Nephroprotective effect of turmeric on oxidative stress, renal histopathology and toxicity induced by gentamicin

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ABSTRACT

Background: Commonly used aminoglycosides have frequent side effect of nephrotoxicity, still are preferred by clinicians because of efficacy against gram negative bacteria, resistant bacteria, nosocomial infections and cost effectiveness. Gentamicin produces oxidative stress; substances ameliorating stress are used to reduce toxicity. Turmeric has multiple medicinal properties including potent antioxidant activity, hence study was undertaken.

Methods: Eight groups containing six animals in each group, treated for 15 days. First group treated with normal saline. Second, fourth and sixth group treated with only gentamicin- sacrificed at 16, 22, 29th day. Third, fifth and seventh group treated with gentamicin and turmeric simultaneously and sacrificed on 16, 22, 29th day. Eighth group was pre-treated with turmeric for thirty days and concurrently treated with gentamicin and turmeric for 15 days and sacrificed on 16th day. Levels of blood urea, serum creatinine, superoxide dismutase and histopathological grades were assessed each time.

Results: Severe renal dysfunction (146±9.2, 2.03±0.26), highest renal injury grading (3.66±0.24) was observed in only gentamicin treated groups followed by spontaneous recovery after withdrawal of drug but with higher levels of oxidative stress (0.04±0.01). Gentamicin and Turmeric treated groups maintained renal function and had lower level of renal damage grades and oxidative stress. Turmeric pre-treated group was having lowest oxidative stress (0.12±0.03), histopathology grade (0.60±0.06) with normal renal functions.

Conclusions: Turmeric has potent antioxidant property which effectively protects kidney from damage induced because of gentamicin.

Keywords: Gentamicin, Nephrotoxicity, Oxidative stress, Turmeric

INTRODUCTION

Aminoglycoside antibiotics are widely used as first or second choice antibiotics in the treatment of complex gram negative infections like, urinary, intestinal and pulmonary due high stability of drug and cost effectiveness.

This group is known for their otootoxicity and nephrotoxicity leading to renal impairment in about 10 to 25 percent of the patients. Increased toxicity is observed if other risk factors are present; like hypothyroidism, hepatic dysfunction, reduced renal mass and concomitant use of other commonly used nephrotoxic drugs like NSAIDs, diuretics and cephalosporins. It has confirmed by both in vitro and in vivo studies that gentamicin produce oxidative stress which is key reason for its toxicity.¹,²

Toxicity of gentamicin is related to its preferential accumulation in proximal convoluted tubules, pathologically characterised by granulovacuolar debris, epithelial necrosis, desquamation and clinically manifested as azotemia.²,³

Various chemical compounds and reactions produce toxic oxygen species called as pro-oxidants or free radicals. Free radicals may be oxygen derived like superoxide (O₂⁻), Hydroxyl (HO⁻), hydrogen peroxide (H₂O₂) or nitrogen derived like Nitric oxide (NO), peroxinitrite (ONOO⁻).
Electromagnetic radiation, cigarette smoke, UV light are exogenous sources while mitochondrial electron transport chain, beta oxidation of fat, auto oxidation of catecholamines are endogenous sources. Excess production of pro-oxidants cause damage to cell. Normally in cell there is a balance between pro-oxidants and antioxidants. Excess reactive oxygen species (ROS) cause increased peroxidation of membrane phospholipids, oxidative modification of proteins and mitochondrial DNA damage leads cell aging and toxicity.4

Antioxidants are the substance when present in low quantity significantly delays or prevents oxidation of substrate like lipids, proteins, carbohydrates and DNA. They may be enzymatic, non-enzymatic or plant derived. Enzymatic antioxidants are superoxide dismutase (SOD) which catalyse breakdown of superoxide, catalase (CAT) involved in reduction of H2O2. Non-enzymatic antioxidants include Vitamin-E, Vitamin-C. Plant derived antioxidants include polyphenol, phenolic acid, carotenoids. Curcumin- a bioactive ingredient of turmeric is having multiple properties like antisepctic, anti-inflammatory, analgesic, anti-cancerous and most importantly antioxidant activity. It acts as scavenger for ROS, as well it protects haemoglobin from oxidation. Study supported that antioxidant property reduces the toxicity of lead by significant increase in superoxide dismutase, reduced glutathione and catalase activities. Hence the present study was undertaken to study the effect of turmeric on oxidative stress and nephrotoxicity induced by commonly used drug gentamicin.5,6

Aim of the study was to evaluate antioxidant property of turmeric and its protective effect on renal histopathology and nephrotoxicity induced by gentamicin in albino rats.

Objectives of the study were a) to study oxidative stress due to gentamicin. b) to study histopathological grade of renal injury. c) to study spontaneous recovery of kidney function in gentamicin toxicity. d) to study short term and long term effect of turmeric treatment on gentamicin toxicity.

METHODS

Study was conducted at SRTR medical college Ambajogai. Healthy albino rats of weights between 150-200 gms were used for the study.

Animals were kept on standard pellet diet after acclimatization to environment. Animals were divided in to eight groups having six animals in each group. Nephrotoxicity was induced by injecting gentamicin Intraperitoneally (IP) at dose of 80mg/kg/day for 15 days.

Turmeric was collected locally and powder was prepared. Suspension of 3% was prepared in gum acasia. A calculated dose of test drug 200mg/kg/day per orally (PO) and GM was given as per protocol of the groups.

Study design

Control group

Group I: Received normal saline 2ml IP for 15 days. Animals were sacrificed on 16th day for biochemical investigations and histopathological examinations.

Gentamicin toxicity

Group II: This group received only gentamicin 80mg/kg/d IP for 15 days. Animals were sacrificed on 16th day for biochemical investigations and histopathological examinations.

Concurrent GM and turmeric

Group III: This group received gentamicin 80mg/kg/d IP and turmeric 200mg /kg/d/PO for 15days. Animals were sacrificed on 16th day for biochemical investigations and histopathological examinations.

GM recovery

Group IV: This group received gentamicin 80mg/kg/d IP for 15. Animals were sacrificed on 22nd day for biochemical investigations and histopathological examinations.

Concurrent GM and turmeric recovery

Group V: This group received gentamicin 80mg/kg/d IP and turmeric 200mg /kg/d/PO for 15 days. Animals were sacrificed on 22nd day for biochemical investigations and histopathological examinations.

GM recovery

Group VI: This group received gentamicin 80mg/kg/d IP for 15. Animals were sacrificed on 29th day for biochemical investigations and histopathological examinations.

Concurrent GM and turmeric recovery

Group VII: This group received gentamicin 80mg/kg/d IP and turmeric 200mg /kg/d/PO for 30 days followed by turmeric in same dose and gentamicin 80mg/kg/day IP for 15 day. Animals were sacrificed on 46 days for assessment of renal function and histopathological examination. In the study, all animals were sacrificed under ether anaesthesia. Blood collected from heart by cardiac puncture, both kidneys were dissected out from retroperitoneum.

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**Assessment of renal toxicity**

Renal dysfunction was assessed by estimation of blood urea levels by DAM method, serum creatinine levels by picric acid method and Histopathological grade of renal damage assessed and highest grade of renal damage was considered for statistical comparisons.

Protective effect of Turmeric was assessed by estimation of superoxide dismutase (SOD).

**Histopathological grades of renal damage**

Both kidneys from all animals were removed, processed and embedded in paraffin wax. Sections stained with Haematoxylin and Eosin (H and E) were examined under light microscope and score was given as per damage as follows.

- Score-0: Normal
- Score-1: Areas of focal granulovacuolar debris with or without evidence of tubular epithelial cell desquamation of small foci. (<1% of total tubular population)
- Score-2: Tubular epithelial necrosis and desquamation easily seen but involving less than half of cortical tubules
- Score-3: More than half of proximal tubules showing desquamation and necrosis but involved tubules easily found
- Score-4: Complete or almost complete tubular necrosis.7

**Statistical analysis**

All values were expressed as mean with SD. For estimation of total variation present in a set of data, all groups were first subjected to ANOVA for measurement of significance between two groups, the student’s t test was used and p Value <0.05 was considered significant.

**RESULTS**

Gentamicin at a dose 80mg/kg for 15 day had induced severe renal dysfunction, depletion of superoxide dismutase and marked damage to renal parenchyma (3.66 ± 0.24). There was spontaneous recovery in renal function to normal in all animals within one week duration after withdrawal i.e. at 21st day and maintained thereafter i.e. at 28th day. These results suggest that nephrotoxicity due to gentamicin is transient and reversible after drugs has been discontinued. Oxidative stress was decreased after withdrawal of gentamicin which hallmark by increased levels of SOD at 21st and 28th day compared to 15th day (Table 1, Table 2).

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Mean with SD</th>
<th>Gentamicin treated group-2 (after 15 days)</th>
<th>Gentamicin treated group-4 (after 21 days)</th>
<th>Gentamicin treated group-6 (after 28 days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood urea</td>
<td></td>
<td>20.68±1.35</td>
<td>24.17±2.63</td>
<td>23.92±2.23</td>
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<tr>
<td>Serum creatinine</td>
<td></td>
<td>0.99±0.06</td>
<td>0.99±0.11</td>
<td>0.99±0.13</td>
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<td>Superoxide dismutase</td>
<td></td>
<td>0.31±0.01</td>
<td>0.05±0.01</td>
<td>0.05±0.01</td>
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<tr>
<td>Renal histopathological grade</td>
<td></td>
<td>3.66±0.24</td>
<td>1.45±0.08</td>
<td>1.16±0.08</td>
</tr>
</tbody>
</table>

In the groups treated with turmeric and gentamicin simultaneously, all animals maintained normal renal functions at 15th, 21st, and 28th day, though there was renal parenchymal damage (1.98±0.07), grades of renal parenchymal damage were less than that of only gentamicin treated groups to extent that these groups maintained normal renal functions and there was higher serum concentration of SOD at all the phases of assessment i.e. at the 16th, 22nd, and 29th days of assessment (Table 3).

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Gentamicin+turmeric treated group-3 (after 15 days)</th>
<th>Gentamicin+turmeric treated group-5 (after 21 days)</th>
<th>Gentamicin+turmeric treated group-7 (after 28 days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood urea</td>
<td>27.68±4.33</td>
<td>23.94±1.35</td>
<td>26.56±2.16</td>
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<tr>
<td>Serum creatinine</td>
<td></td>
<td>0.86±0.05</td>
<td>0.44±0.09</td>
</tr>
<tr>
<td>Superoxide dismutase</td>
<td></td>
<td>0.09±0.01</td>
<td>0.09±0.01</td>
</tr>
<tr>
<td>Renal histopathological grade</td>
<td></td>
<td>1.98±0.07</td>
<td>0.86±0.16</td>
</tr>
</tbody>
</table>
The group which received long term treatment with turmeric for thirty days prior to administration of gentamicin and turmeric for 15 days, all animals maintained normal kidney functions and grade of renal parenchymal damage of lowest in this group (0.60±0.06). When compared with short term treated group of turmeric it showed that there were highest levels of super oxide dismutase. (0.12±0.03 V/S 0.11±0.06) (Table 4). All the parameters except blood urea level were at statistical significant difference in gentamicin and turmeric treated group (Table 5).

### DISCUSSION

The present study was conducted in the albino rat to explore the nephroprotective effect of turmeric on the biochemical changes in the renal function, oxidative stress and extent of histopathological changes in the renal parenchyma after administration of gentamicin.

Serum level of super oxide dismutase (SOD) was used to determine increase in the defence mechanism as oxidative stress induced by gentamicin markedly reduces levels of SOD. It was observed in the present study that gentamicin at a dose 80mg/kg for 15 days had induced severe renal dysfunction. Functional derangements are multifactorial and mainly due to:

- Preferential accumulation to proximal convoluted tubules
- Induction of inflammatory response
- Production of ROS
- Decreased renal blood flow
- Derangements of antioxidant due to its ability to alter mitochondrial respiration.

Gentamicin is transferred through megalin ligands by endocytosis which then transferred to lysosome, and by ultimately binding to acid phospholipids ending into production of phospholipids metabolites.8

It is observed that toxicity of the gentamicin is dose dependent and increasing dose produces more severe dysfunctions, concentration of gentamicin can as high as 30 folds in proximal convoluted tubules than plasma concentration and continuous infusion is more toxic than intermittent dosing, hence in clinical practices lower dose or frequency of the dose administration is reduced to safeguard the kidney functions. This is due to accumulation of drugs in the proximal convoluted tubules and decreased clearance. Co-administration of the other drugs that compete with gentamicin through inhibition of megalin ligands and small proteins or administration of antioxidant reduces its toxicity.9 Gentamicin induces inflammatory response, activation of cells and cytokine production due to production of ROS; drugs that scavenger these damaging superoxide ions will ultimately protect kidney from adverse effects.1

In gentamicin mediated nephrotoxicity; there is significant reduction of renal blood flow and insulin clearance. It was observed that superoxide dismutase has significant increase in both parameters suggesting its protective role.10

In our study, there was evident depletion of superoxide dismutase which was more severely in the animals that were not treated with turmeric and relative higher levels of SOD in pre-treated and simultaneously treated with turmeric; this result highlights the antioxidant property of turmeric. Similar potent antioxidant effects of turmeric comparable to Vitamin C, anti-inflammatory property and other protective actions like antimicrobial as well inhibition of platelet aggregation is explored by different researchers. It is also confirmed that turmeric is non-toxic even at higher doses and protective in condition like atherosclerosis, it acts as hepatoprotective and anti-carcinogenic. Superoxide dismutase (SOD), Catalase (CAT) and Glutathione reductase are first line enzymatic defence whose levels are reduced in oxidative stress. In present literature, there is consensus about role of oxidative stress due to reactive oxygen and nitrogen species in gentamicin toxicity.11

### Table 5: Comparisons of means and SD of different groups.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Group-1 V/s group-2</th>
<th>Group-2 V/s group-3</th>
<th>Group-1 V/s group-3</th>
<th>Group-4 V/s group-5</th>
<th>Group-6 V/s group-7</th>
<th>Group-1 V/s group-8</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood urea</td>
<td>0.0001</td>
<td>0.0001</td>
<td>0.0032</td>
<td>0.85</td>
<td>0.0639</td>
<td>0.69</td>
</tr>
<tr>
<td>Serum creatinine</td>
<td>0.0001</td>
<td>0.0001</td>
<td>0.0097</td>
<td>0.02</td>
<td>0.0001</td>
<td>0.03</td>
</tr>
<tr>
<td>Superoxide dismutase</td>
<td>0.0001</td>
<td>0.0001</td>
<td>0.0094</td>
<td>0.0001</td>
<td>0.0001</td>
<td>0.0001</td>
</tr>
<tr>
<td>Renal histopathological grade</td>
<td>-</td>
<td>0.0001</td>
<td>-</td>
<td>0.0001</td>
<td>0.0021</td>
<td>-</td>
</tr>
</tbody>
</table>

### Table 4: Group 8 pre-treatment with turmeric for 30 days followed by gentamicin and turmeric treatment for 15 days.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Mean with SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood urea</td>
<td>21.15±2.50</td>
</tr>
<tr>
<td>Serum creatinine</td>
<td>0.85±0.13</td>
</tr>
<tr>
<td>Superoxide dismutase</td>
<td>0.12±0.03</td>
</tr>
<tr>
<td>Renal histopathological grade</td>
<td>0.60±0.06</td>
</tr>
</tbody>
</table>
In present study marked damage to renal parenchyma (3.66±0.24) was observed in animals only treated with gentamicin. We observed that there was spontaneous recovery in renal function parameters to normal and lower grade of renal parenchymal injuries in all animals within one week duration after withdrawal i.e. at 21st day (1.45±0.08) and maintained thereafter i.e. at 28th day (1.16±0.08). These results suggest that nephrotoxicity due to gentamicin is reversible after drugs has been discontinued. This extensive damage is due to preferential accumulation of gentamicin in the proximal convoluted tubules which produces variable changes in epithelial cells and graded accordingly. Intracellular changes like hydropic changes with cytoplasmic vacuolation are also seen in some area, similar results i.e. damage and regeneration of epithelium was observed on histopathological examination in kidney after gentamicin administration. Acute renal failure, oxidative stress its scavenging was reported in other studies.1,7,12-24

In our study, oxidative stress was gradually reduced after withdrawal of drug with decreasing grades of parenchymal damage when tested at set duration of interval which was predominant in turmeric treated group. Group which was pre-treated with turmeric for a month shown a lowest oxidative stress, renal tubular damage and maintained kidney function. Statistically significant differences (p <0.01) were observed in renal functions but are of clinical irrelevance as they are within normal range.

CONCLUSION

Gentamicin produces severe but reversible nephrotoxicity at selected doses in albino rats. Underlying oxidative stress evident by decreased levels of SOD whereas turmeric was effective in reducing the oxidative stress in short term treatment but more predominant in long term treated group.

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Conflict of interest: None declared
Ethical approval: The study was approved by the Institutional Ethics Committee

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5. Turmeric-centerchem. V. 01-06/05. 40560, 40690, 47740, 47870-1. Available at www.centerchem.com