Ameliorative Effects of Ascorbic Acid and *Allium sativum* (Garlic) Ethanol Extract on Renal Parenchyma of Gentamicin-induced Nephropathic Rats

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Authors’ contributions

This study was carried out in collaboration between both authors. Author DRO designed the study, performed the statistical analysis and wrote the study protocol. Author JMO wrote the first draft of manuscript. Authors DRO and JMO managed the analyses of the study and the literature searches. Both authors read and approved the final manuscript.

**Article Information**

DOI: 10.9734/JOCAMR/2020/v9i430146

Editor(s):
(1) Dr. Aditi Singh, Amity University, India.

Reviewers:
(1) Wilson Obidah, Modibbo Adama University of Technology, Yola, Nigeria.
(2) Ochieng O. Anthony, Sumait University, Tanzania.
(3) Eshak Mourad EI-Hadidy, Cairo University, Egypt.

Complete Peer review History: [http://www.sdiarticle4.com/review-history/56640](http://www.sdiarticle4.com/review-history/56640)

**Original Research Article**

**ABSTRACT**

To assess ameliorative effects of Ascorbic acid (AA) and *Allium sativum* ethanol extract (ASEE) on renal parenchyma of gentamicin-induced nephropathic rats. Thirty Wistar rats (weighing between 180-205 g) were randomly divided into five groups (A-E). These include Group A administered with 0.9% Normal Saline (0.5 ml/kg body weight (b.w.)), Group B administered with gentamicin (GM, 200 mg/kg b.w.) intraperitoneally (i.p.), Group C administered with GM (200 mg/kg b.w.) i.p. and AA (200 mg/kg b.w.) orally, Group D administered with GM (200 mg/kg b.w.) i.p. and ASEE (200 mg/kg b.w.) orally and Group E administered with GM (200 mg/kg b.w.) i.p. and AA (200 mg/kg b.w.) orally and ASEE (200 mg/kg b.w.) orally. All administrations were done once daily for a period of ten (10) days. The body weight of study
animals was recorded at the beginning and end of study period. After the study period, renal tissue of study animals was harvested, weighed, processed, stained using H & E technique. Stained sections were examined under microscope for histopathological changes within the renal parenchyma and were scored using image-J software.

The results of this study showed that exposure to GM results into significant ($P < 0.05$) reduction in body and renal tissue weight. However, therapeutic exposure to AA and ASEE either as individual or combined treatment regimen culminated into relatively null body and renal tissue weight loss among treatment groups C-E. In addition, exposure to GM precipitates prominent histopathological changes within renal parenchyma of study animals. As observed with body and renal tissue weight changes, treatment with AA and ASEE also comparatively ameliorate GM-induced nephropathy within renal parenchyma of study animals in treatment groups.

The findings of this study therefore showed that AA and ASEE exhibit ameliorative effect on the renal parenchyma of gentamicin-induced nephropathic rats either as distinct or combined treatment regimen.

**Keywords:** Ascorbic acid; Allium sativum; gentamicin-induced nephropathy; renal parenchyma; wistar rats.

### 1. INTRODUCTION

The renal tissue is among most versatile body tissues that perform or significantly contribute to several vital body functions which include excretion of metabolic waste products and foreign chemicals, body fluid and electrolyte homeostasis, regulation of acid-base balance, regulation of arterial pressure as well as hemopoietic and endocrine functions [1-3]. These functions can predispose the renal tissue to exposure and possible accumulation of potential nephrotoxins that may in turn trigger nephropathic changes within the renal parenchyma. Histopathological changes that characterized nephropathy, which may result into renal functional impairment or renal failure, include epithelial necrosis, glomerular congestion, interstitial oedema and inflammation, tubular dilatation and so on [3-5].

Nephropathy has been associated with application of some chemotherapeutic agents such as aminoglycosides antibiotics like gentamicin (GM) [6]. In addition, experimental models of gentamicin-induced nephropathy are commonly used to probe ameliorative or protective potential of chemical or phytochemical compounds against nephropathic effects of the nephrotoxin. As a potential nephrotoxin, GM can induces nephropathy by stimulating production of reactive oxygen species (ROS) leading to oxidative damage of renal parenchyma [7,8]. In essence, chemical or phytochemical compounds with antioxidant activity may exhibit ameliorative or protective effect against nephropathic effects of nephrotoxins such as GM.

Ascorbic acid (AA) has been described as a natural antioxidant which exhibits its potent antioxidant activity through down-regulation of ROS production and suppression of free radicals reactivity [9-12]. Similarly, Allium sativum is a common food spice that possesses antioxidant property and has been extensively applied for various therapeutic purposes such as anti-diabetic, anticancer and hepatoprotective [13-15].

In this study, ameliorative effects of AA and A. sativum ethanolic extract (ASEE) on renal parenchyma of gentamicin-induced nephropathic Wistar rats were assessed either as distinct or combined therapeutic agent.

### 2. MATERIALS AND METHODS

#### 2.1 Chemical Reagents and Plant Material

Chemical reagents used in this study were procured from Bristol Scientific Co. Ltd. (Lagos, Nigeria) while fresh bulbs of A. sativum were obtained from commercial sources in Benin City, Nigeria.

#### 2.2 Extract Preparation

Fresh bulbs of A. sativum were air-dried at room temperature (22-24°C) for four weeks and pulverized into powder form. 200 g of powdered A. sativum was dissolved in absolute ethanol for 72 hours with intermittent agitation. The preparation was then filtered and evaporated to produce 22 g yield (percentage yield of 11%) used as extract for treatment.
2.3 Study Animals

Thirty Wistar rats, weighing between 180–205 g, were employed for this study. The animals were housed in plastic cages at room temperature and exposed to 12 hour light/dark cycle. They were fed with standard animal feed and allowed access to drinking water ad libitum. This study was carried out in the Central Animal House, Igbinedion University, Okada, Edo State, Nigeria.

2.4 Design of Study

After the 14-day period of acclimatization, study animals were weighed and randomly divided into five groups (A–E) comprising of 6 animals each (n=6). Group A represented normal control; group B represented experimental control while groups C–E represented treatment groups. Group A was administered with 0.9% Normal Saline (NS; 0.5 ml/kg body weight (b.w.)), Group B was administered with GM (200 mg/kg b.w.), Group C was administered with GM (200 mg/kg b.w.) and AA (200 mg/kg b.w.), Group D was administered with GM (200 mg/kg b.w.) and ASEE (200 mg/kg b.w.) while Group E was administered with GM (200 mg/kg b.w.) and AA (200 mg/kg b.w.) and ASEE (200 mg/kg b.w.). Administrations of NS and GM were done intraperitoneally while AA and ASEE done orally; once daily for ten days. Dosages employed in this study were considered safe and without toxic effects [13,16-17].

2.5 Collection and Processing of Study Tissue

After the treatment period, each study animal was weighed, sacrificed via cervical dislocation and their right and left renal tissues harvested, weighed and processed. The processing of renal tissues to produce tissue blocks for histopathological study involved the following protocol: fixation using 10% Neutral-Buffered Formalin, dehydration using ascending grades of Alcohol (from 70% to absolute Alcohol), clearing in xylene and embedding using paraffin wax. Alcohol to 70% alcohol), stained in Haematoxylin, rinsed in water, differentiated in 1% Acid Alcohol, blued in Scott’s tap water, rinsed in water, stained in Eosin, rinsed in water, dehydrated with ascending grades of Alcohol, cleared in xylene and mounted with Distrene polystyrene xylene DPX [18].

2.6 Sectioning of Tissue Blocks and Staining of Tissue Sections

Renal tissue blocks were sectioned at thickness of 5 μ, mounted on microscope slides and stained using Haematoxylin and Eosin (H & E) staining technique. The H & E staining protocol employed in this study is given as follows: tissue sections were dewaxed in xylene, hydrated with descending grades of Alcohol (from absolute alcohol to 70% alcohol), stained in Haematoxylin, rinsed in water, differentiated in 1% Acid Alcohol, blued in Scott’s tap water, rinsed in water, stained in Eosin, rinsed in water, dehydrated with ascending grades of Alcohol, cleared in xylene and mounted with Distrene polystyrene xylene DPX [18].

2.7 Histopathological Study

Stained renal tissue sections were viewed under microscope to assess the histomorphology of renal parenchyma of all study animals. Photomicrographs were generated for stained sections and observable renal histopathological features such as interstitial inflammation, epithelial necrosis, glomerular congestion and tubular dilatation within the renal parenchyma of study animals were evaluated using image-J software (NIH, Bethesda, MA, USA).

2.8 Statistical Analysis

Data obtained during this study was analyzed using IBM-SPSS version 20 (IBM Corp, NY, USA) and results presented as Mean ± standard error of mean (SEM). Comparison of statistical results was done by t-test and multiple comparison by one way analysis of variance (ANOVA) with the probability level of p <0.05 regarded as statistically significant.

3. RESULTS

3.1 Evaluation of Body Weight of Study Animals

The mean values of total body weight of study animals in Groups A – E recorded at the beginning and end of the study period were given in Fig. 1. In comparison with normal control group A, mean values of body weight of group B animals showed significant (p < 0.05) reduction while those of treatment groups C – E showed no significant weight loss.

3.2 Evaluation of Renal Tissue Weight of Study Animals

The mean values of right and left kidney weight of study animals in Groups A – E recorded at the end of study period were given in Fig. 2. Similar to body weight changes, mean weight values of renal tissue of group B animals showed significant (p < 0.05) reduction in comparison with normal control group A. However, treatment groups C – E showed no significant renal tissue weight loss.
3.3 Histopathological Results

Histomorphology of renal parenchyma of study animals in Groups A – E were revealed in Fig. 3. Evaluation of renal histopathological features such as inflammation, necrosis and glomerular congestion showed significant ($p < 0.05$) prominence within renal parenchyma of group B relative to normal control group A. However, renal parenchyma of treatment groups C – E showed no significant prominence of renal histopathological features (Fig. 4).
Fig. 3. Representative photomicrograph of renal tissue showing histopathological changes within renal parenchyma of study animals in Groups A – E (H & E X100)

Group A = Normal control (NS), Group B = Experimental control (GM), Group C = GM+AA, Group D = GM+ASEE, Group E = GM+AA+ASEE

Fig. 4. Evaluation of histopathological changes within renal parenchyma of study animals in Groups A – E

(*,+ indicate significant difference from normal and experimental controls respectively at P < 0.05).

Group A = Normal control (NS), Group B = Experimental control (GM), Group C = GM+AA, Group D = GM+ASEE, Group E = GM+AA+ASEE
4. DISCUSSION AND CONCLUSION

4.1 Discussion

The therapeutic application of aminoglycoside antibiotics such as GM has been reported to cause nephrotoxicity or nephropathy in about 30% of cases [19,20]. In principle, the activation of oxidative stress has been described as primary mediator of nephropathic effects of GM and potent antioxidants or ROS scavengers can be of therapeutic value in preventing or ameliorating deleterious effect of GM exposure [21,22]. GM-induced nephropathy is often characterized by tubular damage, necrosis, fibrosis, oedema, inflammation and glomerular congestion which can in turn lead to renal dysfunction [23,24].

According to results of this study, the exposure to GM results into significant ($P < 0.05$) reduction in body and renal tissue weight (Figs. 1 and 2). This can be linked with deleterious effects of GM exposure on the study animal body in general and their renal tissue in particular. However, it can be observed that therapeutic exposure to AA and ASEE as individual or combined treatment regimen culminated into relatively null body and renal tissue weight loss among treatment groups C-E. In addition, exposure to GM precipitates prominent histopathological changes within renal parenchyma of study animals (Figs. 3 and 4). As observed with body and renal tissue weight changes, treatment with AA and ASEE also comparatively ameliorate GM-induced nephropathy within the renal parenchyma of study animals in treatment groups C-E.

As a naturally occurring vitamin, AA can be readily sourced from citrus fruits, strawberries and vegetables and as antioxidant; it acts to suppress ROS production or activation in tissues [25]. Each molecule of AA has in its structure two hydrogen atoms that bear two high-energy electrons which can be readily donated to stabilize ROS [26,27]. Based on the results of this study and in comparison with previous studies, AA potently acts to suppress ROS reactivity thereby functioning as free radical scavenger that ameliorates ROS-induced oxidative damage caused by GM exposure in renal tissue [28-30].

Furthermore, A. sativum is a member of Liliaceae family commonly used as food spice. It is one of the most popular herbs with numerous therapeutic properties and widely used in modern folkloric medicine. Its characteristic strong odour is due to its sulphur-containing compounds (especially S-allylcysteine sulphoxide or allicin) which also confer the potent therapeutic properties [15,31-33]. A. sativum has been reported to contain phytochemicals with potent antioxidant property which constitute the primary basis of its therapeutic activity against oxidant-mediated tissue pathologies like GM-induced nephropathy [33-34]. In comparison with previous studies, ameliorative effect of ASEE against GM exposure observed in this study can be related with antioxidant activity of its phytochemical constituents which suppresses oxidative renal tissue damage that follows GM exposure.

4.2 Conclusion

The findings of this study showed that AA and ASEE both exhibit ameliorative effect on the renal parenchyma of gentamicin-induced nephropathic rats either as distinct or combined treatment regimen. The ameliorative effect of each therapeutic agent against the nephropathic effect of the toxicant can be associated with their distinct antioxidant property.

DISCLAIMER

The research was wholly funded by personal efforts of the authors.

CONSENT

It is not applicable.

ETHICAL APPROVAL

This study was approved by the Research and Ethics Committee of Igbinedion University, Okada, Edo State, Nigeria. The study protocols were in compliance with International guidelines for handling and care for experimental animals.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

REFERENCES


