Effect Of Some Aflatoxins On A Lymphatic Organ (Spleen) Of Male Albino Rats (Histopathological Study)  
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ABSTRACT  
Background: The present study was planned to compare between two kinds of aflatoxins (AFB2 and AF Mix) on spleen of male albino rats. Fifty young male albino rats, each weighing 50g, were fed on diets containing aflatoxins at concentration of 1.0 ppm either of AFB2 or AF mix. for 2, 4, 6 weeks followed by a withdrawal period of 2 weeks.

Material and Methods: Rats were randomly divided into 3 groups: the 1st group of 10 rats was fed on the standard diet. The 2nd and the 3rd groups of 20 rats each were maintained on the standard diet plus either AFB2 or AF mix, respectively. Ten animals from the latter 2 groups as withdrawal period. Pieces of spleen were subjected to histological procedures and the obtained sections (6 µm thick) were stained with the haematoxylin and eosin, also, mercuric bromphenol blue stain for total protein were used in this study.

Results: Marked histopathological alterations were observed in the studied sections under the influence of AFB2 and AF mix. It was found that AFB2 induced more alterations. The most common changes were lymphocytic degeneration, fatty changes with numerous hemorrhagic areas. The two weeks withdrawal period showed a partial recovery of the developed changes.

Conclusion: This study indicated that AFB2 has a toxic effect on spleen than the equivalent level of AF mix. Great caution must be followed to prevent the possible contamination of our food with such mycotoxins.

Key words: Aflatoxins – Spleen – Histopathology.

INTRODUCTION  
Aflatoxins are metabolites of Aspergillus flavus that grow on ground nuts and other food stuffs (Borker, 1966; Barnes, 1967), and were confirmed to be toxic (Newberne et al., 1964).

There are two species of Aspergillus which are known to produce toxins. The isolated toxigenic Aspergillus flavus produces aflatoxin B1 and B2, whereas the isolated A. parasiticus generally produces aflatoxins B1, B2, G1 and G2. Aflatoxin B1 was found to be a potent hepatotoxic and hepatocarcinogenic mycotoxin (Cole and Cox, 1981; Rastogi et al., 2001) and caused severe histopathological alterations in liver (Rati et al., 1991). At low level of aflatoxin B1, the immune function and growth performance in pig were greatly inhibited (Chang and Pan, 1992) and induced liver tumors in rats (Angsubhakarn et al., 1990).

The effects of long-term treatment of three dose levels of aflatoxin B1 (AFB1) on lymphoid cells of weanling rats showed a marked reduction in the population and phagocytic capacity of macrophages due to AFB1 administration (Raisuddin et al., 1990). Aflatoxin B1 was toxic to the systemic immune system in various animal species (Watzl et al., 1999). Also, AFB1 had an an immunosuppressive effect on macrophages after in vivo exposure (Moon et al., 1999). Low level of it inhibited the immune function (Blaney and Williams, 1991; Chang and Pan, 1992).

In rabbits, the addition of aflatoxin B1 to normal diet produced hyperglycaemia and decreased liver glycogen (Verma and Raval, 1992). This was similiary met with in chickens (Abdelhamid et al., 1995) and rats (Choi et al., 1981; Rastogi et al., 2001). In this respect, the elevation of blood glucose level and the decrease in glycogen in liver by aflatoxin was attributed to lowering insulin level and raising of cortisol (Abdelhamid and Dorra, 1990). Also, giving rats intraperitoneal...
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dose of 5 and 7 mg/kg of aflatoxin B₁ for 6-72 hrs and 5-10 days was found to decrease acid and neutral mucopolysaccharides in liver of rats (Dutu and Maltezeanu, 1976). Several investigations have shown the serious effects of aflatoxins on liver, lymphocytes, macrophages and lung.

In chicks, AFB₁ treatment decreased both humoral and cell-mediated immune response in growing chicks at dose level of 1mg/Kg diet (Sirajudeen et al., 2011).

Toxic effects of human on T- lymphocyte (Dugyala and Sharma, 1996) and/or other lymphoid cells, the cytotoxic T-cells and natural killer cells (Methenitou et al., 2001), which impair the function of direct or indirect killing of tumor cells, can have pronounced tumorigenesis effects. The inflammatory mechanisms are initiated when various organs have been damaged by toxic assault (Batey and Wang, 2002). AFB₁ effects on the immune system were dependent on dose and time (Hinton et al., 2003). On the other hand, the reticular fibers in spleen cat, see small rings in the periphery of white pulp with toxin effect (Osuji et al., 2005).

MATERIAL AND METHODS

50 male albino rats (Rattus norvegicus) weighing about 50g were obtained from the animal station in Helwan, Cairo, Egypt. Animals were offered the standard diet (Meyer et al., 1980) and water were daily provided ad libitum. For accommodation, rats were kept in the animal house for one week before experimentation. Then, the animals were randomly divided into 3 groups. The first group, of 10 rats, served as control and was given the standard diet only. The second group, of 20 animals was fed on the standard diet containing aflatoxinB₂ (AFB₂) (1.0 ppm AFB₂). 10 animals of this group were sacrificed after 6 weeks to examine the effect of AFB₂, the other 10 animals were left for 2 weeks feeding only on the standard diet to examine the effect of the withdrawal period. The 3rd group, of 20 animals was given on aflatoxin mixture (AF mix) (AFB₁, AFB₂, AFG₁ and AFG₂) mixed with the diet which contained 1.0 ppm of AF mix for each Kg of diet. From this group, 10 rats were sacrificed after 6 weeks to analyze the effect of AM mix, while the other 10 rats were left for extra 2 weeks as withdrawal period feeding only on the standard diet. AF mix contains AFB₁, AFB₂, AFG₁ and AFG₂ in a ratio 1: 1/2: 1/4: 1/8 respectively. Pure AFB₂ and AF mix were obtained from Sigma Company, USA.

After 2, 4 and 6 weeks, pieces of spleens were taken, fixed in aqueous Bouin for 24 hrs and Carnoy's fluid, dehydrated, cleared and embedded in paraplast. Sections of 6µm in thickness were subjected to histopathological demonstration and stained with haematoxylin and eosin(Drury and Wallington, 1980). Also, we stained total proteins by mercuric bromophenol blue method (Mazia et al., 1953).

RESULTS

The Histopathological Changes:-
In spleen of control untreated rat, the typical structure of spleen was microscopically evident. The red pulp, white pulp and eccentric artery (Fig. 1).

In rats fed on AFB₂ & AF mix for 2 weeks, the spleen showed degeneration and blood hemorrhage, pools of RBCs, numerous degenerated blood cells, megakaryocyte and bizarre arrangement of WBCs in the white pulp (Figs. 2&3).

In rats fed on AFB₁ & AF mix for 4 weeks, the spleen showed numerous necrotic area, highly congested blood sinuses, numerous areas which contained debris of ruptured degenerated cells, highly distorted white pulp, bizarre arrangement of T & B lymphocytes in it, thickened arterial wall with narrow lumen of it (Figs. 4 &5).

In rats fed on AFB₁ & AF mix for 6 weeks, the spleen revealed numerous highly atrophied white pulp, haemosidrin granules, haemolysed RBCs with highly dilated and congested vein and some necrotic areas (Figs. 6a,b &7).

After 2 weeks withdrawal period showed somewhat normal appearance of the white pulp, but the red pulps were contained congested blood sinuses (Fig. 8) and somewhat normal appearance of the splenic tissue, but some degenerated areas were still detected (Fig. 9).

In the figures stained with mercuric bromophenol blue for total proteins, the splenic tissue of a control rat detected normal distribution of it with densely stained RBCs in the red pulp with less stained in white pulp (Fig. 10).

In rats fed on AFB₁ for 2, 4 & 6 weeks, the spleen showing highly dilated and congested trabecular vein which contained faintly stained with highly reduced total
protein in white and red pulps, poorly stained white pulp with negatively stained numerous degenerated areas, haemosidrin granules acquired black coloration and dilated wall of the congested trabecular vein contained haemolysed RBCs and numerous haemosidrin granules (Figs. 11a,b, 12a,b, &13a,b,c,d.).

After 2 weeks withdrawal period in this treatment AFB2, showed somewhat normal appearance of total protein was observed (Fig. 14).

In rats fed on AF mix for 2, 4, & 6 weeks, there were increased stain affinity in the white pulp, dark coloration of haemosidrin granules and necrotic areas were poorly stained. Thickened, branched and elongated trabecula were moderately stained with a slight increased stain affinity in the red pulp (Fig. 15, 16, & 17).

After 2 weeks withdrawal period treatment with AF mix showed less stained white pulp, moderately stained thickened arterial wall with numerous faintly stained areas in the red pulp, but some congested blood sinuses were deeply stained (Fig. 18).

**Explanations of Figures :-**

1- Haematoxylin and eosin stains:-

*Fig. 1.* Photomicrograph of spleen of the control rat, the typical structure of spleen was evident. T&B lymphocytes, the red pulp (RP), white pulp (WP) and eccentric artery(EA). (H&E, 200X).

*Fig. 2.* Photomicrograph of spleen of a rat treated with AFB: for 2 weeks, the spleen tissue displayed degeneration (d), congested blood sinuses (BS) and highly dilated arterial wall (arrow). (H&E, 400X).
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**Fig. 3.** Photomicrograph of spleen of a rat treated with AF mix. for 2 weeks, the spleen showing congested blood sinuses (BS), pools of RBCs, numerous degenerated blood cells, megakaryocyte (M) and bizarre arrangement of WBCs in the white pulp (arrow). (H&E, 400X).

**Fig. 4.** Photomicrograph of spleen of a rat after 4 weeks treatment with AFB2, the spleen showed numerous degenerated areas (d) highly congested blood sinuses (BS), numerous vacuolated cells (V). (H&E, 400X).

**Fig. 5.** Photomicrograph of spleen of a rat after 4 weeks treated with AF mix. Numerous degenerated areas which contained debris of ruptured degenerated cells, highly distorted white pulp, bizarre arrangement of T & B lymphocytes in the white pulp, thickened arterial wall with narrow lumen of it, numerous degenerated and necrotic areas. (H&E, 400X).
Figs. 6 a&b. Photomicrographs of spleen of a rat after 6 weeks, in rat treated with AFB₂, the spleen tissues showing numerous atrophied white pulps, haemosidrin granules (arrow), haemolysed RBCs with highly dilated and congested trabecular vein (tv) and some necrotic areas(N). (H&E, 400X).

Fig. 7. Photomicrograph of spleen of a rat after 6 weeks treated with AF mix. spleen tissue showed fatty degeneration(Fd) in the lymphocytes of the white pulp with numerous(Fd). (H&E, 400X).

Figs. 8 & 9. Photomicrographs of spleen of a rat after 2 weeks as a withdrawal period from AFB₂ and AF mix. The spleen tissue showed some clear signs of spleen tissue repair. The red and white pulps appeared with more defined outlines, but some degenerated areas were still detected. (H&E, 400X).
2-The results of total protein with mercuric bromophenol blue stain: -

*Fig. 10.* Photomicrograph of a spleen showing normal distribution of total protein in the splenic tissue of a control rat. Notice densely stained RBCs in the red pulp with less stained white pulp. (mercuric bromophenol blue, 100X).

*Figs. 11a,b* AFB2 after 2 weeks, the photomicrographs showed highly dilated and congested trabecular vein which contained faintly stained, haemolysed protein of RBCs with highly reduced total protein in white and red pulps. The degenerated areas were negatively stained with less stained arterial wall. (mercuric bromophenol blue, 100X).

*Figs. 12a,b* AFB2 after 4 weeks, the photomicrographs showed deeply stained pools of RBCs in the red pulps, numerous aggregated haemosidrin granules, necrotic and degenerated areas which poorly of negatively stained, highly elongated trabecula was negatively stained. (mercuric bromophenol blue, 100X).
Figs. 13a,b,c &d. AFB: after 4 weeks, the photomicrographs showed poorly stained white pulp with negatively stained numerous degenerated areas in it, congested blood sinuses in the red pulp appeared deeply stained, haemosidrin granules acquired black coloration, degeneration and branched trabeculae were moderately stained, highly atrophied white pulps were poorly stained, dilated wall of the congested trabecular vein contained haemolysed RBCs and congested haemosidrin granules. (mercuric bromophenol blue, 100X).

Fig. 14. AFB: after 2 weeks of withdrawal period, the photomicrograph showing somewhat normal appearance of the white and red pulps. (mercuric bromophenol blue, 100X).

Fig. 15. AF mix after 2 weeks, the photomicrograph showing thickened arterial wall of the white pulp and congested blood sinuses acquired increased stain affinity, numerous aggregations of haemosidrin granules showed dark coloration, necrotic areas were poorly stained. (mercuric bromophenol blue, 100X).
Fig. 16. AF mix after 4 weeks, the photomicrograph showing numerous aggregation of dark haemosidrin with deeply stained congested blood sinuses. (mercuric bromophenol blue, 100X).

Fig. 17. AF mix after 6 weeks, the photomicrograph showed that thickened branched and elongated trabecula was moderately stained with a slight increase stain affinity in the red pulps. (mercuric bromophenol blue, 100X).

Fig. 18. AF mix after 2 weeks withdrawal period, the photomicrograph showed that highly atrophied white pulps were less stained, also moderately stained arterial wall was detected. Fatty vacuoles showed pale stain affinity but some congested blood sinuses were deeply stained. (mercuric bromophenol blue, 100X).

DISCUSSION
The histopathology evaluation of the spleen presented here suggested that AFB caused damage to splenic cells as exemplified by the vacuolar degenerated. This change was more pronounced after treatment by AF mixture. It was known that AFB1 affects immune function in various animal species (Gaylor et al., 1992). There were early reports of a slight inflammatory response in rats due to AFB1-induced injury in the liver (Butler, 1970). Suppression of the inflammatory response via suppression of kupffer cell activation in the liver by AFB1 is in agreement with suppression of macrophage function as seen in the splenic histopathology (Hinton et al., 2001 & 2003). Also, the
changes with histopathology evaluation of the spleen with emphasis on the cell populations involved in the inflammatory response (Kodell et al., 1987, Murdoch et al., 1992). Immunotoxic effects of aflatoxin have been well documented in poultry (Celik et al., 2000 ; Ortatati et al., 2005 ; Sur and Celik, 2005). Sur et al., (2012) demonstrated that the dose caused slightly lymphoid cell depletion in lymphoid organs. Dietary aflatoxin induced immunosuppression in broilers and affected the thymus, bursa of Fabricius and spleen (Celik et al., 2000 ; Ortatati et al., 2005). The hampered functioning of macrophages may be due to the cytotoxic action of AFB1 (Raisuddin et al., 1993). By used mercuric bromophenol blue stain for demonstration total protein in the present study, it showed pale stained affinity in splenic tissues in both treatments. After a withdrawal period for 2 weeks from AFB2 and aflatoxin mix, the present data showed a partial recovery of splenic cells with some clear signs of repair. The red and white pulps appeared with more defined outlines.

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تأثر بعض الأفلاتوكسينات على عضو ليفي طحال ذكر الجزء الأبيض (دراسة نسجية)

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تمت الدراسة لمقارنة نوعين من السموم الفطرية من الأفلاتوكسينات (ألفلاتوكسين ب و خليط الأفلاتوكسينات) على طحال ذكور الجزء الأبيض.

تم تغذية عدد 50 من ذكور الجزء الأبيض بعلبة متوازنة محتوية علية 1 مجم/كم من وزن الجسم ألفلاتوكسين (ألفلاتوكسين ب و خليط الأفلاتوكسينات).

تم تجهيز قطاعات شمعية من الطحال لفحصها هستولوجيا بصبغة الهيماتوكسلين والإيوسين وكذلك صبغة البروموفينول الأزرق لفحص البروتين الكلي بانسجة الطحال بعد أسبوعين و 4 أسابيع و6 أسابيع وذلك مقارنة بمجموعة ضابطة (العلبة المتوازنة فقط). تلي ذلك أسبوعين لإنسحاب الألفلاتوكسينات من الجسم.

وقد أدت المعالجة بألفلاتوكسين ب و كذلك بخليط الأفلاتوكسينات إلى تغيير الشكل الطبيعي للطحال وظهور اضحلال في خلايا الطحال وظهور نزيف دموي وفجوات دهنية وكذلك ظهور اختلافات عديدة يكيمياء الأنسجة في محتمي البروتينات الكلية بخلايا الطحال مقارنة بالمجموعة الضابطة.

أما انحصال المواد السامة من الجزء الأبيض لمدة أسبوعين فقد أظهر تحسنا واضحا في أنسجة الطحال وأثبتت نتائج الفحوصات النسجية أن المعالجة بألفلاتوكسين ب منفردا قد أدت تأثيرات أقوى على النسيج من خليط الأفلاتوكسينات المكافئ له في الجرعات المعطاة على أنسجة الطحال.