Research article

Pathological effects of mercury chloride on reproductive system in white rats

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Abstract

The present study was undertaken to know the reproductive toxicity of mercury chloride in male and females of white rats. This study was used twelve (6 male and 6 females) white rats of approximately of the same body weight (200-220 g) divided equally in to 3 groups; the first group (T1) was received mercury chloride (1mg/kg B.W intraperitoneally once daily for 30 days). While the second group (T2) was received, mercury chloride (1.5mg/kg B.W intraperitoneally once daily for 30 days).Third group was received only 0.2ml of Distilled water considered as control group. At the end of experiment, the animals were sacrificed and small pieces of (2cm³) were taken from ovary, uterus and testis of all animals to histopathology. Histopathological sections of these organs of (T2) group was showed severe pathological changes characterized by vaculation in epithelial cells of uterus, cystic dilatation of uterine glands with degeneration of epithelial lining of its, hemorrhage and decrease in number of growing follicles in ovary also there were sever pathological changes in the testes. While (T1) group was showed less pathological changes characterized by hyperplasia of epithelial lining with few and small uterine gland in uterus, presence of large secondary follicles in ovary and there were less pathological changes in testes. 

Keyword: Mercury Chloride, Pathological effects, White rats.

Introduction

Mercury is one of the most dangerous environmental pollutants to living organisms and it is considered as one of the heavy metal, which is the chief component of various medicines (1). It Its discharge can occur through natural sources such as volcanoes or anthropogenic activities such as industrialized processes, agriculture and metallization (2). The release of mercury can cause increase in the amount of atmosphere mercury, which enters to the soil-water distribution cycles where it can remain in circulation for many years. Mercury poisoning is occur due to exposure to mercury or mercury compounds lead to various toxic effects depending on route of exposure and chemical form of its (3). Animals and humans reacts with their environment daily and are exposed to a lot of chemicals and heavy metals present in the environment by food, air and water (4). It causes a variety of health effects including respiratory, neurological, reproductive, renal, immune and dermatological effects (5). Mercury is an element found in the environment causing oxidative stress in the exposed living organisms leading to tissue damage (6). It is toxicity is linked to its high affinity for sulfhydryl groups (-SH), forming stable complexes causing several changes, like structural changes of sulfhydryl enzymes and inactivation of their active sites (7).
Mercury influences antioxidant mechanisms in the cell lead to cell degeneration, lack of membrane safety and then cellular necrosis (8). Most of the studies with experimental animals are undertaking on male adult rats or pups exposed for short periods to mercury (9), (10) and (11). However, few studies using acute treatments on female’s reproductive system have been carried out. So my present study is make to investigate the pathological changes induced by difference dosage of mercury chloride on male and female reproductive system of white rats for 30 days.

Materials and Methods

Ethical approval
The Animal Ethical Committee of Veterinary Medicine College, University of Al-Qadisiyah, Iraq, has approved the present study under permission No: 416

Experimental animals: The present study was conducted on (12) white rats (6 male and 6 female) of approximately the same age and body weight (200-220 gm). The animals were housed in plastic cages in an air conditioned room with temperature maintained at 25±2 C in animals house of Veterinary Medicine College/ University of Al-Qadisiyah under 12 hours light/12 hours dark. Rats were given food pellets and water ad libitum and divided into three groups 4 rats each (2 male and 2 females) and this study continued for 30 days.

Chemicals: Mercury chloride is a heavy metal obtained from central laboratory in University of Al-Qadisiyah. Mercury chloride (BDH chemical Ltd (England)). The rats administered 1.5mg/kg B.W and 1.0 mg/kg B.W (12) as chronic doses.

Experimental design: Twelve white rats, both sexes were divided into 3 groups (4 rats each) and were treated as following:
1st group was injected with (1mg/kg B.W) intraperitoneally.
2nd group was injected with (1.5mg/kg B.W) intraperitoneally.
3rd group was injected with (0.2ml) distal water as control group.

Tissue samples:
The rats were sacrificed and the ovary, uterus and testis were dislocated by sterile scissor. Then 10% formalin fixed, small pieces (2cm³) were taken from these organs of all groups for histopathology. Processed routinely in histokinette, cut at 5 Mm thickness by microtome (Juny 4291, West Germany) and stained with Haematoxylin &Eosin stain then examined under light microscope (13).

Results
Histopathological section from uterus, ovary and testis from treated rats were examined under light microscope. Uterus: Uterus of second group (high dose group) show vaculation of endometrium cells, cystic dilatation of endometrium gland with degeneration of epithelium which lining these glands Figures (1, 2), while the first group(low dose group) show less pathological changes characterized by hyperplasia of endometrium with infiltration of inflammatory cells also there were few and small of uterine glands Figures (3, 4).

Ovary. The present study revealed that the severity of pathological changes on the ovary was increased with increased of doses, in second group (high dose group). We showed decreased number of growing follicles, also we showed small ovarian follicles at different stages of maturation including primary and secondary follicles also there were hemorrhage in stroma of ovary Figure (5). While in first group (low dose group) these changes is less compared with high dose group and can we showed ovarian follicles at different stages of maturation including
primary, growing and mature follicles. The mature follicle appeared nearer to the surface of the ovary and has the ova Figure (6). Testis: Histopathological examination of testis in second group (high dose group) showing extensive degeneration in some tubules characterized by vaculation of Spermatogonia, severe suppression of spermatogenesis Figure (7) and There are few Leydig cells Figure (8). The severity is depending on the dose whereas these changes is mild in first group (low dose group) which characterized by slight degenerations in spermatids and spermatozoa Figure (9).

Figure (1): Histological section of uterus in rats treated with mercury chloride (1.5mg/kgB.W) show of endometrium cells vaculation. 10XH&E.

Figure (2): Histological section of uterus in rats treated with mercury chloride (1.5mg/kgB.W) show cystic dilatation of endometrium gland (red arrow) with degeneration of epithelium which lining these glands (blue arrow).10XH&E.

Figure (3): Histological section of uterus in rats treated with mercury chloride (1mg/kgB.W) show hyperplasia of endometrium (red arrow) with infiltration of inflammatory cells (blue arrow). 10XH&E.

Figure (4): Histological section of uterus in rats treated with mercury chloride (1mg/kgB.W) show few and small of uterine glands. 10XH&E.

Figure (5): Histological section of ovary in rats treated with mercury chloride (1.5mg/kgB.W) show small and decreased number of growing follicles (blue arrow) and haemorrhage (red arrow).10XH&E.

Figure (6): Histological section of ovary in rats treated with mercury chloride (1mg/kgB.W) show normal section of ovary of characterized by present of mature follicle.10XH&E.
Figure (7): Histological section of testis in rats treated with mercury chloride (1.5mg/kgB.W) show vacuolation of spermatogonia.10XH&E

Figure (8): Histological section of testis in rats treated with mercury chloride (1.5mg/kgB.W) show vacuolation of Spermatogonia (red arrow) and There are few Leydig cells (blue arrow).40XH&E

Figure (9): Histological section of testis in rats treated with mercury chloride (1mg/kgB.W) show slight degenerations in spermatids and spermatozoa.10XH&E.

Discussion
Reproductive problem like abortion, congenital malformation, infertility and decrease in ovulation are the most signs conducted with mercury toxicity (14). The present study demonstrated that mercury chloride has toxic effects on reproductive system of rats in high doses due to histopathological changes, which showed in these organs. The changes in uterus of second group (high dose group) show vacuolation of endometrium cells, cystic dilatation of endometrium gland with degeneration of epithelium which lining these glands. while the first group(low dose group) show less pathological changes characterized by hyperplasia of endometrium with infiltration of inflammatory cells also there were few and small of uterine glands. These changes occur may be due to altering of levels of follicle-stimulating hormone (FSH), luteinizing hormone (LH), estrogen and progesterone (14, 15) and the results agreed with (16, 17) who recorded that high dose of mercury has been shown to inhibit the release of FSH and LH from the anterior pituitary which in turn can effect estrogen and progesterone levels leading influential on endometrium. The present study revealed that pathological changes on the ovary in second group (high dose group) showed decreased in number of growing follicles also we showed small ovarian follicles at different stages of maturation including primary and secondary follicles also there were hemorrhage in stroma of ovary. While in first group(low dose group) these changes is less
comparing with high dose group and can we showed ovarian follicles at different stages of maturation including primary, growing and mature follicles. The mature follicle appeared nearer to the surface of the ovary and has the ova. These changes are agreed with (18). Who were revealed that chronic nonlethal levels of mercury induced histopathological changes in the ovary of rats interfering with their breeding. In accordance (19) revealed that oral given of mercuric chloride in a dosage of 1.5 mg/kg BW using in vivo and in vitro fertilization of mouse could damage the ovary function and reduce the number of ovulation oocytes. These pathological changes in the present study may be explained by the direct toxic effect of mercury on the ovarian tissues by its high affinity for sulphhydryl groups (-SH), forming stable complexes also its reacts with phosphoryl, carboxyl, and amide groups resulting in many dysfunction of enzymes and structural proteins(20). However, it is possible that mercury chloride accumulating in the ovari and having little effect on ovulation (21). or indirectly via the hypothalamus– pituitary– ovarian axis (22). While the changes is less and may be near to normal is may be due to the less effect of mercury in ovary and ovulation in low dose (23). Histopathological examination of testis in second group (high dose group) showing extensive degeneration in some tubules characterized by vacuolation of Spermatogonia, severe suppression of spermatogenesis and There are few Leydig cells. The severity is depending on the dose whereas these changes is mild in first group (low dose group) which characterized by slight degenerations in spermatids and spermatozoa. The result of our study is in agreement with the results obtained from previous study (24) and (25) They mention HgCl₂–induced testicular damage in animals is commonly associated with Spermatogonia damage, spermatid degeneration, Leydig cell dysfunction, testicular disorder and giant cell formation. Various mechanisms have been suggested to explain Hg induced cellular toxicity. Among these mechanisms, lipid peroxidation has been considered a primary initiating mechanism during Hg injuries (26). Many studied recorded that an increase in free radical formation relative to loss of antioxidant defense system (27). Also (27) suggested that HgCl₂ generates free radicals by interacting with DNA that interferes with antioxidant defense system and results in the tissue injury, testis and epididymis are more susceptible to oxidative damage leading to their functional inactivation (28).

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