RAPID DETECTION OF AFLATOXIGENIC PRODUCING ATRAINS OF ASPERGILLUS FLAVUS FROM POULTRY FEES BY UV LIGHT AND AMMONIA

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ABSTRACT

A total 180 samples of pellet poultry feed were collected from poultry feed stores and local markets .They were cultured on potato dextrose agar(PDA) and malt extract agar (MEA) for isolation and identification of *Aspergillus flavus*.They were Sub-cultured on sabouraud dextrose agar(SDA) .Then the isolates were cultured on coconut agar medium(CAM) . The rapid detection of 50 selected isolates of contaminated poultry feed samples with *A.flavus* were achieved by blue –green fluorescence under UV light and by ammonia vapor on CAM medium to determine the aflatoxigenic isolates of *A.flavus*. The detection by UV light revealed that 26 (52%) of isolates were aflatoxigenic (positive) by produce blue-green fluorescence under UV light at 356nm , and also 26 (52%) of isolates were aflatoxigenic (positive) by turned the colony reverse to pink color by exposure to ammonia vapor.

INTRODUCTION

Aspergillus flavus is widely distributed in nature and is largely found at cereal and grains. Before harvest or during storage, *A. flavus* grows on agricultural crops (1). Its growth is affected by the environmental condition such as temperature and relative humidity (2).

Aflatoxins are difuranceumarin derivatives. They are very slightly soluble in water (10–30 μ g/mL), in non-polar solvents they are insoluble , and soluble in moderately polar organic solvents (e.g. chloroform and methanol) and extremely soluble in dimethyl sulfoxide (3) .Under the influence of ultraviolet light they are unstable and in the presence of oxygen, to extremes of pH (< 3, > 10) and to oxidizing agents (4).

Aflatoxins are produced only by a closely related group of Aspergilli: *A. flavus, A. parasiticus* and *A. nomius* strains (5). Other species such as *A. bombycis, A. pseudotamari* and *A. ochraceoroseus*

are also aflatoxin-producing species, but they are found less frequently (6).Aflatoxins cause a problem concerning many commodities also aflatoxin B1(AFB1) act as carcinogenicity, mutagenicity and acute toxicology and determined it to be a human carcinogen. Aflatoxins are common occurrence in feedstuffs, feeds. Aflatoxicoses , the disease caused by exposure to aflatoxin have made severe economic losses in the poultry industry, affecting ducklings, broilers, layers, quail and turkeys to cause clinical signs include anorexia, decreased weight gain, decreased egg production, hemorrhage, embryotoxicity, and increased susceptibility to environmental and microbial stressors (7).

Blue fluorescence a method used for developing qualitative cultural methods for detecting aflatoxigenic *Aspergillus* species grown on suitable media. This techniques use either solid media, such as coconut agar medium (CAM) and potato dextrose agar (PDA) or liquid medium, like aflatoxin producing-ability medium(APA), and a medium with steep liquor (8,9), and achieved by cut a small plugs from *Aspergillus* colonies on medium to culture on the other media. The aflatoxins producer *Aspergillus* were detected under long-wave UV light (365nm). This rapid identification to determine aflatoxigenic isolates from non-aflatoxigenic by appear blue to blue – green fluorescent to aflatoxigenic , and nonaflatoxigenic is non-produce fluorescent (10). Ammonium hydroxide vapor-induced color change a rapid and sensitive method , also for detection of aflatoxigenic and nontoxigenic strains of *Aspergillus* (11). In this method a single colony was grown in the centre of Petri dish. The reverse of colony of aflatoxigenic *Aspergillus* strains turned to pink color when their medium were exposed to ammonia vapor by dropped of ammonia hydroxide on it but nonaflatoxigenic is no color produce (12). This study aims to detect the aflatoxigenic *Aspergillus flavus* by UV light test and by ammonia vapor test which were isolated from poultry feed.

MATERIALS AND METHODS

Collection of samples

A total 180 samples of pellet poultry feed were collected from poultry feed stores and local markets at Basrah governorate during one year from Sep. 2014 to Apr. 2015. Samples were stored for 2-3 days in sterile plastic containers at room temperature (22-25°C) in laboratory. After stored, they were prepared for fungi isolation and identification.

Isolation and identification of A.flavus

Suspension of 20 g of the poultry feed samples with 180 ml of saline solution (0.85% Sodium Chloride) with 0.05% Tween 80 on a horizontal shaker for 30 minutes. Then 0.1 ml of suspension

was inoculated on PDA and MEA(13,14). The pure culture was incubated at 25 °C \pm 2 and after 7 days (15). After incubation , the macroscopic and microscopic distinct colonies by using lactophenol cotton blue were done. The morphological characteristics of *A.flavus* isolates were described microscopically according to Domsch and Gams (16) and Klich (17) .Sub-culture on SDA was done , and also on CAM according to Davis *et al* (18). Then the cultures were incubated at 25°C \pm 2 for 7 days (19).

Coconut based medium test

It was done by fluorescence on coconut agar medium(CAM). A preliminary screen for aflatoxin producer *Aspergillus* was performed on the basis of emission of blue to blue – green fluorescence after UV light excitation at 365 nm after growing the isolates on coconut agar medium , because this agar is inductive of aflatoxin production (20). Producer isolates can be identified by fluorescence in the reverse side of the culture CAM in glass Petri dishes(18, 21).Five millimeter diameter sterile cork borer was used to make a hole in the center of CAM medium in petri dish .The isolate was inoculated of a mass of conidia by cork borer to the hole at the central point Petri dish of CAM , then they were incubated at 28 °C for 7 days. nonaflatoxigenic *Aspergillus niger* was used as control (20,22).

Ammonia vapor test

The fungal isolates were inoculated on CAM as single colonies by cork borer (5mm) diameter in the center of plate and incubated in the dark at 28 °C. for 7 days. The dish was inverted and 1 or 2 drops of concentrated ammonium hydroxide solution are placed on the inside of the lid of Petri dish. Then the Petri dish inverted over the lid containing the ammonium hydroxide. A control as was mentioned in previous test was prepared.

RESULTS

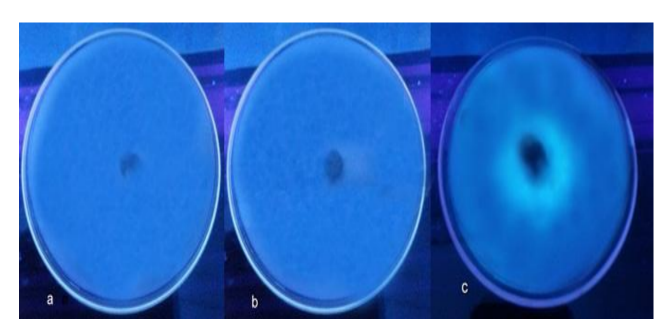
Fifty *A.flavus* isolates were considered out of 180 samples. The detection of aflatoxigenic and nonaflatoxigenic *Aspergillus flavus* by using UV light and ammonia vapor revealed that 26 (52%) of isolates were aflatoxigenic (positive) and 24(48%) of isolates were nonaflatoxigenic (negative) for both methods (table1,2). The detection by UV light 365nm recognized aflatoxigenic by produce blue-green fluorescent colonies in the center of glass Petri dish of CAM in the reverse, from nonaflatoxigenic which were nonproducing fluorescent colonies, similar to the control isolates of nonaflatoxigenic *A.niger*, (figure 1).

Table(1): Number of aflatoxigenic -producing isolates of A.flavus on the CAM

Detection method	Number of detected isolates	Positive isolates	Positive isolates(%)	Negative isolates
Coconut based medium detection	50	26	52	24
Ammonia vapor detection	50	26	52	24

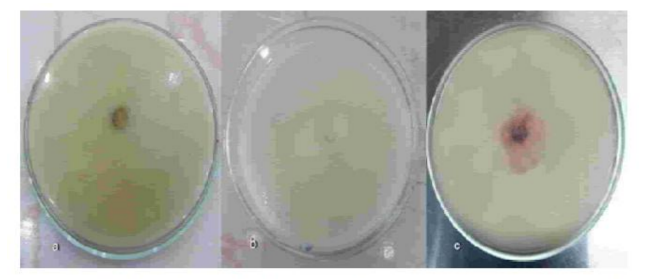
Table (2) : The results of aflatoxigenic A.flavus isolates recovered from poultry feed samples which was obtained by UV light detection , ammonia vapor detection on CAM.

No. of isolate	Coconut based medium test	Ammonia vapor test	No. of isolate	Coconut based medium test	Ammonia vapor test
1	+	+	26	+	+
2	+	+	27	-	-
3	+	+	28	+	+
4	+	+	29	+	+
5	-	-	30	+	+
6	-	-	31	-	-
7	-	-	32	+	+
8	-	-	33	+	+
9	+	+	34	+	+
10	+	+	35	+	+
11	+	+	36	+	+
12	-	-	37	-	-
13	-	-	38	-	-
14	+	+	39	+	+
15	+	+	40	-	-
16	+	+	41	-	-
17	+	+	42	+	+
18	+	+	43	-	-
19	-	-	44	-	-
20	-	-	45	-	-
21	+	+	46	-	-
22	-	-	47	-	-
23	+	+	48	-	-
24	-	-	49	-	-
25	+	+	50	-	-



Figure(1): Showed the detection of aflatoxigenic *A.flavus* by CAM under UV light at 365nm .(a) control of nonaflatoxigenic isolate of *A.niger*, (b) nonaflatoxigenic *A.flavus* (negative) isolate, and (c) aflatoxigenic *A.flavus* (positive) isolate, showing a blue-green fluorescent ring around the colony.

Aflatoxin detection by ammonia vapor to characterize as aflatoxigenic isolates *A.flavus* which produced pink to red color colonies in inverted petri dish by applying 1 or 2 drops of concentrated ammonia hydroxide solution on the inside of the lid, but no color change occurred in nonaflatoxigenic isolates, (figure 2).



Figure(2): Showed the detection of aflatoxigenic *A.flavus* by ammonia vapour.(a) control of nonaflatoxigenic isolate of *A.niger*, (b) nonaflatoxigenic *A.flavus* (negative) isolate, and (c) aflatoxigenic *A.flavus* (positive) isolate, showing a pink-red ring around the colony.

DISCUSSION

By using UV light technique in this study, 26 isolates (52%) from a total 50 isolates of *A*,*flavus* were aflatoxigenic can colored with blue –green fluorescence (positive) on reverse of glass Petri dish of CAM with compared to nonaflatoxigenic showed no color (negative) results and considered as negative. The same results above by ammonia vapor detection were reported in which the colony of aflatoxigenic *A*,*flavus* turned to pink color , while no change in color with non aflatoxigenic isolates. This mean that the number of aflatoxigenic isolates of *A*,*flavus* were equal by UV light at 365 nm and ammonia vapor on CAM. This result is similar with those obtained by Yazdani *et al* (11) , Saito *et al* (12) , Zarari *et al* (24), Nair *et al*(25) and Sudini *et al* (26). While there was difference with Riba *et al* (23) whose confirmed that the cultures of aflatoxigenic *Aspergillus* were tested for 365 nm UV light fluorescence and for bright orange-yellow colony reverse coloring, and also this study disagrees with study of Fani *et al* (27) , which reported only (25.6 %) positive isolates of aflatoxigenic *A*,*flavus* by fluorescence detection on CAM , while less isolates(12 %) were identified as aflatoxigenic using ammonium vapor detection.

CONCLUSIONS

Aflatoxin is a major problem in developing countries where contaminated food commodities may readily reach food stores and homes . It is important to know that the effect of aflatoxin on animals extend beyond the symptoms. There are high percentage of aflatoxigenic *A*,*flavus* in poultry feed product.

الخلاصة

تم جمع 180 عينة من اعلاف الدواجن المركزة من مخازن أعلاف الدواجن و الأسواق المحلية . تم زرعها على الوسطين تم جمع (PDA) و (MEA) لعزل وتحديد سلالات الفطر Aspergillus flavus . وقد تم اعادة زرعها على وسطين ثانويين (SDA) و (MEA). تم الكشف السريع لى 50 سلالة مختارة من عينات علف الدواجن المتلوثة بـ Aflavus بواسطة ضوء (SDA) و (SDA). تم الكشف السريع لى 50 سلالة مختارة من عينات علف الدواجن المتلوثة بـ Aflavus بواسطة ضوء (SDA) و (SDA). تم الكشف السريع لى 50 سلالة مختارة من عينات علف الدواجن المتلوثة بـ Aflavus المنتوبة ضوء (SDA) و (SDA). تم الكشف السريع لى 50 سلالة مختارة من عينات علف الدواجن المتلوثة بـ Aflavus المنتجة الأشعة فوق البنفسجية بطول موجي 356 نانومتر وبخار الأمونيا على الوسط CAM لتحديد سلالات الفطر Aflavus المنتجة الأشعة فوق البنفسجية أن 26 (25٪) من العزلات كانت Aflatoxigenic (إيجابية) من خلال انتاج الضوء الفلورسنتي الاخضر المزرق تحت ضوء الأشعة فوق البنفسجية فوق البنفسجية في 356 نانومتر ، وأيضا لوحظ

ان 26 (52٪) من العزلات كانت aflatoxigenic (ايجابية) من خلال تحول لون ظهر المستعمرة الى اللون الوردي عند التعرض لبخار الأمونيا.

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