NARINGIN PREVENT CISPLATIN-INDUCED NEPHROTOXICITY BY ABROGATION OF OXIDATIVE STRESS AND INFLAMMATION IN RATS

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Nephrotoxicity remains one of the most dangerous effect relevant to cisplatin use in chemotherapy. Rat injection with cisplatin in a single dose of 7 mg/kg intraperitoneally resulted in a significant increase in serum level of urea and creatinine. Also, cisplatin caused marked increase in renal content of malondialdehyde (MDA), while depletion in reduced glutathione (GSH). In addition, cisplatin administration notably increased kidney/body weight ratio, renal contents of nitric oxide (NO), tumor necrosis factor alpha (TNF-α) and cyclooxygenase-2 (COX-2) protein level as well as renal myeloperoxidase (MPO) activity. Histopathological examination confirmed the biochemical and molecular results which revealed several pathological alteration in the renal tissues following cisplatin. Oral pretreatment of rats with naringin (NAR) (80 mg/kg) for 14 days before and 7 days after cisplatin injection significantly reduced the pathological level of serum urea and creatinine and restored oxidative stress parameters. In the same manner, the inflammatory markers as well as kidney/body weight ratio show great improvement following the treatment. The histopathological examination confirms fit with the biochemical and molecular results. In conclusion, NAR showed a great protective effect against cisplatin-induced nephrotoxicity in rats via its antioxidant, anti-inflammatory roles.

INTRODUCTION

The kidneys are the most organ concerning with drug elimination. Approximately 90% of drug excreted mainly by the kidneys. Acute kidney injury (AKI) one of the most complicated factor of chemotherapy use against human malignancies hence, drugs induced renal toxicity remains the most problems that limit the beneficial roles of chemotherapeutic agents. Cisplatin (CP) is a key chemotherapeutic drug used in the treatment of many solid tumors and hematological malignancies. Unfortunately, CP cannot effectively differentiate healthy cells from malignant cells; as a result, it accumulates in healthy tissues, leading to severe clinical toxicities in different body organs, including kidneys as the kidney is the main route of excretion of CP. Tubular cell injury occurs in one third of CP treated patients and manifests as an increase in serum creatinine and urea concentration as well as imbalanced electrolytes, which limits its use as chemotherapeutic agents. Although, intense efforts over the past decades to find less toxic, but equally effective alternatives, CP continues to be widely prescribed.

It was postulated that CP-induced nephrotoxicity was mediated through the generation of large amount of reactive oxygen species (ROS), which induces lipid peroxidation, the oxidative damage in renal cells, increase renal capillary permeability and tubular atrophy. As, overproduction of ROS causes activation of oxidative stress cascade and depletion of endogenous antioxidants, that trigger an immune response and mediate...
inflammation. This inflammation is characterized by overproduction and activation of pro-inflammatory cytokines such as tumor necrosis factor-alpha (TNF-α), nitric oxide (NO) and inflammatory cell infiltration. Additionally, CP induces apoptosis and necrosis of renal tubular cells through activation of both intrinsic and extrinsic mitochondrial pathways as well as its direct tubular toxicity. Several strategies applied dosage optimization, the use of analogues or combined therapy, and many other trials have been done to attenuate CP side effects but no promising results have been achieved.

Thus, there is a need for identifying alternative, natural, and safer sources capable of protecting the cells from oxidative injury, as herbal extracts which, possess many properties such as antioxidant, anti-inflammatory, antiproliferative, and anticarcinogenicity.

Naringin (Nar) is a flavanone glycoside, isolated from the grape and citrus fruit species. The therapeutic activity of NAR has been reported to possess biological and pharmacological properties including anti-carcinogenic, lipid-lowering, anti-apoptotic, anti-atherogenic, metal chelating and antioxidant activities. Furthermore, another animal study reported that NAR has anti-inflammatory effects both in vitro and in vivo by modulating the expression of TNF-α, interleukin-6 (IL-6), interleukin-8 (IL-8) and inducible Nitric oxide synthase (iNOS). Also, it was reported that NAR has potent antioxidant properties via abrogation of oxidative stress and high free radical scavenging activity.

Therefore, the current study was done to investigate the possible nephroprotective effect of NAR against CP-induced nephrotoxicity in rats.

MATERIALS AND METHODS

Drugs and chemicals
Cisplatin was obtained from MYLAN United Pharmaceuticals Co., Egypt, and given i. p. in a single dose of 7 mg/kg. NAR was purchased from Sigma–Aldrich (St. Louis, MO). All other chemicals were obtained from local sources with highest analytical grade.

Animals
Male Swiss albino rats weighing 200–230 g were housed in the animal house of the Faculty of Medicine, Assiut University. The animals were kept one week to adapt with the environmental conditions. Rats were fed a standard diet and allowed freely accessed to water. Experiment was carried out according to the accepted guidelines for animal care.

Experimental design
Twenty four male adult Swiss albino rats were allocated into three groups (eight rats each); two rats from each group were used for histopathological examination as follows:

Group 1: Received saline orally and served as controls.
Group 2: Received CP in a single dose of 7 mg/kg i. p.
Group 3: Pretreated with NAR in a daily dose of 80 mg/kg p. o. for 14 consecutive days and NAR in a daily dose of 80 mg/kg, p.o. for 7 consecutive days after CP injection.

Serum and tissue preparation
At the end of the experiment, rats were anesthetized with ketamine (50 mg/kg) and xylazine (10 mg/kg) by i. p., injection, blood samples were withdrawn by a direct cardiac puncture. Sera were collected in non-heparinized tubes and separated by centrifugation for 20 min at 4000xg and stored at −20°C till the time of assay.

The kidneys were rapidly isolated, washed with ice-cold isotonic saline (0.9%) and divided into two portions the first portion was stored at 10% neutral buffered formalin solution and was subjected to histopathological examinations. The second portion was homogenized (20%) with (Cole-Parmer instrument company, USA) in cold phosphate buffered saline. Tissue homogenates were centrifuged at 3000 rpm for 15 min at 4°C. The supernatant was collected, divided into aliquots and stored at −80°C for evaluation of oxidative stress and inflammatory parameters.
Biochemical analysis
Assessment of kidney function testes
Serum urea and creatinine were estimated colorimetrically according to methods of Fawcett and Scott\textsuperscript{21}, Bartles \textit{et al.}\textsuperscript{22}, respectively, using aqueous primary standard urea solution (50 mg/dl), standard creatinine (2 mg/dl).

Evaluation of oxidative stress markers
Renal content of GSH was assayed according to the method described by Ellman\textsuperscript{23}, while Renal thiobarbituric acid reactive substances measured as MDA were assayed according to the method described by Uchiyama and Mihara\textsuperscript{24}, using 1,1,3,3-tetramethoxypropan as standard.

Estimation of inflammatory markers
NO was assayed according to the method described by montgomery and dymock\textsuperscript{25}. Renal content of COX-2 level was determined using ELISA kit according to the manufacturing instruction based on the principle described by Van Weemen and Schuurs\textsuperscript{26}. MPO content was assayed according to the method described by Bradley \textit{et al.}\textsuperscript{27}. Renal expression of TNF-α was determined by immunhistochemical analysis\textsuperscript{28}.

Histopathological examination of the kidney
Autopsy samples were taken from the kidneys of two rats in different groups and fixed at 10% neutral buffered formalin for 24 h. Washing was done with distilled water, and then dehydaration by serial dilutions of alcohol (methyl, ethyl, and absolute ethyl). Specimens were cleared in xylene and embedded in paraffin at 56°C in a hot air oven for 24 h. Paraffin bees wax tissue blocks were prepared for sectioning at 4 μm thickness by a sledge microtome. The obtained tissue sections were collected on glass slides, deparaffinized, stained with hematoxylin and eosin, and then an examination was done through the light electric microscope. The blocks were made using paraffin bees wax by sledge microtome, followed by deparaffinization and staining by hematoxylin and eosin\textsuperscript{28}.

Immunohistochemical analysis
Tissue samples were flushed and fixed in 10% neutral buffered formalin for 72 hrs. Samples were trimmed and processed by dehydration in alcohols, clearing in Xylene, synthetic wax infiltration and blocking out into Paraplast tissue embedding media. 3-5μ sections were cut by rotatory microtome. The sections were stained with
1- Harris Hematoxylin and Eosin as a general staining method as Outlined by Bancroft and Stevens (2010).
2- Tumor necrosis factor-alpha immunohistochemical Staining by using rabbit polyclonal antibody RB-9034-R7 From Thermo scientific Co. Expression Area percentage of TNF-α were obtained from (ex. 10 random fields) according to by using a full HD microscopic camera attached to the Leica application suite for immunoexpression analysis (Leica Biosystems- Germany\textsuperscript{28}).

Statistical analysis of data
Data were presented as means ± standard error of mean (SEM). Statistical analysis was performed using Graph Pad Prism version 5 (Graph pad, San Diego, CA). A comparison between different groups was carried out using one-way analysis of variance (ANOVA), followed by Tukey multiple comparison tests. The difference is considered significant when $P \leq 0.05$.

RESULTS AND DISCUSSION

Results
Effect of NAR on kidney functions test
Data summarized in table 1 show that injection of CP resulted in a significant increase in serum urea (280%) and creatinine (323%) compared to control group. In contrast, oral administration of NAR significantly reduced the elevated levels of urea and creatinine in serum by 52% and 61%, respectively in comparison with a murine model. At the end of the experiment the kidney-body weight ratio were significantly increased in rats challenged with CP respected to control rats. Treatment with NAR significantly reduced kidney-body weight ratio regarded to CP rats.
Effect of NAR on oxidative stress biomarkers

Administration of CP was associated with marked elevation of renal content of MDA (116%), while significant reduction in renal contents of GSH (66%) compared to control group. On the other hand, treatment of rats with NAR resulted in a notable correction of oxidative stress markers as represented in table 2.

NAR effects on inflammatory biomarkers

Data summarized in table 3 and figure 1 show that CP injection was showed a significant increase in kidney contents of TNF-α, COX-2 (193%) and NO (177%) as well as MPO (244%) activity in related to control animals. NAR treatment remarkably ameliorated renal contents of TNF-α, COX-2 and NO as well as MPO activity in related to a murine challenged with CP as illustrated in table 3.

Effect of treatment with NAR on histopathological findings of kidney tissue in CP-treated rats

Histopathological findings of kidney tissues are illustrated in table 4 and figure 2:
(A) Control saline: shows that normal histological structure of renal parenchyma,
(B) CP-treated group: shows that marked vacuolation (v) and necrosis (n) of renal tubular epithelium as well as periglomerular fibroblasts proliferation (P),
(C) NAR + CP-treated group: shows that slight congestion (c) of glomerular tuft.

Table 1: Effect of treatment with NAR on serum urea, creatinine and kidney–body weight ratio in CP-treated rats.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Urea (mg/dl)</th>
<th>Creatinine (mg/dl)</th>
<th>Kidney–Body Weight ratio (1000×)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control saline</td>
<td>30.50±2.95</td>
<td>0.92 ± 0.077</td>
<td>5.6 ± 0.29</td>
</tr>
<tr>
<td>CP</td>
<td>114 ± 5.71a</td>
<td>3.9 ± 0.14a</td>
<td>13.85±0.0a</td>
</tr>
<tr>
<td>CP + NAR</td>
<td>72.75±5.93a,b</td>
<td>1.81 ± 0.102a,b</td>
<td>9.45±0.3a,b</td>
</tr>
</tbody>
</table>

Data are expressed as mean ± SEM of eight rats per group.

Table 2: Effect of treatment with NAR on kidney contents of MDA and GSH in CP-treated rats.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>MDA (nmol/g Tissue)</th>
<th>GSH (µmol/g tissue)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control Saline</td>
<td>20.75±1.54</td>
<td>8.97±0.319</td>
</tr>
<tr>
<td>CP</td>
<td>44.5±2.84a</td>
<td>3.025±0.213a</td>
</tr>
<tr>
<td>CP + NAR</td>
<td>28.75±2.016b</td>
<td>7.47±0.311a,b</td>
</tr>
</tbody>
</table>

Data are expressed as mean ± SEM of six rats per group.

Table 3: Effect of treatment with NAR on inflammatory biomarkers in CP-treated rats.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>COX-2 (pg/ml)</th>
<th>MPO (U/g tissue)</th>
<th>NO (µmol/g tissue)</th>
<th>TNF-α Area%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control Saline</td>
<td>45±2.95</td>
<td>10.98±0.833</td>
<td>2.37±0.228</td>
<td>0.750±0.064</td>
</tr>
<tr>
<td>CP</td>
<td>132.4±3.77a</td>
<td>37.8±1.93a</td>
<td>6.57±0.268a</td>
<td>41.5±1.93a</td>
</tr>
<tr>
<td>CP + NAR</td>
<td>84.75±2.64a,b</td>
<td>22±3a,b</td>
<td>3.87±0.149b</td>
<td>22±2a,b</td>
</tr>
</tbody>
</table>

Data are expressed as mean ± SEM of six rats per group.

Table 1: Effect of treatment with NAR on serum urea, creatinine and kidney–body weight ratio in CP-treated rats.

Table 2: Effect of treatment with NAR on kidney contents of MDA and GSH in CP-treated rats.

Table 3: Effect of treatment with NAR on inflammatory biomarkers in CP-treated rats.
Fig. 1: Immunohistochemical assay of TNF-α in CP-treated rats:
A: Normal control group
B: CP- treated group
C: NAR+CP treated group
Table 4: Effect of treatment with NAR on histopathological findings of kidney tissues of cisplatin-treated rats.

<table>
<thead>
<tr>
<th>Groups</th>
<th>Control Saline</th>
<th>CP</th>
<th>NAR+CP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vacuolation of renal tubular epithelium and glomerular tufts</td>
<td>–</td>
<td>+++</td>
<td>+</td>
</tr>
<tr>
<td>Congestion of glomerular tufts</td>
<td>–</td>
<td>++</td>
<td>+</td>
</tr>
<tr>
<td>Cystic dilatation of renal tubules</td>
<td>–</td>
<td>+++</td>
<td>–</td>
</tr>
<tr>
<td>Necrosis of renal tubular epithelium</td>
<td>–</td>
<td>++</td>
<td>+</td>
</tr>
<tr>
<td>Cellular cast in the lumen of renal tubules</td>
<td>–</td>
<td>++</td>
<td>–</td>
</tr>
<tr>
<td>Chronic interstitial nephritis</td>
<td>–</td>
<td>+++</td>
<td>–</td>
</tr>
</tbody>
</table>

(-) normal  (+) milde  (+++) moderate  (++++) severe

Fig. 2: Effect of Treatment with NAR on Histopathological Findings of Kidney Tissues of CP-Treated Rats
(A) control saline: shows that normal histological structure of renal parenchyma,
(B) CP-treated group: shows that marked vacuolation (v) and necrosis (n) of renal tubular epithelium as well as periglomerular fibroblast proliferation,
(C) NAR + CP-treated group: shows that slight congestion (c) of glomerular tuft.
Discussion

Nephrotoxicity remains the most complicating factor of CP used in chemotherapy that might be attributed to CP-induced oxidative injury, free radical generation and its direct accumulative effect in renal tissues. Many researches have established the ability of antioxidants of natural origin to protect against pathological conditions induced via chemotherapeutic agents. Many researches had been demonstrated that NAR has antioxidant, anti-inflammatory and anti-apoptotic activities.

Therefore, the present study was designed to evaluate the possible protective effects of NAR against CP-induced nephrotoxicity in rats. Our results revealed that CP-induced renal injury manifested by a significant elevation in serum creatinine, urea, and kidney-body weight ratio accompanied with a experiment. These abnormalities could be attributed to reduction in glomerular filtration rate, increase in glomerular volume and cellular degenerative changes. As well as, gastrointestinal toxicity evidenced by severe body weight reduction.

Moreover, our data showed that a single dose of CP has the ability to produce severe changes in oxidative stress parameters confirmed by a significant increase in renal contents of MDA and NO with a significant reduction in GSH levels in the tissue. Additionally, increased renal NO production may induce cellular injury as was confirmed, NO reacts with superoxide anion to generate peroxynitrite radical, a potent prooxidant and cytotoxic intermediate that causes protein nitration and tissue injury. Also, excess NO depletes intracellular GSH thus increasing the susceptibility to oxidative stress. These results are consistent with de Oliveira Mora et al., 2003 and Shimeda et al., 2005 who found that CP generates a large amount of ROS includes superoxide anion, hydrogen peroxide, and hydroxyl radicals which, induces lipid peroxidation causing an oxidative damage in renal cells and tubular atrophy as well as, depletion of endogenous antioxidants.

Furthermore, our study revealed that CP cause inflammation of renal tubules manifested by of a significant elevation in renal contents of inflammatory biomarker, COX-2, TNF-α and MPO activity which is a marker of neutrophil infiltration via induction of oxidative stress cascade that triggers an immune response and mediates inflammation in renal tissue.

On the other hand, the present study revealed that oral treatment of rats with NAR ameliorates CP-induced alterations in serum levels of creatinine, urea, body weight and kidney-body weight ratio. In addition to, it is significantly mitigates the lipid peroxidation measured by a significant decrease in renal content of MDA, and NO as well as, it cause a significant improvement in the GSH levels. These results could be attributed to the potential antioxidant effect of NAR.

Also, our data showed that NAR treatment obviously mitigated CP-induced renal inflammation via a significant decrease in the renal contents of COX-2, TNF-α and MPO. NAR was found to reduce the level of MPO in a dose dependent manner. The inhibition of neutrophil infiltration in renal tissue by naringin is also evident in the histopathological observations of renal tissue. The reversal of neutrophil infiltration by NAR established its anti-inflammatory activity. This finding is consistent with Chtourou et al., who demonstrated that NAR has antioxidant, anti-inflammatory and anti-apoptotic activities.

Our histopathological findings demonstrated that administration of CP produced various degenerative changes in kidney cells which confirmed the biochemical evidence of the oxidative stress. In contrast, the treatment with NAR obviously mitigated the histopathological changes induced by CP.

Conclusion

The present study revealed the nephroprotective effect of NAR against CP-induced renal injury by abrogation of oxidative stress, inflammation which might be attributed to their antioxidant, anti-inflammatory effect.

REFERENCES

نشرة العلوم الصيدلانية
جامعة أسيوط

النارينجين يمنع السمية الكلوية المستحثة بالسيسبيلاتين بواسطة الحد من الأكسدة والإلتهابات في فنران التجارب

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الاسم الكلوي يظل أحد أهم التأثيرات الضارة لاستخدام عقار السيسبيلاتين في علاج الأورام السرطانية. في هذه الدراسة تم حفظ فنران التجارب بجرعة مفردة من عقار السيسبيلاتين (7 مجم/كجم) داخل البطن والتي نجت عنها ارتفاع محول في مستوي اليوريا والكرياتينين بالدم. أيضاً جرعة السيسبيلاتين أدت إلى ارتفاع محول في محتوى الألمان داي ألفيند مع انخفاض في مستوى الجلوتاثيون. بالإضافة إلى أن السيسبيلاتين أدى إلى زيادة محول في نسبة وزن الكلي / وزن الجسم، محتوى أكسيد النيتريزك ومعامل النخر النوري، وإينزيم السيكلوكسيجينز-2، وإنزيم الميلوبيروكسيديز.

التحليل الظاهري للأنسجة الكلوية يؤكد نتائج التحليل البيوكيميائي والجيني والذي أظهر تغيير في الأنسبة الكلوية على حاف جرعة واحدة من السيسبيلاتين.

في المقابل إعطاء عقار النارينجين بجرعة (80 مجم/كجم) لمدة أربعة عشر يومًا عن طريق الفم قبل إعطاء السيسبيلاتين لمدة سبعة أيام بعد ظهور تغيرات محول في الآثار الضارة الناتجة عن السيسبيلاتين، والتي ظهرت في انخفاض محول في نسبة اليوريا والكرياتينين بالدم، وانخفاض محتوى الألمان داي ألفيند، وأكسيد النيتريزك ونسبة وزن الكلي / وزن الجسم، مع زيادة محول في محتوى الجلوتاثيون كمضاد للأكسدة.

أيضاً إعطاء عقار النارينجين قد أدى إلى انخفاض محول في محتوى معاملات الإلتهابات، والتي تتمثل في مقياس النخر النوري، وإينزيم السيكلوكسيجينز-2، وإنزيم الميلوبيروكسيديز.

وهذه النتائج تؤكد أن التحليل الظاهري للأنسجة الكلوية والتي أظهرت التأثير الفعال للنارينجين في الحماية ضد السمية الكلوية المستحثة بالسيسبيلاتين.