## EFFECT OF *TRICHODERMA* ISOLATES, DELIVERY SYSTEMS AND HOST GENOTYPE ON BIOLOGICAL CONTROL OF COTTON SEEDLINGS DISEASE

Asran-Amal Abdel-Mongy\*

Agricultural Research Center, Plant Pathology Research Institute Cotton Diseases Section, Giza, Egypt

Accepted: September 17, 2007

**Abstract:** Six isolates of *Trichoderma* spp. (belonging to species; *Trichoderma harzianum* and *T. longibrachiatum*) were applied as seed or soil treatments to suppress damping-off of seedlings of ten cotton cultivars under greenhouse conditions. In most cases, cultivar x isolate interaction was a highly significant (p < 0.01) source of variation in the tested seedling growth parameters: incidence of disease, seedling height, and seedling dry weight. This interaction implies that a single isolate of *Trichoderma* can be highly effective in controlling the disease on a cotton cultivar but may have minimal efficiency in controlling the disease on another cultivar. It was also found that, in most cases, cultivar x isolate x application method was a highly significant source of variation (p < 0.01) in the tested growth parameters. Cotton cultivars showed differences in the disease reaction to the biocontrol agents. In the experiments evaluating the *Trichoderma* antagonists and their effect on seedling disease, a highly significant (p < 0.01) experimental treatment interaction was found. This interaction suggests that the outcome of cultivar x isolate interaction is markedly affected by the application method. Thus, the application method should be chosen to maximize the outcome of this interaction. The degree of the control of seedling disease in cotton differed according to the isolates of antagonists, the application method and cultivars.

Key words: seed coating, seedling emergence, Trichoderma, cotton, growth parameters

## **INTRODUCTION**

Cotton (*Gossypium* spp.) is the most important fiber crop, grown world-wide in over 80 countries. It is the first crop in terms of economic value in Egypt. Cotton seedling disease is one of the most serious diseases in all cotton-producing areas in Egypt. The soilborne fungi most often isolated from diseased cotton seedlings in-

<sup>\*</sup>Corresponding author:

asran.amal@gmail.com

clude Fusarium, Rhizoctonia, Macrophomina, and Pythium and inflict a significant financial loss for cotton producers (Youssef and Mankarios 1974; Moubasher et al. 1984; Omar 1999; Howell et al. 2002). These pathogens are capable of colonizing seed within hours from planting and can lead to the following effects: (i) seed decay before germination, (ii) pre-emergence damping-off, (iii) post-emergence damping-off, or (iv) generalized blight (Hillocks 1992; Bell 1999). Due to environmental concerns, there is a considerable interest in finding alternatives to chemical pesticides for suppression of soilborne plant pathogens (Larkin et al. 1998). Identification and selection of effective antagonistic organisms is the first and foremost step in biological control (Kamalakannan et al. 2004). Antagonism by Trichoderma species to various fungi has been well documented (Harman et al. 1989, 2004; Kubicek and Harman 1998; Elad 2000; McBeath et al. 2001; Batta 2004). Biocontrol with beneficial microorganisms seems to be a promising approach to managing cotton seedling damping-off (Howell 1982; Howell et al. 1997; Howell and Puckhaber 2005). A number of Trichoderma isolates collected from the cotton rhizosphere were effective in suppressing seedling disease on cotton under greenhouse conditions (Asran-Amal et al. 2005).

Several factors affect the ability of *Trichoderma* spp. to provide systemic disease control (Hoitink *et al.* 2006). Abiotic and biotic environmental parameters may have a negative influence on the biocontrol efficacy of *Trichoderma* strains, therefore it is very important to collect information about the effects of environmental factors on different activities of *Trichoderma* strains with biocontrol potential (Kredics *et al.* 2003).

Additionally, as different isolates of fungal biocontrol agents are known to vary in biocontrol efficacy, mode of action, and physiology, it is important to determine whether isolates of fungal antagonists all respond similarly to changes in the environment and, consequently, help in the selection of isolates most suitable for mass production (McQuilken *et al.* 1997).

The purpose of this study was to evaluate the effects of delivery methods of antagonistic, *Trichoderma* isolates, and host cultivar on the efficacy of biological control of cotton seedling disease.

## MATERIALS AND METHODS

#### **Fungal isolates**

Pathogenic and antagonistic isolates used in this study are listed in Table 1. Isolates of pathogenic soil-borne fungi were isolated from roots of cotton seedlings with damping-off disease symptoms and collected from cotton-growing areas. Isolation of *Trichoderma* spp. was made on potato dextrose agar (PDA) from rhizosphere of healthy cotton grown in agro-climatically different locations. Monosporic cultures were made and stored on PDA slants for further use.

#### Production of pathogen inoculum

Inoculum of the pathogens was prepared by wetting 40g sorghum seeds with 50 ml water, autoclaving at 15 psi for 30 min, infesting with seed-borne pathogens, and incubating at 25°C for 2 weeks. Inoculum was air-dried and stored in a paper bag at 25 to 27°C in the laboratory. Inoculum level for each of the tested isolates was 50 g of fungus-sorghum mixture/kg of soil.

Isolate code	Organism	Characteristics	Geographic origin	Source
Fo	Fusarium oxysporum	cotton root-borne fungi	Beheria	Cotton Disease Depart.
Fs	F. solani	cotton root-borne fungi	Dagahliya	Cotton Disease Depart.
Rs	Rhizoctonia solani	cotton root-borne fungi	Fayium	Cotton Disease Depart.
Мр	Macrophomina phaseolina	cotton root-borne fungi	Gharbiya	Cotton Disease Depart.
Sr	Sclertium rolfsii	cotton root-borne fungi	Beheria	Cotton Disease Depart.
Pu	Pythium ultimum	cotton root-borne fungi	Daqahliya	Cotton Disease Depart.
T1	Trichoderma harzianum	biocontrol agent	Gharbiya	Asran-Amal et al. 2005
T2	T. harzianum	biocontrol agent	Gharbiya	Asran-Amal et al. 2005
T3	T. harzianum	biocontrol agent	Daqahliya	Asran-Amal et al. 2005
T4	T. longibrachiatum	biocontrol agent	Minufiya	Asran-Amal et al. 2005
T5	T. longibrachiatum	biocontrol agent	Minufiya	Asran-Amal et al. 2005
T6	T. longibrachiatum	biocontrol agent	Giza	Asran-Amal et al. 2005

Table 1. Description, characteristics, and sources of biocontrol and pathogen isolates used in the present study

#### Production of antagonist inoculum

The six fungal antagonists were grown on molasses yeast medium (Papavizas et al. 1984) by liquid fermentation for 14 days, and formulated by mixing 200 ml of fermentor broth with 500 g of autoclaved talc powder. Five grams of carboxymethyl cellulose (CMC) was added as a sticker to the powder after air-drying and the final dried formulation had a moisture level of 11%.

#### Application methods of antagonists

#### Seed treatment

In the first series of tests, 10g of cotton seeds (10 cultivars) were mixed with 4.0ml of an aqueous (11%) pelgel (Lipha Tech) solution as a sticker. Ten grams of seeds were mixed with 6 ml of sticker and 1.3g of each powdered biomass for each fungal isolates. The seeds, sticker, and fungal biomass were mixed thoroughly; the seeds were covered with plastic sheets and stored at  $4^{\circ}$ C for no more than 7 days before planting. Seeds contained >10<sup>7</sup> CFU/1g seed for each fungal antagonist.

#### Soil amendment

*Trichoderma* spp. were cultured in sorghum as described by Budge and Whipps (1991). To prepare inoculum for soil treatment, mixtures comprising 2 liters of flaked maize and perlite (15% v/v)and 200 ml tap water in bags were autoclaved twice for 15 min and then inoculated with 100 ml of a suspension of  $10^6$  spores/ml in distilled water. The bags were incubated at  $25^{\circ}$ C for 3 weeks. The bags were shaken periodically to distribute the mycelium evenly. The concentration of inoculum of biocontrol fungi was  $10^7$  colony forming units (CFU) per cm<sup>3</sup> maize colonized with *Trichoderma*. It was mixed with the soil at the rate of 50g/kg soil.

#### Greenhouse assay for biocontrol activity against cotton seedling disease

The antagonistic capacities of the *Trichoderma* spp. isolates against the pathogen, mixtures of six cotton soil-borne fungi were determined. The autoclaved soil was infested with mixture of the tested fungi *F. solani, F. oxysporum, R. solani, M. phaseolina, S. rolfsii* and *P. ultimum* to obtain final concentration of 30, 30, 0.5, 30, 3, and 5 g/kg soil, respectively. Six antagonist isolates were selected, and evaluated for their efficiency in controlling cotton seedling disease on ten commercial cotton cultivars Giza 91 (V1), Giza 89 (V2), Giza 83 (V3), Giza 90 (V4), Giza 85 (V5), Giza 45 (V6), Giza 70 (V7), Giza 80 (V8), Giza 86 (V9), and Giza 88 (V10). These cultivars were selected because they represent important cotton cultivars grown in Egypt. Each experimental unit consisted of pots (15 cm x 20 cm depth) with 10 seeds per pot. Soil treated with fungal pathogens without antagonists was used as control (C1 or positive control). In addition, autoclaved soil treated with CMC was used as control (C2 or negative control).

A completely randomized design with five replications (pots) was used. Irrigation was provided daily. Survival, plant height (cm), and dry weight (mg/plant) were recorded two months after sowing. The temperature regime during cotton-growing period ranged from 23±2 to 38±2.5°C.

#### Data analysis

All data were analyzed using analysis of variance (ANOVA). Least significant difference (LSD) was used to compare treatment means. Statistical computations were performed using the statistical package STATISTICA 6 (StatSoft Inc., Tulsa, OK, USA).

## RESULTS

ANOVA of Table 2 showed that cultivar, isolate, and cultivar x application method interaction were all significant or highly significant sources of variation in disease incidence in 2004 and 2005. The application method was a non-significant source of variation in 2004, while it was a significant source of variation in 2005. Cultivar x isolate interaction was a highly significant source of variation in 2004 and non-significant source of variation in 2005. Isolate x application method was a non-significant source of variation each year. The second order interaction of cultivar x isolate x application method was a highly significant source of variation in 2004 and non-significant source of variation in 2004 and non-significant source of variation method was a highly significant source of variation in 2004 and non-significant source of variation in 2005.

Table 2. Analysis of variance of the effect of cotton cultivar, *Trichoderma* isolate, method of application of *Trichoderma* isolate, and their interactions on incidence of cotton seedling disease under greenhouse conditions

Year and source of variation	D.F.	M.S.	F. value
	2004		
Cultivar (V)	9	4551.6113	20.2632*
Method (M)	1	40.4800	0.1802
Isolate (T)	7	36359.7148	161.8685 **
V x M	9	1363.5577	6.0704 **
V x T	63	450.7857	2.0068**
МхТ	7	126.7886	0.5644
V x M x T	63	350.7935	1.5617**
	2005		
Cultivar (V)	9	3226.1111	17.0412**
Method (M)	1	2414.990	12.7566**
Isolate (T)	7	46686.5352	246.6110**
V x M	9	607.9177	3.2112**
V x T	63	200.9206	
МхТ	7	260.2871	
V x M x T	63	191.4560	

D.F. - degrees of freedom

M.S. - mean square

\*\* highly significant value

Relative contribution of cultivar, isolate, application method, and their interactions with the variation in disease incidence (Fig. 1) revealed that *Trichoderma* isolates were the most vital source of variation in disease incidence as they accounted for 70.9 and 83.7% of the explained (model) variation in 2004 and 2005, respectively.

A significant of cultivar x isolate x application method interaction in 2004 implies that the cultivar x isolate interaction was clearly affected by isolate T2 (Table 3) which reduced disease incidence on cultivar V2 by 78.3% relative to pathogen-infested control when it was applied as seed treatment (AM1). However, when this isolates was applied as soil treatment (AM2) its efficiency in controlling the disease was decrease to 38.1% on the same cultivar. Another example was isolate T4, which significantly reduced disease incidence by 81.8% on cultivar V6 when it was applied as seed treatment. Nevertheless its efficiency was reduced to 55% on the same cultivar when it was applied as soil treatment. A very highly significant interaction of cultivar x application method in 2005 indicates that cultivars responded differently to the two application methods. Least significant difference (LSD) was used to compare he effectiveness of the two application methods used in cultivars (Table 4).

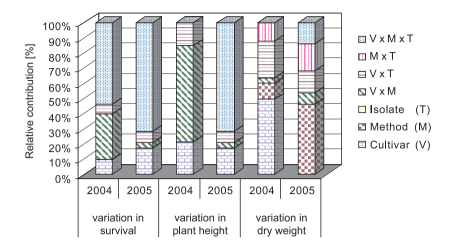


Fig. 1. Relative contribution of the effect of cotton cultivar, *Trichoderma* isolate, method of application of *Trichoderma* isolates, and their interaction to variation in survival, plant height, and dry weight of cotton seedlings under greenhouse conditions

These comparisons showed that the interaction between cultivars and application methods was due to changes in the magnitude of the differences between the application methods within cultivars. For instance, the difference in disease incidence between the two application methods was highly significant on cultivar V1, while it was non-significant in case of cultivar V4. The lack of significance isolate x cultivar and the isolate application method in 2005 (Table 2) indicates that isolate efficiency in controlling the disease was not affected by cultivar nor the application method.

Therefore, LSD was used to compare between the general means of the isolates and this comparison showed that isolate T4 was the most effective isolate in reducing disease incidence since it reduced it by 60.8% (Table 5) relative to the pathogen infested control.

Seedling height was significantly affected by all the sources of variation each year (Table 5). Cultivar was the first in importance as a source of variation in seedling height every year (Fig. 1). Isolates showed almost the same relative contribution to variation in seedling height each year; however, it was the third and the second in importance in 2004 and 2005, respectively. Application method was the second and third in importance in 2004 and 2005, respectively.

The significance of cultivar x isolate x application method interaction implies that the effect of cultivar x isolate interaction on seedling height was markedly affected by the application method. For example, isolate T4 (Table 6) significantly increased seedling height of cultivar V2 by 23.59% when it was applied as seed treatment, while height of seedlings of the same cultivar was increased by 57.05% when the isolate was applied as soil treatment.

An additional example was isolate T1, which was ineffective in increasing seedling height of cultivar V6 when it was applied as seed treatment. However, the isolate increased seedling height of the same cultivar by 32.20% when it was applied as soil treatment.

Appli- cation	Isolate <sup>b</sup>					Culti	varsc					Mean
methodª	Isolate	V1	V2	V3	V4	V5	V6	V7	V8	V9	V10	Wiean
	T1	36	40	47	30	48	26	44	48	34	46	40.1
	T2	40	20	52	56	50	28	38	48	48	36	41.6
	Т3	26	30	54	46	26	32	40	52	38	46	39.0
AM1	T4	52	24	46	32	46	16	50	54	48	32	40.0
AWI	Т5	46	22	62	30	32	34	26	58	32	60	41.2
	Т6	42	32	36	48	40	22	26	38	56	48	38.8
	C1	84	82	90	88	94	88	90	84	92	82	88.4
	C2	20	20	30	18	24	22	14	32	28	28	23.6
	mean	43.2	33.7	52.1	43.5	45	33.5	41	51.7	47.2	47.2	43.9
	T1	40	68	50	60	54	22	46	42	38	38	42.2
	T2	29	48	74	52	52	24	38	40	34	24	41.9
	Т3	45	44	74	58	38	40	52	38	46	18	46.5
AM2	T4	34	68	64	56	46	36	32	36	40	18	39.4
AIVIZ	Т5	42	72	64	48	44	30	24	46	46	32	44.4
	Т6	36	58	60	42	50	24	24	44	40	18	39.8
	C1	86	16	90	94	90	80	88	96	96	90	89.4
	C2	18	82	30	16	22	16	10	28	26	14	19.8
	mean	41.2	44.2	63.3	53.2	40.6	44	55.2	46.2	45.7	31.5	45.4
Overall	mean	42.2	39.6	57.8	38.4	47.2	33.7	40.4	48.9	46.4	39.3	44.7

Table 3. Effect of cotton cultivar (V), *Trichoderma* isolate (T), application method (AM) and their interaction on incidence of cotton seedling disease [%] in 2004

<sup>a</sup> application methods seed treatment (AM1) and soil treatment (AM2)

<sup>b</sup> Trichoderma isolates were T. harzianum (T1, T2, and T3) and T. longibrachiatum (T4, T5, and T6)

C1 pathogen - infested soil and C2 autoclaved soil

<sup>c</sup> cotton cultivars were Giza 91(V1), and Giza 88(V10)

LSD for cultivar x isolate x application method interaction = 24.42 (p < 0.01) or 18.58 (p < 0.05)

Table 4. Effect of cotton cultivar (V), *Trichoderma* isolates (T), application method (AM) and their interaction on incidence of cotton seedling disease [%] in 2005

Appli-	TICh					Culti	vars <sup>c</sup>					
cation method <sup>a</sup>	Isolate <sup>b</sup>	V1	V2	V3	V4	V5	V6	V7	V8	V9	V10	Mean
	T1	40	32	46	46	48	36	42	38	38	26	38.2
	T2	26	16	60	52	46	24	38	40	34	24	36.0
	Т3	30	12	42	46	42	40	48	38	42	38	37.8
AM1	T4	34	40	46	30	40	22	32	36	40	18	33.8
	T5	24	24	58	42	44	30	44	46	42	32	38.6
	T6	18	42	60	42	44	24	38	44	36	38	38.6
	C1	86	88	92	94	88	84	88	90	90	90	89.0
	C2	10	8	26	12	16	8	16	16	12	4	13.0
	mean	33.5	67.2	56.3	44.3	46	33.5	43.3	43.5	47.8	33.8	40.9
	T1	38	70	56	54	52	34	50	50	38	38	44.0
	T2	38	38	50	36	34	22	52	56	38	24	62.2
	Т3	44	30	62	42	38	48	40	40	32	18	47.8
AM2	T4	24	24	48	50	36	48	46	42	28	32	38.0
	T5	40	42	44	60	48	34	48	42	42	36	44.0
	T6	42	34	44	44	54	30	46	42	52	18	40.6
	C1	92	95	96	94	94	96	92	96	92	94	94.1
	C2	22	22	28	16	18	10	10	24	14	12	17.6
	mean	42.5	39.7	51	49.5	46.8	39	48.5	49	42	34	44.2
Overall	mean	38	36.2	53.7	64.9	46.4	36.7	45.9	46.3	41.9	33.9	42.6

<sup>a</sup> application methods seed treatment (AM1) and soil treatment (AM2)

<sup>b</sup> Trichoderma isolates were T. harzianum (T1, T2, and T3) and T. longibrachiatum (T4, T5, and T6)

C1 pathogen – infested soil and C2 autoclaved soil

 $^{\rm c}$  cotton cultivars were Giza 91(V1), and Giza 88(V10)

LSD for cultivar x isolate = 5.01 (p < 0.01) or 3.81 (p < 0.05)

LSD for cultivar x application method = 7.9342 (p < 0.01) or 6.03 (p < 0.05)

# Table 5. Analysis of variance of the effect of cotton cultivar, *Trichoderma* isolate, method of application of *Trichoderma* isolate, and their interaction on height of cotton seedlings under greenhouse conditions

Years and source of variation	D.F.	M.S.	F. value
	2004		
Cultivar (V)	9	590.6174	218.5318**
Method (M)	1	4028.4475	1490.5488**
Isolate (T)	7	376.9914	139.4890**
V x M	9	103.1651	38.1717**
V x T	63	26.0605	9.6426**
M x T	7	41.7557	15.4499**
V x M x T	63	15.3979	5.6973**
	2005		
Cultivar (V)	9	351.1113	112.4282**
Method (M)	1	1255.7887	402.1122**
Isolate (T)	7	289.2909	92.6329**
V x M	9	190.6290	61.0407**
V x T	63	14.3367	4.5907**
M x T	7	84.8411	27.1825**
V x M x T	63	3.1230	6.0330**

D.F. - degrees of freedom

M.S. – mean square

\*\* highly significant value

Table 6. Effect of cotton cultivar (V), *Trichoderma* isolate (T), application method (AM) and their interaction on height of cotton seedlings (cm) in 2004

Appli-						Culti	vars <sup>c</sup>					
cation method <sup>a</sup>	Isolate <sup>b</sup>	V1	V2	V3	V4	V5	V6	V7	V8	V9	V10	Mean
	T1	15.64	15.34	20.75	18.58	20.6	18.54	20.6	15.3	15.04	15.62	17.6
	Т2	16.26	17.86	20.16	18.06	21.5	20.44	20.86	18.9	15.46	15.82	18.53
	Т3	18.54	19.24	20.44	18.8	23.1	19.86	22.5	18.38	15.1	16.5	17.48
AM1	T4	17.68	19.28	18.7	19.62	22.86	21.36	20.94	19.02	15.56	17.5	19.25
	T5	16.94	17.98	17.2	20.3	22.66	21.94	21.34	15.48	15.48	14.16	18.44
	T6	16.28	16.96	17.9	17.9	18.6	21.4	19.42	16.7	17.3	15.14	17.76
	C1	14.52	15.6	17.1	15.32	15.1	18.76	14.3	11.86	14.02	12.26	14.88
	C2	16.2	16.6	17.9	17.02	17.24	18.98	17.48	18.24	14.8	15.86	18.78
	mean	16.63	17.35	18.76	18.2	20.20	20.16	19.68	16.73	15.34	15.40	17.84
	T1	19	21.1	25.3	27.5	25.36	27.1	28.4	19.14	20.1	20.04	23.30
	T2	20.44	26.6	28.3	29.5	25.9	28.5	27.6	17.2	18.9	17.6	24.05
	Т3	19.5	28.66	22.9	28.4	26.4	28.9	28.1	15.72	18.9	18.52	23.6
AM2	T4	20.94	28.74	25.2	29.2	29.4	26.2	28.3	17.3	20	19.08	24.43
AWIZ	T5	21.66	27.2	27.9	22.2	26.88	29.3	27.4	17.6	18.6	18.76	23.75
	T6	20.4	26	24.9	19.2	28.6	33.4	23.9	18.9	21.2	14.5	24.5
	C1	17.4	18.3	16.74	17.4	22.16	20.5	15.6	12.28	18.3	10.2	16.88
	C2	19.3	19.74	19.74	21.4	22.8	20.5	18.28	19.7	19.96	16.2	19.76
	mean	19.83	24.54	23.89	24.35	25.93	26.8	24.69	17.23	19.49	16.92	22.36
Overal	l mean	18.23	20.94	21.32	21.27	23.06	23.48	22.18	16.98	17.4	16.16	20.1

<sup>a</sup> application methods seed treatment (AM1) and soil treatment (AM2)

<sup>b</sup> *Trichoderma* isolates were *T. harzianum* (T1, T2, and T3) and *T. longibrachiatum* (T4, T5, and T6) C1 pathogen – infested soil and C2 autoclaved soil

<sup>c</sup> cotton cultivars were Giza 91(V1), and Giza 88(V10)

LSD for cultivar x isolate x application method = 2.68 (p < 0.01) or 2.04 (p < 0.05)

Appli- cation	Isolate <sup>b</sup>					Culti	vars <sup>c</sup>					Mean
method <sup>a</sup>	Isolate	V1	V2	V3	V4	V5	V6	V7	V8	V9	V10	Mean
	T1	18.74	20.16	22.72	17	22.05	20	20.48	21.06	17.1	17.96	19.72
	T2	17.36	21.66	17.1	18.52	23.5	21.62	20.44	20.76	19.5	17.86	19.83
	Т3	20.16	20.63	19.5	18.62	23.36	21.1	17.96	20.86	20.7	16.98	19.98
AM1	T4	21.66	20.3	18.24	18.52	24.4	21.16	18.9	20.46	18.9	18.3	20.08
AWI	Т5	18.3	21	18.6	19.62	21.4	19.22	19.2	19.26	18	17	19.16
	T6	21.48	21.5	19.6	17.87	23.5	19.74	19.2	19.5	22	17.64	20.20
	C1	17.52	17.5	15.24	15.35	20.2	17.4	17.46	16.98	16.5	15.7	16.98
	C2	18.44	20.8	18.68	19.35	21.3	21.16	19.8	18.12	19.5	17.5	19.46
	mean	19.20	20.8	18.7	18.17	22.46	20.17	19.18	19.62	19.5	17.36	19.51
	T1	18.5	21.2	24.7	26.6	26.7	26.7	27.6	17.6	20.5	20.6	23.07
	T2	19.9	24.8	26.98	28.3	28.3	27.7	27.4	16.7	20.1	18.2	23.8
	Т3	19.1	27	21.6	27.4	28.2	27.3	26.5	16.38	19.4	18.68	23.15
AM2	T4	20.5	27.24	26.58	29.2	28.6	25.6	25.5	17	20.6	19.8	24.06
AWIZ	Т5	21	26.2	25.2	22.8	23.98	28.6	23.9	17.5	19.2	15.82	22.42
	T6	19.9	25.38	24.1	19.6	20.4	30.2	24.28	18.6	21.9	15	21.93
	C1	16	18	18.46	17.4	17	20.2	15.8	13.1	17.1	12	16.49
	C2	19.18	20.12	19.8	22.18	21.18	23.3	20.6	17.1	20.2	16.8	20.04
	mean	19.26	23.74	23.42	24.18	24.37	26.23	23.9	16.73	19.87	17.11	21.88
Overall	mean	19.23	22.27	21.06	21.17	23.41	23.2	21.5	18.17	19.68	17.23	20.69

Table 7. Effect of cotton cultivar (V), *Trichoderma* isolate (T), application method (AM) and their interaction on height of cotton seedlings (cm) in 2005

<sup>a</sup> application methods seed treatment (AM1) and soil treatment (AM2)

<sup>b</sup> Trichoderma isolates were T. harzianum (T1, T2, and T3) and T. longibrachiatum (T4, T5, and T6)

C1 pathogen - infested soil and C2 autoclaved soil

<sup>c</sup> cotton cultivars were Giza 91(V1), and Giza 88(V10)

LSD for cultivar x isolate x application method = 2.88 (p < 0.01) or 2.19 (p < 0.05)

#### Table 8. Analysis of variance of the effect of cotton cultivar, *Trichoderma* isolate, method of application of *Trichoderma* isolate, and their interaction on dry weight of cotton seedling under greenhouse conditions

Years and source of variation	D.F.	M.S.	F. value
	2004		<u>^</u>
Cultivar (V)	9	231480.5781	34.4385**
Method (M)	1	185199.0469	27.5529**
Isolate (T)	7	160276.8906	23.8451**
V x M	9	66445.7891	9.8854**
V x T	63	21377.0313	3.1804**
M x T	7	94026.8594	13.9888**
V x M x T	63	14735.1035	2.1922**
	2005	•	
Cultivar (V)	9	105525.8672	12.9621**
Method (M)	1	2273027.7500	279.2043**
Isolate (T)	7	277453.0625	34.0806**
V x M	9	42903.4023	5.2700**
V x T	63	12440.1826	1.5281**
M x T	7	135355.5313	16.6262**
V x M x T	63	11663.3926	1.4327**

D.F. - degrees of freedom

M.S. – mean square

\*\* highly significant value

The formerly mentioned conclusions concerning cultivar and isolate application method interaction hold true for 2005 data shown in Table 7.

Dry weight of seedlings was significantly affected by all sources of variation each year (Table 8). Cultivar was the main important source of variation each year as it accounted for 30.08 and 28.35% of the explained (model) variation in 2004 and 2005, respectively.

Cultivars and application method was the second in importance as a source of variation in 2004 as it accounted for 19.46% of the explained (model) variation. Isolate was the third and second in importance in 2004 and 2005, respectively. Application method was almost as important as application method x isolate interaction as a source of variation each year (Fig. 1).

The interaction of cultivar and isolates was significantly affected by dry weight. For example, isolate T1 (Table 9) significantly increased dry weight of V7 seedlings by 71.2% when it was applied as soil treatment. Almost the same results were obtained when the experiment was repeated in 2005 (Table 10).

Appli- cation	Isolate <sup>b</sup>	Cultivars <sup>c</sup>										
methoda	Isolate	V1	V2	V3	V4	V5	V6	V7	V8	V9	V10	Mean
	T1	212.8	296.4	347.6	297.4	355.2	254.2	317.6	233	289.4	252.4	285.6
	T2	197.4	269.8	403	332.8	393.8	221	341	238	273.6	290.4	296.08
	Т3	240.2	383.2	380.4	306.2	304.4	229.4	317.4	323.4	286.4	313.8	308.4
AM1	T4	321.4	322.2	375.8	271.1	319.2	235.6	289.4	339.4	341	276.8	309.19
AMI	T5	260.8	275.2	400.6	343.4	329.2	303	302.6	232.6	240.8	264	295.22
	T6	218.4	297	325.4	335	300	249.6	239.2	303.2	313.6	285.4	286.68
	C1	198.2	235.4	225.8	193.4	241	237.6	185.6	129.5	128.6	148.8	182.39
	C2	255.8	308.6	325	213	295	246	246.2	336.8	265.6	262.1	294.36
	mean	238.1	298.4	347.9	286.5	317.22	247.05	279.7	2791.4	278.1	274.6	294.33
	T1	306.8	283.8	481.6	483	478.4	332.4	516.6	373.4	248.4	294	379.84
	T2	249.2	474	609.6	513.8	616.8	488.8	453.8	267.8	265.2	210.8	414.9
	Т3	213.6	451.8	487.2	449.2	363.4	401	454.4	234.6	244.2	244.4	354.38
AM2	T4	312.4	523.4	480.2	587.6	412.8	363.4	400.6	370.8	289.4	268.4	400.9
AWIZ	T5	248.8	306.6	526	340.6	464	344.8	400.6	244.8	213	156.6	324.58
	T6	244.4	448.4	555.2	351.2	487.8	477.6	287	370.2	257.8	176	365.56
	C1	198	239.2	279.4	233	260.2	210.4	141	159	211.6	117.8	204.96
	C2	210.2	248.6	301.4	250.4	299.6	249.2	295.8	275.6	242	147.4	252.02
	mean	247.9	371.9	465.6	401.1	422.6	358.4	368.7	287	246.4	201.9	337.15
Overall	l mean	243	335.1	406.4	343.8	370.03	302.7	324.2	283.2	256	232.0	309.64

Table 9. Effect of cotton cultivar (V), *Trichoderma* isolate (T), application method (AM) and their interaction on dry weight of cotton seedling [mg] in 2004

<sup>a</sup> application methods seed treatment (AM1) and soil treatment (AM2)

<sup>b</sup> *Trichoderma* isolates were *T. harzianum* (T1, T2, and T3) and *T. longibrachiatum* (T4, T5, and T6)

C1 pathogen - infested soil and C2 autoclaved soil

° cotton cultivars were Giza 91(V1), and Giza 88(V10)

LSD for cultivar x isolate x application method = 133.57 (p < 0.01) or 101.63 (p < 0.05)

Table 10. Effect of cotton cultivar (V), *Trichoderma* isolate (T), application method (AM) and their interaction on dry weight of cotton seedling [mg] in 2005

Appli-	Isolate <sup>b</sup>					Cult	vars <sup>c</sup>					Mean
cation method <sup>a</sup>	Isolate	V1	V2	V3	V4	V5	V6	V7	V8	V9	V10	Mean
	T1	307.4	341.6	416.6	307.4	416.4	241.6	274.4	306.6	280	265.6	315.74
	T2	330.4	337.2	432	330.4	432	237.2	336	347.8	268.4	246.4	329.78
	Т3	448.8	336.8	324	448.8	324	336.8	301.2	316.6	241	247.4	332.54
	T4	347.8	349	400.6	347.8	400.6	249	286.8	344.6	284.4	298.6	340.90
AM1	T5	300.6	311.6	335.6	400.6	335.6	211.6	286.2	305.4	325	227.4	293.90
	T6	288.8	315.4	339	288.8	339	215.4	302.8	366.6	298.4	251.4	300.50
	C1	209.2	2008	173.6	209.2	213.6	245.8	167.2	203.4	183.4	177.8	204.90
	C2	294	230.6	312.4	290	212.4	230.6	293.6	319.2	230.6	240.4	284.78
	mean	315	296	341.1	317.8	329.2	246	281.5	308.4	263.9	244.2	290.8
	T1	426	462.2	507.6	451.8	555.8	380.8	387.2	509	372.6	349	440.2
	T2	364	502.6	483.6	562.4	509.8	417.4	372	393.2	445.2	313.2	436.34
	Т3	422.4	502.4	479.8	570.4	482.4	430	374.8	399.4	448.4	308.8	441.8
	T4	454.4	516	467.2	453.7	571	465.6	472	418.8	388.6	322.6	447.2
AM2	T5	426.4	433.8	517.6	549.2	520.4	493.6	326.8	488.2	338.6	400.4	449.5
	T6	459.6	491.2	512	428.4	540.6	374.2	347.2	400	324.4	319.4	419.0
	C1	258.4	237.4	264	249.2	291.8	217.8	210.4	252.8	202.2	206.6	239.0
	C2	357.6	251.6	383.6	347	317.8	365.8	204.8	259.4	329	262.4	307.8
	mean	361.1	424.6	451.9	451.5	473.7	393.1	336.9	390.1	356.1	310.3	394.9
Overall	l mean	338.4	360	396.6	389.6	401.4	319.55	309.2	349.25	310	277.2	345.09

<sup>a</sup> Application methods seed treatment (AM1) and soil treatment (AM2)

<sup>b</sup> *Trichoderma* isolates were *T. harzianum* (T1, T2, and T3) and *T. longibrachiatum* (T4, T5, and T6) C1 pathogen – infested soil and C2 autoclaved soil

<sup>c</sup> cotton cultivars were Giza 91(V1), and Giza 88(V10)

LSD for cultivar x isolate x application method = 147.00 (p < 0.01) or 111.85 (p < 0.05)

## DISCUSSION

Numerous attempts in biological control have resulted in second-rate disease control under varying environmental conditions and sites. This inconsistency possibly was related, at least partially, to a general lack of understanding of how these biocontrol systems work and under which conditions they may or may not be expected to function. This has resulted in the introduction of biocontrol organisms into environments in which they are ecologically unsuitable (Deacon 1991). Any individual biocontrol microorganism can only be expected to perform within a limited set of physical, biological, and environmental conditions. Hitherto, generally, these conditions are inadequately clear (Larkin and Fravel 2002).

For eco-friendly and sustainable management of the disease, 6 isolates belonging to two species of *Trichoderma* (*T. harzianum*, and *T. longibrachiatum*) were applied as seed and soil treatments to suppress damping-off of cotton seedlings on ten cotton cultivars under greenhouse conditions. On the whole, cultivar and isolate interaction was a highly significant source of variation (p < 0.01) in the tested seedling growth parameters (disease incidence, seedlings height, and seedling dry weight).

This interaction implies that a single isolate of antagonist can be vastly effective in controlling the diseases on a cotton cultivar but may have minimal efficiency in controlling the disease on another cultivar. Antagonists also varied in their efficiency as biocontrol agents, and a relative effectiveness of different antagonists varied among growing seasons (Ryan *et al.* 2004). The efficacy of biological control agents can also vary relative to each other and overall when assayed on different host cultivars (Schisler *et al.* 2000). *T. longibrachiatum* conferred varying levels of protection to the cotton seedling disease, depending on isolate, host and pathogen (Sreenivasaprasad and Manibhushanrao 1990).

The interaction also indicates that apparently many genes from both cotton and *Trichoderma* interact to regulate the number of cotton cultivars and *Trichoderma* isolates (Wells and Bell 1983). The methods of applying biocontrol agents to a target area are critical in the development of biocontrol strategies for protection against different diseases (Mao *et al.* 1997).

These findings have an important bearing on antagonism testing methods. Isolates of *Trichoderma* should be tested on as many cotton cultivars as possible, as this will improve the chance of identifying antagonist isolates effective in controlling the disease on more than a few cotton cultivars.

The interaction also suggests that it may be more prudent to evaluate blends of antagonist isolates for wild application on more cotton cultivars. In this investigation, the interaction between cotton cultivars and *Trichoderma* isolates was evaluated under greenhouse conditions favourable for the growth of both cotton cultivars and *Trichoderma* isolates.

Under field conditions, environmental conditions during the different periods of cotton growing season may be more favourable for cotton cultivars or the antagonist isolates. Thus, the findings of this work are not expected to be necessarily related to the degree of biological control that may be observed in the field, but should reflect the capacities and genetic variability of the antagonist isolates and of the various cotton cultivars to response to antagonisms (Bell *et al.* 1982).

It was also found that, in most cases, cultivar x isolate x application method was a highly significant source of variation (p < 0.01) in the tested growth parameters.

This interaction suggests that the outcome of cultivars x isolates interaction is markedly affected by the application method. Thus, application method should be chosen to maximize the outcome of the interaction.

## CONCLUSIONS

Environmental factors play an important role in restricting the activity of potential biological control agents, detailed information on the environmental requirements of *Trichoderma* are required in order to improve its efficacy and also to assist in optimizing large-scale inoculum production. The results obtained here, demonstrated that significant complex interactions occur between methods of application, cultivars and antagonistic isolates. Further tests also are required to improve our understanding for this complex interaction.

#### ACKNOWLEDGEMENTS

Guidance of Prof. A. A. Aly, cotton disease sections, Plant Pathology Research Institute, in planning and conducting the research is also duly acknowledged. Thanks are also due to Dr. K.A. Abd-Elsalam for his valuable suggestions in the manuscript.

#### REFERENCES

- Asran-Amal A., Abd-Elsalam K.A., Omar M.R., Aly A.A. 2005. Antagonistic potential of *Trichoderma* spp. against *Rhizoctonia solani* and use of M13 Minisatellite-primed PCR to evaluate of the antagonist genetic variation. J. Plant Dis. Prot. 112: 550–561.
- Batta Y.A. 2004. Postharvest biological control of apple gray mold by *Trichoderma harzianum* Rifai formulated in an invert emulsion. Crop Prot. 23: 19–26.
- Bell A.A. 1999. Diseases of cotton. p. 553–560. In "Cotton: Origin, History, Technology, and Production (C.W. Smith, J.T. Cothren, eds.). Wiley, New York.
- Bell D.K., Wells H.D., Markham C.R. 1982. In vitro antagonism of *Trichoderma* species against six fungal plant pathogens. Phytopathology 72: 379–382.
- Budge S.P., Whipps J.M. 1991. Glasshouse trials of *Coniothyrium minitans* and *Trichoderma* species for the biological control of *Sclerotinia sclerotiorum* in celery and lettuce. Pl. Pathol. 40: 59–66.
- Deacon J.W. 1991. Significance of ecology in the development of biocontrol agents against soil-borne pathogens. Biocontrol Sci. Technol. 1:5–20.
- Elad Y. 2000. Biological control of foliar pathogens by means of *Trichoderma harzianum* and potential modes of action. Crop Prot. 19: 709–714.
- Harman G.E., Backman, P.A., Turner J.T., Fillatti T., Mccall C., Comai I., Kiser J., Mcbride J., Stalker D., Foudin A.S., Shantharam S. 1989. Symposium: biocontrol and biotechnological methods for controlling cotton pests. p. 15–20. In: "Proceedings of the Beltwide Cotton Production Research Conferences" (J.M. Brown, D.A. Richter, eds.). Memphis, Tennessee, USA.
- Harman G.E. Howell C.R., Viterbo A., Chet I., Lorito M. 2004. Trichoderma species-opportunistic, avirulent plant symbionts. Nature Rev. 2: 43–56.
- Hillocks R.J. 1992. Seedling diseases. p. 1–17. In: "Cotton Diseases" (R.J. Hillocks, ed.). CAB International, Wallingford, Oxon, UK.

- Hoitink H.A.J., Madden L.V., Dorrance A.E. 2006. Systemic resistance induced by *Trichoderma* spp.: Interactions between the host, the pathogen, the biocontrol agent, and soil organic matter quality. Phytopathology 96: 186–189.
- Howell C.R. 1982. Effect of Gliocladium virens on Pythium ultimum, Rhizoctonia solani and damping-off of cotton seedlings. Phytopathology 72: 496–498.
- Howell C.R., DeVay J.E., Garber R.H., Batson W.E. 1997. Field control of cotton seedling diseases with *Trichoderma virens* in combination with fungicide seed treatments. J. Cotton Sci. 1: 15–20.
- Howell C.R. 2002. Cotton seedling preemergence damping-off incited by *Rhizopus oryzae* and *Pythium* spp. and its biological control with *Trichoderma* spp. Phytopathology 92: 177–180.
- Howell C.R., Puckhaber L.S. 2005. Study of the characteristics of "P" and "Q" strains of *Trichoderma virens* to account for differences in biological control efficacy against cotton seedling diseases. Biol. Cont. 33: 217–222.
- Kamalakannan A., Mohan L., Harish S., Radjacommare R., Amutha G., Chiara K., Karuppiah R., Mareeswari P., Rajinimala N., Angayarkanni T. 2004. Biocontrol agents induce disease resistance in *Phyllanthus niruri* Linn against damping-off disease caused by *Rhizoctonia solani*. Phytopathol. Mediterr. 43: 187–194.
- Kredics L., Antal Z., Manczinger L., Szekeres A., Kevei F., Nagy E. 2003. Influence of environmental parameters on *Trichoderma* strains with biocontrol potential. Food Technol. Biotechnol. 41: 37–42.
- Kubicek C.P., Harman, G.E. 1998. *Trichoderma* and *Gliocladium*, Basic Biology, Taxonomy and Genetics Vol. 1. Taylor and Francis, London, 278 pp.
- Larkin R.P., Roberts D.P., Gracia-Garza J.A. 1998. Biological control of fungal diseases p. 141–191. In: "Fungicidal Activity-Chemical and Biological Approaches to Plant Protection" (D. Hutson, J. Miyamoto, eds.). Wiley, New York, NY.
- Larkin R.P., Fravel D.R. 2002. Effects of varying environmental conditions on biological control of Fusarium wilt of tomato by nonpathogenic *Fusarium* spp. Phytopathology 92: 1160–1166.
- Mao W., Lewis J.A., Hebbar P.K., Lumsden R.D. 1997. Seed treatment with a fungal or a bacterial antagonist for reducing corn damping-off caused by species of *Pythium* and *Fusarium*. Plant Dis. 81: 450–454.
- McBeath J.H., Gay P.A., Yokogi T. 2001. Biological control of pink rot by *Trichoderma atroviride*. Phytopathology 91: S5 (Abstract).
- McQuilken M.P., Budge S.P., Whipps J.M. 1997. Effects of culture media and environmental factors on conidial germination, pycnidial production and hyphal extension of *Coniothyrium minitans*. Mycol. Res. 101: 11–17.
- Moubasher A.H., Mazen M.B., Abdel-Hafez A.I. 1984. Studies on the genus *Fusarium* in Egypt. Mycopathologia 85: 161–165.
- Omar M.R. 1999. Studies on susceptibility of cotton to *Macrophomina phaseolina*. M. Sc. Thesis, Al-Azhar Univ. Cairo, 139 pp.
- Papavizas G.C., Dunn M.T., Lewis J.A., Beaggle-Ristaino J. 1984. Liquid fermentation technology for experimental production of biocontrol fungi. Phytopathology 74: 1171–1175.
- Ryan A.D., Kinkel L.L., Janet L.S. 2004. Effect of pathogen isolate, potato cultivar, and antagonist strain on potato scab severity and biological control. Biocontrol Sci. Technol. 14: 301–311.
- Schisler D.A., Slininger P.J., Hanson L.E., Loria R. 2000. Potato cultivar, pathogen isolate, and antagonist cultivation medium, influence the efficacy and ranking of bacterial antagonists of *Fusarium* dry rot. Biocontrol Sci. Technol. 10: 267–279.
- Sreenivasaprasad S., Manibhushanrao K. 1990. Biocontrol potential of fungal antagonists *Gliocladium virens* and *Trichoderma longibrachiatum*. J. Plant Dis. Prot. 97: 570–579.

- Wells H.D., Bell D.K.1983. Antagonism in vitro between isolates of *Trichoderma harzianum* and *Rhizoc-tonia solani* AG4. Phytopathology 73, p. 507.
- Youssef Y.A., Mankarios A.T. 1974. Studies on the rhizosphere mycoflora of broad bean and cotton, IV. The influence of the rhizosphere fungi on plant growth. Mycopath. Mycol. Appl. 54: 173–180.

## POLISH SUMMARY

## WPŁYW IZOLATÓW TRICHODERMA, SYSTEMÓW ICH APLIKACJI I GENOTYPU ROŚLINY ŻYWICIELSKIEJ NA BIOLOGICZNE ZWALCZANIE CHOROBY SIEWEK BAWEŁNY

W celu ograniczenia zgorzeli siewek 10 odmian bawełny wykorzystano 6 izolatów Trichoderma spp., zaliczanych do gatunków T. haraianum i T. longibrachiatum. Przy ich użyciu stosowano zaprawianie nasion bawełny, lub wprowadzano je do ziemi w doświadczeniu prowadzonym w warunkach szklarniowych. W większości przypadków źródłem występującej, wysoce istotnej (p < 0,01) zmienności badanych parametrów wzrostu siewek (występowanie choroby, wysokość roślin i ich sucha masa), było współdziałanie odmiany z izolatem grzyba antagonistycznego. To współdziałanie pokazuje, że izolat Trichoderma może być wysoce efektywny w zwalczaniu choroby na jednej odmianie, ale może wykazywać minimalną efektywność zwalczania na innej odmianie. Stwierdzono także, że w większości przypadków współdziałanie: odmiana x izolat x metoda jego aplikacji było wysoce istotnym (p < 0.01) źródłem zmienności ocenianych parametrów wzrostu. Wykryto różnice w reakcji chorobowej odmian bawełny na zastosowane izolaty Trichoderma. Oceniając wpływ antagonistycznych izolatów oraz sposób ich zastosowania na zgorzel siewek bawełny stwierdzono wysoce istotne (p < 0.01) współdziałanie tych czynników. Sugeruje to, że współdziałanie: odmiana x izolat jest w dużym stopniu zależne od zastosowanej metody ich aplikacji.

Należy więc wybierać metodę pozwalającą na maksymalizację korzystnego aspektu tego współdziałania. Stopień zwalczania zgorzeli siewek bawełny różnił się zależnie od izolatu grzyba antagonistycznego, metody jego aplikacji oraz odmiany bawełny.