امیر حسین محمدی، ضیاءالدین بنی هاشمی^{*} و معصومه حقدل بخش گیاه پزشکی، دانشکده کشاورزی، دانشگاه شیراز و موسسه تحقیقات پسته کشور

دريافت: ۱۳۸۷/۶/۲۵ پذيرش: ۱۳۸۸/۳/۶

گونه های آسپرژیلوس از جمله عوامل تخریب کننده محصولات کشاورزی در مراحل قبل و پس از برداشت می باشند. طی سالهای ۱۳۸۳ تا ۱۳۸۶ با نمونه برداری از خاک مزارع و باغ های مختلف در استان فارس و باغ های پسته استان کرمان گونه های مختلف جنس Aspergillus از خاک جدا شده و جمعیت آن ها نیز محاسبه گردید. نمونه های خاک از عمق صفر تا ۴۰ سانتی متری تهیه شده و جدا سازی جدایه ها با استفاده از محیط کشت های مفر تا ۴۰ سانتی متری تهیه شده و جدا سازی جدایه ها با استفاده از محیط کشت های مفر تا ۴۰ سانتی متری تهیه شده و جدا سازی جدایه ها با استفاده از محیط کشت های مفر تا ۴۰ سانتی متری تهیه شده و جدا سازی جدایه ها با استفاده از محیط کشت های بخــش (Arpergillus انجام گردید. براساس خصوصیات میکروسکوپی و ماکروسکوپی پنج مشناسایی گردید که عبارت بودند از: Fumigati ، Circumdati ، Nigri ، Flavi شناسایی گردید که عبارت بودند از: Funigatus می مال ۱۳ گونـــه Aspergillus مناسایی گردید که عبارت بودند از: Aniger var. niger A. auricomus A. alliaceus مناسایی گردید که عبارت بودند از: As موماند ما مال ۱۳ گونــه As idav مناسایی گردید که عبارت بودند از: Arpa ما مال ۱۳ گونــه عمر مالا شناسایی گردید که عبارت بودند از: مین می مال ۱۳ گونــه که ماکروسکوپ مناسایی گردید که عبارت بودند از: مین مال ۲۵ گونـ معند مالا مالا گونــه کون مالاسایی گردید که عبارت بودند از: مین مالا مالای مالا مالای مالای که مالای مالای مالای می مالان مالای مالا مالای می مالان مالای محیط مالا مالای که با ستاره مشخص شدهاند برای اولین بار از ایران گزارش می شوند. از میان محیط کشت های مورد استفاده، محیط های App مالا مالا وکارین باری ولین ماز داند. کشت های مورد استفاده، محیط های App کارایی بهتری برای جاین دادند.

(E-mail: ziabani@shirazu.ac.ir) * مسئول مكاتبه

در استان فارس کرمان به ترتیب، A. flavus A. flavus بیشترین فراوانی را در استان کرمان به ترتیب A. parasiticus A. flavus A. niger var. niger بیشترین فراوانی را در مقایسه با سایر گونه ها داشتند. در استان فارس، خاک باغ های پسته و خاک های بکر، بیشترین و کمترین جمعیت گونه های Aspergillus را نشان دادند. در استان کرمان، خاک باغ های پسته رفسنجان و زرند، به ترتیب بیشترین و کمترین جمعیت را گونه های باغ های پسته رفسنجان و زرند، به ترتیب بیشترین و کمترین جمعیت را گونه های sepergillus داشتند. همچنین از میان ۲۰ و ۵۲ جدایه Alavus محمیت تولید کنند. فارس و کرمان، به ترتیب ۱۴ و ۳۳ جدایه توانستند آفلاتوکسین تولید کنند.

واژه های کلیدی: جمعیت قارچی، میکوتوکسین، پسته

نشانی نگارندگان: امیر حسین محمدی، دانشکده کشاورزی، دانشگاه شیراز و موسسه تحقیقات پسته کشور، دکتر ضیاءالدین بنی هاشمی، دانشکده کشاورزی، دانشگاه شیراز و معصومه حقدل، موسسه تحقیقات پسته کشور، رفسنجان.

IDENTIFICATION AND PREVALENCE OF ASPERGILLUS SPECIES IN SOILS OF FARS AND KERMAN PROVINCES OF IRAN AND EVALUATION OF THEIR AFLATOXIN PRODUCTION

A.H. MOHAMMADI, Z. BANIHASHEMI^{*} and M. HAGHDEL

Department of Plant Protection, College of Agriculture, Shiraz University and Iranian Pistachio Research Institute

Received: 16.09.2008

Accepted: 27.05.2009

Abstract

Aspergillus species are major causes of pre- and post-harvest degradation of agricultural products. During 2004-07, the presence and population of Aspergillus species were studied in various fields and orchards of Fars and Kerman Provinces. Soil samples were collected from 0-40 cm depth. Isolates were recovered from soil using modified Czapek, AFPA and SPDA media. Based on macroscopic and microscopic criteria, five sections (*Flavi*, *Nigri*, *Circumdati*, *Fumigati*, *Terrei*) and 13 following species were identified: A. alliaceus, A. auricomus, A. carbonarius*, A. flavus, A. fumigatus, A. japonicus var. japonicus, A. niger var. niger, A. ochraceus, A. oryzae*, A. parasiticus, A. sclerotiorum, A. sojae and A. terreus. The species with asterisk are new to Iran. SPDA and AFPA were more efficient for isolation and enumeration of Aspergillus colonies especially aflatoxigenic

^{*} Corresponding author (E-mail: ziabani@shirazu.ac.ir)

aspergilli. In Fars Province: *A. flavus, A. parasiticus, A. niger* var. *A. niger,* and in Kerman Province: *A. flavus, A. parasiticus* and *A. niger* var. *A. niger* were predominant, respectively. In Fars, pistachio orchard and uncultivated soils showed the highest and lowest *Aspergillus* population, respectively. In Kerman Province, soils collected from Rafsanjan and Zarand had the highest and lowest *Aspergillus* population, respectively. Out of 20 and 52 isolates of *A. flavus* obtained from soils in Fars and Kerman Provinces, 14 and 33 isolates produced aflatoxin, respectively.

Key words: Population density, Pistachio, Mycotoxin

Introduction

Aspergillus is a hyphomycetous genus with approximately 250 recognized species (GEISER *et al.* 2007). Members of the genus inhabit various niches, but mainly occur as saprophytes in soils, stored food and feed (DOMSCH *et al.* 1980, KLICH & PITT 1988). The genus was primarily described by MICHELI in 1729. GAMS & SAMSON (1985) have considered LINK (1809) as the validating author (KLICH 2002). After 1965, teleomorph names for sexual states of *Aspergillus* have become firmly established (MALLOCH & CAIN 1972, WILEY & SIMMONS 1973). At present, the genus has been divided into seven subgenera, each containing one or more sections to fit the Botanical Code (KLICH & PITT 1988a, KLICH 2002). These sections are equal to groups established by RAPER & FENNELL (1965). KLICH (2002) has questioned the validity of some groups and the species involved.

Although the majority of aspergilli are saprotrophs, some are parasitic on insects, plants and animals. Aspergilli are rarely directly pathogenic to plants in fields. *Aspergillus flavus* Link and *A. niger* Van Tieghem have been reported as seedling pathogen and crown rot of peanut, respectively (KLICH 2002). *Aspergillus* species are one of the major causes of pre- and post- harvest degradation of agricultural products. The greatest positive economic impact of aspergilli has been in the exploitation of the enzymes and acids produced by a number of species. For example, amylase and citric acid are two of the most important products produced

by various *Aspergillus* species (KLICH 2002). The negative economic impact of aspergilli involves the production of harmful mycotoxins by some species. Mycotoxins are products of secondary metabolism which are harmful to humans and/or animals. Aflatoxin is the most economically important mycotoxin throughout the world that is produced by *A. flavus*, *A. parasiticus* Speare and *A. nomius* Kurtzman, B.W. Horn & Hesseltine. The most toxic form of aflatoxin is aflatoxin B1 (DIENER *et al.* 1987). Other species of *Aspergillus* produce various mycotoxins (KLICH 2002).

Thirteen species of *Aspergillus* have been reported from Iran by MOJTAHEDI *et al.* (1979). They reported *A. niger* as the predominant species on pistachio kernels. HEIDARIAN *et al.* (2005) also isolated *A. phoenicis* (Corda) Thom & Currie and *A. puniceus* Known-Chung & Fennel from pistachio kernels in Iran. Eleven species of *Aspergillus* were isolated by RAHIMI *et al.* (2007) from pistachio nuts and litter.

Materials and Methods

During 2004-06, the frequency of occurrence of Aspergillus species was investigated in various fields and orchards in Fars and Kerman Provinces. In each orchard/field (= plot), four soil samples (500 g) taken from 0-40 cm depth and were pooled as one samples per orchard/field after passing through 2 mm, 40 and 60 mesh sieves in Fars Province. Likewise, four soil samples in each orchard were collected from eight sampling locations (four orchards per each) in Rafsanjan, Sirjan and Zarand of Kerman Province. In each plot, four samples were mixed together and finally, four subsamples (10 g) were suspended in 90 ml of 0.1% water-agar containing 100 ppm NPX (STEINER & WATSON 1965) and shaken for 30 min. Four replicates (0.5 ml) of diluted suspension $(10^{-2}, 10^{-3})$ in each subsample were flooded on the plates containing Czapek (RAPER & FENNEL 1965) with 250 ppm ampicillin. Plates were incubated at 25-27° C for 3-5 days and recovered colonies were purified using single-spore method. In this study, two other media including AFPA (Aspergillus flavus-parasiticus agar) (GOURAMA & BULERMAN 1995), and SPDA (sucrose-pepton-dichloran agar) (DHINGRA & SINCLAIR 1985) were also used for isolation and enumeration of toxigenic Aspergillus species from soil.

For identification of Aspergillus species, four media and two temperatures were applied: CYA (PITT 1973) at 25 and 37° C, CYA20S (CYA+20% sucrose) and MEA (KLICH & PITT 1988a) at 25° C. Inoculated plates were incubated in the dark for seven day. Isolates were examined both macroscopically and microscopically. For macroscopic examination, colony diameter, exudates and soluble pigment, surface and reverse colors and texture were considered. Diameter, shape, color and surface texture of conidia, phialides, metulae, vesicle, stipes, sclerotia and seriation (uniseriate or biseriate) were examined using microscope. All isolates were identified at species level (KLICH & PITT 1988a, KLICH 2002). In Fars Province, 20 isolates of A. flavus were randomly selected and examined for production of aflatoxin B1 and B2. In Kerman Province, 20, 16 and 16 isolates (2 isolates from orchards) of A. flavus from Rafsanjan, Sirjan and Zarand were also evaluated for their ability to produce aflatoxins, respectively, using WEI & JONG (1986) method. Ten ml of sterile distilled water was added to Petri dishes containing seven-day-old colonies of A. flavus and spore suspension was added to 500 ml flask containing 50 g of sterilized rice flour at 25° C. The flasks were incubated at 25° C in darkness for two weeks. To extract aflatoxin, four g of NaCl, 250 ml methanol 55 % and 100 ml n-hexan were added to the flasks. After 45 min shaking, the mixture was passed through the Whatman paper (No. 4). Fifty ml of the suspension was added to the same amount of chloroform and mixed thoroughly. The chloroform phase was transferred into new flasks and evaporated in rotary (40° C) with vacuum condition. Samples were dissolved in 200 µl of banzen-acetonitril (98:2) and 5 µl were spotted on silicagel TLC plate (60F254 Merck). Standard aflatoxin samples were spotted on the edges of the plates. Plates were then placed in developing tank containing developing solvent (chloroform-aceton, 9:1 v/v) for 15-20 min. Plates were air-dried in dark cabinet. Samples were quantified by comparing to standard aflatoxin spots using CAMAG TLC Scanner 3.

Results

All species of *Aspergillus* and other soil fungi such as *Penicillum, Rhizopus, Fusarium* and *Rhizoctonia* were easily isolated on Czapek medium; nevertheless discrimination of *Aspergillus* species was difficult on the medium. Growth and development of soil fungi except *Aspergillus* species was very low on SPDA and AFPA. *A. flavus* and *A. parasiticus* were easily identified on Czapek by the production of yellow to olive green spores three days after inoculation on AFPA and SPDA media. *A. niger* colonies developed dark brown to black conidia after three days and the reverse of colonies became yellow, but not orange. *A. ochraceus* grew slowly at 30° C and yellow colors appeared after three days. Applying these media shortened the time required for isolation and identification of potentially aflatoxigenic *Aspergillus* species. SPDA and AFPA displayed lower population of other fungi than Czapek medium. SPDA also showed lower population of soil bacteria compared with AFPA and Czapek.

In the present study, 340 isolates of *Aspergillus* were recovered from soils in Fars and Kerman Provinces from which 280 isolates were selected and identified to species. Based on macroscopic and microscopic examinations, 13 different species were identified in *Aspergillus* subgenera *Circumdati*, *Fumigati* and *Nidulantes* (Table 1).

Subgenus	Section	Soil	Species	Soil
Circumdati	Circumdati	88.23, 100 ^a	A. alliaceus	41.18, 55.26 ^b
			A. auricomus	29.41, 44.74
			A. ochraceus	64.71, 60.53
			A. sclerotiorum	29.41, 44.74
	Flavi	100, 100	A. flavus	100, 92.11
			A. oryzae	52.94, 63.16
			A. parasiticus	94.12, 86.84
			A. sojae	41.18, 50
	Nigri	100, 100	A. carbonarius	58.82, 81.58
			A. japonicus var. japonicus	41.18, 44.74
			A. niger var. niger	100, 94.74
Fumigati	Fumigati	76.47, 65.79	A. fumigatus	76.47, 65.79
Nidulantes	Terrei	23.53, 36.84	A. terreus	23.53, 36.84

Table 1. Identification of the recovered isolates in Aspergillus subgeneraCircumdati, Fumigati and Nidulantes in Fars and Kerman Provinces

^{a,b} Percentage of soils in which sections and species of *Aspergillus* were isolated in Fars and Kerman Provinces, respectively.

Description of selected Aspergillus species

Aspergillus carbonarius (Bain.) Thom, J. Agric. Res. 7: 12 (Fig. 1)

Thirty-one isolates of the species were used for identification.

Macroscopic characteristics: Colony diameter on CYA, 78 mm; conidia black and mycelium white; reverse pale yellow; sclerotia were not observed. Colony diameter on CYA20S, 85 mm, other characters similar to those on CYA except the stipes that was longer than those on CYA with usually yellow reverse. Colony diameter on MEA 55 mm, conidia black and uncrowded; mycelium white; reverse uncolored, Sclerotia were not observed. Colony diameter 45 mm at 37° C.

Microscopic characteristics: Conidial heads radiate; stipes thick-walled, finely roughened and colorless; vesicles globose 72-87 μ m; biseriate; phialides 10-16 × 5-7 μ m; metulae over the entire vesicle surface 23-45 × 6-14 μ m; conidia very large 6-10 μ m in diameter, walls extremely rough with tubercles and globose.

Aspergillus oryzae (Ahlburg) Cohn, Jahresbericht Schles. Ges. Vaterl. Kult. 61: 226 (Fig. 2)

Twenty isolates of the species were used for identification

Macroscopic characteristics: Colony diameter on CYA 63 mm; conidia olive to brown and mycelium white; reverse uncolored; dark sclerotia occasionally formed; colony texture generally floccose. Colony diameter on CYA20S 67 mm; conidia olive brown to olive yellow, in some isolates conidia were abundant; mycelium white, dense; reverse uncolored and colonies were more floccose compared to other media. Colony diameter on MEA 58 mm; conidia olive or occasionally green; mycelium white, not as dense as on CYA; reverse colorless; colony floccose. Colony diameter 70 mm at 37° C.

Microscopic characteristics: Conidial heads radiate; stipes colorless, walls finely rough; vesicles subglobose 15-62 μ m; aspergilla in some isolates predominantly biseriate, other predominantly uniseriate; phialides 7-13 \times 2.5-4.5 μ m; metulae cover more surface of the vesicle surface measuring 6-15 \times 3.5-5 μ m; conidia globose, 3.5-5.5 μ m in diameter, with smooth to finely roughened walls.

Aspergillus sclerotiorum Huber, Phytopathology 23: 306 (Fig. 3)

Twenty four isolates of the species were used for identification Macroscopic characteristics: Colony diameter on CYA 55 mm, conidia light yellow, mycelium white, reverse brownish yellow. Colony diameter on CYA20S 70 mm, characters similar to those on CYA except reverse that was lake red. Colony diameter on MEA 55 mm, conidia light yellow, mycelium white, not dense, reverse light yellow. Colony diameter 26 mm at 37° C.

Microscopic characteristics: Conidial heads radiate; stipes rough-walled, yellow; vesicles spherical 25.5-46 μ m; biseriate; phialides 5-7.5 \times 2.5-4 μ m; metulae usually cover the entire vesicle surface, 11-14 \times 3-4 μ m; conidia spherical 2-4 μ m in diameter, with smooth to finely roughened walls.

Flavi and *Nigri* sections were found in 100% whereas *Terrei* in 23.53% of collected soils in Fars Province (Table 1). In Kerman Province, *Circumdati, Flavi* and *Nigri* sections were recovered from 100% of soils and *Terrei* in 36.84% of the soils (Table 1). Among the *Aspergillus* species, *A. flavus* and *A. niger* var. *niger* were found in all collected soils of Fars Province whereas in Kerman Province, *A. niger* var. *niger* and *A. flavus* were recovered from 94.74 and 92.11% of the soils. *A. terreus* had the lowest frequency in soil samples of both provinces (Table 1).

Frequency of occurrence (%) of the *Aspergillus* sections are shown in Table 2. For each case, *Flavi* and *Terrei* sections had the highest and lowest frequency, respectively (Table 2). *A. flavus* (21.35%) and *A. parasiticus* (19.45%) in *Flavi* section and *A. niger* var. *niger* (18.97%) in *Nigri* section were the most prevalent species in Fars Province. Other species including *A. oryzae* (7.72%), *A. carbonarius* (6.86%), *A. fumigatus* Fres. (5.65%), *A. sojae* (4.76%), *A. ochraceus* Wilhelm (4.63%), *A. sclerotiorum* (3.30%), *A. japonicus* Saito var. *japonicus* (2.26%), *A. auricomus* (2.02%), *A. alliaceus* (1.75%) and *A. terreus* Thom (1.29%) showed low frequency in various fields and orchards. *Aspergillus* population was variable in various soils of Fars Province. Pistachio orchard soils with 5280 and uncultivated soils with less than 1000 propagules/g of soil, showed the highest and lowest *Aspergillus* population, respectively.

		Sect	ions of A <i>spergillus</i>		
Substrate (Location)	Flavi	Nigri	Circumdati	Fumigati	Terrei
Pistachio (Neiriz)	$2550^{a} (5.8)^{b}$	1910 (4.4)	570 (1.3)	250 (0.6)	0 (0.0)
Cotton (Kheer-Darab)	2185 (5)	900 (2.1)	215 (0.5)	200 (0.5)	0 (0.0)
Sugarbeet (Rooniz)	1800 (4.1)	800 (1.8)	330 (0.8)	180 (0.4)	360 (0.8)
Corn (Saadatshar-Bajgah-Abadeh)	1563 (3.6)	1023 (2.3)	437 (1.0)	160 (0.4)	33 (0.4)
Grapevine (Kavar)	1470 (3.4)	400 (0.9)	850 (1.9)	160 (0.4)	0 (0.0)
Alfalfa (Bajgah-Abadeh)	1355 (3.1)	465 (1.1)	400 (0.9)	165 (0.4)	70 (0.2)
Citrus (Darab)	1370 (3.1)	850 (1.9)	150 (0.3)	250 (0.6)	0 (0.0)
Sesame (Kheer)	1520 (3.5)	600 (1.4)	350 (0.8)	0 (0.0)	100 (0.2)
Tomato (Neiriz)	1250 (2.9)	690 (1.6)	500 (1.1)	100 (0.2)	0 (0.0)
Fig (Estahban)	1140 (2.6)	760 (1.7)	280 (0.6)	260 (0.6)	0 (0.0)
Melon (Maharloo)	1590 (3.6)	400 (0.9)	250 (0.6)	200 (0.5)	0 (0.0)
Almond (Maharloo)	1330 (3.1)	560 (1.3)	250 (0.6)	250 (0.6)	0 (0.0)

Table 2. Colony counts and frequency of occurrence (%) of Aspergillus sections in field and orchard soils of Fars Province

	Sections of Aspergillus					
Substrate (Location)	Flavi	Nigri	Circumdati	Fumigati	Terrei	
Potato (Safashahr)	950 (2.2)	740 (1.7)	250 (0.6)	200 (0.5)	0 (0.0)	
Wheat (Safashahr-Sepidian)	1050 (2.4)	475 (1.1)	175 (0.4)	100 (0.2)	0 (0.0)	
Ornamental (Shiraz-Bajgah)	760 (1.7)	610 (1.4)	110 (0.3)	0 (0.0)	0 (0.0)	
Apple (Bajgah)	900 (2.1)	750 (1.7)	0 (0.0)	0 (0.0)	0 (0.0)	
Uncultivated soil (Kavar-	574 (1.3)	364 (0.8)	0 (0.0)	0 (0.0)	0 (0.0)	
Abadeh-Safashahr-Sepidan)						

^a Data are the means of soil samples of four plots.
^b Frequency of occurrence (%) of the Aspergillus sections are shown in parentheses. The data are based on the proportion of the total isolates of recovered Aspergillus (175166).

Flavi and *Terrei* sections showed the highest (44.97%) and lowest (2.69%) frequency of occurrence in Kerman Province, respectively (Table 3). *A. niger* var. *niger* (22.43%) and *A. carbonarius* (10.38%) in *Nigri* and *A. flavus* (18.42%) and *A. parasiticus* (18.18%) in *Flavi* sections dominated in various soils. *A. oryzae* (4.37%), *A. fumigatus* (4.05%), *A. sclerotiorum* (3.78%), *A. ochraceus* (3.76%), *A. sojae* (3.61%), *A. alliaceus* (3.10%), *A. auricomus* (2.73%), *A. terreus* (2.69%) and *A. japonicus* var. *japonicus* (2.11%) showed low population. Results also showed that soils collected from Rafsanjan and Zarand had the highest and lowest *Aspergillus* population, respectively. *Aspergillus* population in soils with routine cultural practices collected from Rafsanjan, Sirjan and Zarand was 4860, 4332 and 3777 propagules/g soil, respectively (Table 3). Pistachio soils with low manure application in Rafsanjan, Sirjan and Zarand had lower *Aspergillus* population compared to other pistachio soils (1784, 1326 and 1269 propagules/g soil). In these soils, no manure or low amounts of manure had been used by the growers.

In Fars Province, 14 (70%) isolates out of 20 *A. flavus* isolates produced aflatoxin, 13 isolates producing AFB1+AFB2 and one isolate producing AFB1 (Table 4). *A. flavus* isolates of corn (Bajgah), alfalfa (Abadeh), wheat (Saadatshahr), uncultivated soil (Sepidan) and citrus (Darab) produced no aflatoxin. Isolates capability in aflatoxins production was inconsistent and varied from 4.13 to 625.32 ppb for AFB1 and 1.54 to 17.89 ppb for AFB2.

In Kerman Province, 17 and 16 isolates out of 52 isolates of *A. flavus* produced AFB1 + AFB2 and AFB1, respectively (Table 3). Fifteen (75%) isolates out of 20 *A. flavus* from Rafsanjan, produced aflatoxins. Among these, seven isolates (35%) produced 353-226 ppb AFB1 and eight (40%) AFB1 + AFB2 ranging from 382 to 4536 and 15.6 to 225 ppb.

From 16 isolates of *A. flavus* from Sirjan, four isolates (25%) produced 256-1254 ppb AFB1 and five isolates (31.25%) AFB1 + AFB2 varying from 514 to 2346 and 49 to 563.2 ppb (Table 4).

Location		Sect	ions of Aspergil	llus	
Location	Flavi	Nigri	Circumdati	Fumigati	Terrei
	Orchard	soils with rou	tine manure app	lication	
Rafsanjan	2128 ^a (13.6) ^b	1508 (9.7)	701 (4.5)	283 (1.8)	240 (1.5)
Sirjan	1940 (12.4)	1443 (9.3)	651 (4.2)	159 (1)	139 (0. 9)
Zarand	1784 (11.4)	1394 (8.9)	475 (3)	95 (0.6)	29 (0.2)
	Orchard soils	s with low or i	rregular manure	application	
Rafsanjan	803 ° (3.9)	708 (3.4)	178 (0.9)	78 (0.4)	17 (0.1)
Sirjan	588 (1.9)	600 (1.9)	88 (0.3)	50 (0.2)	0 (0.0)
Zarand	538 (1.7)	543 (1.7)	163 (0.5)	25 (0.1)	0 (0.0)

Table 3. Colony counts and frequency of occurrence (%) of *Aspergillus* sections in pistachio orchard soils with routine and low manure application in Kerman Province

^a Data are the means of four plots per location. Twenty to 40 tones of manure are used every other year.
 ^b Frequency of occurrence (%) of the *Aspergillus* sections are shown in parentheses. The data are based on the proportion of the total isolates of recovered *Aspergillus* (124815).

376 (13.4)

115 (4.1)

71 (2.7)

1033 (35)

1297 (45)

Average

^c In soils with low manure application, data are the means of six, four and four locations in Rafsanjan, Sirjan and Zarand, respectively. (in each location, four plots were considered as replicates).

In Zarand, five isolates (56.25%) produced only 989 to 775 ppb AFB1 and four (25%) produced AFB1 +AFB2 ranging from 1942 to 3654 and 87 to 652ppb (Table 4). In Kerman Province, 19 isolates of *A. flavus* (5, 7 and 7 isolates from Rafsanjan, Sirjan and Zarand, respectively) were unable to produce aflatoxin.

Isolate No.	Host Location	Aflatoxin Concentration (ppb)		
Isolate No.	Host Location	(ppt	B2	
1	Corn-Saadatshahr	389.95 ^a	4.32	
2	Corn-Bajgah	ND	ND	
3	Corn-Abadeh	415.3	9.45	
4	Alfalfa-Abadeh	215.67	6.8	
5	Alfalfa-Abadeh	ND	ND	
6	Wheat-Safashahr	ND	ND	
7	Wheat-Sepidan	447.35	12.31	
8	Uncultivated soil-Sepidan	ND	ND	
9	Cotton-Darab	4.13	ND	
10	Cotton-Darab	240.15	5.32	
11	Citrus-Darab	202.47	9.3	
12	Citrus-Darab	ND	ND	
13	Grapevine-Kavar	219.2	8.79	
14	Grapevine-Kavar	489.65	12.356	
15	Fig-Estahban	165.89	5.78	
16	Fig-Estahban	510.32	10.68	
17	Sugarbeet-Rooniz	105.96	1.54	
18	Pistachio-Neiriz	79.7	1.73	
19	Pistachio-Neiriz	625.32	17.89	
20	Pistachio-Neiriz	ND	ND	
21	Pistachio-Rafsanjan	353	ND	
22	Pistachio-Rafsanjan	ND	ND	
23	Pistachio-Rafsanjan	986	ND	
24	Pistachio-Rafsanjan	382	15.6	
25	Pistachio-Rafsanjan	ND	ND	
26	Pistachio-Rafsanjan	1784	ND	
27	Pistachio-Rafsanjan	2236	ND	
28	Pistachio-Rafsanjan	1887	203.4	
29	Pistachio-Rafsanjan	ND	ND	
30	Pistachio-Rafsanjan	578.3	ND	

 Table 4. Aflatoxin production by Aspergillus flavus isolates from various soils in Fars and Kerman Provinces

Isolate No.	Host Location	Aflatoxin Concentration (ppb)		
1501ate 110.	Host Location	B1	B2	
31	Pistachio-Rafsanjan	ND	ND	
32	Pistachio-Rafsanjan	1369	87.9	
33	Pistachio-Rafsanjan	421.5	ND	
34	Pistachio-Rafsanjan	1056	ND	
35	Pistachio-Rafsanjan	1425.6	92	
36	Pistachio-Rafsanjan	986.8	87.9	
37	Pistachio-Rafsanjan	4536	225	
38	Pistachio-Rafsanjan	1026	103	
39	Pistachio-Rafsanjan	ND	ND	
40	Pistachio-Rafsanjan	985.6	87.9	
41	Pistachio-Sirjan	514	85.3	
42	Pistachio-Sirjan	1483	563.2	
43	Pistachio-Sirjan	587.3	49	
44	Pistachio-Sirjan	ND	ND	
45	Pistachio-Sirjan	ND	ND	
46	Pistachio-Sirjan	ND	ND	
47	Pistachio-Sirjan	1542	215.6	
48	Pistachio-Sirjan	256.4	ND	
49	Pistachio-Sirjan	ND	ND	
50	Pistachio-Sirjan	2346	207.8	
51	Pistachio-Sirjan	ND	ND	
52	Pistachio-Sirjan	1225	ND	
53	Pistachio-Sirjan	ND	ND	
54	Pistachio-Sirjan	475	ND	
55	Pistachio-Sirjan	ND	ND	
56	Pistachio-Sirjan	1254	ND	
57	Pistachio-Zarand	756	ND	
58	Pistachio-Zarand	ND	ND	
59	Pistachio-Zarand	ND	ND	

Isolate No.	Host Location	Aflatoxin Concer	Aflatoxin Concentration (ppb)		
Isolate 110.	Host Location	B1	B2		
60	Pistachio-Zarand	1942	87		
61	Pistachio-Zarand	98	ND		
62	Pistachio-Zarand	546	ND		
63	Pistachio-Zarand	ND	ND		
64	Pistachio-Zarand	3098	201		
65	Pistachio-Zarand	ND	ND		
66	Pistachio-Zarand	2457	146		
67	Pistachio-Zarand	ND	ND		
68	Pistachio-Zarand	ND	ND		
69	Pistachio-Zarand	775	ND		
70	Pistachio-Zarand	3654	652		
71	Pistachio-Zarand	598.6	ND		
72	Pistachio-Zarand	ND	ND		

ND: Not Detected

^a Isolates were grown on 50 g rice flour and the presence of aflatoxin was determined by TLC.

Discussion

It seems that variation among the species and population of *Aspergillus* spp. in soil samples is due to variation of plants and cultural practices such as applying different manures and soil tillage. *A. flavus*, *A. parasiticus* and *A. niger* var. *niger* were ranked as first, second and third highest frequent species in Fars Province. These species were also found in the majority of soils (Table 1). Whereas, the order of ranks changed to *A. niger* var. *niger* > *A. flavus* > *A. parasiticus* in Kerman Province. These species were isolated from all pistachio orchard soils with routine manure application. In some soils with low manure application, a number of *Aspergillus* species were absent. These pistachio soils had lower *Aspergillus* population compared with soils with routine manure application. *Aspergillus alliaceus* produced no sclerotium on MEA as reported by KLICH & PITT (1988) and KLICH (2002). Isolates of *A. auricomus* produced many orange sclerotia in small clusters on MEA. The average size of vesicle (14.5 μ m) was in agreement with KLICH & PITT (1988a) and KLICH (2002). Isolates of *A. carbonarius* obtained from Fars and Kerman Provinces had long stipes (up to several millimeters) and large and tuberose conidia as reported by KLICH & PITT (1988a) and KLICH (2002). Most of the isolates of A. flavus were distinguished from A. oryzae by less floccose texture and greener color of colonies. Smaller and smoother conidia were the most important criteria to discriminate A. flavus from A. parasiticus. Based on other reports (KLICH and PITT 1988a,b, GOURAMA & BULLERMAN 1995, KLICH 2002), conidial ornamentation is the most diagnostic criterion to differentiate these fungi. Isolates of A. fumigatus had uniseriate conidial heads with curving phialides that were parallel to each other and stipe axis. The color of the colonies in this species quickly turned into turquoise to dark green (KLICH & PITT 1988a, KLICH 2002). Uniseriate conidial heads with black spinose conidia were a good criteria to identify A. japonicas var. japonicus that was in agreement with KLICH & PITT (1988a) and KLICH (2002). Based on KLICH & PITT (1988a), A. japonicas var. japonicus is distinguished from A. japonicus var. aculeatus primarily by the globose conidia and smaller vesicle in the former. Nevertheless, both varieties have been elevated to one species by KLICH (2002). Isolates of A. niger var. niger had roughened conidia and dark brown to black colonies. Based on KLICH (2002) two former described varieties (KLICH & PITT 1998a), A. niger var. awamori and A. niger var. niger, have been elevated to two species. The average colony diameter at 37° C (zero to 30 mm) and yellow buff colonies of A. ochraceus was in agreement with the KLICH & PITT (1988a) and KLICH (2002) and made a distinction between this species and A. alliaceus. Colonies of A. oryzae were usually floccose with olive colors. These characters in addition of producing large conidia distinguished this species from A. flavus. Isolates of A. parasiticus had predominantly uniseriate conidial heads, more olive colonies and rough-walled conidia. These criteria differentiated this species from A. flavus. Isolates of A. sclerotiorum produced no sclerotia on CYA at 25° C as reported by KLICH & PITT (1988a) and KLICH (2002). Isolates of A. sclerotiorum had also yellow stipes with pale yellow and smooth-walled, small conidia (2.5 µm). Isolates of A. sojae had also predominantly uniseriate conidial heads with roughwalled conidia. A. sojae is differentiated from A. parasiticus by its larger conidia (7.5 µm compared to 4.2 µm) and more floccose colonies. Isolates of A. terreus had compact columnar conidial heads and very small (2.5 µm) and smooth-walled

conidia. This species also possessed hyaline aleurioconidia and lateral cells on hyphae. These characters were in accordance with KLICH & PITT (1988a) and KLICH (2002).

MOHAMMADI & BANIHASHEMI (2006) previously isolated *A. alliaceus*, *A. auricomus*, *A. carbonarius*, *A. flavus*, *A. niger* var. *niger*, *A. oryzae*, *A. parasiticus*, *A. sclerotiorum*, *A. sojae* and *A. terreus* from soil in Fars Province. Therefore, *A. carbonarius* and *A. oryzae* are new species to the mycoflora of Iran. *A. japonicus* var. *japonicus*, *A. carbonarius*, *A. oryzae*, *A. sclerotiorum* and *A. sojae* are new for pistachio orchard mycoflora in Kerman Province. MOJTAHEDI *et al.* (1979) isolated *A. niger*, *A. flavus*, *A. fischeri* Wehmer, *A. tamari* Kita, *A. terreus*, *A. nidulans* (Eidam) G. Winter, *A. umbrosus* Bainier & Sartory, *A. vesicolor* (Vuill.) Tiraboschi, *A. sydowii* (Bain. & Sart.) Thom & Church, *A. ochraceus*, *A. petrakii* Voros-Felkai, *A. fumigatus* and *A. parasiticus* from pistachio kernels. They reported that among the isolated species, *A. niger* had the highest population (MOJTAHEDT *et al.* 1979). HEIDARIAN *et al.* (2005) reported seven *Aspergillus* species from dry pistachios which *A. phoenicis* and *A. punvieus* were new to pistachio mycoflora of Iran.

RAHIMI *et al.* (2007) isolated 11 *Aspergillus* species from pistachio nuts in Kerman, Rafsanjan and Isfahan. They reported that *A. alliaceus*, *A. unguis* and *A. wentii* are new for mycoflora of Iran. *A. alliaceus*, *A. candidus*, *A. niveus*, *A. unguis* and *A. wentii* were also reported as new species for pistachio mycoflora (RAHIMI *et al.* 2007).

It seems that, SPDA and AFPA media are more efficient for isolation and enumeration of *Aspergillus* species especially *A. flavus* and *A. parasiticus*. In addition to quick identification and enumeration of these fungi, population of other soil mycoflora was low due to dichloran. Applying antibiotics, Rosebengal and streptomycine, in SPDA resulted in lower population of soil bacteria compared to AFPA and Czapek media.

Aspergillus population was totally variable in soil samples from Fars Province. Among the field and orchard soils, pistachio, cotton, sugar beet, corn and wheat, ornamental plants, apple, rose had the highest (3180 to 5280) and lowest (1080 to 1950) *Aspergillus* population/g soil, respectively. The lowest population of Aspergillus occurred in uncultivated soils with a mean less than 1000 propagules/g soil. Aspergillus species were diverse in soils of Fars Province. Species related to sections *Flavi* and *Nigri* were found in all collected soils whereas section *Circumdati* and *Fumigati* were not isolated from ornamental, apple and uncultivated soils. *Terrei* section was only isolated from sugarbeet, corn, alfalfa and sesame soils and had the lowest frequency in Fars Province. *A. flavus, A. parasiticus (Flavi section)* and *A. niger* var. *niger (Nigri section)* were three species with highest frequency in Fars Province and showed variation in the population in various soils.

Other studies showed variation in populations of *A. flavus/A. parasiticus* in soils. GRIFFIN & CARREN (1974) found that *A. flavus* population in Virginia field soils was around 0.5-57.3 propagules/g soil. BELL &CRAWFORD (1967) reported significantly greater amount of propagules in naturally infested soils in Georgia $(1.5 \times 10^4 \text{ propagules/g soil})$. MARTINUK & WAGNER (1978) and SHEARER *et al.* (1992) reported 2.8 × 10³ and zero to 256 propagules of *A. flavus/A. parasiticus* from soils that were fertilized and cropped to continuous corn and from cultivated fields on three years rotation of wheat, red clover and corn in Missouri, respectively.

Aspergillus species obtained from Kerman Province were placed in three groups based on their population pattern in soils. First group included A. niger var. niger (Nigri section), A. flavus and A. parasiticus (Flavi section) representing more than 590 propagules/g soil (736, 616 and 597 propagules/g of soil, respectively). Second group included A. carbonarius (342 propagules/g soil) and third group included other Aspergillus species (86 to 141 propagules/g soil). Among the third group, A. terreus (Terrei section) and A. oryzae (Flavi section) had the lowest and highest population, respectively. Species belonging to the first group are important because of colonization of pistachio nuts, production of aflatoxins and decreasing pistachio quality. Among members of this group, A. niger var. niger may be potentially important due to its role in biological control of aflatoxigenic species. DOSTER & MICHAILIDES (1994) reported that A. niger var. niger, A. japonicus, A. flavus, A. parasiticus, A. tamari, A. melleus, A. ochraceus and A. wentii can colonize pistachio litter in pistachio growing areas in California. Among identified species, members of Nigri section (A. niger and A. japonicus) were more common in pistachio litter and soil (DOSTER & MICHAILIDES 1994) but in Kerman Province, *Flavi* and *Terrei* sections showed the highest (1297) and lowest (71) *Aspergillus* population, respectively. Although *A. niger* var. *niger* had a higher population than *A. flavus* or *A. parasiticus*, but *Nigri* section had a lower population compared with *Flavi* section.

Population of Aspergillus species in soil samples of Rafsanjan was globally higher than Sirjan and Zarand (Table 3). These results were not in agreement with MORADI et al. (2004) who reported that population density of A. flavus and A. niger groups were higher in Zarand than Rafsanjan and Sirjan soils. In the present study, we identified five Aspergillus sections with 13 species, whereas Moradi et al. (l.c.), only studied two sections of Aspergillus. Moradi et al. (l.c.) also indicated that sheep and poultry manure had the highest and lowest A. niger/A. flavus groups population, respectively, and cow manure was intermediate. These manures are mixed with the soils and increased the level of organic substances and subsequently Aspergillus population. Cultural practices such as tillage can distribute Aspergillus inoculum in manure and soil and increase the number of spores. These spores could be important by reaching to the aerial parts of the pistachio trees through dust (DOSTER & MICHAILIDES 1992). In some orchards of Kerman Province, low amounts of manure had been used or the application of manure was irregular. Very little or no pistachio debris was observed in some of the orchards. These soils supported lower Aspergillus population than others. DOSTER & MICHAILIDES (1994) reported that spores of A. flavus on male inflorescences are evenly distributed in orchards that can be important in increasing A. flavus population in pistachio orchards in California. In pistachio growing areas of Iran, male inflorescences fall from late March to April but male trees are few compared to California pistachio orchards. Therefore, the male inflorescences may not be important in increasing Aspergillus population in Iran. Nearly 10% of female inflorescences can be fertilized by pistachio pollen in orchards and the rest are distributed on the ground (CRAINE & EVAKIRI 1981). These female inflorescences can be colonized by Aspergillus species and increase fungus population. In pistachio orchards of Iran, pistachio debris is distributed in the vicinity of processing terminals or on orchard ground. Colonization of debris by Aspergillus species can build up the fungus population in

the pistachio orchards. Other reports (DOSTER & MICHAILIDES 2004, SHEARER *et al.* 1992) also showed that pistachio litter and cob and stalk pieces of corn may play an important role in infection of pistachio and corn by increasing the amount of *Aspergillus* inocula in soils. Since *Aspergillus* species are able to decompose numerous substrates including plant debris, it is not surprising to find high population of the fungus in soils of various fields and orchards (DOSTER & MICHAILIDES 1994). It seems that type of manures (cow, sheep and poultry), manure application procedure (burying under the soil or spreading on soil surface), presence of plant debris on field and orchard ground and tillage or cultural practices are important factors for increasing population of *Aspergillus* species in Fars and Kerman Provinces.

In corn fields of America, sclerotia of *A. flavus/A. parasiticus* are an important source of primary inoculum. These sclerotia can not disperse the fungus at harvest, but remain as a long-term survival structure in soils (WICKLOW 1987, WICKLOW *et al.*1984). The method of survival of *Aspergillus* spp. particulary aflatoxigenic species in Fars and Kerman soils still is not well understood. MIRABOLFATHY *et al.* (2005) reported that 42% of soil isolates, 33% of air isolates and 27% of nut isolates of *A. flavus* recovered from Kerman and Semnan Provinces can produce sclerotia. They also observed a high correlation between production of aflatoxin and sclerotium by *A. flavus* whereas, RAHIMI *et al.* (2007) showed that 10% of pistachio isolates of *A. flavus* produced sclerotia and there is not a direct relationship between sclerotium and aflatoxin production among *A. flavus* isolates.

The majority of *A. flavus* isolates (70%) recovered from soil of Fars and Kerman Provinces produced aflatoxin. It seems that the soil is the major source for contamination of crops to aflatoxins. Based on other studies, 41-95% of *A. flavus/A. parasiticus* can produce aflatoxins (DIENER & DAVIS 1966, KLICH & PITT 1988b, TABER & SCHROEDER 1967), but DOSTER & MICHAILIDES (1994) found that most isolates of *A. flavus* from pistachio litter could not produce detectable amounts of aflatoxins. Although many reports revealed that some isolates of *A. flavus* isolates do not produce aflatoxins, the exact percentage of these aflatoxigenic *A. flavus* isolates vary substantially. In the ATCC, only 34-41%

(depending on substrate) of A. flavus and 85% of A. parasiticus produced aflatoxin (WEI & JONG 1986). RAHIMI et al. (2007) reported that 50 and 100% of A. flavus/A. parasiticus isolates recovered from pistachio orchards produced aflatoxin. Isolates of A. flavus recovered in the present study did not produce aflatoxin G. This result was in accordance with DOSTER & MICHAILIDES (1992). COTTY (1989) and TABER & SCHROEDER (1967) who also reported none of the isolates of A. flavus produced aflatoxin G but isolates in ATCC (WEI & JONG 1986) and 26% of the A. flavus isolates from Africa and Thailand (Cotty 1997), produced the G aflatoxins. In addition to A. flavus/A. parasiticus, other mycotoxigenic aspergilli were also isolated and identified in various soils of Fars and Kerman Provinces. A. ochraceus, A. auricomus, A. sclerotiorum and A. alliaceus can produce an important group of carcinogenic toxins such as ochratoxin and less important mycotoxin penicillic acid (BAYMAN et al. 2002). The occurrence of A. alliaceus was correlated with the presence of ochratoxin A in figs, against A. ocharceus (BAYMAN et al. 2004). Although ochratoxin producing species showed low population in Fars and Kerman Provinces, but the presence of these species and the possibility of ochratoxin production must be regarded in field and orchards of Iran.

References

- BAYMAN, P., BAKER, J.L., DOSTER, M.A. MICHAILIDES, T.J. and MAHONEY, N.E. 2002. Ochratoxin production by the *Aspergillus ochraceus* group and *Aspergillus alliaceus*. App. Environ. Microb. 68: 2326-2329.
- BELL, D.K. and CRAWFORD, J.L. 1967. A Botran-amended medium for isolating *Aspergillus flavus* from peanuts and soil. Phytopathology 57: 939-941.
- COTTY, P.J. 1989. Virulence and cultural characteristics of two *Aspergillus flavus* strains pathogenic on cotton. Phytopathology 79: 808-814.
- COTTY, P.J. 1997. Aflatoxin-producing potential of communities of *Aspergillus* section *Flavi* from cotton producing areas in the United States. Mycol. Res. 101: 698-704.
- CRAINE, J.C. and EVAKIRI, B.T. 1981. Morphology and reproduction of pistachio. Hort. Rev. 3: 376-393.

- DHINGRA, O.D. and SINCLAIR, J.B. 1985. Basic Plant Pahtology Methods. CRC Press.
- DIENER, U.L. and DAVIS, N.D. 1966. Aflatoxin production by isolates of *Aspergillus flavus*. Phytopathology 56: 1390-1393.
- DIENER, U.L., COLE, R.J., SANDERS, T.H., PAYNE, G.A., LEE, L.S. and KLICH, M.A. 1987. Epidemiology of Aflatoxin formation by Aspergillus flavus. Ann. Rev. Phytopathol. 25: 249-270.
- DOMSCH, K.H., GAMS, W. and ANDERSON, T.H. 1980. Compendium of Soil Fungi. Academic Press, London. 859 pp.
- DOSTER, M.A. and MICHAILIDES, T.J. 1994. Development of *Aspergillus* molds in litter from pistachio trees. Plant Dis. 78: 393-397.
- GEISER, D.M., KLICH, M.A., FRISVAD, J.C., PETERSON, S.W., VARGA, J. and SAMSON, R.A. 2007. The current status of species recognition and identification in *Aspergillus*. Stud. Mycol. 59: 1-10.
- GOURAMA, H. and BULERMAN, L.B. 1995. Aspergillus flavus and Aspergillus parasiticus: Aflatoxigenic fungi of concern in foods and feeds, a review. J. Food Protect. 58: 1395-1404.
- GRIFFIN, G.J. and GARREN, K.H. 1974. Population levels of *Aspergillus flavus* and the *A. niger* group in Virginia peanut field soils. Phytopathology 64: 322-325.
- HEIDARIAN, R., JAVAN-NIKKHAH, M., ORMAZ, B. and PEYAMBARI, M. 2005. Study of fungal contamination of pistachio seeds in Kerman Province, Iran and some new fungi for Iranian pistachio mycoflora. IV International Symposium on Pistachio and Almond-ISHS, Tehran, Iran, p. 180.
- JONES, R.K. 1979. The epidemiology and management of aflatoxins and other mycotoxins. pp. 381-392. *In*: J.G. Horsfall & E.B. Cowling (eds). Plant Disease. An Advanced Treatise. Vol. 4. How Pathogens Induce Disease?. Academic Press, New York.
- KLICH, M.A. 2002. Identification of Common *Aspergillus* Species. CBS, Utrecht, The Netherland.

- KLICH, M.A. and PITT, J.I. 1988a. A Laboratory Guide to Aspergillus species and their Teleomorphs. CSIRO, Division of Food Processing, North Ryde, NSW, Australia. 116 pp.
- KLICH, M.A. and PITT, J.I. 1988b. Differentiation among Aspergillus flavus and Aspergillus parasiticus and other closely related species. Trans. Br. Mycol. Soc. 91: 99-108.
- MALLOCH, D. and CAIN, R.F. 1972. The Trichocomaceae: Ascomycetes with *Aspergillus, Paecilomyces* and *Penicillium* imperfect states. Can. J. Bot. 50: 2613-2628.
- MARTYNIUK, S. and WAGNER, G.A. 1978. Quantitative and qualitative examination of soil microflora associated with different management systems. Soil Sci. 125: 343-350.
- MIRABOLFATHY, M., MORADI GHAHDARIJANI, M. and WALIYAR, F. 2005. Variability in aflatoxicogenic potential and sclerotial production of *Aspergillus flavus* in pistachio of Iran. IV International Symposium on Pistachio and Almond-ISHS. Tehran, Iran. p. 188-189
- MOHAMMADI, A.H. and BANIHASHEMI, Z. 2006. Isolation and identification of *Aspergillus* species from soil in Fars Province. 17th Iran. Plant Protect. Cong. Tehran, Iran. p. 452.
- MOJTAHEDI, H., RABIE, C.J., LUBEN, A., STEYN, M. and DANESH, D. 1979. Toxic *Aspergillus* from pistachio nuts. Mycopathologia 67: 123-127.
- MORADI, M., ERSHAD, D., MIRABOLFATHI, M. and PANAHI, B. 2004. The role of plant debris, soil and manure on population density of *Aspergillus flavus* and *Aspegillus niger* groups in pistachio orchards of Kerman Province (in Persian). Iran. J. Plant Pathol. 40: 221-234
- PITT, J.I. 1973. An appraisal to identification methods for *Penicillium* species: novel taxonomic criteria based on temperature and water relations. Mycologia 65: 1135-1157.
- RAHIMI, P., SHARIFNABI, B. and BAHAR, M. 2007. *Aspergillus* species isolated from pistachio and determination of their aflatoxin production. Rostaniha 8: 30-42 (In Persian).

- RAPER, K.B. and FENNEL, D.I. 1965. The Genus *Aspergillus*. Williams and Wilkins, Baltimore.
- SHEARER, J.F., SWEETS, L.E., BAKER, N.K. and TIFFANY, L.H. 1992. A study of Aspergillus flavus/A. parasiticus in Iowa crop fields: 1988-1990. Plant Dis. 76: 19-22.
- STEINER, G.W. and WATSON, R.D. 1965. The effect of surfactants on growth of fungi. Phytopathology 55: 1009-1012.
- TABER, R.A. and SCHROEDER, H.W. 1967. Aflatoxin-producing potential of isolates of the Aspergillus flavus-oryzae group from peanut (Arachis hypgaea). Appl. Microbiol. 15: 140-144.
- WEI, D. and JONG, S. 1986. Production of aflatoxins by strains of *Aspergillus flavus* group maintained in ATCC. Mycopathologia 93: 19-24.
- WICKLOW, D.T. 1987. Survival of Aspergillus flavus sclerotia in soil. Trans. Br. Mycol. Soc. 89: 131-134.
- WICKLOW, D.T., HORN, B.W., BURG, W.R. and COLE, R.J. 1984. Sclerotium dispersal of Aspergillus flavus and Eupenicillium ochrosalmoneum from maize during harvest. Trans. Br. Mycol. Soc. 83: 299-303.
- WILEY, B.J. and SIMMONS, E.G. 1973. New species and a new genus of *Plectomycetes* with *Aspergillus* states. Mycologia 65: 934-938.

Addresses of the authors: A.H. MOHAMMADI, Dr. Z. BANIHASHEMI and Department of Plant Protection, College of Agriculture, Shiraz University, Shiraz and M. HAGHDEL, Iranian Pistachio Research Institute, Rafsanjan, Iran.