Effect of Sesamol on Arsenic Induced Hepato and Nephrotoxicity in Rats

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Arsenic is considered to be toxic when it is in an inorganic state and basic sources are contamination of water, smoking tobacco and irrigation of food crops. Sesamol is liposoluble lignans extraction that is used in rats to reduce skin papillomas. The main aim of the study is to study the effect of toxic arsenic on rats, usage of sesamol in treating hepato and nephrotoxicity in rats and analyze kidney tissue and liver tissues of the rats. The study primarily focuses on the effect of injecting arsenic and sesamol to the group of animals or injecting both and analyzing them to understand hepatotoxicity and nephrotoxicity. Sesamol along with arsenic powder have been used and the rats were kept in standard condition. The laboratory experiment has been carried out where both single and double oral treatments were provided. Animals were grouped into four groups where every 4 groups have eight rats each. SPPS software has been used to analyze the data collected. It has been shown treated rats with sesamol, counteract the toxicity effect of arsenic upon liver and kidney tissues. It has been found that sesamol has a protective effect upon arsenic induced liver and nephrotoxicity in rats.

Keywords: Arsenic; Hepato; Immunohistochem Kidney; Sesamol; Toxicity.

Arsenic is considered to be a natural element in the crust of the earth and is distributed in an overall environment of air, water as well as land. It is toxic when it is inorganic and the most toxic inorganic forms include Arsenate and Arsenite that have various mechanisms depending on the state of valence. Though homicides of arsenic receive publicity from the media, the basic source of toxicity of arsenic to the public is through contamination of water and utilizing that water in preparation of food and irrigation of food crops, eating food that is contaminated and smoking tobacco. Exposure in a longer period to toxic arsenic through drinking water and food can lead to chronic arsenic poisoning. Characteristic effects include skin lesions and cancer. Arsenic release into the environment propagates via weathering and the process of mining along with other phenomena like volcanic activity. Compounds of inorganic arsenic are toxic at a high level whereas compounds of organic arsenic tend to be less harmful to human health. Due to the toxicity of arsenic, the symptoms of acute poisoning of arsenic are vomiting, abdominal
pain along with diarrhea, muscle cramping and sometimes death in extreme cases. In case of long term exposure, higher degrees of arsenic in the inorganic form is detected in skin and have changes in pigmentation, skin lesion and patches on palm and foot sole. Along with skin cancer, it also leads to lungs cancer and bladder cancer. Other adverse effects of toxic arsenic include developmental impact, diabetes, pulmonary disease and disease of cardiovascular. It also leads to adverse pregnancy and infant mortality affecting child health and in early childhood, it is linked with enhancement in mortality in young people because of various cancer, lung cancer, failure in kidney and heart attacks.

Sesamol is a liposoluble lignans extraction along with a prominent fragrance component in sesame oil that is used to demonstrate activities of potential anticancer. The water-soluble lignin sesamol which is derived from Sesamum Indicum seed oil and is considered as a potent antioxidant along with a significant potent anticancer. This sesamol protects against atherosclerosis, hypertension and ageing along with healing of the wound, antioxidant, anti-inflammatory and free radical activity of scavenging. In the case of rats, it was evident that sesamol provides a 50% reduction in skin papillomas within 20 weeks after providing 12-O-tetradecanoylphorbol 13-acetate. After treatment of sesamol, plasma cholesterol along with the level of triacylglycerol were reduced and thus it was possibly inhibited HMG CoA reductase activity or activity of lipoprotein lipase. Sesamol has significantly reduced tumor burden along with lipid peroxidation and raised the level of antioxidants. This leads to inhibition of the development of a tumor on the skin along with promotion. Apoptosis in cells of the tumor was evident to cause by downregulation of Bcl-2 and stimulation of Bcl-2 that is aligned with an expression of X protein and is administered with free sesamol and encapsulated sesamol. The main objectives of the study are-

- The effect of toxicity of arsenic on Sprague Dawley rats.
- The usage of sesamol in treating hepatotoxicity and nephrotoxicity induced by arsenic in rats.
- To examine Histopathology in the liver and kidney of rats.

**MATERIAL AND METHOD**

**Drugs**

The powders along with Sesamol and arsenic were collected from Sigma Chemical Co., USA. In 1% of an aqueous solution of Tween 80, Sesamol was made whereas arsenic was made in normal saline which was stabilized through gum of 0.2%.

**Animals and treatments**

Male rat Sprague Dawley having weight approx. 200 g were collected from Animal House, College of Medicine, King Faisal University. These rats were preserved in the standard condition of approx. 24°C temperature, approx. 45% humidity and 12hr light or dark cycle. They were then supplied along with standard laboratory chow as well as water ad libitum. The rats were kept being acclimatized for a week before the conduction of experiments.

Ten hours before the experiment, animals were grouped in random process in four groups with every group having eight rats. A single dose of oral treatment of normal saline is stabilized through 0.2% gum that was provided to the first group and is served as control. In the group of the second, hepato and nephrotoxicity was induced by injected arsenic in rats. 3rd group was provided with arsenic with twice the dose of oral which is about 10mg/kg. Moreover both the 2nd and 3rd group of rats received two injections of either sesamol of dose of 50mg/kg or vehicle of sesamol which is of 1% aqueous solution of Tween 80 after administration of arsenic. The 4th rats’ group were provided with sesamol in absence of arsenic hepato and nephrotoxicity induction.

The protocol of the experiment was approved through the Local Animal Care committee along with procedures conducted aligning with international guidelines based on care and utilization of laboratory species.

**Sample preparation and biochemical studies**

The animals were killed a day after the administration of arsenic. Gathering blood samples was conducted and they were left for 1 hour to clot. After that, blood was centrifuged for 10 minutes at 5000 rpm to get a pure serum that was then stored at 20-degree temperature. To evaluate serum aspartate aminotransferase as well as alanine aminotransferase degree based on
manufacturer named Randox Laboratories Ltd. UK, recommending, colourimetric kits there were utilized.

Firstly the removal of the liver along with kidney tissue has been done, they were washed with ice-cold saline and were stored at 80-degree temperature. Utilizing cold potassium phosphate buffer, the liver and kidney were homogenized. Homogenates were separated at 5000 rpm for 10 minutes at 4°C and the supernatant was utilized to determine the malondialdehyde and to minimize the glutathione degree and to view the activity of catalase utilizing colourimetric kits of the assay as framed by manufacturer’s instruction of bio-diagnostic®. Depending on the manufacturer, Cayman Chemical Co, USA, instruction the degree of Nitric Oxide (NO) was assayed through utilizing a kit of the colourimetric assay.

Exam of Histopathological of liver tissue and kidney tissue

The samples of tissues of liver and kidney from every rat were fixed in 10% formalin that were subjected to dehydration in grades of ascending of alcohol and after that, they were embedded in paraffin. Sections were cut into 4m, which were then marked with hematoxylin along with eosin and investigated through utilizing a light microscope via a pathologist not aware regarding treatment protocol®.

Statistical analysis

To determine the data gathered, every value is expressed in mean ±S.E.M. one-way variance analysis was conducted followed by a Tukey test for multiple comparing. SPSS version 21 have been utilized and the difference is analyzed at the significance degree p<0.5.

RESULTS

Impact of sesamol on the measured biochemical parameters

The table above illustrates the effect of sesamol on antioxidants degree in arsenic which induced hepatotoxicity in animals. Glutathione in the case of treating with arsenic is found to be 108±5.3a and sesamol is about 137.4±5.9. While in

<table>
<thead>
<tr>
<th>GSH (mg/g tissue)</th>
<th>NO (nmol/100mg tissue)</th>
<th>MDA</th>
<th>CATALASE (u/g tissue)</th>
<th>Dose (mg/kg)</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>117.6±1.6</td>
<td>90.4±0.4</td>
<td>24.20±0.66</td>
<td>5.7±0.37</td>
<td>Control</td>
<td></td>
</tr>
<tr>
<td>108±5.3a</td>
<td>105.3±1.15a</td>
<td>53.11±1.13a</td>
<td>3.2±0.52a</td>
<td>10mg/kg Arsenic</td>
<td></td>
</tr>
<tr>
<td>137.4±5.9</td>
<td>142.7±1.8</td>
<td>34.8±0.87</td>
<td>4.8±0.6</td>
<td>50mg/kg Sesamol</td>
<td></td>
</tr>
<tr>
<td>115.6±1.9b</td>
<td>119.7±1.04b</td>
<td>45.26±0.92b</td>
<td>5.5±0.34b</td>
<td>Arsenic+sesamol</td>
<td></td>
</tr>
</tbody>
</table>

*MALD: malondialdehyde, NO: nitric oxide, GSH: glutathione.

All the values are expressed as mean ±S.E.M., n = 8 in each group.

a P < 0.05 vs. control group.
b P < 0.05 vs. arsenic group.

Table 2. Impact of sesamol on renal malondialdehyde (MDA), reduce glutathione (GSH) and NO level and catalase in rats which exposed to arsenic nephrotoxicity.

<table>
<thead>
<tr>
<th>Arsenic+sesamol</th>
<th>Sesamol</th>
<th>Arsenic</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>25.19±0.37b</td>
<td>30.78±0.24</td>
<td>37.6±0.33a</td>
<td>27.21±0.39 Mda(nmol/g tissue)</td>
</tr>
<tr>
<td>3.74±0.06b</td>
<td>4.01±0.15</td>
<td>3.2±0.12a</td>
<td>4.7±0.08 GSH(nmol/g tissue)</td>
</tr>
<tr>
<td>0.114±0.016b</td>
<td>0.118±0.013</td>
<td>0.129±0.02a</td>
<td>0.102±0.011 No(nmol/g tissue)</td>
</tr>
<tr>
<td>0.77±0.05a</td>
<td>1.28±0.05</td>
<td>0.54±0.07a</td>
<td>1.33±0.05 Catalase(U/g tissue)</td>
</tr>
</tbody>
</table>

All the values are expressed as mean ±S.E.M., n = 8 in each group.

a P < 0.05 vs. control group.
b P < 0.05 vs. arsenic group.
the case of treating both arsenic and sesamol, the Glutathione level is 115.6± 1.9b. This reveals that the level of effect of sesamol treatment is higher on an antioxidant degree in arsenic that leads to hepatotoxicity among animals.

In table 2 it has been evident that while injecting arsenic, the level of MDA is 37.6± 0.33, in the case of sesamol it is 30.78±0.24 and while injecting both Arsenic and sesamol it is 25.19± 0.37b. The level of GSH while injecting arsenic is 3.2±0.12a, in the case of sesamol it is 4.01±0.15 and injecting both arsenic and sesamol is 3.74±0.06a.

From the above graph, it has been found through ‘t’ test that while injecting arsenic to rats exposed to hepatotoxicity, an inclination in weight has been observed in the liver than that of injecting sesamol and both arsenic and sesamol altogether.

The above graph depicts that in the case of injecting arsenic along with sesamol, the serum enzyme level of alanine aminotransferase increased

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**Fig. 1.** Impact of sesamol on variation in weight of liver of arsenic induced hepatotoxicity in rats

**Fig. 2.** Effect of treatment of sesamol on the level of serum of alanine aminotransferase and aspartate aminotransferase in rats
and in the case of aspartate aminotransferase while injecting arsenic, the serum enzyme level increased.

**Effects of sesame oil on liver and kidney histopathology**

In figure 4 the animals falling under the control group (figure 4, A) shows normal glomerulus and normal renal tubules of the kidney. There exists no dilation in renal tubules and no such cytoplasmic vacuolization of renal tubules was observed (fig4, A) [12]. The animals falling under the group of arsenic-treated show abnormal dilation of renal tubules and there is a swelling of epithelial cells [13]. Cytoplasmic vacuolization of epithelial cells was observed (fig4, B). The animals falling under the group treated with arsenic along with sesame oil show the recovery of kidney parenchyma (fig4, C). The animals falling under the sesame oil treated group exhibit normal glomerulus along with tubules of the kidney (fig4, D).

**DISCUSSION**

The results shows that while treating the rats with sesame oil on renal malondialdehyde. There was a reduction in the level of glutathione as well as nitric oxide and catalase in rats that were exposed to arsenic nephrotoxicity [10]. Table 3 shows the effect of sesame oil on BUN and serum creatinine levels in rats when they were exposed to arsenic nephrotoxicity. It has been seen that when injecting arsenic, Blood urea

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**Fig. 3.** A) Photomicrograph of the liver from the control group, shows a normal parenchyma of the liver. Hepatocytes (black arrow) are radiating from the central vein (blue arrow). No hepatic pathology was observed. H&E stain. X 40. B) Photomicrograph of the liver from the arsenic treated group, showing vacuolization of cytoplasm in hepatocyte (black arrow). The central vein became congested with inflammatory cells (orange arrow). H&E stain. X 40. C) Photomicrograph of the liver from the group treated with (arsenic plus sesame oil), shows a recovery of the parenchyma of the liver (blue and black arrows). H&E stain. X 40. D) Photomicrograph of the liver from the sesame oil treated group, shows a normal parenchyma of the liver (black arrows). H&E stain. X 40.
nitrogen (BUN) increases and in the case of serum creatinine level it also increased. When injected with Sesamol both BUN and Serum Creatinine are higher among rat when exposed to arsenic nephrotoxicity. When both Arsenic and sesamol are injected to rats, both the level of BUN and Serum are increased than the standard range. Treatment of acute arsenic has led to marked damage of the liver like ballooning degeneration, centrilobular necrosis along with cytoplasmic vacuolation of hepatocytes with sinusoidal congestion (fig-3B). Markedly the treatment of sesamol attenuated the injury of tissue when induced with arsenic and restored the same kind of histopathological figure which was seen in the control group (fig-3C). In the control group (fig3A) animals shows normal liver parenchyma. It was observed that there was radiation of hepatocytes from a central vein and no such hepatic pathology is being observed. Among the animals belonging to the group treated by arsenic, it was observed cytoplasmic vacuolation in hepatocytes. Moreover, it has been observed that the central vein has become congested with that of inflammatory cells. In the group, the animals are treated with both arsenic along with sesamol that indicates a recovery of liver parenchyma. Furthermore, the rats under the group of sesamol treated, exhibit normal liver parenchyma. The animals of hepatocyte were generally compared with the animals from the group of control.

From the figure, it has been depicted that while treating acute arsenic there is marked damage

Fig. 4. A) Photomicrograph of the kidney from the control group, shows a normal glomerulus (black arrow), and normal tubules of the kidney (orange arrow). No renal pathology was observed. H&E stain. X 40. B) Photomicrograph of the kidney from the arsenic-treated group, shows an abnormal dilatation of renal tubules (black arrow), and swelling of the epithelial cells (blue arrow). Note the vacuolation of epithelial cells in the tubules of the kidney (orange arrow). H&E stain. X 40. C) Photomicrograph of the kidney from the group treated with (arsenic plus sesamol), shows a recovery of the parenchyma in the kidney (Black and blue arrows). H&E stain. X 40. D) Photomicrograph of the kidney from sesamol treated group, illustrates a normal glomerulus and tubules of the kidney (black arrows). H&E stain. X 40
of the liver such as ballooning degeneration, centrilobular necrosis and cytoplasmic vacuolation of hepatocytes as well as sinusoidal congestion. While treating with sesamol, the injury of tissue when injected with arsenic restores a similar kind of histopathological figure in the control group. Normal liver parenchyma has been seen along with the radiation of hepatocytes from a central vein and no hepatic pathology is observed. The group of animals who are treated with arsenic suffered from cytoplasmic vacuolization in hepatocytes20. In this case, the central vein is congested along with inflammatory cells. Animals, when treated with both arsenic and sesamol, recover from liver parenchyma but when treated with only sesamol it leads to normal liver parenchyma. The animals of the control group having normal glomerulus and renal tubules of the kidney do not face dilation in renal tubules and also cytoplasmic vacuolization of renal tubules. The animals who are being treated with arsenic face from kidney parenchyma whereas when treated with sesamol normal glomerulus were found along with kidney tubules.

CONCLUSION

From the findings, it has been evident that there is a higher effect of sesamol treatment on the level of antioxidants in arsenic which leads to hepatotoxicity among the animals. This leads the liver to develop inflammation as the animal is exposed to toxic substances like arsenic. In such cases, the liver normally removes and breaks down the drugs along with chemicals from the bloodstream14-16. The level of MDA after injecting arsenic tends to be higher leading to the decrease in the level of GSH and is lower in the case of injecting sesamol leading to the increase in GSH level14. It has also been evident that at the time of treating with sesamol on renal malondialdehyde leads to the reduction of the degree of glutathione along with nitric oxide as well as catalase in rats which is exposed to arsenic nephrotoxicity17. This leads to a lower concentration of Glutathione in plasma from the vein of hepatic that was not higher than that of concentration in plasma from the portal vein and from the aorta that indicates depleted liver was not releasing glutathione into plasma 18-19. It was also evident that when the rat is exposed to nephrotoxicity while injecting arsenic BUN and serum creatinine is increased as well as injecting arsenic and sesamol both lead to an increase in the level of BUN and serum Creatinine. Through ‘t’ test it has been evident that while injecting arsenic to a rat that is exposed to hepatotoxicity they gained in weight of liver than that of injecting either sesamol or both arsenic and sesamol. There is also a rise in serum enzyme level of alanine aminotransferase when injecting arsenic and an increase in the level of serum enzyme of aspartate aminotransferase is seen in the case of injecting arsenic.

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REFERENCES


