Protective Effects of Copper (I) Nicotinate Complex Against Aflatoxicosis

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Abstract: Aflatoxicosis mostly leads to serious clinical signs of malnutrition, growth retardation, and impairment of utilization of nutrients, immunosuppression, liver diseases and cancer. Detoxification or prevention of such toxicity is interesting to researchers in change field. In pursuit for an approach, the synthetic copper(I)- nicotinate complex has been evaluated against the traditional Butylated hydroxy toluene (BHT) for possible protective effects as a food additive in induced aflatoxicosed animals. Induction of toxicity in adult rats by AFB₁ 60 µg/kg body weight divided on three times a week for consecutive five weeks resulted in marked hepatic necrosis, collagen fibers around portal tract and iron deposits as well as features of general cellular collapse and cirrhosis. The protective effects were tested on two animal groups by associate doses of BHT (0.05 mg / kg B.W.) in one group and copper (I) nicotinate complex (0.4 mg/kg B.W.) in the other. Results showed, in both groups, almost normal histological as well as biochemical markers especially those treated with the copper complex, which suggests that anti-inflammatory copper complex can be used as a protective agent against aflatoxicosis when used in doses resemble those used in pharmaceutical vitamin supplements.

Keywords: Aflatoxicosis, Copper (I) Nicotinate Complex, Antioxidant chemoprotectant, Butylated hydroxytoluene (BHT), Malondialdehyde (MDA).

INTRODUCTION

Biological mycotoxicosis depends mainly on the type of mycotoxin [1], as the acute or chronic intoxication by AFB₁ is reflected according to the level of metabolite 8,9-epoxide [2]. The bioactivation product, epoxide, is the most known genotoxic product of AFB₁ [3]. Proteins that bind with aflatoxin B₁ or its metabolites act as reservoir that sustains exposure for aflatoxicosis [4-6]. Evidence has been accumulating regarding the development of human hepatocellular carcinoma HCC with respect to AFB₁-epoxide where the P53 tumor suppressor gene and more specifically, codon 249 of this gene were seriously affected [7]. Butylated hydroxy toluene is one of the most known food additives antioxidant that showed a versatile chemoprotective agent against mammary and liver cancer induced by several carcinogenic chemicals including AFB₁ [8-10]. In mammals, the most important mechanism of AFB₁-epoxide detoxification is attributed to the high affinity of glutathione-S-tranferase towards the more potent exo-epoxide derivation [11]. Such an affinity is proven to be promoted by the antioxidant butylated hydroxy derivatives [12, 13]. Generally, antioxidants may impact their anti-carcinogenic property by inducing cellular detoxification pathway [14, 15]. With regard to many of the biologically active copper chelating complexes [16] was the copper (I)- nicotinate that exhibiting antioxidant activity as well as therapeutic pharmaceutical activity against Newcastle diseases [17], anti-inflammatory effect on gastric ulcer [18], reduction of adverse effects of 5-Fluorouracil in patients with hepatocellular carcinoma [19] and curative effect against induction of fatty liver in experimental animals [20].

AIM OF THE WORK

This work has been designed in order to evaluate the prophylactic effect of a bioactive copper (I) nicotinate complex as a food additive against aflatoxicosis. For this purpose the complex was given within the daily requirement of copper element, versus a commercially used phenolic derivative BHT.

MATERIALS AND METHODS

Fifty five Wister albino male rats (120–150 g B.W.) were accommodated in appropriate healthy conditions and grouped into 4 groups; group I contained 15 animals for control measurements G₁, and 3 groups; 15 animals each, for experiment. The experimental groups were aflatoxicosed by orally ingested AFB₁ (60 µg/kg B.W.), three times weakly for consecutive five weeks. The toxin as a pure sample was a gift from the Department of Botany, Mycology Center, Faculty of Science, Assiut University, Egypt. The First experimental group aflatoxicosed received no associate
treatment; experimental control GII. The second experimental group GIII animals were supplemented by BHT (0.05 g/kg B.W.) concomitant with AFB1 [21]. The phenolic BHT was purchased from Oxford Laboratory – Mumbai- 400002, CAS No. (128-73-0). The third experimental group GIV were treated by the synthesized copper(I)-nicotinate complex (400 µg/kg B.W.) concomitant with AFB1 [22]. Experimental groups were sacrificed at the end of the time course of the experiment (5 weeks). Blood serum and liver tissue samples were collected for chemical analysis and histopathological examination. The tested biochemical parameters on the liver tissue were performed on homogenates (one gm/9 ml phosphate buffer at pH 7.4).

The biochemical tests included total protein [23], amino transferases ALT and AST (diagnostic kit by spectrum diagnostics, Egyptian company for biotechnology S.A.E.), nitric oxide [24], total thiols [25], superoxide dismutase [26], reduced glutathione [27], glutathione peroxidase [28], glutathione-S- transferase [29], serum ceruloplasmine [30] and lipid peroxide [31]. Histological examination including iron staining was done by Perls’ Prussian blue reaction according to Wang and Liao (2003)[32]; Wang, Zhou et al. (2004)[33]; Masson's trichrome was done by Drury R and Wallington E.A. (1980)[34].

RESULTS

Serum total protein (g/dl): Table 1, shows: the animals in GII showed dramatic reduction in serum total proteins that was highly significantly reduced vs normal control animals in GI (p<0.001). Intoxicated animals co-treated with BHT in GIII showed that was non-significantly higher than normal. Similarly intoxicated animals co-treated with Cu-nicotinate complex in GIV showed also non-significantly lower than normal controls. Serum ALT activity (U/L): Table 1, showed dramatic increase in GIII than normal, (p<0.001). The level of serum ALT increased in GIV which was still significant (p<0.05). Aflatoxicosed animals of GIV was non-significantly than normal (p<0.05). Serum AST activity (U/L): Table 1 shows the aflatoxicosed animals in GII that was dramatically significant increased over normal, (p<0.001). AST level of GIII it was still significantly higher than normal but is much less than control. The GIV group showed a reduced level of AST but it was still significantly higher than normal, (p<0.05). Serum total thiols concentration (nmol/ml): Table 1 illustrates the GI that was significantly reduced from normal control, (p<0.001). The GII that was still lower significantly than normal control, (p<0.001). Similarly GIV showed a level of total thiols that was significantly lower than normal control, (p<0.05). Serum and hepatic tissue GSH (nmol/ml), (nmol/mg protein): from Table 1, GII showed prominent reduction in serum GSH value that was highly significantly reduced than normal, (p<0.001). Animals co-treated with BHT GIII showed also reduction that was statistically not different from normal. Similarly GIV showed the same response that was also non-significantly deviated from normal value. In the hepatic tissue Table 2, GII animals showed significant reduction in hepatic tissue GSH value from normal, (p<0.001). Animals co-treated with BHT GIII showed slightly reduction in the level GSH from the normal. While GIV animals showed also nonsignificantly different level from normal, (p<0.05). Total thiols of hepatic tissue (nmol/g): From Table 2, Aflatoxicosed untreated animals, GII showed significant reduction from normal animals (p<0.001). The co-treated BHT animals in GIII that was significantly lower than normal, but was about three times the value in untreated GII. Similarly GIV showed improvement in hepatic tissue total thiols that was nonsignificantly different from normal, (p<0.001). Hepatic tissue lipid peroxidation level (nmol/g): from Table 2, it was found that significantly increased over normal in GII, (p<0.001). This level in GIII group was significantly elevated than normal (p<0.05). GIV didn’t show any significant elevation over

Table 1. Changes in Serum Total Protein Concentration (g/dl), Serum Albumin (g/dl), Serum ALT (U/L), serum AST (U/L), Serum GST (nmol/min/ml), Serum Nitric Oxide (µmol/ml), Serum GSH (nmol/ml), Serum SOD (U/ml), Serum Ceruloplasmine (mg/dl) and Serum Total Thiols (nM/ml) in Control G1, Aflatoxicosed Untreated GII and BHT GIII and Copper Nicotinate GIV Aflatoxin co-Treated Rats

<table>
<thead>
<tr>
<th>Parameters in Serum</th>
<th>Group I (Control), n=10</th>
<th>Group II (AFB1), n = 15</th>
<th>Group III (AFB1+BHT), n = 15</th>
<th>Group IV (AFB1 + copper complex), n = 15</th>
</tr>
</thead>
<tbody>
<tr>
<td>total protein (g/dl)</td>
<td>7.77±0.33</td>
<td>4.17±0.96***</td>
<td>7.31±0.39 N.S.</td>
<td>7.48±0.22</td>
</tr>
<tr>
<td>Albumin (g/dl)</td>
<td>3.49±0.3</td>
<td>1.94±0.8***</td>
<td>3.07±0.9 N.S.</td>
<td>3.28±0.09</td>
</tr>
<tr>
<td>ALT (U/L)</td>
<td>22.5±1.22</td>
<td>50.25±0.95**</td>
<td>26.45±1.42*</td>
<td>23.5±0.33</td>
</tr>
<tr>
<td>AST (U/L)</td>
<td>26.45±0.69</td>
<td>70.08±2.31***</td>
<td>35.38±0.96***</td>
<td>32.5±1.82*</td>
</tr>
<tr>
<td>total thiols (nM/ml)</td>
<td>1.33±40.428</td>
<td>1.305±0.55***</td>
<td>3.312±0.37***</td>
<td>3.827±1.62*</td>
</tr>
<tr>
<td>nitric oxide (µmol/ml)</td>
<td>206.4±0.32</td>
<td>238.74±0.99***</td>
<td>205.7±0.67</td>
<td>198.6±1.19***</td>
</tr>
<tr>
<td>SOD (U/mL)</td>
<td>5.5±8.7</td>
<td>3.125±8.83***</td>
<td>5.75±9.45</td>
<td>8 ±0.001***</td>
</tr>
<tr>
<td>GSH (nmol/ml)</td>
<td>33.51±0.47</td>
<td>21.12±1.33***</td>
<td>32.32±1.05</td>
<td>33.57±0.88</td>
</tr>
<tr>
<td>GST (nmol/min/ml)</td>
<td>450.5±2.36</td>
<td>325.3±2.98***</td>
<td>435.6±1.55*</td>
<td>446.2±3.72</td>
</tr>
<tr>
<td>ceruloplasmine (mg/dl)</td>
<td>50.28±1.36</td>
<td>7.06±0.323***</td>
<td>49.63±0.254</td>
<td>56.3±1.22***</td>
</tr>
</tbody>
</table>

Values are expressed as the mean ± SE for (15) animals in each group.

*p<0.05; **p<0.01; ***p<0.001 represent significant differences as compared to control animals (group I).
Table 2. Changes in Hepatic Tissue Lipid Peroxide (nmol/g wt Tissue), Hepatic Tissue GsPx (nmol/min/mg Protein), Hepatic Tissue Total Thiols (nM/g Protein), Hepatic Tissue Nitric Oxide (µmol/g Protein), Hepatic Tissue GSH (nmol/mg Protein), and Hepatic Tissue GST (nmol/min/mg Protein) in Control G_I, Aflatoxicosed Untreated G_II and BHT G_III and Copper Nicotinate G_IV Aflatoxin Cotreated Rats

<table>
<thead>
<tr>
<th>Parameters in Hepatic Tissue</th>
<th>Group I (Control), n=15</th>
<th>Group II (AFB1), n=15</th>
<th>Group III (AFB1+BHT),n=15</th>
<th>Group IV (AFB1 + Copper Complex), n=15</th>
</tr>
</thead>
<tbody>
<tr>
<td>lipid peroxide (nmol/g wt tissue)</td>
<td>11.9±0.88***</td>
<td>38.94±2.03***</td>
<td>12.89±1.29*</td>
<td>12.64±0.99</td>
</tr>
<tr>
<td>GsPx (nmol/min/mg protein)</td>
<td>280.5±2.33</td>
<td>195.8±0.28***</td>
<td>260.3±2.65*</td>
<td>277.2±1.98</td>
</tr>
<tr>
<td>total thiols (nM/g protein)</td>
<td>3.29±1.32***</td>
<td>1.026±1.53***</td>
<td>2.844±0.97***</td>
<td>3.195±0.33</td>
</tr>
<tr>
<td>nitric oxide (µmol/g protein)</td>
<td>151.2±0.39***</td>
<td>190.5±1.35**</td>
<td>159.3±0.85</td>
<td>146.8±1.03</td>
</tr>
<tr>
<td>GSH (nmol/mg protein)</td>
<td>28.2±1.22</td>
<td>18.1±0.48***</td>
<td>27.5±1.33</td>
<td>29±2.14</td>
</tr>
<tr>
<td>GST (nmol/min/mg protein)</td>
<td>430.3±3.98***</td>
<td>305.2±5.12***</td>
<td>410.8±3.55*</td>
<td>420.3±2.45</td>
</tr>
</tbody>
</table>

Values are expressed as the mean ± SE for (15) animals in each group.

*p<0.05; **p<0.01; ***p<0.001 represent significant differences as compared to control animals (group I).

normal; (p<0.001). Serum nitric oxide concentrations (µmol/ml): in Table 1, The G_II showed dramatically significant increase higher than normal, (p<0.001). The level of serum nitric oxide concentration was non-significantly changed from normal in G_III animals, (p<0.001). The G_IV showed also non-significant lower value, (p<0.001). Hepatic tissue nitric oxide concentrations (µmol/g proteins): from Table 2, The G_II liver NO was significantly elevated than normal (p<0.01). While G_III was non-significantly higher than normal value (p<0.001). Similarly, the animals co-treated with Cu-nicotinate complex; G_IV changed to significant lower value. Serum superoxide dismutase (SOD) activity (U/ml): From Table 1, the aflatoxicosed untreated rats G_II showed significant lower than normal (p<0.01). The level of serum SOD was increased in G_III non-significantly higher than (p>0.001). Aflatoxicosed animals with simultaneously co-treatment with Cu-nicotinate complex G_IV showed significantly higher than normal (p<0.001). HEPATIC animals measurement showed improvement that was non-significantly different from normal (p<0.001). Serum glutathione-S-transferase (GST) concentrations (nmol/min/ml): From Table 1, G_II aflatoxicosed without treatment showed remarkable reduction in serum GST value that was significantly different from normal (p<0.001). G_III group showed more or less reduced level from the normal, that change was significant (p<0.05). G_IV showed improvement that was non-significant change from normal (p<0.001).

Hepatic tissue GST concentration (nmol/min/mg protein): it is seen from Table 2, G_I that was a significant reduction in the enzyme activity than normal (p<0.001). G_III measurement showed improvement that was a significant change from normal (p<0.05). G_IV animals measurement reflected great improvement that was non-significantly different from normal (p<0.01). Serum ceruloplasmine (mg/dl): Table 1 illustrates the level was dramatically reduction in G_II that was a significant reduction from normal (p<0.001). G_III animals value that was non-significantly different from normal (p<0.05). The G_IV group exceeded the normal level to significantly greater value (p<0.01).

Histological Examination: Fig. (1) shows histological features of health control G_I animals. Fig. (2a through e) shows
Fig (2). Stained sections with Heamatoxyllin and eosin for aflatoxicosed animals:

(a) Shows a liver section of GII animals with markedly dilated congested central vein occupied by large thrombus (star). Note that the multiple iron deposits in the field. (Hx. & E. ×400).

(b) Shows liver focal hepatic necrosis (arrow head), predominantly perivascular with infiltration of the hepatic parenchyma by lymphocytes are characterized (Hx. & E. ×400).

(c) Shows liver localized aggregation of cells (star); some appear with vesicular nuclei, others appear with dense nuclei and necrotic cells with vacuolated cytoplasm are also detected. (Hx. & E. ×400).

(d) Shows a liver GII animals, bile ducts hyperplasia as well as mononuclear inflammatory cells formed mainly of lymphocytes, hyperplasia of kupffer cells (Hx. & E. ×400).

(e) Shows a liver of GII group, the portal tracts are thickened by cell proliferation of bile ducts as well as mononuclear inflammatory cells formed mainly of lymphocytes. (Hx. & E. ×400).

Microscopic examination of aflatoxicosed livers of GII animals. The liver appeared with markedly dilated congested central vein occupied by large thrombus and multiple iron deposits as well as hepatic cellular necrosis (Fig 2a). Fig. (2b) shows liver focal hepatic necrosis (arrow head), predominantly perivascular with infiltration of the hepatic parenchyma by lymphocytes are characterized. Liver localized aggregation of cells (star); some appear with vesicular nuclei, others appear with dense nuclei and necrotic cells with vacuolated cytoplasm are also detected in Fig. (2c). In Fig. (2d), the same group liver of aflatoxicosed untreated GII animals, bile ducts exhibit hyperplasia as well
Fig (3). Stained sections with Heamatoxyllin and eosin for co-treated rats by BHT or copper(I)-nicotinate complex animals:

(a) A liver section of G_{III} animals; perivascular infiltration of the hepatic parenchyma with lymphocytes (star), some hepatocytes exhibit vesicular nuclei others appear with dense nuclei (Hx. & E. ×400).

(b) A liver section of G_{III} a minimal cellular infiltration in the portal tract area, multiple small bile ducts are present hepatocytes more or less normal(Hx. & E. ×400).

(c) Shows a liver of G_{IV} group, prevention of the effect of AFB_{1} on the liver where hepatic architecture had almost normal appearance. (Hx. & E. ×400).

as mononuclear inflammatory cells formed mainly of lymphocytes, the portal tracts appear thickened by cell proliferation of bile ducts as well as mononuclear inflammatory cells formed mainly of lymphocytes, Fig. (2e).

Fig. (3a), shows liver section of G_{III} animals; perivascular infiltration of the hepatic parenchyma with lymphocytes (star), some hepatocytes exhibit vesicular nuclei others appear with dense nuclei, in Fig. (3b) of the same group, we find minimal cellular infiltration in the portal tract area, multiple small bile ducts are present, hepatocytes have more or less normal appearance. In Fig. (3c), of G_{IV} group, remarkable modulation in the effect of AFB_{1} on the liver where hepatic architecture returned to normal appearance. Fig. (4a through d) shows cross sections of liver of all groups stained with Masson's trichrome for collagen fibers and it depicts; (a) liver of control animals G_{I}, where delicate collagen fibers are around central vein and portal tract. (b) Liver section of G_{II} animals with markedly increased collagen fibers around portal tract (star). (c) Shows a liver section of G_{III} animals, with reduced thickness collagen fibers around portal tract (star) (d) Seen modulation of the effect of AFB_{1} on the liver where reduction in the collagen fibers around portal tract in G_{IV} animals comparison to those treated with G_{III} group. Fig. (5a through d) shows liver sections of all groups stained with Prussian blue for detection of Iron deposits in tissue. (a) Shows a liver of negative control with normal appearance of liver parenchyma with no observed iron deposits, (b) Shows a liver section of G_{II} animals, multiple iron deposits in the central vein. The liver section of G_{III} animals appeared without any observed iron deposits in the liver parenchyma, see Fig. (4c), while in Fig. (5d), G_{IV} liver section showed no signs of iron deposits in the central vein or liver parenchyma.

Fig. (3a), shows liver section of G_{III} animals; perivascular infiltration of the hepatic parenchyma with lymphocytes (star), some hepatocytes exhibit vesicular nuclei others appear with dense nuclei, in Fig. (3b) of the same group, we find minimal cellular infiltration in the portal tract area, multiple small bile ducts are present, hepatocytes have more or less normal appearance. In Fig. (3c), of G_{IV} group, remarkable modulation in the effect of AFB_{1} on the liver where hepatic architecture returned to normal appearance. Fig. (4a through d) shows cross sections of liver of all groups stained with Masson's trichrome for collagen fibers and it depicts; (a) liver of control animals G_{I}, where delicate collagen fibers are around central vein and portal tract. (b) Liver section of G_{II} animals with markedly increased collagen fibers around portal tract (star). (c) Shows a liver section of G_{III} animals, with reduced thickness collagen fibers around portal tract (star) (d) Seen modulation of the effect of AFB_{1} on the liver where reduction in the collagen fibers around portal tract in G_{IV} animals comparison to those treated with G_{III} group. Fig. (5a through d) shows liver sections of all groups stained with Prussian blue for detection of Iron deposits in tissue. (a) Shows a liver of negative control with normal appearance of liver parenchyma with no observed iron deposits, (b) Shows a liver section of G_{II} animals, multiple iron deposits in the central vein. The liver section of G_{III} animals appeared without any observed iron deposits in the liver parenchyma.
Fig (4). Stained sections with Masson's trichrome for normal control, aflatoxicosed, co-treated animals with BHT or copper(I)- nicotinate complex:

(a) Shows a liver of negative control animals where delicate collagen fibers are around central vein and portal tract (Masson trichrome X 400). (b) Shows a liver of aflatoxicosed animal for five weeks, the markedly increase in the collagen fibers around portal tract (star). (Masson trichrome x400).

(c) Shows a liver of aflatoxicosed animal for five weeks co-treated by BHT, with reduced collagen fibers around portal tract (star) (Masson trichrome×400).

(d) Shows a liver of aflatoxicosed animal for five weeks co-treated by copper(I)- nicotinate complex amelioration in the effect of AFB$_1$ on the liver with reduction in the collagen fibers around portal tract in comparison to those treated with BHT group (Masson trichrome × 400).
Fig (5). Stained sections with Prussian blue for normal control, aflatoxicosed, co-treated animals with BHT or copper(I)- nicotinate complex: (a) Shows a liver of negative control animal of normal looking appearance of liver parenchyma with no observed iron deposits. (Prussian blue stain X 400) (b) Shows a liver of aflatoxicosed animal for five weeks multiple iron deposits in the central vein (star). (Prussian blue stain X 400). (c) Shows a liver of aflatoxicosed animal for five weeks co-treated by BHT without any observed iron deposits in the liver parenchyma (Prussian blue stain ×400). (d) Shows a liver of aflatoxicosed animal for five weeks co-treated by copper(I)- nicotinate complex, no signs of iron deposits in the central vein or liver parenchyma (Prussian blue stain ×400).

see Fig. (4c), while in Fig. (5d), GIV liver section showed no signs of iron deposits in the central vein or liver parenchyma.

**DISCUSSION**

The assayed biochemical parameters in circulating systemic blood as well as hepatic tissue homogenates were in consistency with those findings of the microscopic examination characters. The total serum protein or serum albumin levels were highly reduced by aflatoxicosis due to inhibition of protein synthesis in hepatic tissue and other organs [35, 36]. The antioxidant BHT prevented such reduction in protein synthesis due to chemoprevention of aflatoxin B1 metabolism in biological systems [37]. Similarly, treatment with copper nicotinate complex showed improvement that confers a hepatocellular protective as well as antioxidant effect of copper (I) nicotinate complex [18-19]. Concomitantly, the changes in transaminases gave a positive sign for the protective effect of either BHT or the copper (I)-nicotinate complex against hepatic tissue collapse by aflatoxicosis. Elevated levels of measured hepatocellular
oxidative stress markers (NO and lipid peroxide) were prominent by aflatoxosis [38]. Co-treatment by either BHT or copper nicotinate complex that reduced such biochemical parameters indicates that both of them are real able hepatocellular protective agent.

Reduction of accumulated levels NO and lipid peroxides in hepatic tissue by BHT and the copper complex co-treatment that was regulated and promoted by elevation of antioxidant cellular chemicals such as GSH and total thiols [39], this could be attributed frankly to the antioxidant effect of BHT [20, 40] and the cellular protective antioxidant Cu(I)-nicotinate complex [41]. Since, induction of protective GST has been posited possible chemoprotective mechanism by BHT and other phenolic antioxidant as enzyme promoters has been confirmed before by Hayes et al., (1996)[42]. Herein, results could be rationally accepted. Inasmuch as parallel enzymatic induction for the GST-mediated detoxifying enzyme against AFB1 by using the copper (I)- nicotinate complex, in this study is the first time to be deduced. The blood circulating levels of such parameters that were parallel to those of hepatic tissue could be interpreted similarly due to the antioxidant character of BHT and the predicted cellular protectant anti-inflammatory and antioxidant effect of the Cu(I)-nicotinate complex on the cellular levels.

Since, aflatoxosis is among the widely encountered features of environmental pollution that deteriorates the hepatic tissue; induction of aflatoxins in the tested experimental animals was associated with cellular damage that was characterized by, focal hepatic necrosis and increased collagen fibers around central vein and portal tract as it was detected by microscope. These characters were mostly absent by co-treatment with BHT. These findings are simply accepted due to previous success of the use of BHT in preventing aflatoxosis [43]. In addition, these cellular improving characters were microscopically obvious by utilization of the copper complex under consideration as a protective agent. This improvement was predicted since this complex has been confirmed previously as a therapeutic agent against induced non-alcoholic fatty liver in experimental animals [20].

Herein, after ingestion, AFB1 might be converted into its epoxide or the epoxide of its major metabolites AFM1 and these derivatives produced DNA adducts causing DNA strand breaks and point mutations [44]. Under this pathological condition, the active process of cellular self destruction, apoptosis may occur. The MDA is the end product of lipoperoxidation, considered as a late biomarker of oxidative stress and cellular damage [45, 46]. Hence, the aflatoxicosed animals that were characterized by a marked increase in MDA level in the liver (Table 2), the oxidative damage is considered to be the main mechanism leading to the subsequent hepatotoxicity as it was reported before [47]. Protective activity of the antioxidant Thymoquinone (TQ) also was evaluated in a similar study against liver toxicity induced by AFB1 in mice. The serum levels of hepatic enzymes primarily reflect the degree of liver damage and have been commonly used as a diagnostic marker for hepatotoxicity. As shown in Table 1, AFB1 significantly elevated, AST and ALT which is in agreement with results of other studies [48, 49]. This finding is consistent within histopathological examination in which necrosis and degeneration, hyperplasia of kupffer cells and infiltration of mononuclear cells in liver sections that were observed. BHT prevented the histopathological changes as well as those co-treated by the copper complex decreased the number of inflammatory cells that was in parallel to the biochemical findings. However co-treatment with the copper (I) nicotinate complex initiated complete amelioration of the AFB1 associated changes where the liver tissues looked more or less similar to those of control. Markedly increase in the collagen fibers around portal tract in aflatoxin intoxicated animals in the present study are similar to those previously described for aflatoxosis [50, 51] especially that was published by Bastianello et al., (1987)[50] who described the pathology of acute, subacute, and chronic aflatoxicosis. In this review the subacute form was typically characterized by extensive bile ductule hyperplasia with varying degrees of fibrosis and bridging fibrosis [50, 52, 53]. Typically, the chronic form was also characterized by extensive fibroplasia. Additionally, observed severe bile ductule proliferation and visible regenerative hepatocellular nodules could be frankly defined [50] as it was reported. BHT cotreated animals in this study exhibited reduced collagen fibers around portal tract, while animals co-treated by copper(I)-nicotinate complex exhibited amelioration in the effect of AFB1 on the liver with complete reduction in the collagen fibers around portal tract in comparison to those treated with BHT group. The current study shows that, induced aflatoxicosis dramatically reduced the level of the main blood circulating enzymatic free radical scavenger SOD, which was previously confirmed by others [54, 55]. Such a reduction was elevated and normalized by the antioxidant co-treating agents BHT or the copper nicotinate complex. Notably, the copper complex was significantly more potent than that of the co-treating BHT. This prominent observation could be attributed to the biochemical behavior of the copper complexes that was confirmed to be similar to the SOD activities such as Cu(II)(3,5-diisopropysalicylic acid)2 (CuDIPS) [56], Cu(II) histidine complexes [57], Cu(II) complexes of macrocyclic polyamine derivatives [58], Bis(2,9-dimethyl-1,10-phenanthroline)- Cu(I)nitrate (Cu(I) (DMP)2) [59] and Cu(II)-oligopeptide [60].

Similarly, the blood ceruloplasmin that was highly consumed by aflatoxicosis was improved by association of the co-treatment of the anti-inflammatory agents BHT or the tested copper-nicotinate complex. The latter was also highly significantly more effective than that of the BHT. It is worthy noted that ceruloplasmine is one of the acute phase transport proteins and that after intoxication; copper and zinc are depleted and tend to leave the body with urine, this depletion of copper may be a causative parameter for the decrease of ceruloplasmine [61].

CONCLUSION

Basically, these obtained results by using copper (I)-nicotinate complex as a proelactive therapeutic agent against aflatoxicosis not only acts like BHT but also it was more prominently effective on measured parameters and biologically saver as food additive than the synthetic BHT.

CONFLICT OF INTEREST

The authors confirm that this article content has no conflicts of interest.
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REFERENCES


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