DETERIORATION IN COTTON FIBERS CAUSED BY SOME CELLULOSE-DEGRADING FUNGI

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ABSTRACT

Deterioration in fibers of the cotton cultivars Giza 80, Giza 85, Giza 86, Giza 89 and Giza 90 (long staple), and Giza 88 (extra long staple) caused by Alternaria sp., Fusarium semitectum, Trichothecium sp., Penicillium sp., Trichoderma sp., Rhizopus, Aspergillus flavus, Cladosporium herbarum, F. monoliforme, and Nigrospora sp. Was evaluated under pure culture conditions. The tested properties were upper half mean, uniformity index, short fiber index, Brightness, Yellowness, trash no., maturity, micronaire value, fiber strength, elongation, cellulose, reducing sugar content, and fiber damage index. Cultivars, fungi, and cultivars x fungi interaction were all very highly significant sources of variation in all the tested properties. Cultivars were the most important source of variation in most of the tested properties. Due to the significance of the cultivars x fungi interaction, a least significant difference (LSD) was used to compare the individual fungal means within cultivars for each of the tested properties. These comparisons showed that most of the tested properties tended to decline as a result of fungal infection; however, the magnitude of decline varied from one cultivar to another. The present study clearly demonstrated that cotton cultivars were much more important than fungal isolates in determining the level of deterioration in most of the tested properties. This result implies that the deleterious effects of the cellulose-degrading fungi on quality of fibers could be considerably reduced if resistant of cotton cultivars to these fungi is effectively enhanced.

INTRODUCTION

Many of the fungi associated with lint contamination are capable of producing cellulolytic enzymes in sufficient quantity to degrade cotton fibers if they are able to grow on the seedcotton for long enough. The most efficient cellulose degradets are *Alternaria* spp., *Curvularia*, *Fusarium moniliforme* and *Glomerella* species isolated from cotton bolls. Most species of *Aspergillus*, including *A. niger*, are poor cellulose degraders. It should be noted however, that there is considerable variation with respect to production of cellulolytic enzymes between species of the same genus and, indeed, among isolates of the same species (Hillocks, 1992).

Fungal microflora associated with deterioration of cotton fibers are classified into two groups: field and storage fungi. Field fungi usually invade the maturing seed cotton on the developing plants in the field before harvest of bolls. These fungi require a moisture content in equilibrium with a relative humidity of more than 90 % to grow. The storage fungi are those that grow on stored lint. Most of them are able to grow without free water, and on media with high osmotic pressure (Amer, 1986).

Under proper invironmental conditions, cellulose degrading fungi may lower substantially the quality of cotton fibers. For example, Badr (1980)

studied the effect of infection with rhizopus nigricans, Aspergillus niger and Fusarium oxysporum f.sp. vasinfectum on fiber properties of some Egyptian cotton varieties. She found that all the tested fungi affected fiber tensile strength and elongation at 1/8 inch gauge length, stiffness, and toughness causing substantial losses, which varied from one fungus to another. She also reported that all fiber damage index values of the infected fibers significantly increased than those of the control. in most cases, pH aqueous extracts of infected fibers tended to increase. A highly significant increase in reducing sugars content was also observed. Rao et al. (1989) reported that properties of cotton fibers such as length, strength, fineness, and maturity were adversely affected by high investations of whitefly and sooty mold. Abd El-Rehim et al. (1993) evaluated deterioration in cotton fibers of Giza 75 cv. Caused by Nigrospors, Aspergillus, Fusarium, Alternaria, Rhizopus. Trichothecium, Cladosporium, and Penicillium. Floating fiber index and number of neps tended to increase as a result of fungal infection, micronair reading showed no change, the other properties tended to decrease. Within each of the tested properties, except the micronair reading, the observed magnitude of change varied from one fungus to another; however, none of the tested fungi was able to affect all properties. Abd El-Rahim et al. (2001) evaluated deterioration in quality of Giza 86 cv. Caused by the sooty mold (SM) fungus Cladosporium herbacium (CH) under field conditions in 14 samples, obtained from 7 different locations in 4 governorates. They found that all fiber length parameters were highly affected by SM incidence. Micronaire reading was significantly decreased as a result of SM incidence in all locations. CH caused significant decrease in maturity ratio in all the tested locations. Hair weight was adversely affected by CH in all the tested locations. CH significantly reduced fiber strength in all the tested locations. All chemical properties significantly increased as a consequence of CH infection. However, the greatest magnitude of increase was observed in the case of fiber damage index.

In the present study, we reported on the deterioration effects of 10 cellulose-degrading fungi on physical, mechanical, and chemical properties of 6 cotton cultivars under pure cultural conditions.

MATERIALS AND METHODS

Fungal isolates

Isolates of fungi used in this study (Table 1) were isolated, purified, and identified at Cotton Pathology Lab., Plant Path. Res. Inst., Agric. Res. Cent., Giza, Egypt.

Inoculation of cotton fibers

Substrate for growth of fungal isolates was prepared in 500-ml glass bottles, each bottle contained 10 g of cotton fibers to which 20ml of tap water was added. The bottles were autoclaved for 30 minutes, and the fungal inocula, taken from one-week old cultures grown on potato dextrose agar, were aseptically introduced into the bottles and allowed to colonize fibers for 3 weeks at 26±3°C. the uninoculated controls were autoclaved for 30 minutes.

Congo red test

The Congo red test was achieved according to the method of Celegg(1940). Sample of cotton fibers (which contained about 100 fibers) . was drawn on a glass side and immersed in a solution of 11 % caustic soda for 3 min., then washed for several times with distilled water. A drop of Congo red solution (0.19% in 95% ethyl alcohol) was placed on the fibers and left for 10 min. at room temperature, followed by washing with water for several times. A drop of 18 % caustic soda solution was again placed on the fiber and left for few sec. for swelling fibers, then microscopically examined (G208 projection microscope was used according to ASTM: D 2130-1986)

and sorted into four classes according to the degree of damage as follows:

0= Non deteriorated fibers, were indicated by appearance pink color.

1= Slightly deteriorated fibers, were appeared with several celluloid spiral lines. 2=moderately deteriorated fibers, which regularly appeared with red color.

3= severally deteriorated fibers which appeared with several red colored sloughing off parts.

The fibers damage index was calculated as follows:

Total numbers of damage fibers (classes 1+2+3)
_____ X 100

Total number of tested fibers

Fiber physical properties

Micronaire value, and fiber maturity ratio were determined using Micromate instrument according to (ASTM: D3818-) While, Spain Lab 900B HVI instrument system was used to determine fiber length, fiber uniformity Index, short fiber index, fiber reflectance percentage (Rd%), fiber yellowness degree (+b), fiber strength and fiber elongation according to ASTM: D4605-1986.

Fiber chemical properties

1- Fiber cellulosic materials content:

Cellulosic materials % in cotton fiber was determined according to the methods described by Jenkis (1930).

2- Fiber reducing Sugar content:

Reducing Sugar content in cotton fiber was determined by using Soxhlet extraction according to the methods of Smith (1956).

3- Total wax content of fiber:

Total wax content was determined according to the methode described by Conrad (1944).

All tests were performed at the laboratories of Cotton Research institute, Agricultural Research Center, under constant conditions of temperature. (20 \pm 2 °c) and (65% \pm 5%) of relative humidity.

Statistical analysis of the data

A completely randomized block design with three replicates was used in the present study. The least significant difference (LSD) was applied for comparing treatment means. Analysis of variance (ANOVA) of the data was performed with the MSTAT-C statistical package.

RESULTS AND DISCUSSION

In the present study, 10 cellulose-degrading fungi (Table1) were used to inoculate fibers of 6 Egyptian commercial cotton cultivars. These cultivars were chosen because they are differing in fiber quality. Giza 88 belongs to the extra long staple category, while the remaining cultivars belong to the long staple category.

Table 1. Fungi used in the study.

No.	Fungus	Geographic origin
1	Altrernarid sp	Giza
2	Fusarium semitectum	Daqahiia
3	Trichothecium sp.	Sharqiya
4	Penicillium sp	Sohag
5	Trichoderma sp	Assiute
6	Rhizopus sp	Giza
7	Aspergillus flavus	Gharbiya
8	Cladosporium herbarum	Beheira
9	Fusarium moniliforme	Daqahliya
10	Nigrospora sp	Sharqiya

ANOVA of Table 2 showed very highly significant effects of cultivars, fungi, and cultivars × fungi interaction on fiber length parameters. Cultivars were the first in importance as a source of variation in upper half mean and short fiber index. Fungi were the most important source of variation in uniformity index.

Table 2. Analysis of variance of the effect of some cellulose-degrading fungi on fiber length parameters of six Egyptian cotton cultivars.

Cultival	<u></u>									
Parameters and source of variation	D.F	M.S	F. values	P> F						
Upper half mean (mm)										
Replications	2	0.021	0.0989							
Cultivars (C)	5_	164.649	772.9187	0.0000						
Fungi (F)	10	34.924	163.9437	0.0000						
CxF	50	4.685	21.9922	0.0000						
Error	130	0.213								
	U	niformity Index (%)							
Replications	2	5.329	7.1987							
Cultivars (C)	5	174.399	235.6027	0.0000						
Fungi (F)	10	161.841	218.6371	0.0000						
CxF	50	17.937	24.2313	0.0000						
Error	130	0.740								
		Short Fiber Index	K							
Replications	2	54.874	12.4895							
Cultivars (C)	5	2138.739	486.7829	0.0000						
Fungi (F)	10	866.824	1972916	0.0000						
CxF	50	117.409	26.7226	0.0000						
Error	130	4.394								

Cultivars and cultivars × fungi interaction were almost equally important as sources of variation in uniformity index (Table 3). The very highly significant interaction of cultivars × fungi for all the tested properties (Table 2) indicated that cultivars responded differently to fungi regardless of the tested property. Due to the significance of this interaction, a least significant difference (LSD) was used to compare the individual fungal means within cultivars for each of the tested properties.

Table 3. Relative contribution of fungi, cotton cultivar, and their interaction to variation in fiber length parameters.

Source of	Relative c	Relative contribution to variation in ^a							
variation	Upper half mean (mm)	Uniformity Index (%)	Short Fiber Index						
Cultivars (C)	58.52	25.66	42,20						
Fungi (F)	24.83	47.63	34.21						
CxF	16.65	26.39	23.15						

^{*} Calculated as percentage of sum of squares of the explained (model) Variation.

These comparisons showed that the upper half mean of all cultivars significantly decreased as a result of fungal infection (Table 4); however, the upper half mean of Giza 86 and Giza 88 were notable exceptions because they were resistant to infection with *A. flavus*. Uniformity index of all cultivars significantly increased by all fungi. Short fiber indexes of Giza 80 and Giza 88 were not affected by *A. flavus* and *F. semitectum* respectively. Also, Giza 89 was not affected by *Penicillium* sp., *Trichoderma* sp., and *A. flavus* short fiber index of all the remaining cultivars significantly increased due to fungal infection. The observed decline in the fiber length parameters could be attributed to the deleterious effects of cellulolytic enzymes produced by the fungi. These enzymes weaken the fibers, which become more susceptible to breakage during testing the fiber length. Consequently, the short fiber content increases. These results are in harmony with those of Abd El-Rehim *et al.* (1993) and Mahmoud (1996).

Cultivars, fungi, and cultivars × fungi were all very highly significant sources of variation in brightness, yellowness, and trash no. (Table 5). Cultivars were the most important source of variation in brightness and trash no., while isolates were the most important source of variation in yellowness (Table 6). Brightness of Giza 80, Giza 85, Giza 86, Giza 88 and Giza 90 was significantly reduced by all the fungi (Table 7). However, brightness of Giza 89 showed variable responses to fungal infection. Thus, *F. semitectum, Penicillium* sp., *Trichoderma* sp., *Nigrospors* sp., significantly increased it, while *Alternaria* sp., *Trichothecium* sp., *Rhizopus* sp., *Cladosporium herbarium* and *F. moniliforme* had no effect. On the other hand, *A. flavus* was the only fungus, which reduced it. *Trichoderma* sp., did not effect yellowness of Giza 80, while the remaining fungi sifnificantly increased it. yellowness of Giza 85, Giza 86, Giza 88 and Giza 90 increased by all the tested fungi. *Trichothecium* sp., *Cladosporium herbarium* and *F. moniliforme* did not affect yellowness of Giza 89, while other fungi decreased it..

Table 4. Effects of some cellulose-degrading fungi on fiber length parameters of six Egyptian cotton cultivars.

•									Cu	itivars								
Fungus *	Upper half mean (mm)							Uniformity Index (%)				Short Fiber Index						
	G80	G85	G86	G88	G89	G90	G80	G85	G86	G88	G89	G90	G80	G85	G86	G88	G89	G90
1	24.45	24.30	28.35	30.20	26.63	22.20	70.45	74.90	77.35	76.20	77.40	69.60	38.83	40.57	29.20	26.30	27.50	50.4
2	26.60	24.70	29.05	31.50	27.43	23.33	75.50	74.63	78.93	81.20	77.50	69.07	32.20	36.80	23.53	14.43	27.13	51.9
3	26.25	23.90	27.03	25.50	24.33	22.50	76.65	72.67	75.42	73.42	70.73	69.80	32.27	45.90	33.10	36.40	42.77	50.6
4	28.20	23.80	28.65	28.73	28.27	21.67	79.93	73.10	75.80	77.20	82.07	72.27	25.30	44.07	26.20	20.57	19.23	47.7
5	24.45	24.57	29.40	26.88	29.13	22.67	70.23	73.13	77.50	76.97	80.20	69.93	44.37	41.53	25.20	22.63	18.77	50.4
6	24.45	25.53	29.80	28.50	25.33	23.10	70.67	74.77	77.70	74.75	72.90	70.13	41.85	36.50	25.10	28.17	23.40	51.5
7	28.50	26.03	31.10	31.90	28.50	26.07	79.97	76.10	80.27	80.40	80.30	78.50	20.70	31.87	18.93	18.70	21.30	29.0
В	24.10	23.60	30.60	26.50	26.00	23.30	68.35	71.60	78.67	76.60	75.70	70.17	49.50	45.87	22.70	37.60	31.63	50.00
9	23.55	26.03	27.00	29.30	24.93	23.87	70.02	74.57	77.70	74.65	74.27	71.73	51.10	33.63	32.73	28.93	23.83	43.67
10	25.90	24.07	27.65	29.90	27.40	23.60	76.52	72.50	75.78	76.60	77.87	73.03	35.70	44.80	33.00	24.87	28.77	42.70
Cont.	29.05	30.60	30.60	32.60	34.17	27.30	86.68	85.93	85.20	86.80	84.00	84.70	17.93	10.83	12.33	14.93	19.33	19.93
LSD for LSD for Identificat	fungus(F) x Cul	tivar (C	<u>)</u> (p <l< td=""><td></td><td>0.75 0.99</td><td></td><td></td><td></td><td></td><td></td><td>5)=1.39 1)=1.84</td><td></td><td></td><td></td><td></td><td></td><td></td></l<>		0.75 0.99						5)=1.39 1)=1.84						

Table 5. Analysis of variance of the effect of some cellulose-degrading fungi on fiber brightness (Rd %), yellowness (+b) and trash no.

of six Egyptian cotton cultivars. Parameters and D.F F. values P> F source of variation Brightness (Rd %) Replications 1.203 1.2438 0.2917 5 638.481 659.9382 0.0000 Cultivars (C) Fungi (F) 10 204.152 211.0128 0.0000 32.519 50 CxF 33.6118 0.0000 130 0.967 Error Yellowness (+b) Replications 2 0.079 1.2910 0.2785 5 Cultivars (C) 18.698 306.1839 0.0000 10 29.025 475.2842 0.0000 Fungi (F) CxF 5Ō 2.769 45,3361 0.0000 <u>130</u> 0.061 Error Trash no. Replications 2 2.399 0.3013 5 425.0590 Cultivars (C) 3384.784 0.0000 10 17.5113 0.0000 Fungi (F) 139,444 CxF 50 195.864 24,5965 0.0000 Error 130 7.963

Table 6. Relative contribution of fungi, cotton cultivar, and their interaction to variation in fiber brightness (Rd %), yellowness (+b)and trash no.

Source of	Relative contribution to variation in ^a								
variation	Brightness (Rd)%	Yellowness (+b)	Trash no.						
Cultivars (C)	46.52	17.90	60.19						
Fungi (F)	29.45	55.57	4.96						
CxF	23.69	26.50	34.83						

^a Calculated as percentage of sum of squares of the explained (model) Variation.

The decrease in brightness as well as the increase in yellowness could be accounted for by the production of pigments by fungi; however, it is difficult to account for the increase in brightness and the decrease in yellowness in Giza 89 under the effects of some of the tested fungi. As to trash no., the responses of the cultivars to fungal infection were too variable to draw meaningful conclusions

Cultivars, fungi, and cultivars × fungi interaction were all very highly significant source of variation in maturity, micronaire value, strength, and elongation (Table 8). Cultivars were the most important source of variation in maturity and micronaire value, isolates were the most important source of variation in strength, and the interaction was the most important source of variation in elongation (Table 9).

Table 7. Effects of some cellulose-degrading fungi on fiber color and trash no. of six Egyptian cotton cultivars.

									Cul	tivars								
Fungus	•	Brightness (Rd %)				-	Į .	Yellowness (+b)						Trash no.				
	G80	G85	G86	G88	G89	G90	G80	G85	G86	G88	G89	G90	G80	G85	G86	G88	G89	G90
1	52.75	64.00	64.00	55.30	62.47	57.13	17.80	16.30	15.45	16.80	14.80	17.43	89.33	9.00	14.00	6.67	3.67	16.67
2	53.25	62.73	66.60	57.90	66.27	55.97	17.50	17.03	15.60	15.90	13.13	17.83	39.33	11.33	15.67	10.67	5.67	6.00
3	53.25	62.32	65.40	55.50	62.40	58.47	17.20	15.77	15.65	15.70	15.07	16.23	49.00	10.00	5.00	7.67	4.67	16.00
4	57.70	63.07	64.10	55.50	63.33	56.50	14.75	16.43	15.10	15.75	14.77	16.47	39.00	8.33	14.00	11.67	7.33	9.00
5	52.80	63.93	59.70	58.15	66.73	54.67	14.40	14.93	16.45	15.40	14.37	16.63	28.00	11.67	9.00	7.00	6.00	14.00
Ŝ.	54.20	64.87	60.85	53.75	61.00	57.27	17.55	15.83	16.55	17.20	14.10	16.90	21.67	13.67	13.33	9.67	9.33	16.0
7	52.25	62.43	59.95	47.80	59.20	55.13	15.50	14.30	13.85	14.45	13.77	14.53	12.00	16.33	15.33	21.67	12.67	13.67
3	52.65	61.13	56.35	54.45	62.13	55.67	17.25	15.93	16.65	17.60	15.73	16.93	48.00	6.00	12.90	20.00	12.67	15.00
3	50.55	64.90	61.45	55.05	62.47	53.63	17.10	15.07	16.35	16.40	15.60	17.60	22.00	9.67	12.33	15.00	6.67	21.00
10	51.10	61.33	60.15	53.52	62.83	56.27	17.50	15.83	15.40	15.95	13.90	15.67	60.33	7.33	15.67	20.00	6.33	10.00
Cont	67.57	81.00	77.30	62.90	60.93	68.60	14.30	9.77	10.50	10.75	15.43	13.03	22.33	15.33	6.33	7.00	21.00	5.00
SD for fu						L			F) x Cult F) x Cult				LSD for LSD for					5)= 4.50 1)= 6.03

^{*} Identification of fungl is shown in Table 1

Table 8. Analysis of variance of the effect of some cellulose-degrading fungi on fiber maturity %, micronaire value and fiber mechanical properties of six Egyptian cotton cultivars

IIIECIIAIIR	ai pi upei	cies or six Egyp	uan couon cu	iuvais.
Parameters and source of variation	D.F	M.S	F. values	P> F
		Maturity %		
Replications	2	0.000	5.5418	0.0826
Cultivars (C)	5	0.009	91.9685	0.0000
Fungi (F)	10	0.007	77.7982	0.0000
CxF	50	0.002	20.456	0.0000
Error	130	0.000		
		Micronaire value		
Replications	2	0.013	2.2166	
Cultivars (C)	5	1.609	272.6799	0.0000
Fungi (F)	10	0.046	7.8608	0.0000
CxF	50	0.087	14.7170	0.0000
Error	130	0.006		
		Fiber Strength		
Replications	2	0.737		T T
Cultivars (C)	5	297.950	0.9764	0.0000
Fungi (F)	10	334.811	394.6204	0.0000
CxF	50	25.864	443.4412	0.0000
Error	130	0.755	34.2551	
		Elongation %		
Replications	2	0.001	0.0199	
Cultivars (C)	5	3.336	50.7882	0.0000
Fungi (F)	10	0.940	14.7931	0.0000
CxF	50	0.387	6.0926	0.0000
Error	130	0.064	T	

Table 9. Relative contribution of fungi, cotton cultivar, and their interaction to variation in fiber maturity %, micronaire value and fiber mechanical properties.

	m	P. OP O. 1	_							
Source of	Ţ	Relative contribution to variation in *								
variation	Maturity %	Micronaire value	Fiber Strength	Elongation %						
Cultivars (C)	62.47	62.46	24.28	35.94						
Fungi (F)	3.60	3.57	54.60	20.94						
CxF	33.71	33.77	21.09	43.12						

^{*} Calculated as percentage of sum of squares of the explained (model) Variation.

Maturity of Giza 80 and Giza 85 showed variable responses to fungal infection, while maturity of all remaining cultivars was reduced by all the fungi except that of Giza 89, which was not affected by A. flavus (Table 10). All the tested fungi significantly reduced the micronaire value of all the cultivars except *Trichothecium* sp., which did not affect micronaire value of Giza 86 and A. flavus which did not affect micronaire value of Giza 85, and Giza 89. Fiber strength of all the cultivars was significantly reduced by all the fungi (Table 11).

Elongation of Giza 88 was not affected by any fungus, while that of Giza 90 was significantly reduced by all the fungi. Elongation of the other

cultivars showed variable responses. The decrease in fiber strength could be ascribed to the deleterious effects of cellulolytic enzymes secreted by the fungi, these enzymes attack the amorphous regions of cellulose, which are located between the crystalline regions. Thus, weak point occur in fiber structure. The deterioration in fiber strength by the fungi is coincided with that previously reported by Abd El-Rehim et al. (1993) and Mahmoud (1996).

Table 10. Effects of some cellulose-degrading fungi on Maturity % and micronaire value

		IIICI O	Idne	raiue								
						Cuit	ivars _					
Fungus "			Matu	ity %			Micronaire value					
	G80	G85	G86	G88	G89	G90	G80	G85	G86	G88	G89	G90
1	0.85	0.81	0.85	0.86	0.85	0.82	4.00	3.47	4.10	3.55	4.00	3.70
2	0.87	0.83	0.88	0.86	0.83	0.82	4.20	3.60	4.11	3.65	4.00	3.73
3	0.85	0.81	0.85	0.84	0.91	0.82	3.90	3.63	4.30	3.40	3.87	3.77
4	0.89	0.83	0.87	0.87	0.90	0.81	4.05	3.50	4.05	3.65	4.17	3.63
5	0.85	0.83	0.85	0.82	0.81	0.81	4.05	3.53	3.85	3.70	4.30	3.77
6	0.83	0.84	0.86	0.82	0.87	0.83	4.00	3.77	3.95	3.65	4.15	3.73
7	0.89	0.85	0.88	0.93	0.96	0.85	4.35	3.79	4.16	3.78	4.50	3.78
8	0.82	0.81	0.86	0.84	0.84	0.80	3.85	3.57	4.03	3.60	4.03	3.60
9	0.82	0.80	0.84	0.80	0.83	0.80	3.80	3.57	3.82	3.55	3.83	3.63
10	0.84	0.82	0.85	0.93	0.85	0.81	3.85	3.50	3.90	3.75	4.17	3.63
Cont	0.87	0.92	0.93	0.88	0.96	0.89	4.45	3.90	4.30	4.05	4.53	3.97
LSD for	SD for fungus(F) x Cultivar (C) (p<0.05)=0.015LSD for fungus(F) x Cultivar (C) (p<0.05)=0.13											
LSD for	fundus	F) x C	iltivar (d	C) (p<0.	.01)= 0.	021LS) for fu	naus(F)	xCultiva	ır (C) (ı	o<0.01)	ė 0.17

Identification of fungi is shown in Table 1

Table 11. Effects of some cellulose-degrading fungi on mechanical properties

-		Cultivars											
Fungus	Fiber Strength							Elongation %					
1	G80	G85	G86	G88	G89	G90	G80	G85	G86	G88	G89	G90	
1	22.70	21.93	24.30	30.25	25.20	22.2	6.60	7.17	6.55	6.15	6.80	7.13	
2	25.25	22.37	30.02	33.85	26.80	20.2	6.90	7.77	6.17	6.40	6.50	6.70	
3			25.85					6.40	6.40	6.35	7.10	6.50	
4	29.60	21.60	26.95	27.73	29.22	20.00	7.55	6.93	6.00	6.25	7.90	6.73	
5 6	24.73	22.70	26.82	33.03	32.47	20.00	7.55	6.50	6.55	6.15	6.97	6.60	
6	20.60	23.30	26.70	22.95	21.70	18.60	6.45	6.53	6.50	6.30	6.17	6.77	
7	26.90	27.17	33.20	29.35	33.40	19.73	7.15	7.70	6.50	6.65	7.80	6.97	
8	20.15	23.93	29.35	25.25	23.23	25.50	6.65	6.37	6.15	6.00	6.73	6.47	
9	19.95	20.10	25.82	18.55	19.07	18.80	6.85	6.17	6.48	6.00	6.00	6.83	
10			25.40					6.97	6.40	6.00	6.43	6.57	
Cont	38.40	39.88	39.57	46.75	37.47	39.30	7.07	7.30	6.45	6.25	7.73	7.63	
LSD	for fung	us(F)x(Cultivar	(C) (p<().05)= 1	.40 LS	D for fu	ngus(F)	x Cultiv	ar (C)	(p<0.05	5)= 0.41	
LSD f	or fungi	ıs(F) x	Cultivar	(C) (p<	0.01)=	1.86 LS	D for fu	ngus(F)	x Cultiv	ar (C)	(p<0.01	i)= 0.54	
1 14	and files	tion of	funci le	- house	In Tak	Ja 1							

Identification of fungi is shown in Table 1

ANOVA of Table 12 showed very highly significant effects of cultivars, fungi, and cultivars × fungi interaction on cellulose, reducing sugar, and damage index. Cultivars were the most important source of variation in reducing sugar. Fungi and the interaction were almost equally important as sources of variation in damage index (Table 13).

Table 12. Analysis of variance of the effect of some cellulose-degrading fungi on fiber chemical properties and fiber damage index of

six Egyptian cotton cultivars.

SIX Egy	otian co	otton cultivar	<u> </u>	
Parameters and source of variation	D.F	M.S	F. values	P> F
		Cellulose	%	
Replications	2	0.087	0.2344	
Cultivars (C)	5	268.169	725.1134	0.0000
Fungi (F)	10	16.179	43.7468	0.0000
CxF	50	7.252	19.6087	0.0000
Error	130	0.370		
	Re	educing sugar	content %	
Replications	2	0.000	1.1635	0.3156
Cultivars (C)	5	0.001	27.5695	0.0000
Fungi (F)	10	0.009	197.8962	0.0000
CxF	50	0.003	67.1382	0.0000
Error	130	0.000		
		Fiber damage	Index	
Replications	2	6.173	0.6006	
Cultivars (C)	5	501.201	48.7693	0.0000
Fungi (F)	10	5414.013	526.8101	0.0000
CxF	50	1039.424	101.1411	0.0000
Error	130	10.277		

Table 13. Relative contribution of fungi ,cotton cultivar , and their Interaction to variation in fiber chemical properties and damage index.

Source of	Relative contribution to variation in *								
variation	Cellulose %	Reducing sugar content %	Fiber damage Index						
Cultivars (C)	71.88	2.54	2.31						
Fungi (F)	8.67	36.02	49.84						
CxF	19.44	61.44	47.84						

^{*} Calculated as percentage of sum of squares of the explained (model) Variation.

The fungi significantly decreased cellulose and increased each of reducing sugar and damage index. However, *F. moniliforme* was a notable exception because it significantly decresed reducing sugar of Giza 85 (Table 14).

It is believed that fungi decompose cellulose by a three-enzyme system. The first endo- 1,4-B-D glucanase (EC 3.2.1.4) randomly cleaves internal glucosidic bonds within unbroken glucan chain. The exposed non reducing chain ends become substrate for 1, 4-B-D- glucan cellobiohydrolase (EC 3.2.1.91), which cleaves the cellobiose dimmers from the chain releasing them into the environment.

]	Cultivare																	
Fungus *	Cellulose %						Reducing sugar content %						Fiber damage Index						
	G80	G85	G86	G88	G89	G90	G80	G85	G86	G88	G89	G90	G80	G85	G86	G88	G89	G	9 0
1	80.40	81.41	86.42	85.55	81.50	80.13	0.30	0.27	0.24	0.27	0.26	0.26	47.27	87.30	70.40	79.37	50.23	70	.50
ž	81.60	85.70	88.12	85.83	83.75	80.71	0.25	0.24	0.27	0.27	0.25	0.27	59.37	48.77	54.60	55.63	59.07	58-	60
3	81.53	82.22	89.00	86.30	82.00	80.20	0.25	0.26	0.23	0.23	0.25	0.26	59.00	52.57	83.33	85.17	70.80	49	93
4	81.15	81.17	85.70	86.00	81.00	81.40	0.25	0.23	0.24	0.24	0.27	0.27	79.27	38.13	33.63	70.50	39.33	68==	10
5	81.44	81.40	86.61	87.96	81.86	80.50	0.24	0.26	0.27	0.26	0.24	0.21	71.26	83.50	33.30	86.87	70.20	72	2.90
6	80.89	81.21	89.00	85.40	88.54	80.70	0.21	0.22	0.29	0.27	0.25	0.20	76.83	59.03	46.37	88.77	74.77	72	= .27
7	83.10	86.37	89.45	88.26	89.00	81.81	0.20	0.19	0.22	0.23	0.25	0.22	34.63	17.00	30.00	29.91	36.37	36	.47
В	81.06	81.73	87.80	85.40	84.15	76.90	0.26	0.25	0.26	0.23	0.24	0.24	83.03	54.23	76.87	53.93	43.30	69	₽.93
9	80.63	82.51	85.64	87.23	79.00	80.00	0.26	0.11	0.26	0.25	0.25	0.21	70.53	94.47	90.40	88.88	70.74	76	5.60
10	80.03	79.38	83.00	85.00	82.00	78.00	0.27	0.27	0.29	0.28	0.28	0.28	89.03	65.53	70.30	35.23	37.70	58	3.83
Cont	95.00	96.00	89.33	98,00	95.00	93.00	0.11	0.12	0.11	0.10	0.11	0.12	13.00	6.50	7.60	8.43	9.53	11	.73
LSD for fu LSD for fu				(p<0.05 (p<0.01	,			fungus((p<0.05)							5)====	5.18 6.8

Identification of fungi is shown in Table 1

The hydrolysis of cellobiose to glucose is accomplished by B-glucosidase (EC 3.2.1.21) (Goodman et al. 1986). Therefore, the increase in reducing sugar content could be ascribed to the decomposition of fiber cellulose to glucose. The increase in damage index suggests that the fungi caused breaks in the primary wall of the fiber.

Thus, only the interior of the fiber became more accessible for staining with congo red stain. Therefore, the amount of this stain in the fiber is an indication for the degree of deterioration (Abd Ei-Rehim and Aly, 1999).

The present study clearly demonstrated that cotton cultivars were much more important than fungal isolates in determining the level of deterioration in most of the tested properties. This result implies that the deleterious effects of the cellulose-degrading fungi on quality of fibers could be considerably reduced if resistance of cotton cultivars to these fungi is effectively enhanced.

REFERENCES

- Abd El-Rehim, S.A.; A.A. Aly; H.A. Eisa and Z. M. Askalany (1993). Deterioration of cotton fibers caused by some cellulolytic fungi isolated from rotted cotton bolls. Menofiya J. Agric. Res. 18: 2095-2110.
- Abd El-Rehim S.A. and A.A. Aly (1999). Deterioration in cotton fibers caused by isolates of the sooty mold fungus *Cladosporium herbarum*. J. Agric. Sci. Mansoura Univ., 24:2223-2238.
- Abd El-Rehim S.A.; M.A. Abdel Aziz and E.M. Hussein (2001). Effects of sooty mold disease caused by Cladosporium herbarum on quality of cotton fibers and seeds. J. Agric. Sci. Mansoura Univ., 26(2):805-817.
- Amer, M.A.A. (1986). Studies on cotton-seed infection by fungi. M.Sc. Thesis, Helwan Univ., Alexandeia, 127p.
- American stander for testing materials (ASTM) (1986). D: 1578-67, D: 1425-84, D: 2130, D: 3818 and D: 4605.
- Badr, F.A. (1980). Effect on infection with different fungi on fiber of some Egyptian cotton cultivars under field and laboratory conditions. M.Sc. Thesis, Cairo Univ., Egypt, 123p.
- Conrad, C.M. (1944). Determination of wax in cotton fiber. Anew alcohol extraction method. Ind. Eng. Chem. Anal. 16 pp 745.
- Goodman, R.N.; Z. Kiraly, and K.R. Wood (1986). The biochemistry and physiology of plant diseases. University of Missouri Press, Columbia. 433p.
- Hillocks R. J. (1992). Microbial contamination of the lint. Pp. 263-273. in: *Cotton Diseases*, R J Hillocks Ed.) C. A. B. International, Wallingford.
- Jenkis, S.H. (1930). Determination of cellulose in straw. Biochem . J. 24:1428.
- Mahmoud, A.S.M. (1996). Biochemical studies on toxic Fusarium moniliforme and Fusarium roseum effects on cotton fiber properties. M.Sc. Thesis, Cairo Univ., Egypt,.

Rao, N.V.; A.S. Reddy; R. Ankaiah and S. Mukundan (1989). Effect of whitefly, Bemisia tabaci Genn. On cotton yield and associated components. Insect Science and its Application. 10:685-690.

Smith, F.G. (1956). The sources content of the Western Australian Honey. J. Agric. Res. 2:177-184.

Sundaram, V. (1979). "Handbook of Methods of Tests for Cotton Fibers, Yarn, and Fabrics". Indian Council for Agric. Res., P. 71-76.

التدهور في ألياف القطن الناجم عن الإصابة ببعض القطريات المحللة للسليولوز معوض رجب عمر أ، عبير سمير عرفة و سلوى أحمد عبد الرحيم أ . 1. معهد بحوث أمراض النباتات، مركز البحوث الزراعية ، الجيزة ، مصر. ٢. معهد بحوث القطن ، مركز البحوث الزراعية ، الجيزة ، مصر.

القحت ألياف القطن الأصناف جيزة ٨٠ و جيزة ٥٠ و جيزة ٢٠ (تيلة طويلة) و جيزة ٨٨ (تيلة فائقة الطول) بعشرة فطريات محالة السليولوز، ونلك تحت ظروف المعمل. قيم التدهور في صفات الألياف الناجم عن نشاط هذه الفطريات من خلال دراسة التغيرات التي طرات على الصفات التالية: طول التيلة و معدل انتظام الطول ونسبة الشعيرات القصيرة ودرجة انعكاس الضوء ودرجة الاصفرار ونسبة العادم ونسبة النضج وقراءة الميكرونيتر ومتانة التيلة والنسبة المثوية للاستطالة ونسبة السليولوز والمحتوى من السكريات المختزلة و معدل تدهور الألياف. كانت الأصناف والفطريات وتفاعل الأصناف × الفطريات مصادر عالية المعنوية للتباين في جميع الصفات موضع الدراسة . نظرا المعنوية تفاعل الأصناف × الفطريات فإن أقل فرق معنوي استعمل لمقارنة تأثير الفطريات على كل صنف ، وذلك لكل صفة . أظهرت هذه المقارنات أن أغلب الصفات كانت تميل إلى التدهور بتأثير الإصابة الفطرية ، إلا أن حجم بكثير من عزلات الفطر حكمصدر للتباين في أغلب الصفات موضع الدراسة وعلى ذلك فإن المناف القطن أهم فإن التدهور في صفات الجودة بالنيلة الناجم عن الإصابة بهذه الفطريات - يمكن الحد منه بدرجة ملحوظة عن طريق زيادة مستوى مقاومة الأصناف لهذه الفطريات.

كلية الزراعة – جامعة المنصورة خارجي قام بتحكيم البحث أ. د/ عايدة حافظ عفيفي أ. د/ عبد الودود زكي عاشور