Phytoprevention of aflatoxin production

Rasooli I (Ph.D.)^{1*}, Fakoor MH (M.Sc.)¹, Allameh AA (Ph.D.)², Rezaee MB (Ph.D.)³, Owlia P (Ph.D.)⁴

1- Medicinal Plants Research Center, Shahed University, Tehran-Qom Express

Way, Opposite Imam Khomeini's shrine, Tehran - 3319118651, Iran

2- Faculty of Medical Sciences, Tarbiat Modarres University, Tehran, Iran

3- Department of Medicinal Plants, Institute for Research in Forests and Rangelands, Tehran, Iran

4- School of Medicine, Shahed University, Tehran, Iran

*Correspondence author: Tel: +98-21-51212600, Fax: +98-21-51212601

P.O.Box: 18155/159, Tehran, Iran E-mail: rasooli@shahed.ac.ir

Receive: 1 Jul. 2008 Acceptance: 14 Mar. 2009

Abstract

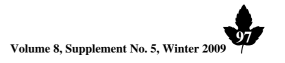
Background: Consumption of mycotoxic foods is associated with several cases of human poisoning, or mycotoxicosis, sometimes resulting in death.

Objective: Phytopreventive inhibition of *Aspergillus parasiticus* growth and its aflatoxin production by the essential oils extracted from *Thymus kotschyanus* Boiss & Hohen and *Zataria multiflora* Boiss. is reported in this study.

Methods: Minimal inhibitory concentration (MIC), minimal fungicidal concentration (MFC) and fungicidal kinetics of the oils were determined and compared with each other. Results: The oils from the above mentioned plants were found to be strongly fungicidal and inhibitory to aflatoxin production. Both oils inhibited aflatoxin B₁ (AFB₁) production by A.parasiticus. T. kotschyanus and Z. multiflora oils at 25 ppm concentration, reduced AFB₁ levels by 100% and 47.87% respectively. Aflatoxin production was significantly inhibited at lower than fungistatic concentration of both oils. The analysis of oils by GC and GC/MS led to identification of 27 and 22 components in T. kotschyanus and Z. multiflora Boiss. respectively which were very similar to each other.

Conclusion: Prevention of fungal growth and aflatoxin production by natural compounds is recommended.

Keywords: Thymus kotschyanus, Zataria multiflora, Essential oil, Antifungal, Aflatoxin



Introduction

Aflatoxins are secondary metabolites produced by certain strains of A. flavus, A. parasiticus and A. nomius which is phenotypically similar to A. flavus but with a distinctive bullet shaped sclerotia [1]. Other species that produce aflatoxins in minute quantities include Α. peudotamarii, A. bonbysis and A. ochraceoroseus [1]. These fungi grow rapidly on a variety of natural substrates and consumption of contaminated food can pose serious health hazards to human and animals. Aflatoxin B₁ (AFB₁) is a highly toxic and carcinogenic metabolite produced by Aspergillus species agricultural on commodities [2]. Natural products may regulate the cellular effects of aflatoxins and evidence suggests that aromatic organic compounds of spices can control production of aflatoxins [3]. Spices occupy a prominent place in the traditional culinary practices and are indispensable part of daily diets of millions of people all over the world. They are essentially flavouring agents used in small amounts and are reported to have both beneficial effect and antimicrobial properties, if properly stored [4]. Their antimicrobial properties have been found to be mostly due to the presence of alkaloids, phenols, glycosides, steroids, essential oils, coumarins and tannins [5, 6]. Allameh et al. [7] reported more than 50% inhibition of aflatoxin production at 50% (v/v) neem extract concentration. Bhatnagar and McCormic [8] have demonstrated that addition of neem leaf extract above 10% (v/v) effectively inhibited aflatoxin production by A. parasiticus and A. flavus. Under such conditions the mycelial dry weight was not affected [8, 9]. Thyme oils present high antimicrobial effect [10, 11, 12] compared to the oils of other plants [13]. Thyme oils may be utilized in the food industry and as a dietary supplement. Various species of thyme have been reported to possess antifungal properties [14, 15]. The aim of this study was to investigate the effect of essential oils of *Thymus kotschyanus* and *Zataria multiflora* Boiss. on *A. parasiticus* growth and AFB₁ production. Moreover the chemical compositions of the oils were studied as well.

Materials and methods

Chemicals, Cultures and Media

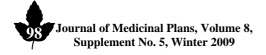
Α. parasiticus (NRRL-2999) was maintained on Sabouraud Dextrose Agar (E. Merck) slants. Spore suspensions were prepared and diluted in sterile yeast extract sucrose (YES) broth to a concentration of approximately 10⁶ spores/ml. Spore population was counted using haemocytometer. YES broth also served as aflatoxin production medium. Subsequent dilutions were made from the above suspension, which were then used in the tests. Incubation temperature was 28 + 2°C. Aflatoxin standards were the products of Sigma Chemical Co. St. Louis, Mo. USA. TLC silica gel 60 F254 were purchased from E. Merck, Germany. All other solvents and reagents were of analytical grade obtained from E. Merck, Germany.

Oil extraction

The plants were collected from National Botanical Garden of Iran in May 2006. The fresh aerial parts i.e. the leaves, were subjected to hydrodistillation for 90 min using a Clevenger. The extraction was carried out for 2 h after a 4-h maceration in 500 ml of water. The oils were stored in dark glass bottles in a freezer until they were used.

Oil analysis

GC analyses were performed using a Shimadzu-9A gas chromatograph equipped



with a flame ionization detector, and quantitation was carried out on Euro Chrom 2000 from Knauer by the area normalization method neglecting response factors. The analysis was carried out using a DB-5 fusedsilica column (30 m 0.25 mm, film thickness 0.25 µm, J & W Scientific Inc., Rancho Cordova, CA, USA). The operating conditions were as follows: injector and detector temperature, 250°C and 265°C, respectively; carrier gas, Helium. Oven temperature program was 40 - 250°C at the rate of 4°C/min. The GC/MS unit consisted of a Varian Model 3400 gas chromatograph coupled to a Saturn II ion trap detector was used. The column was same as GC, and the conditions were as above. spectrometer conditions were: ionization potential 70 eV; electron multiplier energy 2000 V. The identities of the oil components were established from their GC retention indices, relative to C7- C25 n-alkanes, by comparison of their MS spectra with those reported in the literature [16], and by computer matching with the Wiley 5 mass spectra library, whenever possible, by co-injection with standards available in the laboratories.

Antifungal analysis

The fresh oils were tested for their antifungal activities. The disc diffusion method was used for antifungal screening as follows: Sterile Sabouraud dextrose agar medium (Merck) was prepared and distributed into Petri plates of 70 mm diameter. This medium was used for antibiogram assays. The disc size used was 6 mm (Whattman no. 1) paper. The spore suspension was streaked over the surface of the Sabouraud dextrose agar plates using a sterile cotton swab in order to get a uniform microbial growth on both control and test plates. Under aseptic

conditions, the discs were placed on the agar plates and then 5 and 20 µl from each of the oils were put on the discs. The plates were then incubated at 28 + 2°C for 10 days in order to get reliable microbial growth. Diameters of microbial inhibition zones were measured using vernier calipers. The minimal inhibitory concentration (MIC) was determined by a broth dilution method in test tubes as follows: 50µL from each of various dilutions of the oils was added to 5mL of nutrient broth tubes containing 10⁶ CFU/mL of spores. The tubes were then incubated on an incubator shaker as to evenly disperse the oil throughout the broth tubes. The highest dilution (lowest concentration), showing no visible growth, was regarded as MIC. Cells from the tubes showing no growth were subcultured on Sabouraud dextrose agar plates to determine if the inhibition was reversible or permanent. The minimal fungicidal concentration (MFC) was determined as the highest dilution (lowest concentration) at which no growth occurred on the plates.

Fungicidal kinetics of the oils

MFC concentrations of the each oil was added to 5 ml of each spore suspension containing 10^6 spores/ml and were then incubated at 28 ± 2 °C for 5 - 60 min at increments of 5 min in an incubator shaker. Samples were taken after the time intervals and were cultured on Sabouraud dextrose agar for 48 h at 28 ± 2 °C. The control tube had not essential oils. Microbial colonies were counted after incubation period and the total number of viable spores per ml was calculated. The calculation was converted to percent dead spores using routine mathematical formulae.

Determination of mycelial weight

Flasks containing mycelia were filtered through Whatman filter no. 1 and then were washed with distilled water. The mycelia were



placed on pre weighed petri plates and were allowed to dry at 50°C for 6 h and then at 40°C over night. The flasks containing dry mycelia were weighed. The net mycelial dry weight was then determined.

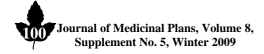
Measurement of aflatoxin B₁

10⁶ spores/ml were added to 100-ml Erlenmeyer flasks containing 20 ml of YES containing the essential oils concentrations lower than the MIC levels (6.25 ppm, 12.5 ppm, 25 ppm and 50 ppm) of T. kotschyanus and Z. multiflora Boiss. Aflatoxin B₁ measurement was determined at the end of the incubation period (28 ± 2 °C for 7 days). Aflatoxin extraction was performed routinely with solvent extraction [7, 17]. Silica gel 60-F precoated TLC sheets (E Merck, Germany) were used for analysis of aflatoxin B₁ produced by the fungal strain. The relative percentage of aflatoxin B₁ was estimated by TLC scanner III (Desaga densitometer CD60, Germany) at 365 nm using thin layer chromatography (TLC) method [18].

Results and Discussion

The inhibitory effects of the two kinds of Thyme essential oils against Aspergillus parasiticus were tested. Preliminary experiments were carried out in vitro using the disc diffusion and broth dilution methods to investigate antifungal action of the essential oils. Two concentrations of essential oils from T. kotschyanus and Z. multiflora Boiss. tested on Sabouraud dextrose agar plates showed a very strong antifungal property. 5 and 20µl from each of the oils caused complete inhibition of the growth of toxigenic A. parasiticus strain for 10 days. Many spices and herbs, such as cassia, basil, sweet basil, cloves, anise, star anise seeds, thyme, cinnamon, marigold, spearmint and caraway

have been reported to inhibit toxigenic and foodborne moulds [15, 19 -22]. MIC and MFC techniques were employed to assess fungistatic and fungicidal properties of the oils. It was found that both oils had static effect at 250 ppm. T. kotschyanus oil at 600 ppm and multiflora Boiss. oil at 750 ppm concentrations exhibited fungicidal property. Nguefack et al. [23] reported complete inhibition of Aspergillus flavus, Aspergillus fumigatus and Fusarium moniliform by the oils of Ocimum gratissimum, thymus vulgaris and Cymbopogon citratus at 800, 1000 and 1200 ppm respectively. Study of fungicidal kinetics of essential oils revealed more than 50% spore death in 15 minutes (Fig. 1). Complete lethal effects by T. kotschyanus and Z. multiflora Boiss. were observed within 30 and 50 minutes respectively (Fig. 1). Both oils inhibited aflatoxin B₁ (AFB₁) production by A. parasiticus. The order of inhibitory activity was T. kotschyanus > Z. multiflora Boiss. A concentration of 12.5 ppm T. kotschyanus in the YES broth resulted in 10.16% reduction of AFB₁ levels. Increasing *T. kotschyanus* concentration to 25 ppm, reduced AFB₁ levels by 100% limits. In the case of Z. multiflora Boiss., 3.34% inhibition of aflatoxin B₁ production was observed at a concentration of 12.5 ppm in the YES broth. When the concentration of Z. multiflora Boiss. was raised to 25 ppm in the YES broth, an inhibition of 47.87% was achieved (Table 1). Aflatoxin production was significantly inhibited at lower than fungistatic concentration of both oils (Table 1). Chemical analysis of the oils led to identification of 27 and 22 compounds in T. kotschyanus and Z. multiflora Boiss. oils respectively (Table 2). The profile of the oil components from kotschyanus was similar to that of multiflora Boiss. Z. in almost all the



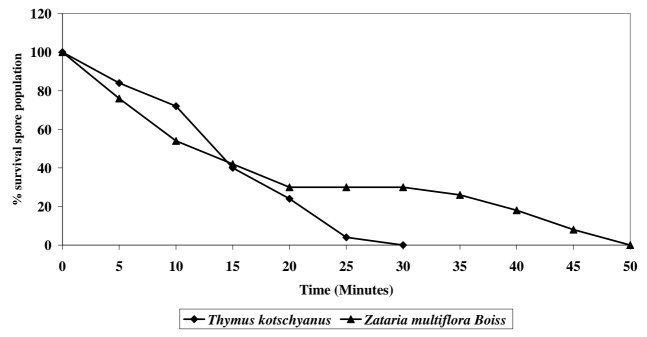


Fig. 1- Kinetics of A. parasiticus spore destruction at MFC levels of the fresh essential oil from Thymus kotschyanus and Zataria multiflora Boiss., Initial spore concentration: 10⁶/ml

Table 1: Effect of essential oils on growth of and aflatoxin production by Aspergillus parasiticus NRRL 2999

Oil and concentration (ppm)	Total Mycelial dry weight (mg)	Aflatoxin (mg/ml)	Aflatoxin (ng):dry weight (mg) ratio per ml	% Aflatoxin inhibition	
Control (0)	450 ± 20	1036	46.04	0	
T. kotschyanus (6.25)	450 ± 20	960	42.66	7.34	
T. kotschyanus (12.5)	440 ± 30	910	41.36	10.16	
T. kotschyanus (25)	450 ± 10	ND	0	100	
T. kotschyanus (50)	430 ± 30	ND	0	100	
Z. multiflora Boiss. (6.25)	470 ± 10	1050	44.68	2.95	
Z. multiflora Boiss. (12.5)	400 ± 30	890	44.50	3.34	
Z. multiflora Boiss. (25)	410 ± 30	492	24	47.87	
Z. multiflora Boiss. (50)	430 ± 20	486	22.60	50.91	

ND: Not detectable



Table 2: Chemical composition of essential oils from Thymus kotschyanus and Zataria multiflora Boiss.

No.	T.kotschyanus	RI*	%	Z. multiflora	RI*	%
1	α – Thujene	930	1.43	α – Thujene	925	0.9
2	α – Pinene	943	2.02	α – Pinene	933	5.0
3	Camphene	952	0.93	Camphene	947	0.3
4	Sabinene	979	0.39	Sabinene	970	0.6
5	β-Pinene	981	1.83	Myrcene	986	1.6
6	α – Phellandrene	1002	0.25	Decane	996	3.9
7	3-Carene	1007	1.5	α – Terpinene	1013	1.4
8	<i>p</i> -Cymene	1018	5.25	<i>p</i> -Cymene	1018	15.0
9	1,8-Cineole	1028	1.5	Limonene	1026	0.8
10	Limonene	1030	0.8	1,8-Cineole	1027	0.7
11	γ-Terpinene	1050	10.34	γ-Terpinene	1057	6.5
12	Transe-Sabinene			-		
	hydrate	1056	1.78	Sabinene hydrate	1064	0.3
13	Cis-Sabinene hydrate	1066	0.43	Terpinolene	1086	0.2
14	Camphor	1126	0.18	Undecane	1098	3.8
15	Borneol	1155	2.32	Dodecane	1198	8.9
16	4-Terpineol	1160	0.25	Thymol (Methyl ether)	1232	0.5
17	α – Terpineol	1165	0.3	Carvacrol		
	-			(Methyl ether)	1242	5.2
18	Carvacrol			,		
	(Methyl ether)	1212	0.3	Thymol	1289	3.3
19	Thymol (Methyl ether)	1221	0.47	Carvacrol	1298	37.0
20	Thymol	1287	19.55	Thymol acetate	1349	0.2
21	Carvacrol	1297	41.49	Tetradecane	1394	1.9
22	Thymol acetate	1340	0.24	β-Caryophylene	1418	1.9
23	Geranyl acetate	1354	0.3	_	_	_
24	β-Caryophyllene	1414	2.5	_	_	_
25	Germacrane D	1484	1.52	_	_	_
26	β-Bisabolene	1497	0.73	_	_	_
27	Germacrane B	1513	1.3			

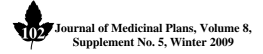
*RI: Retention index in reference to C₇-C₂₅ n-alkanes identified 99.9% in *T. kotschyanus* and *Z. multiflora*

compounds but at different concentrations. The major components of T. kotschyanus and Z. multiflora Boiss. oils were Carvacrol Thymol (41.49,37%), (19.55,3.3%), y-Terpinene (10.34, 6.5%) and P-Cymene (5.25, 15%) respectively (Table 2). The effectiveness of *T*. kotschyanus and Z. multiflora Boiss. on the growth of the A. parasiticus strain and aflatoxin production are probably due to major substances such as thymol and carvacrol showing antifungal effects [24-26] and completely suppressing aflatoxin synthesis [27]. It is well known that a phenolic-OH group is very reactive and can easily form hydrogen bonds with the active sites of enzymes [28].

The present study demonstrated potentials of the essential from oils Thymus kotschyanus and Zataria multiflora Boiss. in the control of growth and aflatoxin productivity of A. parasiticus. The use of these aromatic plants commonly used as spices or in beverage formulations is feasible as they are considered safe. However, further studies on their stability, toxocity and public acceptability as food preservatives are needed.

Acknowledgement

This study was fully supported by the Research Centre of College of Basic Sciences, Shahed University, Tehran-Iran. Authors wish



to express their gratitude to Mr. Mohammad

Habibi for the competent technical assistance.

References —

- **1.** Fente CA, Jaimez OJ, Vizquez BI, Franco CM, & Cepeda A. A new additive for culture media for rapid determination of aflatoxin-producing Aspergilli strains. *Appl. Environ. Microbiol.* 2001; 67 (10), 4858 62.
- **2.** Leontopoulos D, Siafaka A, & Markaki P. Black olives as substrate for *Aspergillus parasiticus* growth and aflatoxin B1 production. *Food Microbiol*. 2003; 20 (1), 119 26.
- **3.** Chatterjee D. Inhibition of fungal growth and infection in maize grains by spice oils. *Lett. Appl. Microbiol.* 1990; 11, 148 51.
- **4.** Oluwafemi F. Correlation between dietary aflatoxins and human male infertility. PhD. Thesis. 2000; University of Benin, Benin City, Nigeria.
- **5.** Ebana RUB, Madunagu BE, Ekpe ED, & Otung IN. Microbiological exploitation of cardiac glycosides and alkaloids from Garcinia kola, Borreria ocymoides, Kola nitida and Citrus aurantiofolia. *J. Appl Bacteriol*. 1991; 71, 398 401.
- **6.** Masako T, Sachis O, Reiko H, & Tadakatsu S. The bacterial activity of tea and coffee. *Lett Appl Microbiol*. 1989; 8, 123 5.
- **7.** Allameh A, Razzaghi AM, Shams M, Rezaei MB, & Jaimand K. Effects of neem leaf extract on production of aflatoxins and fatty acid synthetase, citrate dehydrogenase and glutathione Stransferase in *A. Parasiticus. Mycopathologia.* 2001; 154, 79 84.
- **8.** Bhatnagar D, & McCormic SP. The inhibitory effect of neem (*Azadirachta indica*) leaf extracts on aflatoxin synthesis in *Aspergillus parasiticus*. *J. Am. Oil Chem. Soc.* 1988; 65, 1166 8.
- **9.** Zeringue HJ, and Bhatnagar D. Inhibition of aflatoxin production in *Aspergillus flavus* infected cotton bolls after treatment with neem (Azadirachta indica) leaf extracts. *J. Am. Oil*

Chem. Soc. 1990; 67, 215 – 6.

- **10.** Juliano C, Mattana A, & Usai M. Composition and in vitro antimicrobial activity of the essential oil of Thymus herba-barona Loisel growing wold in Sardinia. *J. Essen. Oil Res.* 2000; 12, 516 22.
- **11.** Karaman S, Digrak M, Ravid U, & Ilcim A. Antibacterial and antifungal activity of the essential oils of Thymus revolutus Celak from Turkey. I. *Ethnopharmacol*. 2001; 76 (2), 183 6.
- **12.** Tzakou O, Verykokidou E, Roussis V, & Chinou I. Chemical composition and antibacterial properties of *Thymus longicaulis* subsp. Chaoubardii oils: Three chemotypes in the same population. *J. Essen. Oil Res.* 1998; 10, 97 9.
- **13.** Benjilali B, Tantaoui-Elaraki A, Ayadi A, & Ihlal M. Method to study antimicrobial effects of essential oils—Application to the antifungal activity of six Moroccan essences. *J. Food Prot* 1984; 47, 748 52.
- **14.** Pina-Vaz C, Gonc alves Rodrigues A, Pinto E, Costa-de-Oliveira S, Tavares C, & Salgueiro L. Antifungal activity of Thymus oils and their major compounds. *J. Eur. Aca. Dermatol. Venereol.* (JEADV) 2004; 18, 73 8.
- **15.** Rasooli I, Rezaei MB, & Allameh A. Growth inhibition and morphological alterations of *Aspergillus niger* by essential oils from *Thymus eriocalyx* and *Thymus x-porlock. Food Control* 2006; 17, 359 64.
- **16.** Davies NW. Gas Chromatographic Retention Index of Monoterpenes and Sesquiterpenes on Methyl silicone and Carbowax 20 M phases. *J. Chromatogr.* 1998; 503, 1 24.
- **17.** Razzaghi AM, Allameh A & Shams M. Screening of aflatoxin producing mould isolates based on fluorescence production on a specific medium under ultraviolet light. *Acta Med. Iranica* 2000; 38, 67 73.



- **18.** AOAC. International Official Methods of Analysis. Gaithersburg, MD, USA Official Method 1995, 970, 45.
- **19.** Atanda OO, Akpan I, & Oluwafemi F. The potential of some spice essential oils in the control of *A. parasiticus* CFR 223 and aflatoxin production. *Food Control* 2007; 18, 601 7.
- **20.** Soliman KM and Badeaa RI. Effect of oil extracted from some medicinal plants on different mycotoxigenic fungi. *Food Chem. Toxicol.* 2002; 40, 1669 75.
- **21.** Rahimfard N, Sabzevari O, Shoeibi Sh, Pakzad SR, Ajdari S, Hajimehdipoor H, Bagheri F and Safaei M. Antifungal activity of the essential oil of the *Eugenia caryophyllata* on *Candida albicans*, *Aspergillus niger* and *A flavus*. *Biomed*. *Pharmacol*. *J*. 2008a; 1 (1), 43 6.
- **22.** Rahimfard N, Sabzevari O, Shoeibi Sh, Pakzad SR, Ajdari S, Hajimehdipoor H, Bagheri F, and Bagheri A. Antifungal activity of the essential oil of the *Cinamoman zeylanicum* on *Candida albicans*, *Aspergillus niger* and *A. flavus. Biomed. Pharmacol. J.* 2008b; 1 (1), 85 8.
- **23.** Nguefack J, Leth V, Amvam Zollo PH and Mathur SB. Evaluation of five essential oils from aromatic plants of Cameroon for controlling food

- spoilage and mycotoxin producing fungi. *Int. J. Food Microbiol.* 2004; 94, 329 4.
- **24.** Sokovic M, Tzakou O, Pitarokili D, & Couladis M. Antifungal activities of selected aromatic plants growing wild in Greece. *Nahrung* 2002; 46 (5), 317 20.
- **25.** Bouchra C, Achouri M, Idrissi Hassani LM, & Hmamouchi M. Chemical composition and antifungal activity of essential oils of seven Moroccan Labiatae against *Botrytis cinerea* Pers: Fr., *J. Ethnopharmacol.* 2003; 89 (1), 165-9.
- **26.** Pinto E, Pina-Vaz C, Salgueiro L, Goncalves MJ, Costa-de-Oliverira S, Cavaleiro C, Palmeria A, Rodrigues A, & Martinez-de-Oliveria J. Antifungal activity of the essential oil of *Thymus pulegioides* on Candida, Aspergillus and dermatophyte species. *J. Med. Microbiol.* 2006; 55 (10), 1367 73.
- **27.** Mahmoud AL. Antifungal action and antiaflatoxigenic properties of some essential oil constituents. *Lett. Appl. Microbiol.* 1994; 2, 110 3.
- **28.** Farag RS, Daw ZY, & Abo-Raya SH. Influence of some spice essential oils on *Aspergillus parasiticus* growth and production of aflatoxins in a synthetic medium. *J. Food Sci.* 1989; 54, 74 6.