Effect of Triclosan on the Renal Cortex of Adult Male Albino Rats and the Possible Protective Role of Ellagic Acid: Histological and Biochemical Study

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Abstract

Background: Triclosan (TCS) is widely used broad spectrum bactericide. Ellagic acid (EA) has radical scavenging properties.

Aim of the work: to assess the structure of the renal cortex after TCS administration and to determine the possible protective role of EA.

Materials and methods: Thirty adult male albino rats were divided into three groups; control group, TCS treated group; TCS was administered 200 mg/kg, once daily for six week, by oral gavages. TCS-EA treated group: received in the same doses of TCS and EA (30 mg/kg) for six weeks. Blood urea nitrogen (BUN), serum creatinine (SC) and uric acids (UA) were measured. Renal cortex samples were processed for light and electron microscope examination.

Results: Examination of TCS treated group revealed increased glomerular cellularity in some corpuscles. Podocytes showed effacement of the foot processes and focal thickening of the glomerular basement membrane. Some tubules showed marked cellular disorganization. Disoriented basal mitochondria, areas of rarified cytoplasm and electron-dense bodies were noticed. Intense positive caspase-3 reaction was expressed in the tubular cells. TCS-EA showed improvement of morphological organization of renal cortex glomeruli, proximal and distal tubules and BUN in TCS treated group in comparison with the control group. Their levels in TCS-EA treated group showed a significant decrease in comparison with TCS treated group

Conclusion: In conclusion, TCS leads to alterations in the histological structure and functions of renal cortex of albino rats and EA supplementation could protect from these changes.

Keywords: Renal cortex; Triclosan; Ellagic acid; Caspase-3

Introduction

Triclosan (TCS) (2,4,4’-trichloro-2’-hydroxydiphenyl ether) is a member of bisphensol compounds [1]. TCS is a broad spectrum bactericide. It is extensively used in many personal care products including soap, deodorants, hand sanitizer, cosmetics and toothpaste as well as household products such as odor-fighting socks and germ-resistant sponges [2]. TCS has antiseptic activity through disrupting bacterial membrane activities and inhibits fatty acid synthase involved in lipid biosynthesis [3].

The widespread TCS use, make the potential to its exposure for the general population through ingestion of food and drinking water contaminated with it or dermal contact with products containing it [3,4]. TCS was detected in the blood [5], breast milk [6] and urine [7]. It is bioaccumulated and concentrated in the fatty tissues of worldwide people [8]. It is also has been found in rivers, streams and in sewage sludge applied to agriculture [9]. Urinary excretion was found to be a major route of elimination [3].

TCS is an endocrine disruptor; previous studies have shown that TCS disrupts thyroid hormone action [10]. TCS also has estrogenic and androgenic hormone properties. TCS exposure could potentially contribute to the development of breast cancer [11].

Flavonoids, which are polyphenolic antioxidants, occur naturally in vegetables and fruits. Ellagic acid (2,3,7,8-Tetrahydroxycromen[5,4,3-cd]chromene-5,10-dione) is also a naturally occurring phenolic constituent in certain fruits and nuts [12]. EA is mostly abundant in berries, walnuts, pecans, pomegranate, cranberries and other plant foods in the forms of hydrolysable tannins called ellagitannins [13]. Ellagic acid (EA) as phytochemical has received particular attention because of its wide array of biological properties, such as radical scavenging, antiviral and antibacterial properties [14].

Programmed cell death occurs following the induction of an intracellular genetically regulated cell death program. A number of physiological and pathological stimuli including lack of nutrients, activation of cell surface death receptors, chemicals, ionizing radiation and direct physical injury can activate the apoptosis [15].

Caspases are a group of enzymes that are involved in the regulation of apoptosis resulting in the classical apoptotic features. Caspase 3 activation occurs in response to variety of apoptotic inducers which activate endonuclease and induce DNA fragmentation [16].

Despite the extensive use of TCS, few studies have investigated the

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toxicological effect of TCS. So, this study aimed to assess the structure of the renal cortex after TCS administration and to evaluate the possible protective role of EA.

**Material and Methods**

**Material**

- **Chemicals:**
  - **Triclosan** (Sigma Aldrich Company, Methyl TCS, C11H9ClO2, CAS Number: 4640-01-1).
  - **Ellagic acid** (Sigma Aldrich Company, E2250, ≥95% (HPLC), powder, from tree bark, CAS-No.: 476-66-4).
  - **Caspase-3:** anti-Caspase 3 (CPP32) Ab-4, 4, rabbit polyclonal antibody (CAT. # RB-1197-P0, ready to use, Thermo Scientific™ Lab vision).

- **Kits:**
  - Biomerieux France kits for estimation of blood urea nitrogen (BUN) and serum creatinine (SC) and uric acid (UA).
  - **Cap3:** anti-Caspase 3 (CPP32) Ab-4, 4, rabbit polyclonal antibody (CAT. # RB-1197-P0, ready to use, Thermo Scientific™ Lab vision).

- **Experimental animals:** Thirty adult male albino rats, with an average weight of 200 g, were used in this study. Rats were housed in wire mesh cages and were fed with pelleted standard chow diet in addition to tap water. Animals were kept for 7 days before beginning the experiment for acclimatization. The experimental protocols were approved by the Ethical Committee of Zagazig University.

**Methods**

- **Experimental design:** Animals were randomly divided into the following groups: Group I (control group): This group included 18 rats that was further equally subdivided into 3 equal subgroups; Group IA (negative control group): were gavaged with 1 ml of distilled water once daily for 6 weeks. Group IB (corn oil positive control group): were orally gavaged 1 ml of corn oil (a vehicle) once daily for six weeks. Group IC (EA positive control group): were gavaged with EA alone. Group II (TCS treated group): included 6 rats that received TCS alone. The dose was selected on the basis of the oral LD50 (5000 mg/kg, in rats) [17]. Group III (TCS-EA treated group): included 6 rats that received the same dose of TCS, 1 hour subsequent to administration of EA.

Rats of groups II and III received TCS in a dose of 200 mg/kg (suspended in 1 ml of corn oil) by oral gavages once daily for five weeks [15]. Rats of groups IC and III received EA at a dose 30 mg/kg (dissolved in corn oil) by oral gavages, once daily for six weeks [18].

At the end of experiment, the rats were anesthetized with 50 mg/kg, bw sodium pentobarbital intraperitoneally [19]. Blood samples were obtained directly from the heart by cardiac puncture. Blood samples were collected into tubes containing ethylenediamine-tetraacetic acid (EDTA) and centrifuged at 3000 g for 10 min. Plasma was separated and then stored at 20°C until analyzed. Then intra-cardiac perfusion was done by 2% glutaraldehyde for fixation.

**Histological study:** The cortex of both kidneys of each animal was dissected, excised and cut into smaller pieces and processed for light (1cm³) and electron microscope (1mm³) study. To prepare paraffin blocks, specimens were immediately placed in 10% buffered formalin. After 10 min, when the tissue was hardened to avoid soft tissue dissipation, specimens were fixed in 10% buffered formalin for 24 h and processed to prepare 5 μm sections stained with haematoxylin and eosin, Masson's trichrome and PAS stains [20].

The immunohistochemical staining for localization of the caspase-3 was carried out by means of the avidin–biotin– peroxidase complex method following the manufacturer's instructions (Dako Company, Wiesentheid/Bavaria, Germany, Biotin Blocking System, Code X0590). Paraffin sections of 4 μm were deparaffinized in xylene and rehydrated in a descending series of ethanol. The specimens were subjected to antigen retrieval in a citrate buffered solution (pH 6.0) for 10 minutes using a microwave. Endogenous peroxidase was eliminated by incubation in 10% H2O2 in phosphate-buffered saline (PBS), pH 7.4 for 10 minutes. After washing, the specimens were blocked in ready-use normal goat serum for 20 minutes at room temperature. Then, the sections were incubated with the specific anti-Caspase 3 primary antibody at room temperature for 30 minutes. Dilutions were done with antibody diluent (TA-125-UD; Lab Vision). Peroxidase activity was demonstrated using an AEC (3-amino-9-ethyl carbazole) substrate kit (TA-004-HAC; Lab Vision). The sections were rinsed in PBS. It was applied overnight (1:200) in a humidified chamber at 4°C then washed in PBS twice. The secondary antibody was anti-rabbit antibody universal kit (code no. K0773, lot). Sections were covered with biotinylated secondary antibody for 30 minutes and then washed in PBS. Then by peroxidase–labeled avidin– biotin solution reaction (Novo Stain Super ABC Kit, Novocastra, Newcastle upon –Tyne, UK) for 45 minutes. Then washed in PBS. Finally, freshly prepared diaminobenzidine (Sigma, St. Louis, MO) was added for 4 minutes. A chromogen, washed with distilled water followed by Mayer's haematoxylin as a counter stain. The sections were washed, dehydrated mounted and examined. For the negative control, the same steps were followed, but the primary antibody was replaced by PBS [21]. Brown caspase 3 positive immunoreaction is predominantly cytoplasmic with some nuclear staining [22].

Specimens for electron microscope were immediately fixed in 2.5% phosphate-buffered glutaraldehyde (pH 7.4). Renal cortex specimens were immediately fixed in 2.5% glutaraldehyde in 0.1 M cacodylate buffer at pH 7.4 for 2 h at 4°C. Then the specimens were washed with the buffer, post fixed in 1% osmium tetroxide in distilled water for 2 h at 4°C. They were dehydrated with ascending grades of ethanol and then put in propylene oxide to prepare Epon-Araldites resin blocks. Ultrathin sections were stained with uranyl acetate and lead citrate [23] and examined and photographed using a JEOL JEM 1010 electron microscope (JEOL Ltd, Tokyo, Japan) in the Electron Microscope Research Laboratory of the Histology and Cell Biology Department, Faculty of Medicine, Zagazig University (Egypt).

**Morphometric study:** Quantitative morphometric measurements were obtained using a Leica Qwin 500 image analyzer computer system (Leica, Hessen, Germany): Faculty of Medicine, Zagazig University, Egypt. The area percentage (area %) of positive immune reaction for caspase-3 was measured using the interactive measure menu. The area percentage and standard measuring frame of a standard area equal to 118 476.6 µm2 were chosen from the parameters measuring 10 readings from five sections from each rat of the randomly chosen five rats in each group. In each randomly chosen field, the section of the renal cortex was enclosed inside the standard measuring frame; the areas where brown positive immune reaction for caspase-3 were measured. The whole data was subjected to statistical analysis.

**Biochemical study:** Assessment of kidney function tests had been carried out using kit of Biormerieux France.

**Blood urea nitrogen (BUN) (mg/dL) and serum creatinine (SC) (mg/dL):** They were done according to the pamphlet of Biormerieux France kit by enzymatic colorimetric method.
Serum uric acid (mmol/L): Serum uric acid was measured according to the pamphlet of Biomerieux France kit, by a carbonate phosphotungstate method.

Statistical analysis: Data obtained from hormonal assay and morphometry were expressed as mean ± SD. They were fed into the computer using statistical package for the social sciences (SPSS, version 20; IBM, Armonk, New York, USA) software package. Statistical analysis was carried out using one-way analysis of variance and the posthoc test (Scheffe) for pairwise comparison. The level of significance was set at P value less than 0.05.

Results

Histological results

Light microscope results: Histological examination of all control subgroups IA, IB and IC showed nearly similar structure. Figures for subgroup IA were used to differentiate with other groups. Examination of H & E stained sections of the control adult male albino rat renal cortex showed normal renal structure. Each renal corpuscle was formed of a glomerular tuft of capillaries surrounded by Bowman’s capsule. Proximal and distal convoluted tubules cells showed acidophilia of proximal (P) and distal (D) convoluted tubules cells with vesicular nuclei. However, few distorted tubules were also detected (Plate 1D). Examination of TCS treated group revealed intense positive PAS reaction in the renal cortex, glomeruli of renal corpuscles appeared with dark stained nuclei. Some tubules showed desquamation of tubular epithelial cells while others show marked cellular disorganization. Intracellular vacuoles and luminal casts were observed in other tubules. Dilated tubules with flattened epithelium were detected. Many inflammatory cells and congested blood vessels were also seen. (Plate 1B and 1C). Examination of TCS-EA showed improvement of morphological organization of renal cortex, glomeruli of renal corpuscles appeared with their capillary tufts surrounded by Bowman’s capsules. Proximal and distal convoluted tubules cells appeared with acidophilic cytoplasm. Most of their nuclei were vesicular. However, few distorted tubules were also detected (Plate 1D).

Examination of Masson’s trichrome stained sections of control group renal cortex group showing thin blue collagen fibers around renal corpuscles and tubules (Plate 2A). On the other hand, examination of TCS treated group revealed excess blue-stained collagen fibers around tubules (arrow), around and within glomeruli (arrowhead). Foci of interstitial fibrosis (curved arrow) are also noticed. C: Triclosan-ellagic acid treated group showing blue collagen fibers (arrow) around the renal corpuscle and tubules. D: Control group showing moderate PAS +ve reaction in the basal lamina (arrow), luminal brush border (arrowhead) of the renal tubules (T) and renal glomeruli (G). E: Triclosan treated group showing strong PAS +ve reaction in the renal glomeruli (G) while renal tubules (T) show moderate reaction in basal lamina (arrow), luminal brush (arrow head) border. F: Triclosan-ellagic acid treated group showing moderate PAS +ve reaction in the basal lamina (arrow), luminal brush (arrow head) border of the renal tubules (T) and glomeruli (G). (scale bar 25 μm, A, B, C; Masson’s trichrome stain & D, E, F; PAS stain).

Examination of PAS stained sections of control group renal cortex group showing thin blue collagen fibers around renal corpuscles and tubules (Plate 2A). On the other hand, examination of TCS treated group revealed excess blue-stained collagen fibers around the tubules, around and within the glomeruli. Foci of interstitial fibrosis were also noticed (Plate 2B). In TCS-EA treated group, blue collagen fibers were observed around the renal corpuscle and tubules (Plate 2C).

Examination of PAS stained sections of control group revealed positive PAS reaction in the basal lamina, luminal brush border of the renal tubules and renal glomeruli (Plate 2D). On the other hand, TCS treated group revealed intense positive PAS reaction in the renal
glomeruli while renal tubules show moderate reaction (Plate 2E). In TCS EA treated group, moderate positive PAS reaction in the basal lamina and luminal brush border of renal tubules and glomeruli were detected (Plate 2F).

Immunohistochemical examination of control group revealed minimal positive cytoplasmic staining for caspase-3 in cells of proximal and distal tubular cells (Plate 3A). However, TCS treated group showed intense caspase-3 positive reaction expressed in the cytoplasm and nuclei of the tubular cells and moderate reaction were also noticed in glomeruli (Plate 3B). In addition, TCS-EA treated group showed moderately expressed caspase-3 reaction in glomeruli, proximal and distal tubules (Plate 3C).

Electron microscope results: Ultrastructural examination of the glomeruli of control group kidney showed podocytes with irregular euchromatic nuclei, primary processes and interdigitating secondary processes. The glomerular basement membranes (GBM) were formed of central electron dense lamina densa and lamina rarae on either side. In TCS treated group, many podocytes showed electron dense nuclei with effacement of the foot processes (arrow head) and focal thickening of the glomerular basement membranes (curved arrow). In TCS-EA treated group, podocytes appeared with irregular electron lucent nuclei. Relatively normal arrangement of secondary foot processes and regular thickening of the GBM with restoration of its trilamellar structure are seen. Fenestrated blood capillaries were also detected (Plate 4E and 4F).

Examination of proximal convoluted tubules (PCTs) of control group showed epithelial cells lining proximal convoluted tubules with...
long apical microvilli and large euchromatic nuclei, some of them had prominent nucleoli. Many elongated basal mitochondria were seen in-between extensive basal infoldings. All cells were rest on basal lamina (Plate 5A and 5B). In TCS treated group, cells showed disoriented basal mitochondria with very few basal infoldings. Multiple cytoplasmic vacuoles, areas of rarified cytoplasm and electron-dense bodies were noticed. Nuclei with clumps of heterochromatin and apoptotic nuclei were apically displaced. Casts with different electron density were found at the tubular lumen. Apical long microvilli were preserved on some cells while others show sparse short microvilli. Irregular basal lamina were also noticed (Plate 5C and 5D). TCS-EA group showed many PCT cells with numerous apical long microvilli, euchromatic nuclei, many mitochondria, electron-dense bodies and apical vacuoles were detected in their cytoplasm. Cells were rest on regular basal lamina (Plate 5E and 5F).

Examination of distal convoluted tubules (DCTs) of control group revealed epithelial cells with oval nuclei and few short apical microvilli. Many elongated basal mitochondria were seen in-between basal infoldings. Small vacuoles were found in the cytoplasm (Plate 6A and 6B). In TCS treated group, cells showed areas rarified cytoplasm, many mitochondria; some of them appeared swollen with disrupted cristae. Some cells show nuclei with multiple peripheral patches of heterochromatin while others have deeply indented nuclear envelope. Apical displacements of all nuclei are also noticed. In TCS-EA group, many cells had euchromatic nuclei, few short apical microvilli, and mitochondria in-between basal infoldings (Plate 6C and 6D).

**Morphometrical Results:** The mean area percentage of caspase-3 expression for all groups is presented in (Table 2): there was a significant increase (P<0.05) in caspase-3 expression in TCS treated group (II) compared with control group (I). There was a significant decrease (P<0.05) in caspase-3 expression in TCS-EA treated group (III) compared with TCS treated group (II).
Biochemical results

The mean values of blood urea nitrogen (mg/dl), serum creatinine (mg/dl), uric acid (mmol/L) for all groups is presented in Table 1. There was no significant difference between the biochemical results of group IA, group IB, and group IC. The biochemical results of group IA were chosen to express the results of control group and compare with the rest of groups. There was significant increase in the levels of BUN, SC and UA in TCS treated group in comparison with the control group. On the other hand, the level of BUN, SC and UA in TCS-EA treated group showed a significant decrease in comparison with TCS treated group.

Discussion

TCS is as a chlorinated aromatic polychloro-phenoxy phenol compound that has functional groups representative of both ether and phenols [24]. Biodegradation and photolytic degradation diminish the availability of TCS. However, TCS by-products such as methyl-triclosan and other chlorinated phenols may be more resistant to degradation and have higher toxicity than the parent compound [10]. Also, TCS irradiation by sunlight can degrade into a form of dioxin, a class of chemicals linked to a broad range of toxicities cispla [25,26].

Examination of TCS treated group revealed increased glomerular cellularity with obliteration of the Bowman's space in some glomeruli while others appeared destructed with dark stained nuclei. Many podocytes had electron dense nuclei with effacement of the foot processes and focal thickening of the glomerular basement membranes. The mechanism of TCS induced cytotoxicity could be explained by Lack and Reed and Mathur and D'Cruz [27,28]. Lack and Reed [27] stated that TCS metabolism have been shown to undergo via a minor pathway of glutathione conjugation. These conjugates formed accumulate within the kidney where they undergo activation to a chemically reactive species which can introduce an acyl group into proteins leading to renal cell necrosis and compensatory cell regeneration. Mathur and D'Cruz [28] stated exposure to TCS caused an imbalance in pro-oxidant/ antioxidant levels and thereby induces the generation of ROS. Tipping [29] postulated that podocytes are generally regarded as targets of reactive ROS by stimulating podocyte production of the proinflammatory cytokine GM-CSF that induced apoptosis in podocytes and directly injure the glomerular basement membrane. Binelli et al. [30] added that TCS acts as a DNA adduct and exerts genotoxic effects leading to apoptotic damage.

In TCS treated group, some tubules showed desquamation of tubular epithelial cells while others show marked cellular disorganization. Dilated tubules with flattened epithelium were detected. Many inflammatory cells and congested blood vessels were also noticed. Tubular cells had many multiple cytoplasmic vacuoles, areas of rarefied cytoplasm electron-dense bodies and luminal casts were detected. These findings were in accordance with the works performed by Dann and Hontela [10] who reported that TCS were primarily changes in the kidneys evidenced by inflammation and tubular degeneration. Moreover, previous reports linked the chronic TCS toxicity with the occurrence of focal interstitial nephritis, fibrosis, tubular dilatation and eosinophilic casts in the tubular lumina [31]. Barlas and Aydoğan [32] stated that phenol induced enlargement of intertubular area with hemorrhage in rats. Tootian et al. [33] attributed that phenol inducing tubular degeneration to vascular stasis in the kidney. The degenerative alterations may be due to its ability to impair epithelial cell membrane integrity. U.S Food and Drug Administration [34] reported that TCS induced renal changes by ROS damaging of variety of transport proteins including Na/K ATPase [35]. Reduced Na+, K+-ATPase activity in the proximal tubular epithelial cells resulted in intracytoplasmic vacuolation, disruption of the arrangement and swelling of the mitochondria and large number of lysosomes [36].

In TCS treated group, disorganization of basal mitochondria with very few basal infoldings were detected in renal tubule cells. Also, nuclei with clumps of heterochromatin and apoptotic nuclei were also detected. These results came in agreement with Zucherbraun et al. [38] who suggested that TCS induced tubular cell death by apoptosis. Also, Newton et al. [1] proved that TCS can impair fluidity of the inner mitochondrial membrane resulting in mitochondrial dysfunction. Tootian et al. [33] stated that inefficiency of renal function and structure.
is attributed to mitochondrial destruction. Previously, Kaneda et al. [39] reported that mitochondrial dysfunction reduces active transport of renal tubules which impairs reabsorption capacity of the tubules that may progress to cell death.

TCS treated group showed excess collagen fibers around the tubules, around and within the glomeruli. These results were supported by Efratiadias et al. [40] who stated that progressive loss of renal function is associated with development of tubulointerstitial renal fibrosis with destruction of renal tubules and interstitial capillaries and accumulation of extracellular proteins. Pozdžik et al. [41] explained that by epithelial to mesenchymal transition (EMT); a process in which of tubular epithelial cells are transformed to mesenchymal fibroblasts migrating to adjacent interstitial parenchyma constitutes main mechanism of renal fibrosis along with local and circulating cells.

TCS administration results in an apparent increase PAS reaction in the renal glomeruli and moderate reaction in renal tubules. Thickened GBM with distorted foot processes of podocytes was noticed. Rateb and Abdel-Hafez [42] attributed this thickening may develop to compensate for the increased glomerular permeability and proteinuria. Oh et al. [43] demonstrated that oxidative stress induced deleterious structural changes, including loss of the brush border membrane, deposition of PAS positive materials, and cast formation.

Examination of TCS-EA group revealed improvement of morphological organization of renal cortex with regular arrangement of secondary foot processes and regular thickening of the GBM with restoration of its trilamellar structure. Cells of proximal and distal convoluted tubules cells appeared nearly normal however few distorted tubules were also detected. These results are in agreement with Al-kharusi et al. [18] who proved that EA ameliorate oxidative stress toxicity and decreased the tubular necrosis and apoptotic cells of the renal cortex in rats treated with EA. Ateşşahin et al. [12] reported that EA protected the kidney from tubular degeneration, desquamation and tubular dilatation. Chirino and Pedraza-Chaverri [44] postulated that introduction of an exogenous antioxidant is thought to help the endogenous antioxidant system in scavenging the ROS produced during an imbalance in redox status. In view of the fact that TCS treatment results in decreasing in kidney glutathione [12], EA increased the activity of three antioxidant enzymes superoxide dismutase, glutathione peroxidase and catalase which help in free radical attack [45]. Also, Han et al. [46] reported that EA exhibited anti-fibrosis effect and increase of free radical scavenging activity and inhibited lipid peroxide production. EA inhibits cytokine-induced ROS, inflammation, and expression of adhesion molecules [47]. Moreover, recent studies have confirmed that EA reducing apoptosis through scavenging ROS [48].

Immunohistochemical examination of TCS treated group showed was a significant increase of caspase-3 expression in TCS treated group in the cytoplasm and nuclei of tubular cells comparing with control group. On the other hand, there was a significant decrease of caspase-3 expression in TCS-EA treated group in cells of proximal and distal tubules comparing to TCS treated group. These changes of cleaved caspase-3 due to activation of apoptosis pathways are in accordance with Newton et al. [1] who stated that mitochondrial depolarization induced by TCS is an early event in the promotion of apoptosis. Loro et al. [15] reported that TCS can induce apoptosis by increased levels of ROS [49]. This can subsequently activate extrinsic (Fas and FasL) and intrinsic (mitochondrial) apoptotic pathways. Intrinsic apoptosis pathway is regulated by pro-survival proteins and pro-apoptotic protein. Dysregulation of these proteins results in the release of cytochrome C (pro-apoptotic factor) from outer mitochondrial membrane to cytoplasm resulting in activation of caspase-9. Activated caspase-9 triggers the effector caspase-3 [50]. Kuhla et al. [51] stated that extrinsic apoptotic pathway by binding of death receptors which, bind to caspase-8 and activates effector caspase-3. The two divergent pathways of apoptosis converge on the common downstream effector, caspase-3. Pozdžik et al. [41] mentioned that progressive tubular atrophy was related to induction of apoptosis secondary to caspase-3 activation and defective regeneration of tubular epithelial cells. Kaushal et al. [52] the extent of cell injury caused by a toxic agent depend on the balance between activation of caspas triggered by a toxic agent and on the induction of survival signals capable of blocking the activation of caspases. It was stated that the anti-apoptotic effects of EA are due to its inhibition of ROS generation, which, in turn, represses the accumulation of intracellular calcium, stabilizes the mitochondrial membrane and prevents the release of cytochrome c which required for the activation of caspase-3 [53].

In the present work, biochemical results revealed a significant increase in the levels of blood BUN, SC and UA in TCS treated group in comparison with control group. On the other hand, TCS-EA showed significant decrease in their blood level in comparison with the group TCS treated group. Our results are similar to those of Tootian et al. [33] who found that increases of BUN, SC and UA metabolites following phenol administration are indicators of the biochemical damage to the kidney. In TCS-EA treated group the finding is consistent with Chao et al. [54] who found that dose dependent intake of EA reduced plasma BUN and elevated SC clearance. Al-kharusi et al. [18] mentioned that the antioxidant EA has a protective effect against oxidative stress. Ateşşahin et al. [12] postulated that alterations in glomerular function may be secondary to ROS which induced mesengial cell contraction, altering the filtration surface area and modifying the ultra filtration coefficient factors that decrease the glomerular filtration rate and elevation of BUN and SC.

Conclusions

In conclusion, TCS leads to alterations in the histological structure and functions of renal cortex of albino rats. EA supplementation protect from these changes and regulating the apoptotic process. So, ellagic can be used as a protective agent against TCS toxicity. Further investigations, as ROS detection in tissue by ROS assay and vitro experiments, are needed to explore the exact mechanisms and the pathogenesis of TCS toxicity.

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