Does Oregano Protect Against Testicular Toxicity Produced by Ethylene Glycol in Adult Male Albino Rat?

¿El Orégano Protege Contra la Toxicidad Testicular Producida por Etilenglicol en Ratas Macho Albinas Adultas?

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SUMMARY: Origanum vulgare Linn has traditionally been used as a diuretic and antispasmodic. Therefore, we investigated the active extract of Origanum vulgare for possible andrological effect and preventive effects against testicular damage using ethylene glycol rat model of testicular damage, to rationalize its medicinal use. Male Wistar rats received lithogenic treatment comprising of 0.75 % ethylene glycol injection twice with one day interval, then in drinking water, active extract of Origanum vulgare treatment (20 mg/kg) was given for 3 weeks to prevent toxic damage including loss of body weight gain and appetite. Following oral administration of EGME, a rapid decrease in testis weight associated with testicular cell damage was observed. Origanum vulgare treatment (20 mg/kg) prevented as well as reversed toxic changes including loss of body weight gain.

KEY WORDS: Origanum vulgare; Ethylene glycol; Testicular damage.

INTRODUCTION

The ability of ethylene glycol mono-n-alkyl ethers to produce testicular damage has been known for some time. Wiley et al. (1938) first demonstrated the testicular effects of ethylene glycol monomethyl ether (EGME) in rabbits in 1938. Stenger et al. (1971) have shown that the monoethyl derivative (EGEE) can also induce testicular damage in dogs and rats. More recently, Nagano et al. (1979), using a variety of ethylene glycol monoalkyl ethers in studies with orally treated mice, have shown EGME, ethylene glycol monomethyl ether acetate (EGMEA) and EGEE to be the most potent of the series examined in producing adverse testicular effects. EGME has also been shown to produce a decrease in testicular weight in the rat and mouse following inhalation exposure for 2 weeks at 1000 ppm.

Glycolic acid is the major metabolite of ethylene glycol responsible for toxicity (Barceloux et al., 1999). The toxic mechanism of ethylene glycol poisoning is mainly due to the metabolites of ethylene glycol. Initially it is metabolized by alcohol dehydrogenase to glycolaldehyde, which is then oxidized to glycolic acid (Maier, 1983). The metabolic effects occur 12 to 36 hours post ingestion, causing primarily metabolic acidosis which is due mainly to accumulated glycolic acid. Additionally, as a side effect of the first two steps of metabolism, an increase in the blood concentration of lactic acid occurs contributing to lactic acidosis. The formation of acid metabolites also causes inhibition of other metabolic pathways, such as oxidative phosphorylation. The rate-limiting step in this cascade is the conversion of glycolic to glyoxylic acid. Accumulation of glycolic acid in the body is mainly responsible for toxicity (Brent, 2001). Ethylene glycol has been shown to be toxic to humans (Friedman et al., 1962) and is also toxic to domestic pets such as cats and dogs. A toxic dose requiring medical treatment varies but is considered more than 0.1 mL per kg body weight (mL/kg) of pure substance. That is roughly 16 mL of 50 %
ethylene glycol for an 80 kg adult and 4 mL for a 20 kg child. Poison control centers often use more than a lick or taste in a child or more than a mouthful in an adult as a dose requiring hospital assessment (Caravati et al., 2005). The orally lethal dose in humans has been reported as approximately 1.4 mL/kg of pure ethylene glycol (Friedman et al.). That is approximately 224 mL of 50 % ethylene glycol for an 80 kg adult and 56 mL for a 20 kg child. Although survival with medical treatment has occurred with doses much higher than this, death has occurred with 30 mL of the concentrate in an adult (Amathieu et al., 2006). In the EU classification of dangerous substances it is ‘harmful’ (Xn) while more toxic substances are classified as ‘toxic’ (T) or ‘very toxic’ (T+). The U.S. Environmental Protection Agency generally puts substances which are lethal at more than 30 g to adults in Toxicity Class III. Ethylene glycol has a low vapor pressure; it does not evaporate readily at normal temperatures and therefore high concentrations in air or intoxication are unlikely to occur following inhalational exposures (Hodgman et al., 1997). There may be a slight risk of poisoning where mists or fogs are generated, although this rarely leads to poisoning as ethylene glycol causes irritation and coughing when breathed in, alerting victims to its presence (Wills et al., 1974). Ethylene glycol is not well absorbed through skin meaning poisoning following dermal exposure is also uncommon (Driver et al., 1993).

**Origanum vulgare** Linn (family, Lamiaceae) is distributed throughout Asia, Europe and North America and is commonly known as Wild Marjoram and Winter Sweet and locally in Pakistan as Mirzanjosh, Sathra. It is widely used in the traditional medicine as lithotriptic, diuretic and antispasmodic along with other medicinal uses, such as stimulant, expectorant, antibacterial, anticancer, anti-inflammatory, antioxidant and laxative (Lemhadri et al., 2004).

Oregano is a light green in color with strong aromatic odor and pleasantly bitter taste Oregano goes well with tomato sauces, pizza and vegetables (Gilani et al., 2005). Some researches provides evidence that oregano improves growth in growth retarded animals and has no specific immunostimulatory effects on porcine immune cells (Williamson, 2001). **Origanum vulgare** Linn (family, Lamiaceae) is distributed throughout Asia, Europe and North America and is commonly known as Wild Marjoram and Winter Sweet and locally in Pakistan as Mirzanjosh (PDR for Herbal Medicines, 2000). It is widely used in the traditional medicine as lithotriptic, diuretic and antispasmodic along with other medicinal uses, such as stimulant, expectorant, antibacterial, anticancer, anti-inflammatory, antioxidant and laxative (Duke, 2002).

**MATERIAL AND METHOD**

**The materials used for this experiment are as follows:**
OHAUS electric weighing balance, Teflon Pyrex tissue ground, Soxhlet apparatus, thimble, extractor, bolt head flask, water bath, condenser, animal cages, hemocytometer, Electronic microscope, Test tubes, measuring cylinder, light microscope, desiccator.

**Chemicals and Reagents:** Anhydrous sodium sulphate, n-hexane, normal saline solution, ethanol, distilled water, formalin, chloroform, eosin, blue in scott’s tap water.

**Animals:** The present study included (50) sexually active albino rats of weight between 250-280 g. The animals were obtained from the Animal House of Zagazig University Egypt. The animals were kept in the Animal House in experimental room for one week prior to the commencement of study, for acclimatization to experimental conditions with 12 h light and dark cycle. The animals was fed ad libitum with commercially prepared rat feeds made up of 21 % protein, 3.5 % fat, 6 % fiber, 0.8 % phosphorus and given fresh water. All the experimental procedures were done following the experimental guidelines of the institutional ethics committee on the care and use of laboratory animals.

**Preparation of plant extracts:** **Origanum vulgare** leaves were bought from Sekam company for medicinal plant (Cairo, Egypt) and identified by taxonomist The aerial part of the plant material was cleaned of adulterants and kept soaked for three days in the aqueous-ethanol (30:70) with occasional shaking, at room temperature. The filtration was carried out using a muslin cloth and then through Whatman qualitative grade 1 filter paper. This procedure was repeated twice and then all the filtrates obtained were combined and concentrated on a rotary evaporator (RE-111, Buchi, Flawil, Switzerland) accompanied with B-700 recirculation chiller and a water bath model 461 at 65 °C under vacuum. Finally, the extract change to a thick pasty mass called as crude extract (Ov.Cr), yielding approximately 12 % which reconstituted in distilled water (100 mg/ml) to be used in the study.

**Preliminary phytochemical analysis:** The crude extract of **Origanum vulgare**, was screen for the presence of different phytochemical groups such as alkaloids, saponins, coumarins, sterols, terpenes, tannins and flavonoids by using different methods of extract: Group A: Ethanolic extract. Group B: Aqueous extract. Group C: Ethyl acetate extract. Group D: Hexan extract (Sarac & Ugur, 2008).
Experimental Protocol

Five groups with 10 rats were selected randomly in each group. Groups from 2-5 were the test groups, while group 1 was used as a control. Group one was fed with 5 ml /kg) orally of distilled water using a poly-ethene catheter while group two, three, four and five were giving 0.5 ml of 0.75 % ethylene glycol injection twice with one day interval then ethylene glycol EG (0.75 %) 50 mg/kg/day: Group 2 Control positive {untreated} received oral feeding of ethylene glycol EG (0.75 %) treatment for 14 days by stomach tube (1 ml/kg), Group 3 Treated group of extract A received oral feeding of ethylene glycol EG (0.75 %) treatment and extract B in a dose of 20 mg / Kg (5 ml/kg) orally for 14 days. Group 4 Treated group of extract B received oral feeding of ethylene glycol EG (0.75 %) treatment and extract B in a dose of 20 mg / Kg (5 ml/kg) orally for 14 days. Group 5 Treated group of extract C received oral feeding of ethylene glycol EG (0.75 %) treatment and extract C in a dose of 20 mg / Kg (5 ml/kg) orally for 14 days.

Sample Collection. The animals were anesthetized by deep ether anaesthesia and then sacrificed. Orchietomy was performed by open castration method. A pre-scrotal incision was made and the testicles was milked out of the incision site and weighed with the aid of OHAUS electric weighing balance. The testicles were exposed by incising the tunica vaginalis. The spermatic cord was exposed, ligated and incised. Semen samples were thereafter collected from the caudal epididymis. The method of collection was similar to that described by Olugbenga & Oniovosa (2005).

Sperm count and motility assay: Immediately after dissection, the epididymal contents was dropped on a glass slide and viewed under the light microscope to determine the motile and non-motile sperm cells. The motile and non-motile sperm cells were distinguished by the movement of the motile and non-motile sperm cells. The motile and non-motile sperm cells were counted by hemocytometer using the improved Neubauer (Deep 1/10 mm, LABART, Germany) chamber as described by Pant & Srivastava (2003).

Histological procedure. After extracting the testes from the animal’s body, the organ was promptly and adequately treated with 10 % formal saline (fixation) in order to preserve its structure and molecular composition. The sections were floated on water, transferred to a glass slide and stained with haematoxylin and eosin stains with the aid of the Leica auto stainer XL. The slide was then viewed under a light microscope with varying magnification. Spermatogenesis was assessed by a method which depends upon scoring ‘cross sectional’ profiles of seminiferous tubules according to Johnson, using Johnson’s score. For each testis, five vertical sections from the polar and the equatorial regions were sampled and an unbiased numerical estimation of the following morphometric parameters was determined using a systematic random scheme; The diameter of seminiferous tubules with profiles that were round or nearly round were measured for each animal and a mean, D, was determined by taking the average of two diameters, D1 and D2 (Perpendicular to one another). D1 and D2 were taken only when D1/D2 ≤ 0.85, others include epithelial height, testicular weight, sustentacular cells (Sertoli cells) to germ cell ratio (elongated spermatids) (Qin & Lung, 2002). Tissues were fixed for 24 hr and processed into paraffin wax. Sections were stained with hematoxylin and eosin and with periodic acid-Schiff technique for the demonstration of the spermatid acrosome and definition of the stages of spermatogenesis was based on that used by Leblond & Clermont (1952).

RESULTS

The control negative group (200x; H&E) has normal cyto-architecture, reveal interstitial tissues (IT) and seminiferous tubules (ST) with sustentacular cells and the normal developing germ cells: spermatogonia (Sp), spermatocytes I (SPI), spermatids (Spt), spermatozoa (Spz) with flagellum. In addition to that no sloughing of the germinal epithelium and the lumen of the epithelium remained patent and intact (Fig. 1).

Table I. Androgenic parameters.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Control</th>
<th>Group 2</th>
<th>Group 3</th>
<th>Group 4</th>
<th>Group 5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sperm Motility (%)</td>
<td>54.1±2.3</td>
<td>30±2.4</td>
<td>45.8±1.6</td>
<td>43.1±1.3</td>
<td>48±1.6</td>
</tr>
<tr>
<td>Sperm count (x 10^6/ml)</td>
<td>19.5±0.5</td>
<td>13.1±6.1</td>
<td>18.6±0.5*</td>
<td>19.2±3.4**</td>
<td>17±0.5</td>
</tr>
<tr>
<td>Sustentacular to Spermatid ratio</td>
<td>8.1±0.2</td>
<td>1.8±0.9</td>
<td>5.9±0.9</td>
<td>7.1±0.4**</td>
<td>5.6±0.5</td>
</tr>
<tr>
<td>Johnsen’s score</td>
<td>8.1±0.9</td>
<td>4.5±0.3</td>
<td>6.8±0.21</td>
<td>7.0±0.6*</td>
<td>5.8±0.2</td>
</tr>
<tr>
<td>Seminiferous tub dia (µm)</td>
<td>344.2±9.9</td>
<td>154.3±10</td>
<td>269.4±13.8</td>
<td>302.4±7**</td>
<td>245±12</td>
</tr>
<tr>
<td>Height of Epith (µm)</td>
<td>189±7.2</td>
<td>66.9±6</td>
<td>133.6±13*</td>
<td>161±1.4**</td>
<td>154±9*</td>
</tr>
<tr>
<td>Testicular Weight (g)</td>
<td>1.42±0.07</td>
<td>0.9±0.3</td>
<td>1.2±0.7</td>
<td>1.3±0.2**</td>
<td>1.2±0.5</td>
</tr>
</tbody>
</table>

*shows the significant difference with control group. (P<0.05) **shows highly significant difference with control group (P<0.01).
In control positive group 2 shows closely packed small round abnormal spermatocyte massive decrease of spermatozoa within the lumen of seminiferous tubule, complete absence of sustentacular cells and secondary spermatocytes included cytoplasmic vacuolation (Fig. 2).

Group 3, Group 4 & Group 5 shows seminiferous tubules with few spermatozoa and interstitial space with interstitial cells (Leydig cells) shows presence of few sperm cell within the lumen of the seminiferous tubules vascular interstitial cells due to vascular interstitial cell moving spermatozoa along the lumen of the tubules EGME Treatment (Fig. 3-5).
DISCUSSION

Following oral administration of EGME, a rapid decrease in testis weight associated with testicular cell damage was observed. Prolonged dosing resulted in continued spermatocyte degeneration as well as maturation depletion of the spermatid population, leaving tubules containing only sustentacular cells, spermatagonia and preleptotene spermatocytes. Although zygote spermatocytes showed slight degeneration with prolonged exposure to high doses, maturation arrest was evident by the increased numbers of cells present. Arrest of spermatocytes at this stage indicated a failure of the cells to accomplish successful transition from zygote to pachytene. At the ultrastructural level the mitochondria of the spermatocytes showed early evidence of disruption. Cessation of treatment indicated that the testicular lesion was reversible in the large majority of tubules. Testis weights returned to control values and the majority of tubules regained full spermatogenesis and spermatoozoa production within one full spermatogenic maturation cycle (7-8 weeks in the rat). The presence of a small proportion of totally atrophic tubules indicated a loss of spermatogonia, as the cell type not protected by the blood-testis barrier (BTB), are the most vulnerable to toxic effects (Meistrich, 1986). Effects of toxic compounds on the testis may be reversible following cessation of compound exposure through seminiferous epithelium in the rat. The presence of a small proportion of totally atrophic tubules indicated a loss of spermatogonia, as the cell type not protected by the blood-testis barrier (BTB), are the most vulnerable to toxic effects (Meistrich, 1986). Effects of toxic compounds on the testis may be reversible following cessation of compound exposure through seminiferous epithelium in the rat. The presence of a small proportion of totally atrophic tubules indicated a loss of spermatogonia, as the cell type not protected by the blood-testis barrier (BTB), are the most vulnerable to toxic effects (Meistrich, 1986).

REFERENCES


RESUMEN: Origanum vulgare Linn se ha usado tradicionalmente como diurético y antiespasmódico. Por lo tanto, investigamos el extracto acuoso del Origanum vulgare (20 mg / kg) durante 3 semanas con el objetivo de prevenir el daño tóxico, la pérdida de peso corporal y el apetito. Palabras clave: Origanum vulgare, Etilenglicol, Daño testicular.