Effects of Potassium Dichromate on Distribution of
Toxocara canis Larvae within Brain of Mice

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Abstract: The frequency of the infective larvae of Toxocara canis, the common parasite of the dog was assessed in brain portions of male Albino mice ingested two doses: 200 mg/Kg and 400 mg/Kg of potassium dichromate-hexavalent (VI) as an indicator. Only 2,000 ova of the parasite were inoculated orally to twenty seven healthy young male mice. Frequency of the parasite was tested 13 weeks post-inoculation. Using bio-statistical analysis (one way analysis of variance) the frequency of detected larvae of both doses in the brain portions of the experimental animals showed an insignificant difference (p ≥ 0.05) in comparison with control. The results indicate the safety of small and mild intoxication doses of the potassium dichromate on distribution of Toxocara canis in brain portions of mice.

Key words: Potassium dichromate, Toxocara canis, brain, immunity.

1. Introduction

The interest in the biochemical and toxicological aspects of chromium is due to its role as environmental pollutants and a nutritional factor in the public as well as a suspected carcinogenic in occupationally exposed workers [1]. Potassium dichromate, K₂Cr₂O₇, a crystalline ionic solid with a very bright, red-orange color, is a common inorganic chemical reagent, most commonly used as an oxidizing agent in various laboratory and industrial applications. As with all hexavalent chromium compounds Cr(VI), it is acutely and chronically harmful to health. The salt is popular in the laboratory because it is not deliquescent, in contrast to the more industrially relevant salt sodium dichromate [2]. The exposure sources of chromium compounds are described in some other works [3-5]. The main use of Cr(VI) is as a precursor to potassium chrome alum, used in leather tanning [2-6].

Side effects of it may extend further to other industries. In 2005-2006, potassium dichromate was the 11th-most-prevalent allergen in patch tests (4.8%) [7] and one of the most common causes of chromium dermatitis [8]. It is highly likely to induce sensitization leading to dermatitis, especially of the hand and fore-arms, which is chronic and difficult to treat. Toxicological studies have further illustrated its highly toxic nature. With rabbits and rodents, concentrations as low as 14 mg/Kg have shown a 50% fatality rate amongst test groups [9]. Aquatic organisms are especially vulnerable if exposed; hence responsible disposal according local environmental regulations is advised. As with other Cr(VI) compounds, is carcinogenic and should be handled with gloves and appropriate health and safety protection. It also is corrosive and exposure may produce severe eye damage or blindness [10]. Human exposure further encompasses impaired fertility, heritable genetic damage and harm to unborn children.

In a previous work the toxic effects of potassium dichromate was assessed on various tissues of mice [5] and the toxicity of the lead monoxide on the activity of immune system in fighting the infectious larvae of Toxocara canis was assessed [11]. The distribution
frequency of *Toxocara canis* larvae was significantly higher in mice fed on lead monoxide indicating the toxicity of lead monoxide. This led us to explore whether the ingestion of potassium dichromate does exert any similar inhibiting effects on immune system as the lead monoxide.

The parasite *Toxocara canis* has a complex ascarid life cycle as it is not immediately infectious when it leaves the definitive host. It must grow and develop into the infective stage, ensheathed, in order to infect the definitive host. When *Toxocara canis* ova are intubated to mice (the normal host), the larvae hatches and invades the wall of small intestine within 2 hours [12]. After one day, they accumulate in the liver, lungs, kidneys and spleen. They may eventually invade the brain and the eyes [13, 14]. They may cause ocular lesions [15] and an epileptic condition in man. Their survival in the brain is interpreted to be as refuge free from the effects of host immunological responses [14]. A 2.5% solution of potassium dichromate at 4 °C could provide an infectivity of *Cryptosporidium* oocytes for immune suppressed adult mice [16]. In the present work, ingestion of potassium dichromate crystals, similarly to lead monoxide, was tested against the distribution of *Toxocara canis* larvae in brain portions.

2. Materials and Methods

Extraction of the *Toxocara canis* egg from the intestine of dead puppies was performed according to method described by Al-Tae [11]. Twenty seven healthy newly weaned male mice were selected to study the distribution frequency of the infective larvae in brain portion. All animals were divided into groups and maintained under identical environmental condition at 25 °C ± 2. Normal laboratory chow and tap water were available *ad lib* and were replaced three times a week. Using gavage animals were intubated with 2,000 eggs of the parasite.

Potassium dichromate crystals were obtained from FLUKA AG, CH-9470 Buschs, Switzerl and as ground powder and mixed with diet to prepare two concentrations (doses) of 200 and 400 mg/Kg body weight. The mice were fed the mixture for 13 weeks as 200 mg/Kg body weight (G1 = 8) or 400 mg/Kg (G2 = 10). After 13 weeks, all animals were sacrificed by cervical head dislocation for *Toxocara canis* distribution test. The brain was dissected out and cut into 4 pieces; the two cerebral hemispheres, and left and right (LCH & RCH); medulla oblongata (MO); and cerebellum. They were then squashed between two clean and dried slides and counting of the larvae was done by direct examination via Olympus BO light microscopy.

3. Results

Upon continuous monitoring system, neither epilepsy-like condition nor any other abnormal behavioral condition was recorded. However, via counting the frequency of the migrant larvae to the brain portions, insignificant differences ($p \geq 0.05$) were seen between group 1 (G1) and (G2) in comparison with control (Table 1).

4. Discussion

The resulting pathology of *Toxocara canis* is

<table>
<thead>
<tr>
<th>Animal groups</th>
<th>Mean density of <em>Toxocara canis</em> larvae in brain portions</th>
<th>Total</th>
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<tbody>
<tr>
<td></td>
<td>MO</td>
<td>LCH</td>
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<tr>
<td>Control (n = 9)</td>
<td>6.9 ± 2.1</td>
<td>30.3 ± 4.3</td>
</tr>
<tr>
<td>Group-1 (n = 8)</td>
<td>8.5 ± 2.1</td>
<td>29.9 ± 3.3</td>
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<tr>
<td>$p$</td>
<td>$&gt; 0.05$</td>
<td>$&gt; 0.05$</td>
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<tr>
<td>Group-2 (n = 10)</td>
<td>11.0 ± 1.3</td>
<td>24.3 ± 2.8</td>
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<tr>
<td>$p$</td>
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dependent on the intensity of infection and the location of the larvae within the body organs [17]. Infection protocols or mice strains also vary considerably making species comparison difficult [18]. It also suggested that *Toxocara canis* larval burdens vary between individual outbred mice receiving the same inoculation may refer to the role of immunity in the establishment of irreversible brain, particularly, cerebral infection [19]. The mice strain used in this study might have been different; and hence interprets the lower concentration of control mice in the present results than those of other studies [11, 15, 20]. Nevertheless, mice remain a useful animal model for significant applicability and manipulation.

With lead monoxide used instead of potassium dichromate, ingestion *Toxocara canis* ova, produced almost 3-4 folds (significant differences ($p \leq 0.05$)) frequency of the infective larvae in brain portions of mice more than in the present results [11]. The lower density of the infective larvae with potassium dichromate could be interpreted as the effect of the Cr(VI) was less toxicity exerted on the immune system in comparison with lead monoxide. These results are concomitant with those of Abdelhameed [20] and the most recent work of Holland and Hamilton [19] in that the greater concentration of the parasites in the brain could result as infection progress.

The results indicates that future systematic immune responses to *Toxocara canis* in the brain has received little attention so far which open the door wide for future immunological researches to link the distribution of the infective larvae to the immune system in both human and animal.

**5. Conclusion**

Perhaps potassium dichromate is much safer than lead monoxide in affecting immune system in mice. Animal species, strains and inoculation methods may yield varying results dependent on susceptibility of each and perhaps to environmental condition too. Further experiments are necessary to link the distribution of *Toxocara canis* larvae with the immunological studies would help in understanding the implication of various heavy metals as well as cerebral toxocariasis.

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**References**

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