in Iraq to reduce the effect of ear-cockle disease of *A. tritici*, [14] used chitin synthesis inhibitor cyromazine (Trigard) in addition to [15] that used fungicides Dividend 030FS, Vitavax 200WP, Dithane S-60D, and herbicide Granstar 75DF. Due to the massive effect of this disease on wheat, causing a reduction in the yield, and in order to manage such challenge, the study aimed to evaluate the efficiency of some bioagent of *Trichoderma* spp. in addition to some plant extracts for use as alternative substances to control seed gall nematode *A. tritici* under laboratory and field conditions.

2. MATERIALS AND METHODS

Galls of *A. tritici* were collected from infested samples of wheat (Ibaa-99) obtained from the Department of Seed Test and Certification and Center of Receiving Wheat Grain in Wasit Province, Iraq. *Anguina tritici* J1 suspension was prepared by immersing galls in distilled water for 2 hours. J1 suspension was transferred to a beaker and the nematode population was counted under a stereomicroscope using slid count nematodes. 1 ml of J1 suspension was prepared to obtain 50 ± 5 J1 ml−1.

The study was conducted in the laboratory and field of the College of Agriculture, Wasit University, during the 2017–2018 agricultural season. The bioassay was conducted in the following ways.

2.1. Lab-Bioassay

First, in vitro as lab-bioassay, by preparing 1 ml of the J1 suspension in a 50 mm Petri dish and 1 ml of substances, the five treatments (Table 1) were added to a Petri dish separately with the recommended concentration in triplicate. The control treatment contained J1 with distilled water only. All Petri dishes were covered with plastic bags and incubated at 26°C ± 2°C for 1, 3, 5, 7, 9, and 11 days, respectively. This experiment was repeated to confirm the results.

The effects of treatment were evaluated by the J1 moving test during the incubation period. A sample of dead J1 was differentiated based on the method in Ref. [16]: (1) straightness of juveniles and color alternation to brown and (2) making sure that juveniles are not moving when transferred to the water for 2–3 hours. The mortality percentage J1 was calculated during the period of incubation and corrected according to the following equation [17]

Corrected mortality (%) = \( \frac{\text{number of living juveniles in the treatment} - 100}{\text{number of living juveniles in the control treatment}} \) (1)

2.2. Field-Bioassay

Field-bioassay was conducted by preparing plastic pots (25 × 25 × 25 cm) filled with soil infestation by 10 galls/pot and sowing five seeds of wheat (Ibaa-99) per pot after treatment with the five treatments separately (Table 1) with three replicates. The pots were also sprayed with all treatments at leaf age of two of planting. The control treatments were sprayed with distilled water only. Pots were placed under field conditions and irrigated when needed.

The efficiency of treatments was evaluated, after harvesting and collecting wheat’s spike, according to wheat growth, yield, and infection attributes: germination percentage, number of branches/plants, plant length (cm), spike length (cm), spike weight (g), number of seed spike⁻¹, and number of galls spike⁻¹. The infection percentages were calculated according to the following equation:

Infection percentage (%) = \( \frac{\text{number of infected plants in pot}}{\text{total number of plants in pots}} \times 100 \) (2)

2.3. Statistical Analysis

The study was conducted in a factorial complete randomized design. The data were analyzed using Genstat statistical software and Microsoft Excel. Means were compared using Least significant differences (L.S.D) at the probability level of \( p \leq 0.01 \) for laboratory experiment and \( p \leq 0.05 \) for pot experiment. The Regression analysis is used to determine the relationship between treatments and accumulated mortality percentage during incubation periods [18].

3. RESULTS AND DISCUSSION

The results of the lab-bioassay indicate that all treatments caused J1 mortality of *A. tritici* particularity Biocont-T with 40.56% as compared to other treatments (Table 2). The biopesticide Biocont-T had better efficiency than isolate T-113 in average accumulated J1 mortality percentage of %40.56–9.01%, respectively. The results of the plant extract also showed that Nematic was more efficient on accumulated J1 mortality than licorice extract (12.52% and 8.22%, respectively) compared to control treatment (0.33%).

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Formulation</th>
<th>Compound</th>
<th>Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Biocont-T (biopesticide)</td>
<td>WP</td>
<td><em>Trichoderma harzianum</em></td>
<td>19 × 10⁶ spores ml⁻¹</td>
</tr>
<tr>
<td>T-113</td>
<td>Spore suspension</td>
<td><em>Trichoderma hamatum</em></td>
<td>19 × 10⁶ spores ml⁻¹</td>
</tr>
<tr>
<td>Bio Atlantis Nematic</td>
<td>EC</td>
<td>Seaweed extract</td>
<td>2.5 ml l⁻¹</td>
</tr>
<tr>
<td>Licorice extract</td>
<td>WP</td>
<td>Licorice root extract</td>
<td>1 g l⁻¹</td>
</tr>
<tr>
<td>Rugby 100 (negative control)</td>
<td>ME</td>
<td>Cadusafos</td>
<td>1 ml l⁻¹</td>
</tr>
</tbody>
</table>
The J₂ mortality percentage of *A. tritici* in all treatments during the incubation periods’ 1st to 11th days has shown a significant effect (Table 2). The means of the accumulated J₂ mortality percentage demonstrated that gradual increase was highest on the 11th day with 39.01%. The interaction between treatments and incubation periods revealed a significant effect, while Biocont-T was more toxic to J₂ and reached maximum mortality of 92.8% on the 11th day of incubation compared to other treatments, especially nematicide Rugby with 88.05%.

Regression analysis results showed a positive relation between all treatments and J₂ mortality during incubation periods. Y demonstrated that incubation periods at 26°C ± 2°C will increase the percentage of J₂ mortality, while linear relation for treatments (Fig. 1) referred to a significant relation between the accumulated J₂ mortality and incubation period. This was shown by the positive value of factor $R^2$ which reached the highest value in treatments T-113 and Biocont-T with 0.987 and 0.954, respectively, and without any significant differences from treatment with nematic with 0.926, in addition to licorice extract.

The results of the field-bioassay in the pots revealed that treatments had a significant effect on the growth and yield of wheat plants that were inoculated with galls (Table 3). It was found that the Biocont-T showed a significant effect on the J₁ mortality reflex on the growth of wheat attributes such as germination percentage, number of plant branches, plant length, spike length, and seed weight and number of seeds in each spike of 60.1%, 2.3 branch plant⁻¹, 66.8 cm, 8.3 cm, 0.73 g, and 36.5 seed spike⁻¹, respectively. Meanwhile, there was limited improvement in germination percentage as a result of using *Trichoderma* in both treatments with Biocont-T and isolate T-113 with 60.1% and 60.5%, respectively.

The results of treatment effect on wheat infection with *A. tritici* in plastic pots (Table 4) revealed that it has a significant decrease of infection percentage of wheat plant with J₁ compared to the control treatment. Biocont-T has the highest effect by decreasing the infection percentage 77.78% and then T-113 with 80.55%. The other treatments have variable effects compared to the control which infected 100%.

The infection percentage results were reflected on average galls per spike. This has shown that all plants whose seeds were treated with Biocont-T contain the lowest number of galls in their spikes of 1.36 (Table 4). Nematic was also shown to be effective with 1.93 galls per spike. The other treatments had variable effects compared to the control treatments which were 5.66 galls per spike.

Many researchers pointed out that *Trichoderma* is used to inhibit the growth of plant-parasitic nematodes. The secondary metabolism of *Trichoderma* includes secretion of chitinase, which is an effective component against many pathogenic microorganisms; furthermore, chitin is one of the cell wall components of nematode eggs. El-Hasan et al. [19] reported that *Trichoderma* species also produced some chemicals, harzianic acid, alamethicins, tricholin, peptaibols, antibiotics, and enzymes, which may oppose nematode feeding.

The biopesticide Biocont-T has been distinguished by an accumulated effect on J₁ of *A. tritici*. It was also pointed out by Pandey et al. [20] that *Trichoderma* provide essential nutrients to plants in addition to their nematode-inhibiting ability and possessing a promising potential of nematode antagonism as well. They release some toxic substances/metabolites against a number of pathogenic fungi and nematodes [21,22].

Ozberk et al. [23] reported that grain yield losses due to *A. tritici* were reported to be up to 32%. Seed gall also reduced the number of grains spike⁻¹, grain weight spike⁻¹, test weight (kg hour⁻¹), and 1,000-kernel weight (g) but had no effect on sodium dodecyl sulfate (SDS) protein content (%). Mohamedova et al. [24] reported that there was significant decrease in yield between varieties of wheat caused by nematode *A. tritici*.

Some other studies showed that certain plant parts and extracts possess nematocidal properties; this may be due to the contents of these extracts such as saccharides, proteins, growth regulators, and nutrients. Musa et al. [25] reported that licorice extract contains saccharides, protein, and nutrients (P, K, Cu, Mg, Mn, Fe, and Zn). Al-Jawary [26] also reported that licorice extract is similar to gibberellin Gibberellic acid 3 (GA₃) because it contains mevalonic acid, the bioprimer for GA3, which is responsible for cell division and elongation.

Application of plant parts or extracts to nematode-infested soil can affect nematodes directly and stimulates soil microbes that may
Figure 1: Linear relation between treatments and J₂ mortality of *A. tritici* during incubation.

Table 3: Effect of treatments on wheat inoculated with galls of *A. tritici*.

<table>
<thead>
<tr>
<th>Attributes Treatment</th>
<th>Germination %</th>
<th>No. of branches plant⁻¹</th>
<th>Plant length (cm)</th>
<th>Spike length (cm)</th>
<th>Spike weight (g)</th>
<th>No. of seeds spike⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Biocont-T</td>
<td>60.1</td>
<td>3.2</td>
<td>66.8</td>
<td>8.3</td>
<td>0.73</td>
<td>36.5</td>
</tr>
<tr>
<td>T-113</td>
<td>60.5</td>
<td>3.0</td>
<td>54.5</td>
<td>7.8</td>
<td>0.65</td>
<td>23.5</td>
</tr>
<tr>
<td>Nematic</td>
<td>59.0</td>
<td>3.1</td>
<td>52.2</td>
<td>7.8</td>
<td>0.56</td>
<td>18.7</td>
</tr>
<tr>
<td>Licorice extract</td>
<td>66.6</td>
<td>2.0</td>
<td>56.5</td>
<td>7.2</td>
<td>0.61</td>
<td>20.2</td>
</tr>
<tr>
<td>Rugby</td>
<td>63.0</td>
<td>2.7</td>
<td>53.0</td>
<td>7.9</td>
<td>0.70</td>
<td>26.7</td>
</tr>
<tr>
<td>Control</td>
<td>45.7</td>
<td>1.8</td>
<td>45.7</td>
<td>6.9</td>
<td>0.49</td>
<td>22.8</td>
</tr>
<tr>
<td>L.S.D. 5%</td>
<td>25.16</td>
<td>0.48</td>
<td>8.08</td>
<td>0.71</td>
<td>0.14</td>
<td>11.2</td>
</tr>
</tbody>
</table>
reduce nematode population. Under this context, the use of plant extracts with nematicidals can prove to be an effective, safer, and cheaper control measure [27].

4. CONCLUSION
Ear-cockle disease causes great loss of wheat crop. The present study was carried out to evaluate the interactive effect of Trichoderma spp. and plant extracts to be used as alternative substances to control the nematode. It was revealed that biopesticide Biocont-T has proved to be effective in suppressing the J1 of A. tritici with a smaller number of galls per spike. This study will be helpful for studying new bioagents and plant extracts as alternatives to expensive and harmful chemical nematicides against nematodes in a long-term control program in a wheat plant.

5. ACKNOWLEDGMENT
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6. AUTHOR CONTRIBUTIONS
All authors made substantial contributions to conception and design, acquisition of data, and analysis and interpretation of data; took part in drafting the article or revising it critically for important intellectual content; agreed to submit to the current journal; gave final approval of the version to be published; and agree to be accountable for all aspects of the work. All the authors are eligible to be an author as per the international committee of medical journal editors (ICMJE) requirements/guidelines.

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This study does not involve experiments on animals or human subjects.

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REFERENCES

Table 4: Effect of treatment on wheat infection with A. tritici.

<table>
<thead>
<tr>
<th>Attributes</th>
<th>Treatment</th>
<th>Infection %</th>
<th>No. of galls spike-1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Biocont-T</td>
<td>77.78</td>
<td>1.36</td>
<td></td>
</tr>
<tr>
<td>T-113</td>
<td>80.55</td>
<td>3.37</td>
<td></td>
</tr>
<tr>
<td>Nematic</td>
<td>82.89</td>
<td>1.93</td>
<td></td>
</tr>
<tr>
<td>Licorice extract</td>
<td>89.22</td>
<td>4.80</td>
<td></td>
</tr>
<tr>
<td>Rugby</td>
<td>45.0</td>
<td>0.93</td>
<td></td>
</tr>
<tr>
<td>Control</td>
<td>100.00</td>
<td>5.66</td>
<td></td>
</tr>
<tr>
<td>L.S.D. 5%</td>
<td>28.11</td>
<td>0.83</td>
<td></td>
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</table>

L.S.D. 5%


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