**Escherichia coli** Generate Oxidative Stress and Enhance Lipid Peroxidation in the Kidney of the Rat

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**Abstract**

The urinary tract is generally infected by *Escherichia coli* bacteria. They commonly originate in the patient's own bowel, and infection occurs mostly via the ascending route. In this study, we have observed the influence of *E. coli* on oxidative stress generation and lipid peroxidation in the kidney of the rat. *E. coli* were taken from the soil, urine, buffalo intestine and goat intestine for this study. Rats were infected with isolated *E. coli* from different sources and Lipid peroxidation, glutathione assay was done to achieve the goal of this study. Percentage survival data showed that the *E. coli* isolated from urine had more lethargy phenomena because their survival present was 66.66% and the mortality rate was higher in this group. Although *E. coli* isolated from Goat intestine has also shown the same mortality, but the *E. coli* isolated from urine sample shown this mortality from the second day whereas the *E. coli* isolated from goat intestine from the third day. By and large data indicated that the *E. coli* isolated from urine sample generated high oxidative stress and damage rat kidney because the kidney is an organ frequently exposed to oxidative stress. Overall, free radical generated due to *E. coli* infection further enhances lipid peroxidation in the rat which is harmful to the physiology because it may cause a renal disorder in the rat.

**Keywords:** *E. coli*, Virulence; Rat; Kidney; Lipid peroxidation

**Introduction**

Our gastro-intestinal tract is a biologically diverse and complicated system which contains numerous bacterial species. Intestinal microbes harbor a diverse microbial community, often containing opportunistic bacteria with virulence potential [1]. Both the host and bacteria are thought to derive benefit from each other. While most of the activities of the normal flora benefit their host, some of the normal flora are parasitic (live at the expense of their host), and some are pathogenic (capable of producing disease) [2]. Diseases that are produced by the normal flora in their host are called endogenous diseases. Most endogenous bacteria are opportunistic in infection [1]. Among endogenous bacteria, *Escherichia coli* are generally harmless for humans but some of them are parasitic and may cause life dangerous complication like a severe kidney damage function or even sepsis. *E. coli* may live for months in water and ground but they get the opportunity they multiply very quickly and cause digestion problems [3,4]. We can describe *E. coli* as opportunistic bacteria only in the specific situation when it arrives from the intestine to other organs and tissues and causes illnesses by producing Shiga toxin [5,6]. The most frequent are urinary tract and sexual organs infections. *E. coli* outbreak has killed many persons in the USA in and other parts of the world [7,8]. Virulence factors of *E. coli* are studied and reported in different population by many research groups [9-13]. *E. coli* infection was also reported in domestic animals like rabbits, cats, dogs, goats and horses [14]. The main site of *E. coli* infection is kidney and their infection is spread through blood [15]. Therefore in this study kidney and serum burden was measured after total dosages regimen (infection time period) and data of their dilution series showed that kidney and blood burden was higher in buffalo having *E. coli* infection. The sources of *E. coli* were soil, urine, buffalo intestine and goat intestine. This study was designed to investigate the antibiotic resistance, serum analysis and virulence of *E. coli* (isolated from the different source) in the rat.

**Materials and Methods**

**Concept design of the study**

The present study was emerged from isolation of *E. coli* from soil sample, urine sample, goat (G) and buffalo (B) intestine. They were collected locally and stored at -20°C until use.

**Isolation of *E. coli* bacteria**

Samples were firstly inoculated in 1% tryptone broth for 24 hours for the isolation of *E. coli* from different sources mentioned above. Then the broth was again inoculated on Mac-konkey Agar media for 24 hrs. Then the pink colonies were selected and again transferred on EMB agar media for 24 hrs. The metallic shine colonies were selected and stored in LB agar media for at 4°C until use.

**Biochemical tests for identification of bacteria**

**Urease test:** Urea agar medium and urea broth were prepared and sterilized by autoclaving at 121°C. Media was poured into the tubes and allows them to solidify in a slanting position to form slopes. The tubes were labeled with the name of the inoculating organism. Inoculate the liquid and agar media with a transfer loop. Incubate the slants/broths for 24-48 hrs at 37°C.

**Carbohydrate catabolism by microorganisms test:** Glucose

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media was used to identify the oxidative and fermentative abilities of microorganisms. Acid end products were generated, pH was lowered and the bromthymol blue indicator was turned yellow.

**IMVIC tests:** Indole methyl red Voges-Proskauer citrate (IMVIC) test was undertaken to identify the \textit{E. coli} in frequent form.

**Animals study**

Swiss albino rats of either sex weighing 100 ± 5 gm were selected randomly from the animal house of Madras Veterinary College, Chennai, India. They were housed in a controlled room with a 12 h light dark cycle at the room temperature of 22 ± 2°C, humidity 30% to 60%. They were kept on standard pellet diet (Lipton India Ltd.). Animal maintenance and handling were in accordance to the internationally accepted standard guidelines of Committee for the Purpose of Control and Supervision of Experiment on Animals (CPCSEA). Experiments were conducted according to the guidelines and ethical norms, approved by ministry of social justice and empowerment, government of India. Six rats were taken for each \textit{E. coli} sample isolated from different sources. 10^9 CFU/ml broth cultures of \textit{E. coli} strains isolated from different sources were given to experimental animals.

**Leucocytes counting**

It was done by using following formula:

\[
\text{WBC/ \mu L} = \frac{\text{Average of cells} \times \text{Correction for dilution}}{\text{No. of squares counted} \times \text{Volume of one square}}
\]

**Biochemical assays**

**Lipid per oxidation (LPO):** MDA, an end product of LPO reacts with TBA to form a colored substance. Measurement of MDA by TBA reactivity is the most widely used method for assessing LPO. Lipid peroxidation was measured via the amount of liberated malondialdehyde (MDA) in the system that assayed the pink product, as reported earlier [16]. The colored end product was read at 540 nm.

**Glutathione (GSH) measurement:** Cellular GSH activity was measured according to method of Paglia and Valentine et al. [17]. The rate of change of absorbance during the conversion of NADPH to NADP was recorded using Shimadzu spectrophotometer at 340 nm for 3 min.

**Statistical analysis**

Results were expressed as mean ± SD (standard deviation). Three sets of experiments were performed and the results were analyzed by One-Way ANOVA (analysis of variance) test. The upper level of significance was chosen as P<0.05 (Significant).

**Results**

Pure culture of isolated \textit{E. coli} from different sources was tested with three antibiotics (Gentamycin, Erythromycin, and Chloramphenicol) to study the inhibition effects. The result is depicted in Table 1. Maximum inhibition was observed by Erythromycin (33.5 mm) followed by Chloramphenicol (26.5 mm) for urine isolates cultures. Gentamycine inhibited more to the goat intestine isolate (24.0 mm). Data surveillance of test animal (Table 2) relieved that out of six animals tests for each group of culture only two were found dead by urine isolate and only one were found dead by goat intestine isolated culture whereas other culture did not cause dead and showed 100% survival. Variation in
Table 3: E. coli burden in blood serum of rat.

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Infection of E. coli isolated from different sources</th>
<th>Serum burden (number of E. coli ± SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>First dilution</td>
</tr>
<tr>
<td>1</td>
<td>Control</td>
<td>1.00 ± 0.29</td>
</tr>
<tr>
<td>2</td>
<td>Soil</td>
<td>5.00 ± 1.31</td>
</tr>
<tr>
<td>3</td>
<td>Urine</td>
<td>2.00 ± 0.48</td>
</tr>
<tr>
<td>4</td>
<td>B. Intestine</td>
<td>69.30 ± 80.27</td>
</tr>
<tr>
<td>5</td>
<td>G. Intestine</td>
<td>17.00 ± 5.24</td>
</tr>
</tbody>
</table>

Table 4: E. coli burden in kidney of rat.

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Infection of E. coli isolated from different sources</th>
<th>LPO level (mM/mg protein ± SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Control</td>
<td>0.54 ± 0.23</td>
</tr>
<tr>
<td>2</td>
<td>Soil</td>
<td>1.36 ± 0.83</td>
</tr>
<tr>
<td>3</td>
<td>Urine</td>
<td>4.09 ± 0.38</td>
</tr>
<tr>
<td>4</td>
<td>B. Intestine</td>
<td>1.69 ± 0.79</td>
</tr>
<tr>
<td>5</td>
<td>G. Intestine</td>
<td>2.52 ± 0.30</td>
</tr>
</tbody>
</table>

Table 5: LPO level in rat infected by E. coli isolated from different sources.

Figure 2: Lipid peroxidation in infected rat showed that there was increase in LPO in treating group and it was found significant from control at P<0.05.

Figure 3: Glutathione measurement showed that there was decrease in GSH level in treated rat and significant from control at P<0.05.
in the infected kidneys may be due to the infiltration of the pelvic epithelium and sub epithelial connective tissue by mononuclear cells and polymorphonuclear leukocytes. With the above mentioned data (result section), the kidney and serum burden of E. coli was measured with regimens showed more virulence to cause infection the antibiotic effect on E. coli are also measured. The increase of LPO and decrease in GSH in treating group may cause increase oxidative stress in the kidney (Figures 2 and 3).

Conclusion

This study showed the effect of E. coli infection in terms of perturbation of LPO and GSH in experimental animal. Peroxidation of lipids disturbs the integrity of cell membranes and leads to rearrangement of membrane structure. They produce adverse modifications to cell components, such as lipids, proteins, carbohydrates, DNA and alter physiological functions.

Acknowledgements

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