Differential Hepatic Gene Expression and Antioxidant Activity in Male and Female Rats Induced by Subchronic Aflatoxicosis B1

Ghada Ali Omran1*; Nagwa Thabet Abo El-Maali2; Mady Ahmed Ismail3; Nashwa Ahmed Mohamad Mostafa4; Nasser Masood Nasser5

ABSTRACT

Aflatoxin B1 (AFB1) is the most toxic mycotoxin that was proven to be deleterious to human and several animal species. Current work aimed at evaluating the sex-based differential hepatic genotoxic effect and the antioxidant activity as implications of subchronic aflatoxicosis B1. Albino rats were used that comprised two equal AFB1 treated groups with AFB1 contaminated olive oil (50µg/kg) and a control group for each gender that received the vehicle only. Parts of animals’ livers were homogenized for gene expression assessment using quantitative RT-PCR and antioxidant activity analyses. Caspase-3 immunohistopathological examination was concomitantly undertaken. Results showed that AFB1 induced significant overexpression in cell cycle proliferation (ODC1), apoptosis (Aen), and antioxidant heme oxygenase (Hmox) genes in males alongside Bax (Bcl2-associated protein) under-expression. Meanwhile, female rats showed significant overexpression for (Hmox) and under-expression of Bax and Tnf (Tumor necrosis factor). Concomitant total hepatic antioxidant activity of liver homogenates showed a reduction in males, contrasting females. Degenerate vacuolated hepatocytes, polymorphic nuclei, cellular infiltration with concomitant Caspase-3 positive cells were profound findings in male rats. Hence, AFB1 is deferentially genotoxic at the given dose especially to male rats towards carcinogenicity, oxidative stress and apoptosis compared to a brief but compensated oxidative stress in females.

Introduction

Aflatoxins are a class of mycotoxins that are natural secondary metabolites of fungi (Aspergillus parasiticus and Aspergillus flavus) commonly contaminating food staff especially cereals. Human exposure comes from ingestions of mycotoxin contaminated food products that may lead to deleterious health effects (Bondy and Pestka 2000; Richard, 2007). Aflatoxin B1 (AFB1) is the most toxic form among other classes and recorded to be hepatotoxic mainly causing toxic hepatitis and hepatocellular carcinoma. Furthermore, its carcinogenicity was proven to many other organs, immunosuppressive, teratogenic and mutagenic in human and several tested animal species (Verma et al., 2004; Kitya et al., 2010; WHO 2011).

Some previous epidemiologic studies have validated aflatoxin-DNA adducts as a risk biomarker of hepatocellular carcinoma (Groopman et al., 2008). Additionally, G
(guanine): C (cytosine) to T (thymine): A (adenine) transversion observed in AFB1-DNA adducts is found frequently in codon 249 of p53 gene of patients highly exposed to AFB1 and having liver carcinoma (Hussain et al., 2007). Aflatoxin induces hepatotoxicity mainly by the formation of reactive oxygen species (ROS) and consequent oxidative damage (Preetha et al., 2006).

Microarray and quantitative real time PCR results from a previous study undertaken by Yarru and associates, revealed that subchronic intake of aflatoxin B1 contaminated diet (2mg/kg of diet) had led to changes in hepatic gene expression of male boiler checks. Genes coding for specific physiological pathways, including oxidation phosphorylation, detoxification, cell proliferation, immune response, metabolism and antioxidant activities were altered (Yarru et al., 2009). However, sexual dimorphism of gene expression patterns in rodents and humans exposed to AFB1 are still scarce. That is why the current study was implemented to explore certain sex-based differential hepatic gene expression changes related to oxidative DNA damage, apoptosis and cellular proliferation. The total antioxidant capability, histopathological and immunohistochemical alterations in livers of albino rats after subchronic consumption of AFB1 were concomitantly considered.

Material and Methods

Aflatoxin B1 Extraction and Quantitation

Aflatoxin B1 was extracted and quantified from a corn strain sample (AUMC 9783) that was contaminated with Aspergillus flavus strains and collected from south Egypt in another previous work. The selected strain was checked for their aflatoxin producing abilities on coconut agar medium (CAM) that proved positive only for AFB1 (Ismail et al., 2016). This was validated using high performance liquid chromatography (HPLC 1200 Series) and fluorescence detector G1321A (Agilent Technologies, Waldbronn, Germany) as described by Abo El-Maali and coworkers (Abo El-Maali et al., 2016).

Briefly, the sample was cultivated on Czapek liquid medium (Samson et al., 2004); composed of (g/l distilled water): 30 g/l sucrose, 1g/l K2HPO4, 3g/l NaNO3, 0.5g/l MgSO4.7H2O, 0.5 g/l KCl, 0.01g/l FeSO4.7H2O; pH 6.2±0.2, that was fortified by yeast extract (2 g/l) and peptone (10 g/l) (El-Kady et al., 1991). Later, 50 ml of the liquid medium were added in each of 250 ml sterile Erleynmyer flasks.

After sterilization, each flask was inoculated with three agar discs made from 7-day-old cultures on CAM plates. The flasks were then incubated at 28 °C for 10 days. At the end of incubation period, Aspergillus flavus culture was extracted with chloroform for 24h at room temperature. The HPLC (High Performance Liquid Chromatography) mobile phase was acetonitrile: methanol: water (30: 55: 15 v/v %) with a flow rate of 1.5 ml/ min under isocratic condition. Thirty microliter injection volume and 30°C column temperature was used. The detection wave lengths were fixed at 365 nm and 455 nm for the excitation and emission respectively prior to AFB1 identification and quantitation using appropriate standards.

Experimental Design

Thirty six adult rats weighing approximately 300 g (mean 300± 55g) males and 200g (mean 200±30g) females were recruited. All rats were obtained from the Faculty of Medicine’s Animal House, Assiut University. Animals were allowed to acclimatize to their environment for one week.
before the experiment. They were housed under controlled environmental conditions at 25 °C with 12 h dark/ light cycles and free access to food and water. Animals handling was undertaken according to the National Institute of Health guidelines for animal care as stated in the institutional animal care and use committee guidebook and appropriate ethical approval from Faculty of Medicine Ethical Committee in Assiut University.

The rats were divided into two main groups (a control group and an experimental group for each gender). The first main group (control) was subdivided into two subgroups (8 male and 8 female rats per group) and the second main group (experimental), were subdivided into two subgroups (10 male rats and 10 female rats per group). The rats were orally gavaged with 50 µg/kg of the AFB1 (Meki et al., 2001). The amount given to each rat was one ml of olive oil only for the control group and one ml of olive oil mixed with aflatoxin B1 to the experimental group matching designated dose. Rats were gavaged for 90 days (5 days/week) and afterwards were sacrificed by neck dislocation for harvesting livers by end of this period. Parts of recovered fresh livers have been processed immediately for the histopathological analyses and the rest were rapidly saved at – 80 °C for other downstream RNA and antioxidant activity applications.

**RNA Extraction and Quantification**

RNA (ribonucleic acid) was isolated and purified from the liver samples using the PureLink® RNA Mini Kit (Ambion, Life Technologies, California, USA) according to the manufacturer recommendations with slight modifications. Briefly, 25-30 mg liver pieces were homogenized by a manual sterilized mortar then 0.6 ml lysis buffer plus 2-mercaptoethanol were added. The mixture was vortexed for at least 40 seconds. After that, one volume 70% ethanol was added to each volume of cell homogenate and vortexed to mix thoroughly and to disperse any visible precipitate that may form after adding ethanol. The rest of the procedure was undertaken as per manufacturer recommendations. The purity and quantity of RNA were determined by a spectrophotometer (Genequant 1300, Biochrom, England) using UV absorbance at 260/ 280 nm.

**Quantitative Real Time PCR**

The primers (Invitrogen, Life Technologies, USA) used in current study were of five genes as members of oxidative stress, DNA damage, immunity, apoptosis and cellular proliferation genes (Lin et al., 2002; Suenaga et al., 2013), in addition to a housekeeping gene as shown in table (1). One step reverse transcription (RNA to cDNA) and quantitative real time PCR (RT-qPCR) with SYBR green technology were performed in a real time PCR machine (MxP3000, Stratagene, Germany) using SensiFAST™ SYBR® HI-ROX one step kit (Bioline, USA) and the above mentioned primers. The manufacturer recommendations were followed with minor modifications as using half volumes of master mix components (total volume is 10µl). A three step cycling conditions was followed (one cycle at 45°C for 10 min for reverse transcription, one cycle at 95°C for 2min for polymerase activation and 40 cycles at 95°C for 5s (denaturation), 60°C for 10s (annealing) and 72°C for 5s (extension). A melt profile analysis was implemented to protect specificity of primers and PCR products. Data were analyzed using the ΔΔCt method (Livak and Schmittgen 2001). Each RNA sample was also analyzed for Gapdh gene expression, which served as an internal control for correcting relative specific gene expression levels.
Table (1): Gene names, functions and real time PCR primers, annealing temperatures used in gene expression analysis in livers of rats exposed to AFB1 along with their controls.

<table>
<thead>
<tr>
<th>Gene name</th>
<th>Symbol</th>
<th>Forward primer</th>
<th>Reverse primer</th>
<th>Gene category</th>
<th>Annealing temperature</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glyceraldehyde-3-phosphate dehydrogenase</td>
<td>Gapdh</td>
<td>5’-atggccctcctggtctcactcc-3’</td>
<td>5’-gcctgctcctctttggtctg-3’</td>
<td>Housekeeping gene</td>
<td>53°C</td>
<td>Suenaga et al., 2013</td>
</tr>
<tr>
<td>Heme oxygenase (decycling) 1</td>
<td>Hmox</td>
<td>5’-caagcacagggtgacagagg-3’</td>
<td>5’-tctgtgagggactctggttttg-3’</td>
<td>Oxidative stress, DNA damage, apoptosis, cell proliferation,</td>
<td>54°C</td>
<td>Suenaga et al., 2013</td>
</tr>
<tr>
<td>Ornithine decarboxylase 1</td>
<td>OCD1</td>
<td>5’-agacagaccggctcggacgat</td>
<td>5’-agacatgggcagagtgccaa-3’</td>
<td>Cell cycle proliferation, rate-limiting step in pathway of polyamine biosynthesis</td>
<td>51°C</td>
<td>Lin et al., 2002</td>
</tr>
<tr>
<td>Bcl2-associated protein</td>
<td>Bax</td>
<td>5’-gagctgctcctcagttggcactcagaaac-3’</td>
<td>5’-ggcgaattggagatgaactgg-3’</td>
<td>Apoptosis, cell cycle, cell proliferation, DNA damage</td>
<td>48°C</td>
<td>Suenaga et al., 2013</td>
</tr>
<tr>
<td>Apoptosis enhancing nuclease</td>
<td>Aen</td>
<td>5’-gcaagtgcctcctcagttggcactcagaaac-3’</td>
<td>5’-ggaatggagctcctcagttggcactcagaaac-3’</td>
<td>Apoptosis, DNA damage</td>
<td>51°C</td>
<td>Suenaga et al., 2013</td>
</tr>
<tr>
<td>Tumor necrosis factor (Tnf superfamily, member 2)</td>
<td>Tnf</td>
<td>5’-ggcctgctcctcagttggcactcagaaac-3’</td>
<td>5’-cccggacagttctgatgtcactcagaaac-3’</td>
<td>Immunity, apoptosis, cell proliferation</td>
<td>53°C</td>
<td>Suenaga et al., 2013</td>
</tr>
</tbody>
</table>

Total Antioxidant Activity Assay

The total antioxidant activity was determined using a colorimetric method via the Total Antioxidant Capacity kit (Biodiagnostic, ARE) according to the manufacturer’s instructions. The assay of the antioxidant capacity is determined by the reaction of antioxidants in liver homogenate samples (like catalase, superoxide dismutase, and glutathione peroxidase; macromolecules such as albumin, ferritin and ceruloplasmin and a collection of small molecules, including β-carotene, ascorbic acid, α-tocopherol, reduced glutathione, uric acid, and bilirubin) with exogenously provided hydrogen peroxide. The antioxidants in the sample eliminate a certain amount of H₂O₂ and the rest is detected colorimetrically by an enzymatic reaction which involves the conversion of 3, 5, dichloro-2-hydroxy benzensulphonate to a colored product (Abdel-Aziem et al., 2014).

Frozen liver tissue samples were thawed, weighed and homogenized by an automated tissue homogenizer in 5ml cold buffer (50 mM potassium phosphate, pH 7.5, containing 0.9% sodium chloride and 0.1% glucose) per gram tissue. The homogenates were centrifuged at 4000rpm for 15 min at 4°C, and then the supernatant was removed for assay on UV-Visible spectrophotometer (Pharmacia LKB Ultraspec III UV/Vis Spectrophotometer, Biochrom, England) at 505nm. Total antioxidant concentration was calculated as mM/L = AB-ASA x 3.33 where, AB: blank absorbance, ASA: sample absorbance.
Histopathological and Immunohistochemical Examination

Approximately 0.5-cm³ pieces from the freshly isolated liver tissue from all rats were prepared by a frozen section technique using a cryostat instrument (Leica Biosystems, Newcastle, UK) for 10 min at −20°C to cut 5-8µm-thick sections. The procedure followed in such technique was according to Bancroft and Gamble (2008). Then, the obtained sections were stained with Hematoxylin and Eosin for the general histopathological structure using a light microscope (Olympus CH, Japan). For immunohistochemical examination, primary antibody Caspase-3 (CPP32) Ab-4: rabbit polyclonal antibody (Thermo Fisher Scientific, USA) in 1:100 dilution was utilized. Another anti-caspase-3 Biotinylated Goat Anti-polyvalent secondary antibody (Abcam, USA) in a dilution of 1:100 was used for the detection of hepatic caspase-3 according to Hofman and Taylor (Hofman and Taylor, 2013). For a negative control staining, some sections were incubated with PBS instead of the primary antibody. For a positive control, a section of a control tonsil was used.

Statistical Analysis

Fold changes in expression of used genes were calculated by ∆∆Ct method; where fold change=Log 2^(-∆∆Ct) and ∆∆Ct=(Ct of target gene - Ct of Gapdh gene) treated- (Ct of target gene - Ct of Gapdh gene) control. A two tailed - Student t test with unequal group variance was used to compare 2-∆∆Ct genetic expression differences (Schmittgen and Livak, 2008) in male and female rats compared to their matched controls. Additionally, the antioxidant capacity differences between controls and fed rats from each gender and gene fold differences between treated male and female rats were analyzed. SPSS IBM Statistics software version 22 (IBM, NYC, New York, USA) was implemented. Data was expressed as means ± SD (standard deviation). Significance values were set < 0.05 level.

Results

Gene Expression Changes

The five genes examined in rats’ livers showed differential expression patterns between controls and AFB1 treated rats from each gender in one side and between gavaged males versus females from the other side. Figure (1) shows gender differentially expressed five genes (Hmox, Tnf, Aen, Bax and ODC1). Three genes were significantly upregulated in male rats (Aen, ODC1 and Hmox) with fold changes of 2.086, 3.016 and 1.5901 respectively. Meanwhile, Bax was significantly down regulated with a ratio of 0.732 (expression is reduced by 1.366 folds due to AFB1 treatment according to Schmittgen and Livak’s formula through taking the negative inverse of 2^(-∆∆Ct)), and the remaining gene (Tnf) showed no significant change relative to their controls. On the other hand, female rats showed considerable overexpression in Hmox gene with ratios of 1.775 and under expression of Tnf and Aen with ratios of 0.719 and 0.775 in that order (fold decrease by 1.39 and 1.29 respectively relative to their female controls).
Fig. (1): Quantitative real time PCR fold changes of hepatic gene expression in male versus female rats treated with AFB1 and relation of each gender to its controls using ΔΔCt method. TNF: Tumor necrosis factor; ODC1: Ornithine decarboxylase 1; Bax: Bcl2-associated X protein; Aen: Apoptosis enhancing nuclease; Hmox: Heme oxygenase. Intersection of Y & X axes was adjusted to one to display gene up or down regulation in relation to control ratios. Error bars represent ± SD (standard deviation). *p <0.05, **p < 0.01, ***p<0.001 were used for significant difference of each gender with its control; for significant gene expression fold changes in male versus female rats, # P<0.05, ##P<0.01, ###P<0.001 were used.

Regarding the differential gene expression pattern in male rats treated with aflatoxin B1 compared to their matching female counterparts, a two tailed Student t-test was implemented. Statistically significant differences were observed in Aen, ODC, Tnf and Hmox genes. Major variation was observed in Aen and Tnf genes. They were up regulated in males in contrary to females while the Bax gene was significantly different from males towards upregulation. Ornithine decarboxylase (ODC1) gene was significantly more expressed in males than females with no considerable differentiation in both sexes in Hmox gene as well.

**Total Antioxidant Activity**

The total antioxidant activity of liver homogenates was analyzed using t-test in comparing AFB1 treated groups to controls from each gender. Figure (2) displayed significantly decreased antioxidant activity in males compared to their parallel controls (p value= 0.0318). On the contrary, females showed a mild increase in antioxidant activity but with no significant difference from controls (p value=0.763).
Histopathology and Immunohistochemistry

The histopathological examination was assessed by a light microscope using Hematoxylin and Eosin stains and Caspase-3 immunostaining in male livers revealed the following:

1- Control: The liver sections showed normal hepatic structure. The liver was formed of lobules. Each lobule was characterized by the presence of a central vein and portal areas at the periphery. The hepatocytes were arranged into plates radiating from the central vein. They were polyhedral with acidophilic cytoplasm. They had large rounded central vesicular nuclei with prominent nucleoli (Figure 3-A).

2- Aflatoxin B1 (AFB1) treated: many liver cells appeared degenerated with vacuolated cytoplasm. The nuclei showed polymorphism and some of them appeared hyperchromatic while others showed irregularly sized nuclei (Figure 3-B). Localized areas of cellular infiltration were also observed (Figure 3-C).

3- Caspase-3 immunostaining for detection of apoptosis in controls showed a mild positive reaction in some liver cells (Figure 3-D) while AFB1 treated males displayed a highly positive reaction in many cells (Figure 3-E and F).

Fig. (2): Total antioxidant activity in liver homogenates of male and female rats treated with aflatoxin B1 compared to their controls. Bars represent mean antioxidant values, Error bars represent +SD (Standard deviation), *p <0.05.
Fig. (3): Hepatic histopathological and immunohistochemical examinations of Aflatoxin B1-treated and controls of male rats. A: controls showed a normal hepatic structure. The liver was formed of lobules. Each lobule is characterized by presence of central vein (C) in the center. The hepatocytes (H) were arranged in plates radiating from the central vein. B: in AFB1 treated rats, many liver cells appear degenerated with vacuolated cytoplasm (V). The nuclei showed polymorphism (arrow). Some nuclei appeared hyperchromatic while others showed irregular sized nuclei. C: Localized area of cellular infiltration was also observed (arrow) in AFB1 treated group. With Caspase-3 immunostaining: in D & F; there is mild positive reaction in some liver cells in controls (arrow). In E & G: a highly positive reaction was observed in many cells in AFB1 treated rats (arrow). A, B, C: stained with H & E x400. D, E, F, G: caspase-3 immunostaining, D & E x 100, F & G x1000.

Light microscopic examination of the female livers revealed the following by use of Hematoxylin and Eosin stains and Caspase 3 immunostaining:

1-Control: The liver sections showed a normal hepatic structure which was nearly the same as the male liver (Figure 4-A).

2-Aflatoxin B1(AFB1) treated: showed minimal vacuolation in liver cells with mild cellular infiltration compared to the male. Nuclear polymorphism was not observed. (Figure 4-B).

3-Caspase-3 immunostaining for detection of apoptosis exhibited a nearly negative reaction in liver cells of controls (Figure 4-C and E), while AFB1 treated females displayed a moderate positive reaction in some liver cells (Figure 4-D and F).
**Fig. (4):** Hepatic histopathological and immunohistochemical examinations of Aflatoxin B1 treated and controls of female rats. A: controls showed normal hepatic structure. The liver was formed of lobules. Each lobule was characterized by presence of central vein (C) in the center. The hepatocytes (H) were arranged in plates radiating from the central vein. B: AFB1 treated rats, showed minimal vacuolation (V) in liver cells with mild cellular infiltration (arrow) compared to males. Nuclear polymorphism was not observed. With Caspase-3 immunostaining: in C&E there was nearly a negative reaction in liver cells of controls. In D&F: a moderate positive reaction was observed in some cells of AFB1 treated rats (arrow). A,B: stained with H&E x400. C, D, E, F: caspase-3 immunostaining. C&D x100, E&F x1000.

**Discussion**

Liver is considered the main target organ for aflatoxicosis (Yildirim et al., 2011). Aflatoxin B1 (AFB1) is a potent hepatotoxic and hepatocarcinogenic mycotoxin. It has been postulated to mediate various deleterious effects in tissues through the induction of oxidative stress (Özen et al., 2009). Investigation of gender based gene expression and antioxidant activity in response to AFB1 exposure were considered in rat liver. Subchronic treatment of albino rats with AFB1 resulted in a well differentiated hepatic gene expression response in each gender.

The observed upregulation of ornithine decarboxylase (ODC1) gene, especially in male rats, could contribute to increased cellular proliferation rates predisposing to hepatocellular carcinomas as could be observed in previous studies of rats (Yarru et al., 2009) and boiler checks livers (Butler et al., 1981). Ornithine decarboxylase (ODC1) was overexpressed in many cancers like human colon, liver (Tomasi et al., 2013), and breast carcinomas (Xu et al., 2015) as well. Overexpression observed in male rats along with hepatic histopathological characterization of possible precancerous foci may potentiate future carcinogenesis. This hepatocarcinogenic potency of AFB1 especially in rats given single or divided doses was described previously in many reports (Haddad, 1998). Noteworthy, this carcinogenic behavior was recorded to be much higher in males in both
experimental animals and humans supporting current findings (Woo et al., 2011). The observed sexual dimorphism in gene expression can be attributed to female estrogenic hormones that can suppress inflammation and reduce cancer risk during early phases of hepatocarcinogenesis (Naugler et al., 2007).

There was an increase in the apoptotic pathway as can be deduced from significant upregulation of Apoptosis enhancing nuclease (Aen). The latter is an exonuclease that mediates p53-induced apoptosis. When induced by p53 following DNA damage, it digests double-stranded DNA into single-stranded DNA and intensifies DNA damage signals, leading to enhancement of apoptosis (Suena et al., 2013). On the other hand, other apoptosis, oxidative stress and DNA damage contributing genes were either significantly elevated comparative to control males (Hmox) or only showed trivial increment like the Tnf gene, paralleling recent studies on male rats and mice that were given similar genotoxic hepatocarcinogens within 48 h of exposure (Watanabe et al., 2007; Watanabe et al., 2009; Watanabe et al., 2012; Suenaga et al., 2013). Noteworthy, the pro-apoptotic Bax gene was downregulated in males despite overexpression of other apoptosis enhancing genes (Aen). This has been reported before as a result of non-functional p53 overexpression in hepatocellular carcinomas (Beerheide et al., 2000). A reduced Bax expression had also been described in many human tumors, such as cervical (Soufla et al., 2005) and ovarian cancers (Tai et al., 1998). Analyses of apoptosis-related genes showed that Aen and Bax were the main targets in hepatic apoptosis induction in response to the hepatocarcinogen aminoanthracene exposure in a time dependent manner, where Bax was observed in short term high dose exposure and Aen was detected late, more with medium doses (Gato et al., 2014). This latter gene upregulation was found important to both the short and long-term apoptotic response to AFB1 in low or high doses given to growing barrows among other screened apoptosis gene expression changes in that study (Rustemeyer et al., 2011).

The genetic effects observed were consistent to the depressed total antioxidant capacity of liver homogenates versus controls. This in part confirmed previous findings as regard enhancement of apoptotic changes and decrease in liver antioxidant enzymes, increase in malondialdehyde and nitric oxide levels in chronically AFB1 treated male rats with a similar dose (Meki et al., 2001; Rastogi et al, 2001; Theumer et al., 2010; Yang et al., 2012).

Aflatoxin B1 treated female counterparts showed a slight elevation of some genes like OCD1, Bax and a significant change in Hmox with concomitant increment in antioxidant levels although statistically insignificant when compared to controls. Heme oxygenase expression was induced in response to oxidative stress, as a protective behavior in animal models (Kikuchi et al., 2005). Meanwhile, Tnf and Aen were downregulated indicating a possible compensated oxidative stress in this species given that dose to resist apoptosis and DNA damage.

To our knowledge, no previous reports available that have considered genotoxic behavior of AFB1 in female rats. Lack of a significant change though increased antioxidant activity in treated females may be attributed to the complexity of endogenous systems involved in correction and repair. This may occur when a brief increase in oxidative stress rapidly induces compensatory different antioxidant activities that limit the stress and the ability to detect a noticeable alteration by chemical testing methods like the absence of modifications in H2O2 release in detection method used (Blumberg, 2004). This could be a consequence of higher scavenging of ROS.
(reactive oxygen species) by antioxidant enzymes induced by persistent aggression of the mycotoxins (Theumer et al., 2010; Abdel-Aziem et al., 2014).

Regarding the immunohistopathological examination, we found vacuolar degeneration in the liver cells of male rats in addition to many caspase-3 positive cells indicating the occurrence of apoptosis. This finding was similar to previous studies, which reported a dose-dependent vacular degeneration with various severity showing irregularly shaped vacuoles in hepatocytes as signs of fatty accumulation (Özen et al., 2009). Moreover, vacular degeneration reported could be related to impaired lipid transport rather than increased lipid biosynthesis (McLean and Dutton, 1995).

Localized areas of cellular infiltration were observed, especially in male rats. This can be explained through the aflatoxins primary affection of cellular immunity process in most animal species (Oğuz et al., 2002). In the AFB1-exposed chickens, Yıldırım et al. (2011) have declared mild mononuclear cell infiltration in the liver’s portal areas. Small inflammatory cell infiltrates encompassing mononuclear cells, lymphocytes, plasma cells, and few segmented neutrophils occur to respond to degenerate vacuolated hepatocytes (Hinton, 2003).

Nuclear polymorphism mainly in male hepatocytes was observed as well, some nuclei appeared hyperchromatic while others showed irregular size. This could be elucidated as aflatoxin is metabolized by hepatic cytochrome P450 enzyme system to produce a highly reactive intermediate, aflatoxin 8, 9-epoxide, which consequently binds to the hepatocyte DNA nucleophilic sites to form adducts (Smela et al., 2001), thus damaging the liver cells structure (Pasha et al., 2007; Kumar and Balachandran, 2009). These DNA adducts subsequently lead to heritable genetic alterations that drive the hepatocyte in the direction of transformation (Ho et al., 2016) and initiation of aflatoxin B1-induced hepatocarcinogenesis (Preston and Williams, 2005).

In female rats, there was mild liver affection in the form of minimal vaculation with mild cellular infiltration and moderate caspase-3 positive reaction in some cells compared to males. Nuclear polymorphism was not observed. Increased expression of caspase-3, especially in males, might be explained by its different functions, which are involved in immune and inflammatory reactions to apoptotic dying cells. Upon caspase-3 activation, different proteins related to cell structural integrity are cleaved (Galluzzi et al., 2016).

The present study had underlined sex-related differences in aflatoxin B1 induced toxicity. A milestone study carried out by Kamdem et al. (1982) reported a difference in drug-metabolizing enzymes in the two sexes when exposed to different doses of AFB1. Those hepatic enzymes showed higher basal activities in males versus females. Notably, after AFB1 treatment, the levels of enzyme activities became higher in females and the detoxification system was more competent and protective against deleterious aflatoxin metabolites, contrasting males. Moreover, it had been reported that the drug-metabolizing and transporter genes in the human liver showed a sex-based differential pattern. The female predominantly expressed CYP3A4, the most essential cytochrome P450 catalyst of drug metabolism in the human liver. This sexual dimorphic expression of hepatic P450s and other genes is regulated mainly by the release of plasma growth hormone (GH) in a temporal pattern. The latter showed a pulsatile release manner in male rats and mice versus nearly a continuous pattern among females (Waxman and Holloway, 2009) which in turn, can introduce differences in drug absorption, metabolism, excretion, and possibly affecting the higher adverse AFB1 reactions in male rats (Yang et al., 2012).
Conclusion

It could be concluded that AFB1 was differentially genotoxic at the given dose especially to male rats towards carcinogenicity, oxidative stress and apoptosis compared to brief but compensated oxidative stress in females. Further studies may focus on different dose regimens in either sex alongside addressing the mechanism underlying such sex-based different response to AFB1 exposure. Moreover, follow up of gene expression associated changes in protein levels or activity, is highly desired concurrently with assessment of genotoxicity endpoints like oxidative DNA damage-related p53 activation.

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د. غادة على عمران، د. نجوى ثابت أبو العلاى، د. ماضي أحمد إسماعيل.

د. نشوى أحمد محمد صطفي، د. ناصر سعود ناصر

قسم الطب الشرقي والسموم الإكلينيكية، كلية الطب، جامعة مرسى مطروح، مصر العربية

قسم الكيمياء، كلية العلوم، جامعة مرسى مطروح، مصر العربية

قسم النبات والأحياء الدقيقة، كلية العلوم، جامعة مرسى مطروح، مصر العربية

قسم علم الأنسجة، كلية الطب، جامعة مرسى مطروح، مصر العربية

قسم الكيمياء، كلية التربية والعلوم، جامعة عين الدinen، اليمن

الأفلاتوكسين B1 هو من السموم الفطرية الأكثر شهرة والتي ثبت أنها ضارة للإنسان والديد من أنواع الحيوانات. هدفت الدراسة الحالية إلى تقييم تباين السمية الكبدية والنشاط المضاد للأكسدة على أساس الجنس وذلك نتيجة التعرض بصورة شبه مزمنة بالأفلاتوكسين B1. استخدمت الجرذان البيضاء التي تضم مجموعتين متساويتين من كل جنس ذكر أو إناث حيث تم تجريعهم 50 ميكروجرام/كم من الأفلاتوكسين B1 المخلوط مع زيت الزيتون، ومجموعة ضابطة لكل جنس تجرع المادة المذنبة فقط. تم تحليل النشاط المضاد للأكسدة. وقد تم إجراء اختبار الهندستلوجيا المناعية لبروتين كاسباس 3 بشكل متزامن. أوضحت النتائج أن الأفلاتوكسين B1 أحدث زيادة مفرطة في التعبير الجيني لجين خاص بالدورة الخلوية (ODC1)، والمضاد للأكسدة عند الذكر إلى جانب واحد كبير Hmox وأخر ناصع بموت الخلايا المرتبطة (Aen)، وفي الفطرية intens الأفلاتوكسين B1 في التعبير الجيني لجين Bax، كما أظهر النشاط الكلي لمضادات الأكسدة في الكبد انخفاض في الذكور بعكس الإناث. وكانت خلايا الكبد المتحملة ذات الفقاعات، والتي تمتد في الأشكال، والنسج الطري مع ما يصاحب ذلك من ظهور الخلايا الكبدية الإيجابية لبروتين كاسباس 3 واضحة بجلاء في ذكور الجرذان. ومن ثم، فإن الأفلاتوكسين B1 يسبب سمية جينية متبادلة بحركة المستخدمة أعلاه، خاصة عند ذكور الجرذان نحو السرطانى، والإجهاد التأكسدي وموت الخلايا المبرمج مقارنة مع الإجهاد التأكسدي العارض فقط في الإناث.