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AFLATOXICOSIS IN BROILERS IN KHARTOUM STATE

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المستخلص

اجرِت هذه الدراسة للكشف عن وجود الأفلاتوكسين وتحديد تركيزه في عَلائِق الدجاج الاحمر وأَبْاده مع توضيح تأثيرات هذه السموم المرضية على الكبد.

تم جمع ١٠٠ عينة عُليقة (العليقة الكاملة النهائية، أمِّيَاز الفول السوداني و الّذرة) و ١٠٠ اَعيِّنة كبد ب بصورة عشوائية من ١٠ مزارع للدجاج الاحمر بولاية الخرطوم.

بعدعملية استخلاص الأفلاتوكسين من العَلائِق وأَباده تم الفحص باستخدام اختبار الإليرة.

وجد أن جميع عَناتَاتَ العَلائِق تحتوي على الأفلاتوكسين، تراوح تركيز الأفلاتوكسين في عِناتَات العَلِيقَة النهائية (١٠٠ عِنِّة) بين ١ - ٩٧ جَزءَ من البيليون بمتوسط ٤٥.٥ جَزءَ من البيليون، بينما كان تركيزه في عِناتَات أمِّيَاز الفول السوداني (٢٠ عِنِّة) بين ٤٨ - ١٠٩ جَزءَ من البيليون بمتوسط ٩١.٤ جَزءَ من البيليون وكان تركيزه في عِناتَات الّذرة (٢٠ عِنِّة) بين ٧ - ١٠ جَزءَ من البيليون بمتوسط ٩٣.٣ جَزءَ من البيليون. وجد أن ١٧ عينة من عِناتَات الكبد تحتوى على أفلاز بأَباده من الأفلاتوكسين تراوحت بين ٢ - ١٢ جَزءَ من البيليون بمتوسط ٤.٩ جَزءَ من البيليون.

أُشتملَت التغييرات المرضية العيانية في عِناتَات الكبد المصابَة على مناطق نزائف كِبيرة، تضخم و شحوب بالاسياف للتِحَقان و النَخْر. التغييرات المرضية السبِّيجية أُشتملت على احْتِقان الأوردة المركزية، نزائف، فَراغات سيتوبلازمية(تَكَسَ استشفات). نَخْر و اِرتِشاح ليماوا في شكل عقدات.
Abstract

The occurrence and concentration of aflatoxins in broilers feedstuff and livers were investigated and their toxicopathological changes in the liver were described. 100 samples of feed including (Finished Total ration, groundnut cake and dura (sorghum)) and 100 liver samples were collected randomly from 10 poultry (broiler) farms in Khartoum state. After extraction, the aflatoxins were detected using ELISA test.

Aflatoxins were detected in all feedstuff samples collected. The concentrations in total finished ration samples (60) varied between 10 and 97 ppb with a mean of 45.5 ppb, groundnut cake samples (20) ranged between 84 and 109 ppb with a mean of 91.4 ppb and in dura (sorghum) samples (20) varied between 7 and 10 ppb with a mean of 9.3 ppb.

Seventeen out of one hundred liver samples examined were positive for aflatoxins residues and the concentrations varied between 2 and 12 ppb with a mean of 4.9 ppb.

The affected liver showed hepatomegaly, paleness, wide areas of hemorrhages and necrosis. Histopathology revealed congestion of central vein, hemorrhages, vacuolar degeneration, necrosis and nodular lymphoid infiltration.

Introduction

Aflatoxins (AF) are a group of heterocyclic toxic metabolites of toxigenic fungi *Aspergillus flavus* and *A. parasiticus*, (Diaz, 2005). They are difuranocoumarin derivatives produced by a polyketide pathway. Four major aflatoxins produced in feedstuffs and foods are aflatoxins B1, B2, G1 and G2. The most potent and the most frequently occurring of the four compounds is aflatoxin B1. Aflatoxin M1 is a metabolite of aflatoxin B1 that occurs in various animal tissues and fluids from animals. Infection and production of aflatoxins in field crops by species is often associated with drought stress and insect damage (Richard et al., 1993). When AF is ingested by animals, it is readily absorbed via the gastrointestinal tract into the portal blood and is carried to the liver where it is metabolized. (Hsieh, 1983). The main lesions of aflatoxicosis in birds appear in the liver which shows perportal necrosis with bile duct proliferation and fibrosis, jaundice, generalized oedema and hemorrhages, and depletion of lymphoid organs. (Charlton, 2006).
Very little information is available regarding poultry aflatoxicosis in Khartoum state. Hence this study is an attempt to evaluate the magnitude of poultry aflatoxicosis and examine the interrelationship of aflatoxins to poultry health and production.

Materials and Methods

Samples collection:
The samples were collected randomly from 10 poultry (broiler) farms of different capacities i.e. Farms A, B and C, are big intensive farms (above 25,000 chicks) located in Khartoum municipality. Farms E, F and G are medium farms (5,000 - 15,000 chicks are located in Khartoum North municipality. Farm D which is an intensive farm and farms H, I and J which are small extensive farms (less than 5,000 chicks) are located in Omdurman municipality.

Feedstuff samples:
100 samples, each weighing 250gm were obtained, i.e. 10 samples were taken from each farm. These samples comprised 6 samples finished ration, 2 samples groundnut cake and 2 samples Dura (Sorghum). Finished ration samples collected included, two samples each, from recently prepared ration, from feeders and from sacs in the stores.

Liver samples:
100 liver samples were collected from the 10 farms (10 samples each) immediately after slaughtering. Each sample was divided into two parts, the first one was blended and stored at -20°C for ELISA test and the other one was kept in 10% formol saline for histopathological examination.

Samples Extraction:

Feedstuff samples:
This was done according to manufacturer instruction (GIPSA FGIS 2005-12, Neogen Corporation USA/Canada). Samples were grounded using high speed blender for 5 min, 25 ml of 70% methanol was added to 5gm of the grounded sample, and shaked vigorously for 3 min. The extract was then filtered using whattman filter paper.
Liver samples:
Liver extracts were done according to Gathumbi et al. (2003). 5 ml of Ice-cold (-18°C) methanol 5%-acetone10% (50:50) were added to 2 gram of blended liver tissue, the mixture was homogenized for 5 min using a magnetic stirrer. The homogenate was then placed at (-18°C) for 20 min, before 5 mL of Phosphate Buffered Saline (PBS) was added and a further 30-min homogenization on a magnetic stirrer was carried out. The homogenate was centrifuged at 1500 rpm for 15 min and the supernatant was recovered. (0.5 mL) of the supernatant was diluted 1:5 in PBS and defatted by homogenization for 30s with(5 mL) 10% n-heptane followed by a 5 min centrifugation at 2000 rpm and then the lower methanol-acetone-PBS layer was recovered This extract was stored at 2 - 8°C until analyzed.

Enzyme linked Immuno-sorbent Assay (ELISA):

Assay principles:
The aflatoxins concentration in both feedstuff and liver extracts was measured using Veratox® aflatoxins quantitative kits (GIPSA FGIS 2005-12, Neogen Corporation USA/Canada).

Histopathological Techniques:
The liver specimens for histopathology were taken soon after slaughtering and kept in 10% formol saline ,processed and sections 4-5µ were prepared and stained with haematoxyline and eosin (Culling, 1974).

Results

Feedstuff:

Finished ration samples:
Aflatoxins were detected in all finished ration samples (60). The concentrations varied between 10 and 97 ppb with a mean of 45.5 ppb (Table 1)..
Recently purchased ration samples:
The mean concentration was calculated as 35±13.4 ppb. Farm J scored the highest concentration i.e. 57 ppb where as farm C showed the lowest concentration of 10 ppb (Table 1).

Stored ration samples:
The mean concentration was calculated as 52.5±25.8 ppb. Farm H and I scored the highest concentration i.e. 97 ppb where as farm C showed the lowest concentration 13 ppb (Table 1).

Samples taken from feeders:
The mean concentration was calculated as 50±21.3 ppb. Farm D scored the highest concentration i.e. 97 ppb where as farm G showed the lowest concentration 12 ppb (Table 1).

Groundnut cake samples:
Aflatoxins were detected in all samples collected (20), and the concentrations ranged between 72 and 120 ppb with a mean of 94.1±13.6 ppb (Table 1).

Dura (Sorghum) samples:
Aflatoxins were detected in all samples collected (20), and the concentrations varied between 7 and 12 ppb with a mean of 9.3±1.3ppb (Table 1).

Liver samples:
Seventeen out of one hundred liver samples examined were found positive to aflatoxins. The positive liver samples were found in all farms examined (10) except farm B and C. Aflatoxins residues varied between 2 ppb in farm D, E and J and 12 ppb in Farm J with a mean of 4.9±2.9 ppb. ix out of 10samples were positive in farmJ (Table 1).
Gross lesions:
Some livers showed wide areas of hemorrhage (Fig 1), hepatomegaly and paleness (Fig 2). Others revealed marked congestion and necrosis.

Histopathological findings:
The liver showed congestion of central vein, diffuse hemorrhages and necrosis (Fig 3), hydropic degeneration of the hepatocytes (Fig 4) and lymphoid nodules.

Fig: 1. Liver showing wide areas of hemorrhages. (Farm H), 5 ppb aflatoxins residue in the liver.

Fig: 2. The right lobe of the liver is larger than the normal left.
- Both of them are pale in color. (Farm I), 10 ppb. Aflatoxins residue in the liver was
Table 1: Concentration of Aflatoxins in feedstuff and liver samples (ppb).

<table>
<thead>
<tr>
<th>Farm</th>
<th>Finished ration</th>
<th>Peanut cake</th>
<th>Dura (Sorghum)</th>
<th>Liver samples</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Purchased recently</td>
<td>Stored</td>
<td>From feeders</td>
<td>S1</td>
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<tr>
<td>Big intensive farms</td>
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<td>A</td>
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<td>47</td>
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<td>B</td>
<td>37</td>
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<td>Medium farms</td>
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<td>E</td>
<td>20</td>
<td>27</td>
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<td>G</td>
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<tr>
<td>Small extensive farms</td>
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<td>D</td>
<td>50</td>
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<td>96</td>
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<td>H</td>
<td>26</td>
<td>16</td>
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<td>I</td>
<td>13</td>
<td>36</td>
<td>97</td>
<td>73</td>
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<td>J</td>
<td>57</td>
<td>37</td>
<td>70</td>
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</table>

Key: S= sample.
Fig. 3 - Liver showing hemorrhages and necrosis as judged by disappearance of nuclei and cytoplasmic changes. (Farm J), 12 ppb aflatoxins residues in the liver H & E 400 X

Fig. 4 liver showing Cytoplasmic vacuolation of hepatocytes. (Farm I), 10 ppb aflatoxins residues in the liver. H & E 400 X
Discussion

Aflatoxins contamination can occur in a wide variety of feedstuffs including corn, sorghum, wheat, groundnuts, soya, rice, cottonseed and various derivative products made from these primary feedstuffs (Busby and Wogan, 1979).

No significant differences in the mean aflatoxins concentration between the samples collected from the feeders (50 ppb) those collected from stores (52 ppb) or that purchased recently (35 ppb). This could be attributed to that the upper limit of the range in some samples collected from feeders and stores were found to be high. In a survey conducted in India in different poultry farms, Banerjee and Shetty, (1992), reported concentrations of aflatoxins in feedstuff samples ranging between 5.5 and 90 ppb which is relatively nearer to our result (10 – 97 ppb). The maximum aflatoxins level in the broiler feedstuff (97 ppb) in this study is higher than the value in poultry feedstuff (50 ppb) reported by Oruc et al., (2007) in Turkey and lower than the value reported by Dawlatana et al., (2002) in Bangladesh (160 ppb).

This study showed that the concentration of aflatoxins in groundnut cake samples ranged between 72 and 120 ppb. This range is relatively narrow compared to the wide ranges (1-244 ppb) obtained by Yoshizawa, (1981) in Philippines. These differences could be attributed to the wide variation in climatic, environmental conditions and in differences in managerial levels between Sudan and Philippines. On the other hand, aflatoxin concentration in sorghum samples obtained in the study is low (7-12ppb) Logrieco et al., (2003) also found little or no aflatoxins in the sorghum samples they analyzed.

In general aflatoxin levels currently reported in feed stuff are considered high according to the internationally accepted rule that growing poultry should not receive more than 20 ppb aflatoxins in the diet (Jones et al., 1994). The liver is the mostly affected organ when poultry is fed aflatoxins (Charlton, 2006). This study showed that the concentrations of aflatoxins residues in liver samples varied between 2 and 12 ppb with a
mean of 4.9 ppb. In an experimental study Mintzlaff et al., (1974) found that when the broiler chicks were fed rations containing aflatoxins at concentration of 15000 µg/kg for 8 weeks, aflatoxins residues in the liver tissue were 15 ppb. In another study Fernandez et al., (1994) reported a range of 0.29 – 0.63 ppb aflatoxins residues in livers of broilers fed on aflatoxins contaminated feedstuff (5 mg/kg) for 32 days. This differs from our results and might be due to differences in exposure duration and concentration of aflatoxins in feedstuff and the breeds used. In this study, hepatomegaly, and paleness of the liver were seen in association with aflatoxins; this has been previously, observed by (Kumar and Balachandran, 2009), congestion and haemorrhage were also evident in most liver which agree with Lamont(1979), and Slowik et al., (1985). As reported by Lawlor and Lynch, (2005) necrotic changes were recognized grossly in most samples examined.

The microscopic lesions of the liver revealed congestion of central vein, hemorrhages, slight to moderate cytoplasmic vacuolation (hydropic degeneration), necrosis and lymphoid nodules. Similar results were obtained by Hussain et al., (2008) and Kumar and Balachandran, (2009) in their studies of experimental aflatoxicosis in broiler chicks. Kohler, (1963) and Lanza, (1980) found fatty change, proliferation of bile ducts and cirrhosis in broiler livers exposed to 200 ppb of aflatoxins. These results are different from our results and could be attributed to the comparatively high levels of aflatoxins doses used in those studies.

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References


