Evaluation of the Protective Effect of *Acacia senegal* Extract against di-(2-ethylhexyl phthalate) Induced Hepato- and Neurotoxicity in Rats

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Authors’ contributions

This work was carried out in collaboration between all authors. All authors read and approved the final manuscript.

ABSTRACT

Di-(2-Ethylhexyl phthalate) (DEHP) is an aromatic diester used to improve plasticity of industrial polymers. It exhibited adverse changes in both liver and brain. The current study was planned to evaluate efficiency of *Acacia senegal* extract to ameliorate against liver and brain toxicity induced by DEHP. In this study, markers of the serum hepatic functions were elevated significantly (P˂0.05) in the DEHP-treated group. In coincide with these results, the antioxidant enzymes declined significantly (P˂0.05) associated with elevation of the lipid peroxidation product (LPO) in liver of DEHP-treated group. Furthermore, DEHP caused decline in activity of the antioxidant enzymes associated with elevation of LPO level in brain tissue. In consistent with these results, DEHP
caused elevation of excitatory amino acids with decrease of inhibitory amino acids and monoamines in that tissue. A. senegal extract showed ameliorative effect by restoring activities of the antioxidant enzymes to normalcy with reducing the LPO level in the both tissues. The electrophoretic protein and lipoprotein patterns in liver tissue presented that the lowest similarity index (SI) values were noticed in the DEHP-treated group (66.67 and 71.43%, respectively). No changes detected in protein and lipoprotein patterns in brain tissue. DEHP caused electrophoretic quantitative mutagenicity by increasing quantity of the α-EST2 band in liver tissue. As regards β-EST enzyme in liver and brain tissues, DEHP caused qualitative mutagenicity leading to decreasing the SI % in liver and brain tissue (66.67 and 25%, respectively). Moreover, it induced cleavage of the genomic DNA in both tissues. A. Senegal extract increased the SI values by restoring the normal bands and hiding the abnormal ones and maintained integrity of the genomic DNA pattern in liver tissue.

Keywords: Di-(2-Ethylhexyl phthalate); Acacia Senegal; electrophoresis; protein; esterases; genomic DNA.

1. INTRODUCTION

Phthalates are consisting of alkyl or dialkyl esters of phthalic acid (benzene dicarboxylic acid). They are metabolized into tomonophthalate by lipases or esterases in the intestines; monophthalate is degraded into oxidative metabolites. The monoesters are generally considered to be responsible for the general toxic effects [1]. Di-(2-Ethylhexyl phthalate) (DEHP) is an aromatic diester that used as plasticizer in polyvinyl chloride (PVC) resins for fabricating flexible vinyl products [2]. It is well known that the DEHP belongs to the peroxisome proliferator compounds which normally found in food packed from plastic materials. These compounds have been known to changes in metabolic pathways and subsequently induce liver enlargement [3].

It is a lipophilic compound that can be absorbed through skin and lungs by both humans and rodents. The general population is susceptible to be exposed to DEHP mainly through oral rout [4]. Dietary exposure to DEHP occurs through migration of the plasticizer from food packaging and from the environment where it exists as a contaminant in drinking water and aquatic food. Fatty foods such as milk, butter and meats are a major source. Because phthalate plasticizers are not chemically bound to PVC, they can easily leach and evaporate into food or the atmosphere. Moreover, low-molecular-weight phthalate esters, including DEHP are used to make coatings for oral medications [5]. DEHP is a well-known hepatotoxin as it stimulates proliferation of the hepatic peroxisomes and produce liver hypertrophy, hyperplasia and tumors. It causes induction of lipid metabolizing enzymes in the liver [6]. Also, it is shown to be hepatocarcinogenic causing hepatocellular carcinomas and adenomas. The incidence of hepatocellular carcinoma is a DEHP dose dependent. DEHP and its metabolites are suspected of producing neurotoxicity and endocrine-disrupting effects [7].

Esterases catalyze the hydrolysis of ester bonds of various compounds (i.e., lipids, oils, and phthalates) and break them down into the corresponding carboxylic acids. The initial step in the degradation of phthalate esters seems to be a de-esterification reaction by specific esterases, to give rise to phthalate anions and alcohol [1,8]. DEHP is able to inhibit acetylcholinesterase activity and up-regulated transcripts of growth associated protein [9]. Moreover, it exhibits adverse effects on the behavior and brain-derived neurotrophic factor of rats. Therefore, DEHP endocrine disruptors may also contribute neurodevelopmental disorders such as disturbances of the neurotransmitters excitatory and inhibitory amino acids that subsequently lead to attention-deficit hyperactivity disorder and autism [10]. For this reason, the present study aims to reveal the deleterious effect of DEHP on both of liver and brains tissues.

The previous published researches emphasize that Acacia species exhibit various biological activities including antioxidant potential, anti-inflammatory, antitumor and neuronprotective [11]. Acacia senegal, commonly known as Gum Arabic, is a drought or arid region tree. It is a native of Africa and is also found in the Indian subcontinent and Pakistan. Official parts of A. senegal such as gum, seeds, fruits, leaves and bark are considered as the richest resources of bioactive tannins, flavonoids, phenolics, alkaloids, saponins, polysaccharides and terpenoids [12,13]. It is postulated that Acacia senegal gum is used for soothing mucous
membranes of the intestine and to cure bleeding, bronchitis, diarrhea, malaria, typhoid fever and respiratory tract infections besides possessing antiplatelet and antifertility activities [14]. *A. Senegal* gum contains neutral sugars (rhamnose, arabinose and galactose), acids (glucuronic acid and 4-methoxyglucuronic acid) and minerals (calcium, magnesium, potassium and sodium) [15]. Furthermore, it is rich in various effective phytoconstituents that are represented by flavone, catechin, polyphenols, tannins, chalcones, alkaloids and flavonoids [16]. It exhibited various pharmacological activities represented by the protection against cyclophosphamide induced urinary bladder cytotoxicity [17], limitation of the hepatotoxicity induced by acetaminophen [18] and antidiabetic activity against streptozotocin induced diabetes in rats [19]. The present study is planned to explore the possible ameliorative or therapeutic effect of *A. senegal* extract against liver and brain toxicity induced by DEHP.

2. MATERIALS AND METHODS

2.1 Extract Preparation

Plant material of matured *Acacia senegal* pod with seeds was purchased from Khedr El Attar, Khan El Khalili, Cairo, Egypt. It was oven dried and subjected to mechanical size reduction. As documented by Kannan et al. [20], the powdered material was soaked for one week at room temperature in ethanol (70%) at the concentration 10% (W/V). On the 7th day, extraction was carried out using ultrasonic sound bath accompanied by sonication (40 minutes) and filtered. The solvent was allowed to evaporate and the extract was concentrated by rotary evaporator (Panchun Scientific Co.) at 35°C. The gummy extract was freeze dried for further use.

2.2 Treatment Dose

Based on the results obtained by Meena et al. [21] and supported by Singh et al. [22] who reported that no deaths or adverse effects were detected during the 24-hour observation period in mice treated with up to 3000 mg/kg.bw. The dose at the concentration of 300 mg/kg.bw was chosen for the therapeutic application during the experiment.

2.3 Ethics Statement

The experimental design and animal handling were carried out according to experimental protocol approved by Institutional Animal Ethics Committee of National Research Centre, Dokki, Giza, Egypt (No: 471/2016). Moreover, it was conducted in accordance with guidelines as per "Guide for the care and use of laboratory animal" and with permission from Committee for the Purpose of Control and Supervision of Experiments on Animals. Every effort was done to minimize the number of animals and their suffering.

2.4 Experimental Design

Healthy thirty six adult male rats (*Sprague Dawley*) of initial weights 200 ± 20 g (9-12 weeks old) were housed in filter top polycarbonate cages. The animals were interned for 28 days and provided with water *ad libitum* and standard food and maintained under normal environmental conditions at 25 ± 2°C. After an acclimatization period of 1 week, animals were equally divided into six groups (6 rats in each). Group I (Control group): Rats were fed with normal diet as *ad libitum* and received corn oil alone as vehicle for the extract via gavage for concessive 28 days. Group II (DEHP-treated group): Rats were fed with normal diet associated with the treatment orally with DEHP (1000mg / Kg bw) via gavage for concessive 28 days. The DEHP dose was selected according to Jain et al. [23]. Group III (*A. senegal* extract post-treated group): Rats received DEHP for the 14 day orally for 14 days followed by oral administration of acacia gum extract for another 14 days. Group IV (*A. senegal* extract simultaneous treated group): Rats received DEHP and treated orally with acacia gum extract at the same time and continue daily for 28 consecutive days via gavages. Group V (*A. senegal* extract treated group): Rats were fed with normal diet associated with the treatment orally with acacia gum extract (300mg / Kg) by stomach tube for concessive 28 days. Group VI (*A. senegal* extract pre-treated group): Rats received acacia gum extract orally for 14 days followed by DEHP for another 14 days.

2.5 Samples Collection

One day after the end of experimental period (i.e., on 29th day), all animals were fasted for 12 h. The animals were anesthetized through slight exposure to diethyl ether. The blood samples which were drawn by retro orbital puncture. The blood samples were allowed for clotting at room temperature and then centrifuged at 3000 rpm for 15 min; the serum was separated and kept in clean stoppered vials at -20°C until the
biochemical assay. The animals were sacrificed by cervical dislocation. Liver and brain tissues were dissected out on ice quickly, cleaned and washed in ice-cold saline then homogenized in 0.01 M Tris–HCl buffer (pH 7.4). Aliquots of this homogenate were used for measuring markers of the oxidative stress and for the different electrophoretic patterns. Also, another brain tissue was homogenized in methanol (75%) in an ice jacket followed by deproteinization then centrifuged in cooling centrifuge at 4°C for 20 min at 5000 rpm for amino acids analysis.

2.6 Biochemical Analysis

2.6.1 Liver functions

Hepatic dysfunction was assessed by measuring activity of liver enzymes such as alanine aminotransferase (ALT), aspartate aminotransferase (AST) [24], alkaline phosphatase (ALP) [25] and acetylcholinesterase (AChE) [26] in serum samples using commercially available kits based on referenced methods.

2.6.2 Oxidative stress markers

The lipid peroxidation (LPO) product was assayed in liver and brain tissue homogenates by spectrophotometric method [27] using a UV-Vis spectrophotometer (Shimadzu uv-2401 pc). Furthermore, the total antioxidant capacity (TAC) was estimated in these tissues by colorimetric method [28], which is based on capacity of the sample to inhibit production of thiobarbituric acid reactive substances from sodium benzoate under the influence of the free oxygen radicals. The antioxidant enzymes such as superoxide dismutase (SOD) [29], catalase (CAT) [30] and glutathione peroxidase activity (Gpx) [31] were measured in liver tissue homogenates.

2.6.3 Brain amino acids and monoamines

Specimen of each brain tissue was weighed and homogenized in 75% aqueous HPLC grade methanol. The homogenate was spun at 4000 rpm for 10 min. and the supernatant was divided into two equal portions; the first one was dried at room temperature using vacuum (70 Millipore) and used for determination of excitatory (Aspartic (ASP) and Glutamic (GLU) and inhibitory amino acids (GABA and Glycine (GLY)) according to method described by Heinrikson and Meredith [32] who used High Performance Liquid Chromatography (HPLC) equipped with the precolumn PITC derivatization technique. Whereas, the second portion was used for determination of brain monoamines (Noradrenalin (NE), dopamine (DA), serotonin (5HT)) and their metabolites (DOPAC, HVA and 5HIAA) by HPLC according to method suggested by Pagel et al. [33]. The resulting chromatogram identified each monoamine position and their metabolites concentration from the sample with respect to that of the standard.

2.7 Statistical Analysis

Significant difference between groups were assessed by one way analysis of variance (one-way ANOVA) followed by least significant difference (LSD) test and confirmed by Bonferroni test to compare the significance among the rat groups. A "P" value of less than 0.05 was considered to indicate statistical significance.

2.8 Histopathological Examination

After sacrifice, autopsy specimens were taken from liver tissues of different experimental groups and immediately fixed in 10% formal saline solution for 24 hr. The tissues were washed in tap water then dehydrated in serial dilutions of alcohol solutions. Tissue fragments were then cleared in xylene and embedded in paraffin and used for histopathological examination. Paraffin wax tissue blocks were prepared for sectioning at 4 microns thickness by sledge microtome. The tissue sections were collected on glass slides and deparaffinized then stained by hematoxylin and eosin (H&E) stain for examination through the light electric microscope [34].

2.9 Samples Preparation

The tissues (liver and brain) were rapidly freezeed with liquid nitrogen and ground then homogenized in water-soluble extraction buffer (0.2 mg / ml). The homogenates were centrifuged at 10,000 rpm for 10 min. The clear supernatant containing water-soluble proteins was transferred to new eppendorf tubes. Individual samples of each group were pooled together and used as one sample and protein concentration was estimated in all pooled samples according to method described by Bradford [35] using bovine serum albumin as standard.
2.10 Electrophoretic Protein and Lipoprotein Patterns

The polyacrylamide gel electrophoresis (PAGE) was carried out at the concentration 10% according to protocol documented by Laemmli [36] using Mini-gel electrophoresis (BioRad, USA). The native and denaturing gels were stained Coomassie Brilliant Blue G-250 for visualizing the protein bands [37]. The relative mobility (Rf), band intensity (Int) and quantity (Qty) of the electrophoretically separated proteins were determined in addition to the molecular weight (Mwt) which was estimated in comparison to marker of standard molecular weights with regularly spaced bands ranging from 6.458 to 195.755 KDa. Moreover, the gel was stained for lipoprotein pattern by Sudan Black B (SBB) [38].

2.11 Electrophoretic Localization of In-gel Enzyme Activity

The native gel was processed for localization of in-gel α- and β-esterase (EST) activities according to method modified recently by Ahmad et al. [39] who postulated that the gel was incubated in reaction mixture containing α, β-naphthyl acetate (5.58 X 10⁻³ mM, pH 7.5) as substrates along with dye coupler Fast Blue RR.

2.12 Genomic DNA Pattern

The genomic DNA was extracted from liver and brain tissues according protocol suggested by Barker et al. [40]. Purity and concentration of the genomic DNA were determined spectrophotometrically using a UV-Vis spectrophotometer (Shimadzu uv-2401 pc). Absorbance was measured at wavelengths of 260 and 280 (A₂₆₀ and A₂₈₀, respectively) nm. The absorbance quotient (OD₂₈₀/OD₂₆₀) provides an estimate of DNA purity. An absorbance quotient value of 1.8 < ratio (R) < 2.0 was considered to be good and accepted, purified DNA. A ratio of <1.8 is indicative of protein contamination, where as a ratio of >2.0 indicates RNA contamination. Integrity of the genomic DNA was assayed using gel electrophoresis (Bio-Rad, USA) by resolving DNA extracts on a 1.5% agarose containing ethidium bromide. The DNA bands were visualized on a UV transilluminator and photographed by Gel Documentation System then analyzed in comparison to DNA molecular weight marker (HyperLadder II) with regularly spaced bands ranging from 50 bp to 2000 bp.

2.13 Data Analysis

The bands in polyacrylamide gel and the DNA fragments in agarose gel were analyzed using Quantity One software (Version 4.6.2). Percent of the similarity index (SI %) was calculated according to formula suggested by Nei and Li [41] to compare all treated groups to control group.

3. RESULTS AND DISCUSSION

Liver is a more sensitive target for DEHP exposure [42]. It is a major organ that is susceptible to be attacked by reactive oxygen species (ROS) [43]. The antioxidant system is required to be developed for maintaining the redox homeostasis in the liver. When the ROS produced in excessive rate, the homeostasis will be disturbed resulting in oxidative stress. Subsequently, this leads to liver injury and other chronic and degenerative disorders [44]. ALT activity is an important index to measure degree of cell membrane damage, while AST is an indicator of mitochondrial damage since it contains 80% of this enzyme [45]. Activity of AChE used as an indicator of liver function. Its activity falls in catabolism and rises in anabolism. Its activity altered in case of hepatocellular lesions [46]. Elevated levels of serum enzymes are considered as indicators of cellular leakage and loss of functional integrity of the cell membrane in liver [47]. As presented in Table 1, levels of hepatic markers (AST, ALT, ALP and AChE) were found to be significantly (P<0.05) elevated in the DEHP-treated group. This was in a consistent with results obtained by Barse et al. [48] who reported that the elevated levels of these markers in sera of DEHP-treated rats correspond to extensive hepatocellular damage and hence release of these in the blood circulation. As compared to DEHP-treated group, A. senegal extract decreased the hepatic markers significantly (P<0.05) and restored its levels to normalcy except in A. senegal extract pre-treated group (GVI). This might be attributed to presence of the higher flavonoid constituents that possess antioxidant properties and was found to be useful in treatment of liver damage. The flavonoids exhibited hepatoprotective activity through free radical scavenging property and its efficiency to inhibit the peroxidation reaction [49].
The LPO produced as a result of peroxidation of polyunsaturated fatty acids in the biological membrane. The LPO level is a measure of alterations and damage in structure of cellular membranes [50]. As revealed in Table 2, activities of enzymatic antioxidants (SOD, CAT and Gpx) decline significantly (P<0.05) in liver tissue of DEHP-treated group. Furthermore, Fig. 1 illustrated that DEHP caused significant (P<0.05) decline in TAC level associated with elevation of LPO level in that tissue. The increased LPO level associated with lowering the TAC level in liver of DEHP-treated group might be attributable to the failure to prevent formation of excess free radicals in a concomitant decline in activities of antioxidant enzymes in liver [20,23]. All these data revealed that DEHP increased oxidative stress through disturbance of cellular redox systems in rat liver [51]. In parallel to results of hepatic markers, A. senegal extract elevated levels of antioxidants with lowering the LPO level in all A. senegal extract treated groups except GVI with respect to the DEHP-treated group.

As illustrated histologically in Fig. 2, DEHP caused irreversible histopathological changes in liver tissue represented by infiltration in bands of fibrosis and inflammatory cells in the portal areas all over the parenchyma associated with dilatation in the congested portal vein. This was in a consistent with results of oxidative stress markers. These alterations might refer to disturbances in the redox system and the oxidative activity of DEHP with subsequent generation of superoxide anion stimulating the peroxidation process. The lipid peroxides accumulate in addition to production of nitric oxide leading to toxic disintegration of cellular organelles and alteration of membrane permeability with subsequent hepatocytes edema and vascular dilatation [6,52]. This was in consistent to results of liver functions and markers of the oxidative stress. A. senegal extract significantly reversed these changes in rats of GIII and GIV groups, while the GVI group showed no LPO restoration. Generally, the body exhibits the ability to neutralize or prevent the toxicity produced by free radicals by using endogenous antioxidant enzymes through an effective defense mechanism. The balance between ROS generation and antioxidant defense mechanism may be lost in DEHP-induced hepatotoxicity as mentioned by Jain et al. [23].

As compared to healthy liver (GI), A. senegal extract lowered severity of the histological alterations to the mild stage of congestion of the hepatocytes in lacunae in all A. senegal extract treated groups except in A. senegal extract treated group (GII). As a result of changes in the redox system, the antioxidant levels in the GIII and GIV groups. The hepatoprotective effect of A. senegal extract was further accomplished by histopathological analysis. Histopathological findings of liver samples were in accordance with the results obtained in biochemical studies, indicating that A. senegal extract is able to inhibit DEHP-induced hepatotoxicity. This might refer to presence of the phenolics and flavonoids (For example, polyphenolic compound Kaempferol, gallic acid, catechin and umbelliferone) which displayed potential scavenging compound against liver toxicity induced by DEHP in rats

<table>
<thead>
<tr>
<th></th>
<th>GI</th>
<th>GII</th>
<th>GIII</th>
<th>GIV</th>
<th>GV</th>
<th>GVI</th>
</tr>
</thead>
<tbody>
<tr>
<td>ALT (U/L)</td>
<td>32.1±0.8</td>
<td>82±0.4^a</td>
<td>35±0.8^b</td>
<td>35±0.6^b</td>
<td>34.4±1.0^b</td>
<td>76.6±0.6^b</td>
</tr>
<tr>
<td>AST (U/L)</td>
<td>38.7±0.6</td>
<td>86.3±0.9^a</td>
<td>38.9±1.5^b</td>
<td>38.9±0.8^b</td>
<td>40.05±0.4^b</td>
<td>74.6±0.8^a</td>
</tr>
<tr>
<td>ALP (U/L)</td>
<td>52.5±0.6</td>
<td>87.7±0.8^a</td>
<td>54.5±1.4^b</td>
<td>54.5±0.9^b</td>
<td>53±1.3^b</td>
<td>86.3±0.9^b</td>
</tr>
<tr>
<td>ACH (U/L)</td>
<td>38.5±2.9</td>
<td>89.1±1.0^a</td>
<td>38.7±1.8^b</td>
<td>38.7±0.7^b</td>
<td>38.5±2.3^b</td>
<td>66.4±0.7^a</td>
</tr>
</tbody>
</table>

Values are expressed as mean ± SE for 6 rats/group. a: significant difference from GI group at P < 0.05, b: significant difference from GII at P < 0.05.

Group I: Control group; Group II: DEHP-treated group; Group III: A. senegal extract post-treated group; Group IV: A. senegal extract simultaneous treated group; Group V: A. senegal extract treated group; Group VI: A. senegal extract pre-treated group
caused significant (p<0.05) elevation of excitatory amino acids (ASP and GLU) associated with decrease inhibitory amino acids (GABA and GLY) in brain tissue. DEHP can pass to brain through the blood barriers due to its chemical structure inducing neurotoxicity by increase excitatory and decrease inhibitory amino acids. In the current study, the excitatory amino acids increased in association with decreasing the inhibitory amino acids. This might occur as a result of neurotoxicity by increase GLU that could be reflecting an alteration of GLU uptake. Moreover, DEHP result in activation of the glutamatergic system and glutamate pathway leading to counter effects on the release of GABA involvement of the Ca²⁺-related excitotoxic process, possibly mediated by the N-methyl-D-aspartate subtype of glutamate receptors [54]. N-methyl D-aspartate, α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid and kainate receptors are the glutamate receptors which activate in case of neurotoxicity. This subsequently leads to increase Ca²⁺ and Na⁺ and decrease K⁺ level in the cell resulting to cellular neuronal death [55]. Additionally, in the present study DEHP caused significant decline in monoamines and their metabolites (DOPAC, HVA and 5HIAA) which declared important pathway implicated to neurotoxicity that mediated by monoamine oxidase. The inhibitory amino acids (GABA and GLY) level in DEHP-treated group was significantly decreased. This may be due to inhibition of GABA synthesis or destruction of GABAergic neurons [56]. A. senegal extract showed amelioration effect for brain amino acids in A. senegal extract treated groups except A. senegal extract post-treated group (GIII) in comparing with DEHP-treated group (GI) and almost recovery in comparing with control (GI) group. On the other hand, GVI
showed slightly amelioration in comparing with GII. A. senegal extract simultaneous treated group (GIV) showed no significant changes in comparing with GI.

GI: Liver section showing no histopathological alteration with normal histological structure of the hepatocytes in the lacunae was recorded (H&E, X 400).

GII: Liver section showing dilated congested portal vein with thickened hyalinized wall. In addition, bands of fibrosis and inflammatory cells infiltration were detected in the portal areas all over the parenchyma (H&E, X 400).

GIII: Liver section showing mild congestion of the hepatocytes in the lacunae (H&E, X 400).

GIV: Liver section showing mild dilated portal vein with thickened hyalinized wall (H&E, X 400).

GV: There was no histopathological alteration and normal histological structure of the hepatocytes in the lacunae (H&E, X 400).

GVI: Liver section showing mild dilated congested portal vein and inflammatory cells infiltration in the portal areas all over the parenchyma (H&E, X 400).

Fig. 2. Histological examination of liver toxicity induced by DEHP and possible prophylactic or therapeutic role of acacia extract

Group I: Control group; Group II: DEHP-treated group; Group III: A. senegal extract post-treated group; Group IV: A. senegal extract simultaneous treated group; Group V: A. senegal extract treated group; Group VI: A. senegal extract pre-treated group.
DEHP caused significant ($P < 0.05$) reduction in monoamines (NE, DA and 5HT) and their metabolites (DOPAC, HVA and 5HIAA) in comparison with control group (GI). As compared to DEHP-treated group, A. senegal extract exhibited showed amelioration effect in groups (GIV, GVI) and almost recovery with respect to control (GI) group. On the other hand, GIII showed slightly amelioration in comparing to GI and showed no significant changes in comparing to GI (Table 4). Earlier researches performed by Keddy et al. [57] confirmed the presence of flavonoids, tannins, flavone, catechin, polyphenols and alkaloids in A. senegal extract. The flavonoids exert the ability to inhibit the enzymes (kinases and phosphatases) which act as critical proteins for signal transduction pathways to regulate oxidative stress and cell survival [57]. The present results postulated that the A. senegal extract has significant neuroprotective and mild treated activity without side effects. This may be probably due to the higher content of tannins and flavonoids that exhibit antioxidant properties through their ability to donate hydrogen and chelate metals [58]. Mean-while, A. senegal extract (phenolic acids) in GIV and GVI groups prevented DEHP induced neurotoxicity by lowering ASP and GLU concentration as well as elevation in GABA, GLY and monoamines (norepinephrine, dopamine and serotonin) in rat brain. Our results in line with Radad et al. [59] who explained the neuroprotective effect of Ginsenosides (flavonoids) as a neuroprotective effects on dopaminergic cells stressed with glutamate. Amelioration and neutralization of excitatory, inhibitory amino acids, monoamines and their metabolites due to co or/ pre treatment of A. senegal extract near normal group may attributed to the antioxidant capacity for phenolic content that decrease LPO that the major marker for amino acids and monoamines disrupted in brain rat intoxicated with DEHP. Obtained data were in agreement with Benavente-Garcia et al. [60] who found that flavanoids and catechins are important antioxidants and superoxide scavengers. Their scavenging efficiency depends on the concentration of phenol and the number and location of the hydroxyl groups.

As compared to the corresponding control (Fig. 3a), several protein bands disappeared in DEHP-treated group associated with appearance of one characteristic band (Rf 0.40, Mwt 30.19 KDa, Int 972.08 and Qty 9.61). Moreover, the lowest SI value was observed in the DEHP treated group (66.67%). Furthermore, it could be observed that 2 abnormal characteristic lipoprotein bands were identified at Rfs 0.18 (Int. 2295.22 and Qty 7.56) and 0.48 (Int. 2905.98 and Qty 9.57) (Fig. 4a). The lowest SI value was observed in the DEHP-treated group (71.43%) and this was constant with results of the proteomic analysis. The treatment with A. senegal extract increased the SI value by restoring the normal bands and hiding the abnormal bands. On the other hand, the electrophoretic protein and lipoprotein patterns in brain tissue (Fig. 3b and 4b) showed no alterations in number or arrangement of the bands in all treated group with respect to control group (the highest SI value 100%). During the current study, DEHP induced changes in the liver electrophoretic protein pattern. This was in accordance with Watanabe et al. [61] who postulated that phthalate caused quantitative changes in about 12 proteins which are considered as part of the pleiotropic response in rat liver. This might be attributed due to peroxisome proliferation, activation of enzymes in the lipid oxidation metabolic pathway.

### Table 3. Prophylactic and therapeutic effect of *acacia* extract on excitatory and inhibitory amino acids against brain toxicity induced by DEHP in rats

<table>
<thead>
<tr>
<th>Groups</th>
<th>ASP</th>
<th>GLU</th>
<th>GABA</th>
<th>GLY</th>
</tr>
</thead>
<tbody>
<tr>
<td>GI</td>
<td>1.13±0.16</td>
<td>0.96±0.10</td>
<td>8.34±0.88</td>
<td>3.79±0.37</td>
</tr>
<tr>
<td>GII</td>
<td>2.23±0.18a</td>
<td>2.13±0.37a</td>
<td>4.51±0.55a</td>
<td>1.77±0.20a</td>
</tr>
<tr>
<td>GIII</td>
<td>2.11±0.23a</td>
<td>2.08±0.27a</td>
<td>5.75±0.46ab</td>
<td>2.08±0.20a</td>
</tr>
<tr>
<td>GIV</td>
<td>1.08±0.10</td>
<td>1.16±0.06</td>
<td>8.21±0.62</td>
<td>3.55±0.20</td>
</tr>
<tr>
<td>GV</td>
<td>1.11±0.10</td>
<td>0.84±0.06</td>
<td>7.79±1.06</td>
<td>3.91±0.37</td>
</tr>
<tr>
<td>GVI</td>
<td>1.58±0.19ab</td>
<td>0.96±0.16b</td>
<td>8.24±0.59b</td>
<td>3.62±0.45ab</td>
</tr>
</tbody>
</table>

Values are expressed as mean ± SE for 6 rats /group. a: significant difference from GI group at $P < 0.05$, b: significant difference from GII at $P < 0.05$.

Group I: Control group; Group II: DEHP-treated group; Group III: A. senegal extract post-treated group; Group IV: A. senegal extract simultaneous treated group; Group V: A. senegal extract treated group; Group VI: A. senegal extract pre-treated group.
Moreover, DEHP caused alterations in the electrophoretic lipoprotein pattern. This might refer to ability of phthalates to alter lipid metabolism through its inhibitory effect on hepatic cholesterologenesis and 3-hydroxy-3-methylglutaryl CoA reductase (EC 1.1.1.34) which is considered as the key regulatory enzyme in hepatic sterol synthesis [62].

As presented in Fig. 5a, it was revealed that DEHP caused no qualitative alterations in number and arrangement of α-EST bands in liver tissue. Otherwise, it caused mutagenicity at the quantitative level by increasing quantity of the α-EST2 band (Rf 0.78, Int. 3148.55 and Qty 16.96) (Fig. 5b). The treatment with A. senegal extract decreased quantity of the α-EST2 band in all extract treated groups. As regards β-EST enzyme, it was observed that DEHP caused qualitative mutagenicity represented by disappearance of β-EST2 band with appearance of 3 characteristic ones at Rfs 0.31, 0.43 and 0.52 (Int. 1131.16, 5391.56 and 1178.68; Qty 0.76, 3.60 and 0.79, respectively). Moreover, the lowest SI % was noticed in DEHP-treated group (66.67%). A. senegal extract increased the SI % to the highest value (100%) through restoring the β-EST2 band associated with hiding the characteristic ones (Fig. 6a).

Table 4. Prophylactic and therapeutic effect of *acacia* extract on monoamines and their metabolites against brain toxicity induced by DEHP in rats

<table>
<thead>
<tr>
<th>Groups</th>
<th>Monoamines concentration (µg/g)</th>
<th>NE</th>
<th>DA</th>
<th>5HT</th>
<th>DOPAC</th>
<th>HVA</th>
<th>5HIAA</th>
</tr>
</thead>
<tbody>
<tr>
<td>GI</td>
<td>0.98±0.07</td>
<td>2.09±0.20</td>
<td>0.87±0.10</td>
<td>0.84±0.04</td>
<td>0.66±0.07</td>
<td>0.71±0.06</td>
<td></td>
</tr>
<tr>
<td>GII</td>
<td>0.43±0.03&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.02±0.17&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.57±0.04&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.49±0.07&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.48±0.07&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.51±0.06&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td>GIII</td>
<td>0.58±0.06&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>1.25±0.14&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>0.65±0.08&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>0.61±0.06&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>0.55±0.05&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>0.53±0.01&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td>GIV</td>
<td>0.70±0.07&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>1.56±0.25&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>0.66±0.05&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>0.76±0.04&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.71±0.06&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.70±0.10&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td>GV</td>
<td>0.81±0.08</td>
<td>1.87±0.13</td>
<td>0.77±0.06</td>
<td>0.76±0.07</td>
<td>0.67±0.09</td>
<td>0.75±0.05</td>
<td></td>
</tr>
<tr>
<td>GVI</td>
<td>0.70±0.10&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>1.98±0.10&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.82±0.02&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>0.74±0.05&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>0.70±0.10&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.73±0.08&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
</tr>
</tbody>
</table>

Values are expressed as mean ± SE for 6 rats /group. a: significant difference from GI group at P < 0.05, b: significant difference from GII at P < 0.05.

Group I: Control group; Group II: DEHP-treated group; Group III: A. senegal extract post-treated group; Group IV: A. senegal extract simultaneous treated group; Group V: A. senegal extract treated group; Group VI: A. senegal extract pre-treated group.

Fig. 3. Electrophoretic protein patterns showing prophylactic and therapeutic effect of daily oral administration of *acacia* extract against tissue toxicity induced by DEHP on a) liver tissue and b) brain tissue of rats

Group I: Control group; Group II: DEHP-treated group; Group III: A. senegal extract post-treated group; Group IV: A. senegal extract simultaneous treated group; Group V: A. senegal extract treated group; Group VI: A. senegal extract pre-treated group.
Moreover, in the brain tissue, α-EST pattern was represented by only one band. DEHP caused no qualitative abnormalities. Otherwise, it caused qualitative alterations by enhancing quantity of the α-EST band (Rf 0.21, Int. 1618.29 and Qty 15.66). The treatment with A. senegal extract decreased quantity of the α-EST band in all extract treated groups (Fig. 5b). As regards β-EST enzyme, it was noticed that phthalate caused mutagenicity represented qualitatively by disappearance of β-EST1, β-EST2, β-EST3 and β-EST5 bands with existence of 2 characteristic bands at Rfs 0.44 and 0.75 (Int. 105.78 and Qty 0.54 and Qty 1.62, respectively). Furthermore, it caused quantitative alterations through decreasing quantity of the β-EST4 band (Rf 0.86, Int. 190.50 and Qty 0.97) with respect to control one (Rf 0.86, Int. 284.94 and Qty 1.61) (Fig. 6b). In addition, the lowest SI value was observed in DEHP-treated group (25%). The SI % increased and reached the highest value (100%) in all A. senegal extract treated groups through restoring the absent bands and hiding the characteristic ones. Moreover, quantity of the β-EST4 band still lowered in all DEHP-treated groups that treated with extract in combination. Esterases belong to the lysosomal lipolytic enzymes that catalyze hydrolysis of the neutral lipids (triglyceride and cholesterol esters) which are introduced into cells as components of lipoproteins and lipid deposits [63]. During the present study, DEHP caused mutagenicity detected electrophoretically in the α- and β-esterases. This might be attributable to its effect on the membrane bound enzymes that are sensitive to changes in membrane fluidity which reflects alterations in the physical state of the membrane-lipids [64]. Moreover, the inhibitory or stimulatory effect of phthalates on various membrane bound enzymes is an indirect effect of their ability to modify membrane fluidity. In addition, phthalates exhibit lipophilic activity and can partition across organelar membranes and disturb the physical composition of the lipid microenvironment surrounding the particulate enzymes [65]. In the brain tissue, DEHP caused quantitative changes in the α- and β-EST patterns. This may be attributed to the inhibitory effect of DEHP on ChE. Consequently, this leads to accumulation of Ach and then activation of glutamatergic neurones by increase GLU that is a major agonist of N-methyl D-aspartate receptors and a major excitatory neurotransmitter in the central nervous system, as well as being a potent excitotoxin [66].

As illustrated in Fig. 7a, it was noticed DEHP induced cleavage of the genomic DNA in liver tissue giving 3 unique fragments at Rfs 0.17, 0.56 and 0.88 (Mwts 2585, 1100 and 1384 BP;
Q % 9.97, 14.05 and 20.49, respectively). With respect to control, the genomic DNA remains integrated in the DEHP-treated groups that treated with A. senegal extract in GIII and GIV groups. While in the GVI group, there was extra band identified at Rf 0.18 (Mwt 2552 BP and Q % 9.54). Furthermore, DEHP induced DNA damage severely in the brain tissue (Fig. 7b) giving 4 fragments at Rfs 0.1, 0.26, 0.51 and 0.80 (Mwts 3465, 1214 and 798 BP; Q % 4.33, 3.60, 3.36 and 9.02, respectively). In addition, the genomic DNA pattern still damaged in brain tissue of A. senegal extract treated groups. It was noticed that 3 fragments were identified at Rfs 0.10, 0.33 and 0.50 (Mwts 3542, 1631 and 1223 BP; Q % 3.88, 5.05 and 6.98, respectively) in GIII group, 2 fragments identified at Rfs 0.10 and 0.30 (Mwts 3516 and 1777 BP; Q % 4.92 and 4.55) in GIV group. In addition 3 fragments were identified at Rfs 0.10, 0.52 and 0.83 (Mwts 3465, 1196 and 766 BP; Q % 8.04, 3.99 and 8.31, respectively) in GVI group.

Fig. 5. Electrophoretic α-esterase pattern showing prophylactic and therapeutic effect of daily oral administration of acacia extract against tissue toxicity induced by DEHP on a) liver tissue and b) brain tissue of rats

Group I: Control group; Group II: DEHP-treated group; Group III: A. senegal extract post-treated group; Group IV: A. senegal extract simultaneous treated group; Group V: A. senegal extract treated group; Group VI: A. senegal extract pre-treated group

Fig. 6. Electrophoretic β-esterase pattern showing prophylactic and therapeutic effect of daily oral administration of acacia extract against tissue toxicity induced by DEHP on a) liver tissue and b) brain tissue of rats

Group I: Control group; Group II: DEHP-treated group; Group III: A. senegal extract post-treated group; Group IV: A. senegal extract simultaneous treated group; Group V: A. senegal extract treated group; Group VI: A. senegal extract pre-treated group
During the current study, DEHP exhibited ability to produce DNA damage. This was in accordance with Kornbrust et al. [67] who suggested that phthalate induced DNA damage, either directly or as a result of the peroxisomes proliferation associated with increased production of hydrogen peroxide and other DNA-damaging oxygen radicals.

Treatment with A. senegal extract reversed these changes in the electrophoretic and genomic patterns. This might be attributed to ability of the extract to inhibit xanthine oxidase and superoxide radicals and hence to normalize the imbalanced antioxidant system in hepatocytes [20,68]. These findings were supported by results of the oxidative stress markers reported earlier. As regards in the brain tissue, phthalate induced cleavage of the genomic brain DNA. This may refer to the severe changes which include alterations in DNA methylation, histone proteins, non-coding RNA, or chromatin arrangement leading to corresponding gene expression changes in the brain and hence modifications in neuronal cells [69]. A. senegal extract could not prevent the DNA damage in the brain. This because dose of the extract was not sufficient to exhibit its beneficial activity on the genomic DNA. In parallel to the study suggested by Sharma et al. [70], A. senegal can be added as an adjuvant to existing therapies for the treatment of brain related disorders. However, further studies required to reveal the most suitable dose targeting different brain areas to prove the benefits conclusively.

4. CONCLUSION

The present study was conducted that DEHP caused elevation of hepatic enzymes (ALT, AST, ALP and AChE) with lowering activities of antioxidant enzymes (TAC, SOD, CAT and Gpx). In coincide with these biochemical results, it was noticed that DEHP caused severe histopathological alterations in the liver tissue. A. senegal extract lowered the hepatic markers and restored activities of the antioxidant enzymes to normal values. Also, the adverse effect of DEHP targeted the brain tissue. This finding was supported by elevation of excitatory amino acids (ASP and GLU) and decline of inhibitory amino acids (GABA and GLY) and monoamines in brain of DEHP-treated rats. A. senegal extract exhibited ameliorative effect on levels of brain amino acids, monoamines and their metabolites. DEHP caused mutagenicity in the native protein, lipoprotein and isoenzyme patterns and in liver tissues. No changes detected in electrophoretic protein and lipoprotein patterns in brain of DEHP-treated group. Furthermore, DEHP induced cleavage of the genomic DNA in liver and brain tissues. A. senegal extract exhibited beneficial activity by restoring the normal bands and hiding the abnormal ones.
COMPETING INTERESTS

Authors have declared that no competing interests exist.

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