NEPHROPHARMACOLOGICAL ACTIVITY OF ETHANOLIC EXTRACT LEPIDIUM SATIVUM L. SEEDS IN ALBINO RATS USING CISPLATIN INDUCED ACUTE RENAL FAILURE

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ABSTRACT
The present study was designed to investigate to possible potential nephrocurative & nephroprotective activity of 200mg/kg ethanolic extract of Lepidium sativum L. seed was used to against cisplatin (5mg/kg, i.p.) induced nephrotoxicity. The experimental protocol designed as the animals were divided into four groups (n=6) like control, model control, curative (200mg/kg) and protective groups (200mg/kg) were received vehicle, cisplatin, cisplatin + extract, and extract + cisplatin respectively. After 6th days, blood collected from retro-orbital sinus of rats and determined urea and creatinine level in serum of each group after then rats were sacrificed for quantitative estimation of various enzymes and ATPase content in kidney tissue. A single dose of cisplatin induced loss in body weight, increase urine excretion, increased urea & creatinine level in serum; it was significantly recovered by 200mg/kg in curative and protective groups. The enzyme estimation in kidney tissue it found that increase malondialdehyde, superoxide dismutase, catalase and reduced glutathione level, it was significantly monitored by 200mg/kg in curative and protective groups. The level of brush border enzymes like Na+K+ATPase, Ca++ ATPase and Mg++ATPase were found significantly reduced after single dose cisplatin injection. It was overcome by treatment of same extract in curative and protective groups. Finally it is concluded that the present study data conformed nephrotoxicity induced by cisplatin due oxidative stress and ethanolic extract of Lepidium sativum L. seeds may have nephroprotective and curative activity.

Keywords: Cisplatin; Nephrotoxicity; urea; creatinine; glutathione; Lipid peroxidation.

INTRODUCTION
A large number of medicinal plants, natural products and dietary components have been evaluated as potential nephroprotective agents1. The Lepidium sativum L. (family-Brassicaceae) is a native shrub. The Lepidium sativum (L.) seeds contain volatile essential aromatic oils, active principle and fatty oils and carbohydrate, protein, fatty acid, Vitamin: β-carotene, riboflavin, and niacin, and ascorbic acid, Flavonoids, Isothiocynates glycoside2. The Lepidium sativum L. seeds are used as aperients, diuretic, good anti inflammatory, demulcent, aphrodisiac, carminative, galactagogue, antiasthematic, antiscorbutic, and stimulant3-4. Cisplatin (cis-diaminedichloroplatinumII) (CDDP) is one of most potent anticancer drug. It is produced dose limiting nephrotoxicity and high dose of CDDP produce the impairment of kidney, causes decrease in renal blood flow, glomerular filtration rate and increases urea and creatinine level in blood5. The cisplatin induced nephrotoxicity was characterized by signs of injury such as changes in urine volume, body weight, increase the products of lipid peroxidation, and change renal clearance6. Kidneys have some antioxidant enzyme like superoxide dismutase (SOD), lipid peroxidase and glutathione (GSH), and catalase which protect kidney from free radicals like nitric oxide and superoxide etc. The cisplatin is inhibited the activity of antioxidant enzyme in renal tissue like glutathione, SOD, GSH and Catalase depletion and increase thiobarbuturic acid – reactive substance (TBARS)7. Thus, the purpose of current study was to investigate whether oral administration of ethanolic extract of Lepidium sativum L. (ELS) seeds has any protective and curative effect against cisplatin induced nephrotoxicity in albino rats. Its region behind Lepidium sativum seeds L. were traditionally used as diuretic and anti inflammatory8. 

MATERIALS AND METHODS
Drug and Reagents
Cisplatin (VHB, Life sciences Inc., India), DTNB (Merck Pvt. Ltd., India). Glutathione (Merck Pvt. Ltd., India), Thiobarbuturic acid (Loba chemicals Pvt.ltd. India).

Plant material
Lepidium sativum L. seeds were purchased from market of Mandsaur city (M.P., India). The plant was identified by Dr. H.S. Chattarjee (Ex professor of botany), P. G. College of Mansdaur, and M.P. And voucher specimen (BRNCP/L/02/2006) was submitted in department of Pharmacognosy; BRNCP, Mandsaur, M.P. The trampled seeds were extracted by soxhlet apparatus using ethyl alcohol as a solvent. The extract was dried by rotator evaporator under reduced pressure.

Animals
Adult male wistar rats having weight around 180-210 g were maintained at 25 ± 2°C and kept in well ventilated animal house under photoperiodic condition in large...
polypropylene cages and were standard food and water ad libitum. The experiment was carried out in accordance to the guidelines mentioned in the CPCSEA, and Institutional Animal Ethical Committee approved the experiment protocols (Reg.No.-947/ac/06/CPCSEA).

**Experimental design**

The acute toxicity study of ethanolic extract of *Lepidium sativum* seeds L. was not occurred at 2000mg/kg (as per the OECD - 420) on male Wistar rats. One tenth dose of 2000mg/kg was selected regarding toxicity study.

Total duration of study was 16 days. The animals were divided into four groups containing six animals in each group. Group I served as control and received normal saline throughout the experiment, Group II (Modal Control) received single dose of cisplatin (5mg/kg i.p.), 1st days, Group III (Protective) received ELS extract (200mg / kg p.o.) for 1st to 10th day and 11th day, single dose (5mg/kg, i.p.) of cisplatin was administered, Group IV (Curative) received same dose of cisplatin on day 11th, and after 6th days ELS extract (200mg / kg p.o.) was administered up to 16th days.

**Biochemical assays**

After the treatment period, blood was collected from retro-orbital sinus of rat under ether anaesthesia and centrifuged using the table top centrifuge (REMI) at 3000 rpm to get serum. Level of urea and creatinine was estimated using Span diagnostic kit on chemical analyzer (microlab3000) for assessment of renal toxicity. [8&9], After then Kidneys were removed, homogenized and centrifuged at 10,000 rpm at 0°C for 20 min. the supernatant was used for estimation of different antioxidant level by calorimetric method using spectrophotometer (Merck thermo spectronic, Model NO. UV-1, double beam), Glutathione reductase (GSH) estimated by Sedlak and Lindsay method10,11, Lipid peroxidation by thiobarbutic acid-reactive substances (TBARS) methods12,13, Superoxide dismutase (SOD) by method developed by Misra and Fridovich (1972). 14 Catalase (CAT) by colorimetric assay15, and the sediment of the centrifuge was used for estimation of the Na`K`ATPase by Bontin methods16, Ca`2`ATPase by Hjerken and Pan17, Mg`2`ATPase by Ohinishi et al. method18.

**Statistical analysis**

Results were expressed as one way analysis of variance (ANOVA) followed by Dunnett’s test and P< 0.05 was considered as significant.

**RESULTS AND DISCUSSION**

In present study rat treated with single dose of cisplatin shown marked reduction of body weight (173.33±4.21) in model control group as compared to control group (195.00±4.28). It was significantly (*P<0.05) recovered with treatment of 200 mg/kg ethanolic extract of *Lepidium sativum* L. seeds in curative group but not significantly in protective groups (Table no.1).

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Groups</th>
<th>Final body weight (gm) at last day of experiment</th>
<th>Urea level in serum (mg/dl)</th>
<th>Creatinine level in blood serum (mg/dl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Control</td>
<td>195.00±4.28</td>
<td>24.16±1.04</td>
<td>0.94±0.05</td>
</tr>
<tr>
<td>2.</td>
<td>Model control</td>
<td>173.33±4.21</td>
<td>76.66±2.24</td>
<td>2.32±0.10</td>
</tr>
<tr>
<td>3.</td>
<td>Protective (200mg/kg)</td>
<td>179.17±3.74</td>
<td>60.83±2.76**</td>
<td>2.00±0.04*</td>
</tr>
<tr>
<td>4.</td>
<td>Curative (200mg/kg)</td>
<td>188.33±2.591</td>
<td>40.83±0.90**</td>
<td>1.81±09**</td>
</tr>
</tbody>
</table>

Each value represents mean± S.D. of six animals

ns, statically different non significant when compare to the model control

**P<0.01, * P<0.05, **P<0.01, as compared to the model Control.

**Table 2:** Effect of treatment with ethanolic extract of *Lepidium sativum* seeds on the lipid peroxidation and antioxidant enzyme of kidney.

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Groups</th>
<th>µmol GSH/gm. Kidney tissue</th>
<th>n Mol MDA/gm. ml</th>
<th>(Unit SOD /gm kidney tissue)</th>
<th>CAT (µ mole of H2O2/gm kidney tissue)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Control</td>
<td>69.50±1.54</td>
<td>14.00±0.57</td>
<td>21.83±0.94</td>
<td>323.33±1.75</td>
</tr>
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<td>2.</td>
<td>Model control</td>
<td>45.33±1.66</td>
<td>24.50±0.61</td>
<td>07.16±0.60</td>
<td>201.67±3.33</td>
</tr>
<tr>
<td>3.</td>
<td>Protective (200mg/kg)</td>
<td>48.16±1.24**</td>
<td>21.50±0.76*</td>
<td>10.00±0.51*</td>
<td>219.17±6.24*</td>
</tr>
<tr>
<td>4.</td>
<td>Curative (200mg/kg)</td>
<td>59.66±1.28**</td>
<td>17.16±0.60**</td>
<td>15.33±0.88**</td>
<td>301.83±2.63**</td>
</tr>
</tbody>
</table>

Each value represents mean± S.D. of six animals

ns, statically different non significant when compare to the model control

**P<0.01, * P<0.05, **P<0.01, as compared to the model Control.
**Figure 1:** Effect of treatment with ethanolic extract of *Lepidium sativum* (L) seeds on urinary volume (ml/24hr.) in protective and curative groups.

Each group represents mean± S.D. of six animals, **P<0.01, * P<0.05, **P<0.01, as compared to the Model Control.

**Figure 2:** Effect of treatment with ethanolic extract of *Lepidium sativum* (L) seeds on Na⁺/K⁺ ATPase (mM of phosphate liberated/mg tissue) in protective and curative group.

Each group represents mean± S.D. of six animals. **P<0.01, * P<0.05, **P<0.01, as compared to the Model Control.

**Figure 3:** Effect of treatment with ethanolic extract of *Lepidium sativum* (L) seeds on Ca²⁺ ATPase (mM of phosphate liberated/mg tissue) in protective and curative group.

Each group represents mean± S.D. of six animals. **P<0.01, * P<0.05, **P<0.01, as compared to the Model Control.
cisplatin induced a significantly serum creatinine in wistar previous findings, we conformed that a single dose indicating induction of acute renal failure. According to accompanied by increase in serum creatinine level reduction of glomerular filtration rate, which is dose 5mg/kg body weight in rabbit caused a mark Jeong et al 20 indicates increase glomerular filtration rate. significantly reduced the level of urea and creatinine that extract of rate level suggests the reductio n of glomerular filtration groups (Table no.1). The increased urea and creatinine was significantly recovered in curative and protective groups. All the groups that received cisplatin (*P<0.05) effect on creatinine recovery in protective groups. (Table no.1). The administration of extract was significantly overcome with both 200mg/kg extract in curative group but 200mg/kg of protective group comparatively shown less significant (fig.1) The change of renal function observed in the rat correlate well with the nephrotoxicity effect with man22. The single dose of cisplatin (5mg/kg) result increased urea and creatinine level in model control group as compare to model control group. That is agreement with Matsushima et al21. The administration of extract was significantly overcome with both 200mg/kg extract in curative group and but less significant(*P<0.05) in protective group animals, the Na`K`ATPase were significantly decrease after damage of kidney, pathophysiological change in occur in proximal tubules cisplatin toxicity by formation of reactive species which cause the redistribution of brush border enzyme27. After damage of kidney, pathophysiological change occur in proximal tubules cisplatin toxicity by formation of reactive species which cause the redistribution of brush border enzyme27. In aspect of kidney tissue estimation, it is shown as significantly (**P<0.01) increase the lipid peroxidase (24.50±0.61) and decrease the level of GSH (45.33±1.66), SOD (07.16±0.60) and CAT (201.67±3.33) after single dose injection of cisplatin in model control group. The lipid peroxidase, SOD and CAT were monitored significantly (** (P<0.01) in curative group. However less significant (*P<0.05) same dose in protective group. (Table2). Our present result data shown that significantly monitored GSH, SOD, and CAT and lipid peroxidation. This is indicate that extract have antioxidant activity. Whereas in present phytochemical study of the extract have revealed the presence of Flavonoids, and amino acids like glutamine, Cysteine, and Glycine. The tannin (Phenolic compound), Flavonoids have antioxidant activity and Glutamate, Cysteine, Glycine were used to synthesis of the endogenous glutathione26. It’s all may contribute synergistic reason to increase GSH level in kidney tissue significantly.

The level of brush border enzymes like Na`K` ATPase, Ca` ATPase and Mg`ATPase were found to reduced significantly (**P<0.01) as compared to model control group animals, the Na`K` ATPase, Ca` ATPase and Mg`ATPase were recovered Significantly (**P<0.01) in protective group and but less significant(*P<0.05) in protective groups with treatment of same extract. (Fig. 3, 4 &5). After damage of kidney, pathophysiological change in occur in proximal tubules cisplatin toxicity by formation of reactive species which cause the redistribution of brush border enzyme27. In present result data reveal that the level of membrane bound enzyme Na`K` ATPase, Ca` ATPase and Mg`ATPase were significantly decrease after cisplatin administration. The extract was administrated cause to significantly increased Na`K` ATPase, Ca`
ATPase and Mg\(^{2+}\)ATPase in protective and curative groups. It is due to antioxidant activity. Finally it is concluded that the present study data conformed nephrotoxicity induced by cisplatin due oxidative stress and ethanolic extract of *Lepidium sativum* L. seeds may have nephroprotective and curative activity. However curative treatment was more significant than protective group. 

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