Evaluation of Mustard oil and temperature in the toxicity reduction of *Aspergillus flavus* in Walnut fruits

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DOI: https://doi.org/10.36077/kjas/2024/v16i1.10507

Received date: 13/11/2022

Accepted date: 28/12/2022

Abstract

The study aimed to isolate and identify fungi that produce mycotoxins from walnut fruits and the possibility of limiting their presence in the fruits and reducing their toxic effects. The molecular diagnosis in this study of the fungus Aspergillus flavus was carried out using the internal transcribed spacer (ITS) genetic marker. As for the results of testing the ability of some A. *flavus* isolates to produce aflatoxins B1 including the ammonia test, the results of this test showed the ability of four out (of six) isolates of A. flavus on the production of aflatoxin B1 by changing the color of the base of the coconut medium on which the fungus isolates are grown, with a percentage of 66.66 %. The results of the chemical analysis using TLC technology for Aspergillus flavus showed the ability of three isolates to produce aflatoxin B1 out of six isolates of A. flavus with a percentage of 50 %. As for the most important results of the effectiveness of mustard oil in inhibiting the growth of the fungus A. *flavus* in walnut fruits, the highest percentage of inhibition was at 15% concentration, which amounted to 57.33%. The most important results of the storage experiment after three months of storage, included the test of the effect of mustard oil and temperature on the growth and density of A. flavus. The test was the effect of mustard oil on the reduction ratio of aflatoxin B_1 as it reached the highest percentage of toxin reduction when adding mustard oil only, which amounted to 98.213%, compared to other treatments, the test was the effect of mustard oil and temperature on the amount of aflatoxin B1 toxin, which gave the highest rate when adding A. flavus to walnuts, which amounted to 13.425 micrograms/ g., At a temperature of 5 °C, it gave the lowest rate of aflatoxin B₁ toxin, which was 0.880 μ g.gm⁻¹. The highest rate was 8.620 μ g.gm⁻¹, at 25°C.

Keywords: mycotoxins, walnut fruits, Aspergillus flavus.



Introduction

diagnosis of fungi using the Polymerase chain reaction (PCR) technique and testing the extent of

walnut (Juglans nigra) is a northern contamination of walnut fruits with fungi and testing The hemisphere fruit tree with a temperate and subtropical the extent of susceptibility to fungi.

climate (30) and is native to Europe and Asia from the Balkans to southwestern China (15). It is widely **Materials and Methods**

cultivated all over the world because of its nutritional value (1). Walnuts are high in mono-saturated fatty

acids, vitamins, proteins, and carbohydrates, and as Nut samples were obtained from local markets well as it is rich in omega-3. People have been aware(American, Iranian and Chinese) for the purpose of of this fruit since the beginning of time. Because its obtaining isolates of toxin-producing fungi and then botanical structure is similar to and nutritious to the transferred to the Laboratory Plant Diseases at the human brain (29), it is associated with a lower risk of College of Agriculture - University of Kufa to isolate vascular and heart injury (21). During storage withand identify the fungi accompanying those samples.

fungal species, including Asperillus spp, Fusarium

spp, and *Pancillium spp*, they produce a wide range of

toxic receptors (27). They produce antioxidants

against these toxins in addition to their resistance to Isolation and diagnosis of fungi infecting walnuts high temperatures. (34) and many methods have been

used to control fungi in the store, including the use of Nus t samples were taken and isolated from them by chemicals such as sterilization using carbon dioxidedilution method and direct cultivation on PDA and others(23, 37), The use of chemicals is notmedium, where the fruits were cut into small pieces (1 without negatives as a result of environmentalcm) and were superficially sterilized with sodium pollution and the resistance shown by the pest towardshypochlorite solution 2% concentration for two these materials Chemicals and damage to human andminutes after which those pieces were washed with animal health as a result of the accumulation of these sterile distilled water and then placed on filter papers substances in the food chain (33), where plant extracts get rid of free water Then the fruit pieces were were used because they have antagonistic activity forplanted in plastic dishes (9 cm in diameter) containing many fungi and have preferred characteristics such as the medium of PDA by placing four pieces at a their rapid decomposition and lack of impact ondistance of 3 cm from the edge of the dish and a fifth human health (17). The use of some essential oilspiece in the middle of the dish. (20), and after the end derived from plants such as mustard oil because it hasof the incubation period, the growing fungi isolates anti-fungal, anti-bacterial, anti-microbial, and foodwere purified and then identified based on the pathogen activity (11 and 8). Therefore, the currenttaxonomic characteristics mentioned by Pitt and study aimed to investigate the fungi contaminatingHocking (24).

walnut fruits and test the toxins production and study. The percentages of appearance and frequency of fungi the possible measurements in reducing infection and appearance and frequency of fungi combined toxicity. It also aimed to confirm the



 $Percentage of appearance (\%) = \underbrace{\frac{\text{The number of samples in which the sex or gender appeared}}{\text{Total number of samples}} x 100$ $Percentage Frequency (\%) = \underbrace{\frac{\text{The number of isolates of one species}}{\text{The number of isolates of all fungi}} x 100$

Molecular diagnosis of isolates of the fungus A. flavus The isolates of the fungus A. flavus on liquid PD

Molecular diagnosis of the fungus *A. flavus* isolated medium by placing three tablets of five mm in diameter, one week old, in a flask of 250 mL volume, diameter, one week old, in a flask of 250 mL volume, at a rate of 100 m. L⁻¹. Thereafter the extract was filtered through Whatman No. 4. Then the filtrate was then placed in a separator funnel with a capacity of genetic marker

Aflatoxin B1 producing by A. flavus isolates

Detection of Aflatoxins by ammonia treatment method:

250 mL, then 30 mL of chloroform was added to it, then the funnel was shaken gently for 30 seconds while expelling the accumulated gases whenever needed (at least twice) and then left on the holder for

one minute in order to The two layers separate, I

The ability of *A.flavus* isolates isolated from theneglected the upper layer and took the lower layer and tested nuts to produce aflatoxin B1 was carried outrepeated the process for three times. Then the filtrate using (10) method, using coconut medium. In dishes was taken and placed in a clean, sterile flask and each with a diameter of 9 cm, then 3 dishes wereplaced in the electric oven at a temperature of 40°C inoculated with tablets of fungi for each isolate, byuntil it dried, then dissolved in one ml of chloroform. placing a disk with a diameter of 5 mm from the The presence of aflatoxin, B1 was detected using thinmedium of the PDA grown on The Fungi, at the age of sheet chromatography (TLC) technology with one week, in the center of the dish. A week later, the dimensions of 20-20 cm. The plates were activated in ability of the mushroom isolates to produce aflatoxins the electric oven was heated at 105 °C for one hour was revealed using a 20% ammonia solution by before use (7).

placing filter papers saturated with ammonia solution A light straight line was made on the TLC plate at a in the cover of the dish containing the isolate of the distance of 1.5 cm from the base of the plate, and 15 fungi growing in the middle of the coconut. The color μ l was taken by capillary tube of the standard toxin of the colony bases from transparent to pink or redAFB1 and placed on the line at a distance of 2 aflatoxins.

Detection of aflatoxins using Thin L Chromatography (TLC):

cm from the spot of the standard poison. *A.flavus* at Layer the same distance and in an amount equal to the amount of the standard poison, then the spots were left to dry and then placed in a separation tank containing



a separation system consisting of a mixture of 1- Preparation of A. flavus inoculum: A. flavus chloroform and methanol at a ratio of 2:98 v/v and inoculum was prepared by growing the fungus on monitored until the solution reached a distance of PDA medium for a week under temperature of $30 \pm$ approximately two cm from the upper end For the2°C. Then the fungal spores were harvested by adding plate, the plates were taken out and dried under 10 mL of sterile distilled water to each of the plates on laboratory conditions for 5 minutes and then which the fungus was grown, and then passed A examined under ultraviolet light at a wavelength ofsterile glass rod on the surface of the colonies to 365 nm and the presence of aflatoxin B1 was detected facilitate the process of separating spores from the by matching the migration factor Rf and fluorescenceconidia carriers. Then, planktonic spores were color of the standard poison with the color and collected for each fungus separately and their numbers migration factor of samples of extracts of A.flavuswere calculated using a hemocytometer slide. The isolates (31) fungal spore concentrations were adjusted at (106

Testing the efficacy of Indian mustard oil extract on spores/mL water). the growth of A.flavus on PDA medium:

2- Design of the experiment: The following treatments Indian mustard oil was obtained from one of the were applied to walnut fruits, where 10 mL of:

each of the repeaters of one treatment and placing it in

local markets, so it was added in proportions 5, 10 and 1-Distilled water only 200 gm of nuts

15 mL / 100 mL of PDA medium and poured into the2-10 mL Indian mustard oil 200 gm walnuts

dishes. A fungus from the tested fungi. Three plates of 3-The spores of A. flavus are stuck in only 200 gm of PDA medium (control treatment) were inoculatednuts

with PDA discs only on which the fungus A. flavus4-Suspended A. flavus spores + 10 mL of Indian was grown. The dishes were incubated at 25 ± 2 °Cmustard oil 200 gm of nuts

for a week. After that, the diameters of A. flavus Each treatment was stored in three bags containing colonies were calculated, and according to the each bag (200 gm) for a period of (30, 60, and 90) implemented treatments, the inhibition percentage was days at temperatures (5, 15, 25, and 35) °C, after extracted according to the Abbott equation reported which the contamination percentage was calculated in by Shaaban and Al-Mallah (35) where: each of the implemented treatments. Taking 10 g of

Inhibition% =
$$\frac{R_1 - R_2}{R_1} \times 100$$

R1 = maximum radial growth of a fungusan electric mixer with the addition of 100 mL of colony (control treatment). sterile distilled water, then it was mixed for 5 minutes,

R2 = maximum radial growth of the studied then a series of dilutions was made until 10⁻⁵, then one fungus colony in the treatment dishes. mL of the last dilution was grown on the PDA

medium. And three replications for each treatment, Evaluation of the efficiency of Indian mustard oil and and thus all treatments were dealt with, after which storage temperature in protecting walnut fruits from the dishes were incubated at a temperature of $25 \pm 2^{\circ}$ infection with the fungus A. flavus: C. After that, the number of colonies in each dish was

This experiment was carried out to find out therecorded after 3-4 days. Then, the number of active effectiveness of both mustard oil and temperature onunits of the fungus was calculated, according to the protecting walnut fruits from infection with A. flavus. coefficients, according to the following equation:



The number of active units = the average number of layer, and after three minutes, the upper and lower spores per replicate x the reciprocal of the dilution. layers were separated, keeping the lower layer. The

process was repeated twice. Then the extract was transferred to a separating funnel with a capacity of the 100 The concentrations of aflatoxin B1 were measured spectrophotometer, by а after extracting 100 mL, 15 mL of 1-N hydrochloric acid, 20 mL of mycotoxins from the samples of the implemented chloroform, and 20 mL of chloroform were added to treatments, the aflatoxin B1 toxin was extracted by it. Shake well and leave the funnel on the holder for taking 30 gm of walnuts from each treatment and then one minute. The lower layer was taken and to the of chloroform and then mixing the mixture for 10th upper layer, 20 mL of chloroform was added and transferring it to an electric mixer containing 100 mL extracted again. The two lower layers were collected Minutes after that, the mixture was filtered by filter and passed through filter paper containing a layer of paper, then the filtrate was taken and placed in a clean anhydrous sodium sulfate (Na₂SO₄) in order to get rid and sterile flask and placed in an electric oven at a of the remaining water. Then the filtrate was taken temperature of 40 ° C until dryness, then it was and placed in an electric oven at a temperature of 40 dissolved in 5 mL of chloroform and then the °C until dryness, after which it was dissolved in 5 mL concentration of the poison was estimated. As for the of chloroform (9) and then the poison concentration ochratoxin a poison, it was extracted by taking 30 gm was estimated by a Spectrophotometer. This method and Then to an electric mixer containing 100 mL of of chloroform. The atmosphere from each treatment depends by its nature, on the property of the the extraction solution consisting of acetonitrile-water compound to absorb light in the ultraviolet or infrared (90:10) mL, then mix the mixture for 10 minutes, then wavelengths, where there is a direct proportion filter The extract through what man No. 4. Then the between the absorption period and the concentration filtrate was placed in a separator funnel with a of the poison. It is possible by drawing a standard capacity of 250 mL, then 25 mL of hexane was added light absorption and the curve between the to it for the purpose of getting rid of fat (Defatting). concentration of the poison and extracting the Then The funnel was shaken gently for 30 seconds concentration value corresponding to the reading needed (at least twice). Then it was left on the holder compared with Standard poison concentrations for one minute in order for the two layers to separate. (Supplements 1 and 2), where the wavelengths were I neglected the top layer and took the bottom layer, 365 and 360 nm for aflatoxin B1 and ochratoxin A, and repeated the process Three times to ensure the respectively. The reduction ratios were calculated removal of fat. Then, 25 mL of distilled water, 8 mL

of saturated sodium bicarbonate solution (NaHCO₃),Using the following equation: and 25 mL of chloroform were added to the lower



Concentration of control treatment - concentration of the model

Reduction ratio = _____

x 100

Control treatment concentration

Statistical analysis:

All experiments were carried out in a completely the appearance of five species of fungi. The most common were *A. tubingensis*, *A. flavus*, *Fusarium* sp, randomized design, C.R.D (Complete Random Penicillium sp, and Trichoderma sp. Its frequency Design) as one-factor experiments, and the averages rates reached 55.17, 20.68, 13.79, 6.89 and 3.44%, were compared according to L.S.D (Less significant respectively. (Table 1) While the incidence of (100, differences) method and under a probability level of 66.66, 33.33, 33.33 and 16.66) %, respectively, 0.05 (4).

Results and Discussion

explains the reason for the dominance of *Aspergillus sp*. The ability to withstand drought and high humidity (2), and withstand high temperatures up to 50 $^{\circ}$ C (12).

The results of isolating fungi from walnuts showed

Isolation and identification of fungi accompanying the nut:

fungus type	(%) frequency	(%) apperance		
A. tubingensis	55.17	100		
A. flavus	20.68	66.66		
Fusarium	13.79	33.33		
Penicillium	6.89	33.33		
Trichoderma	3.44	16.66		

Table 1. Percentages of the frequency and appearance of fungi isolated from walnut

Molecular diagnosis of A. flavus isolates

tree and in the National Center for Biotechnology Information (NCBI) under the entry (Accession number ON394601.1.) Based on the sequences of its

The results of the molecular diagnosis of the nitrogenous bases for the ITS-rDNA region as well as pathogenic fungus *Aspergillus flavus* indicated that the sequences of global strains of the same pathogenic this isolate was established in the Neighbor-joining fungus were obtained from Gen Bank Data



that

are

structures

Repository. The genetic distances were calculatedstrain (16). The biological basis of the ammonia test using the neighbor-joining method. depends on the production of a group of yellow dyes

with

different

chemical

intermediate compounds in the aflatoxin synthesis

pathway, and that these dyes turn red or purple when combined /*with a basic solution - such as ammonium

Testing the ability of some isolates for *A. flavus* to produce aflatoxins B1:

Ammonia test:

The results of this test showed the ability of 4 out of 6 isolates of *A. flavus* to produce aflatoxin B1 by changing the color of the base of the coconut medium. on which the fungus isolates were grown, with a percentage of 66.66 %. The isolates that produced the most aflatoxins were Af4, while isolates Af2 and Af5 showed a medium ability to produce aflatoxins, and the rest of the isolates were weak in their production of aflatoxins (Table 2).

The fungal strains vary in their ability to secreteproduce aflatoxin quantitatively and qualitatively may aflatoxin, some of them may excrete aflatoxin toxins, be due to the different genetic content of the strains, while there are other strains that secrete more than one and this explains the gradient in the red color. (26). type of mycotoxin depending on the type of fungal

medium coc	onut (CEA)	J I	
-	fungul isolation	The ability to produce aflatoxin B1	

Table 2. Testing the ability of some isolates of A. *flavus* to produce aflatoxin B1 in

	J	
Af1	+	
Af2	++	
Af3	-	
Af4	+++	
Af5	++	
Af6	-	

(+): the color of the middle base changed to pink, (-): the color of the middle base did not change.

Detection using thin sheet chromatography (TLC) The results of this test showed the ability of some isolates of *A. flavus* to produce aflatoxin B1, whereas the test showed the ability of 3 isolates to produce



aflatoxin B1 out of 6 isolates of *A. flavus* at aproduce aflatoxin B1. The difference in the ability of percentage of 50%, as in (Table 3). The fungusisolates to produce aflatoxins B1 may be attributed to isolates varied in their toxin production, and isolate the presence of genetic differences between the fungal AF4 was the most toxin-producing isolate based onisolates (18). It is noted that the percentage of isolates the intensity of its fluorescence. And Af5 and Af2 areproducing aflatoxin B1, which were detected by this the least productive of aflatoxin production. Thesetechnique, is less than the number of isolates results initially agree with what one study indicated producing aflatoxin B1 using the ammonia reaction about the ability of 75% of *A. flavus* isolates totechnique. Accordingly, the thin plate produce aflatoxin B1. And an approach to what waschromatography method is the most accurate method mentioned (14). This indicated that 38.88% of *A* for determining the isolates producing mycotoxins in *flavus* isolates isolates are able togeneral, including aflatoxin B1.

Table 3. Testing the susceptibility of a number of A. *flavus* isolates to produce aflatoxinB1 isolated from walnuts by thin plate chromatography (TLC) method

fungul isolation	The ability to produce aflatoxin B1					
Af1	-					
Af2	+					
Af3	-					
Af4	+++					
Af5	++					
Af6	-					

(+) aflatoxin B1-producing isolate (-) non-aflatoxin B1-producing isolate

Evaluation of the effectiveness of differenthighest percentage of inhibition was at the concentrations of mustard oil in inhibiting the Qatariconcentration 15%, which amounted to 57.33%, and growth of the fungus *A. flavus*: the inhibition percentage in concentrations 5, 10 and

The results of the experiment showed the treatment (table 4) This is similar to what previous studies of nuts contaminated with the fungus *A. flavus* with indicated to the effectiveness of many medicinal fungus *A. flavus*, as the percentage of inhibition producing fungi (28).



Table 4. Effect of different concentrations of mustard oil on the growth of A. Flavusgrown on PDA

Transactions	retarding percentage% Colonies diameter			
		average (cm)		
PDA. Comparison	0.00	7.5		
5	48.00	3.9		
10	52.00	3.6		
15	57.33	3.2		
L.S.D.(0.05)	1.631	0.1883		



Figure 1. shows the evaluation of the effectiveness of different concentrations of mustard oil in inhibiting the growth of the fungus A.concentration of 5% and number (3) which different represents a concentration of 10% and number flavus, are three concentrations (5%, 10% and 15%). No. (1) (4) represents a concentration of 15%. represents the comparison, which is A. Effect of mustard oil and temperature on the growth and density of A. flavus after three months of storage. tubingensis only, and No. (2) represents A



The results of this test showed significantis the presence of erucic acid in mustard seed oil, differences in the growth rate of the number of which plays a role in phytoremediation and reproductive units of A. flavus_spores, as it gave theantioxidants (36). The table also indicates that there highest growth rate when A. flavus was added to theare significant differences in the effect of temperature nut, which amounted to 3.200 x 10⁻⁵ spores/gmon the number of reproductive units of A. flavus, so compared to the control treatment (0.25 x 10⁻⁵ spores.the storage treatment at a temperature of 35 °C gave gm⁻¹). In Table (5), while the mustard oil reduced thethe lowest rate in the number of reproductive units of numbers of microbial units of the fungus, where thethe fungus, which amounted to 0.575×10^{-5} spores. treatment of A. flavus + mustard oil gave 0.575×10^{-5} gm⁻¹ compared to the other treatments (5, 15, 25). mL, spores / gm⁻¹, compared with the treatment of addingwhich gave (1.080, 1.140, 1.560 \times 10⁻⁵ spores. g), fungus only, which amounted to 3.200×10^{-5} spores.respectively. The highest rate in the number of Gm-1, while the oil treatment was given Mustard only reproductive units of the fungus was (1.560×10^{-5}) had the lowest growth rate in the number of spores. gm⁻¹) at 25 °C compared with other reproductive units, which amounted to (0.000 x 10⁻⁵treatments, so crop losses were recorded from 20% to spores. gm^{-1}) compared to the control treatment (0,25550% in developing countries due to inappropriate x 10^{-5} spores. gm⁻¹). Whereas, the addition of distilled storage practices (22).

water gave only 1.425 $\times 10^{-5}$ spores / g, and the reason

Table 5. Effect of mustard oil and temperature on the growth and density of A. flavus

Transactions	Number of reproductive units of spores (spore/gm) x 10^5				The average
	5م°	15م°	25م°	°م35	
Comparison	0.0	0.4	0.2	0.3	0.225
distilled water	0.4	1.6	2.6	1.1	1.425
Mustard oil	0.0	0.0	0.0	0.0	0.000
A. flavus	4.2	2.9	4.6	1.1	3.200
A. flavus + Mustard oil	0.8	0.8	0.4	0.3	0.575
The average	1.080	1.140	1.560	0.575	1.080

after three months

L.S.D. 0.05 coefficients = 0.0715, temperatures = 0.0639, coefficients + temperatures = 0.1429

Effect of mustard oil and temperature in reducing theoil, *A. flavus*, mustard oil + *A. flavus*) (98.213, 0.000, toxicity of aflatoxin B1 produced by *A. flavus* after71.188) % respectively, as indicated in Table (6), three months. where the highest percentage of toxin reduction when

This test showed the effect of mustard oil on the compared to other treatments, and the lowest reduction ratio of aflatoxin B1 when it gave (mustard percentage of toxin reduction in the treatment of



adding *A. flavus* only, which amounted to (0.0000%)respectively, and the highest rate was 42.953% at a compared to other treatments because it has many Thetemperature of 25 °C compared to other treatments, characteristics that make mustard a valuable botanicaldue to the conditions that promote the growth of fungi medicine (19).The table also indicates that there arealways may not lead to the production of mycotoxins. significant differences in the effect of temperature onIn general, it water activity above 0.78, relative the percentage of toxin reduction produced by thehumidity between 88% and 95%, and temperature fungus. % compared to the other treatments (5, 15, 25between 25 and 30 °C are favorable conditions for the °C) which gave (42.855, 41.927, 42.953) %,growth of fungi and production of mycotoxins (32).

Table 6. Effect of mustard oil and temperature in reducing the toxicity of aflatoxin B1produced by A. flavus after three months

Transactions			The average			
		Temperatures				
-	5م°	15م°	25م°	35م°		
Comparison	0.0	0.0	0.0	0.0	0.00	
Mustard oil	100.0	92.85	100.0	100.0	98.213	
A. flavus	0.0	0.0	0.0	0.0	0.0000	
A. flavus +Mustard oil	71.42	74.86	71.81	66.66	71.188	
The average	42.855	41.927	42.953	41.665	42.350	

 $\overline{\text{L.S.D. 0.05 Transactions}} = 0.5109$, temperature = 0.5109, Transactions + temperature = 1.0219

Effect of mustard oil and temperature on the amount *flavus* only gave (13.425 μ g.g⁻¹). The addition of of aflatoxin B1 microgram/gm produced by *A. flavus* mustard oil only had the lowest rate, which amounted after three months of storage: to (0.075 μ g/gm) compared to the control treatment

 $(2.708 \ \mu g. g^{-1})$. While the treatment of adding distilled The results of this test showed significant water gave only 8.708 µg/gm. This is similar to what produced by A. *flavus* fungus, as it gave the highest⁽²⁵⁾ said that mustard oil is used as a treatment and rate when adding A. *flavus* to walnuts, which amounted to(13.425 μ g.g⁻¹) compared to the control treatment, which amounted to $(2.708 \ \mu g.g^{-1})$ As temperature on the amount of aflatoxin B1 toxin there were significant differences in the effect of amount of aflatoxin B1, the treatment of A. flavus + flavusindicated in (table 7) While mustard oil reduced the temperature of 5 °C gave the lowest average in the mustard oil gave 3.800 µg.g⁻¹ compared with the amount of aflatoxin B1 toxin, which was 0.880 µg.g⁻¹ treatment of adding A. flavus only, which amounted to compared to the other treatments (15, 25, 35) m $^{\circ}$ (13.425 μ g.g⁻¹), while the treatment of adding A.



which gave (8.307, 8.620, 4.760) μ g.g⁻¹ respectively.on the biochemical, genetic and physiological And the highest rate was (8.620 μ g. g⁻¹), at acharacteristics of the fungal strains and also on temperature of 25°C compared with the otherexternal factors such as temperature, humidity and treatments. Because the production of mycotoxinsspecific environments (12). from *Aspergillus, Pencillium* and *Fusarium* depends

Table 7. Effect of mustard oil and temperature on the amount of aflatoxin B1

Transactions	T	ıg/gm	The average		
_	Temperatures				
-	5م°	15م°	25م°	[°] م35	
Comparison	0.0	4.2	5.4	1.2	2.708
distilled water	2.6	14.1	9.0	6.6	8.075
Mustard oil	0.0	0.3	0.0	0.0	0.075
A. flavus	1.4	18.3	22.0	12.0	13.425
A. flavus +Mustard oil	0.4	4.6	6.2	4.0	3.800
The average	0.880	8.307	8.620	4.760	5.617

microgram/gm produced by A. *flavus* after three months of storage

L.S.D. 0.05 coefficients = 0.3742, temperatures = 0.3347, coefficients + temperatures = 0.7484

Conflict of interest

The authors have no conflict of interest.

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