EXPERIMENTAL STUDY

Ovarian development in Wistar rat treated prenatally with single dose diisobutyl phthalate

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Abstract: Phthalates are a class of industrial compounds with an array of toxicological properties used in day to day life. Diisobutyl phthalate on (DIBP) is used as an additive to keep the plastics soft or flexible (plasticizer) in nitrocellulose plastic, nail polish, explosives, lacquer manufacturing etc. Although DIBP exposure in humans is generally low, people in adhesive industries and pharmaceutical industries are exposed to higher levels. The aim of this study was to determine the effect of single dose of DIBP on developing ovary of Wistar rat. One hundred and eight adult pregnant Wistar rats were divided into control and experimental groups. Rats in experimental group were given DIBP on day 10, 12 and 14 of gestation at 0.375, 0.75 and 1.25 ml/kg body weight dose intraperitoneally in a single dose. Sections of ovaries collected on day 21 of gestation were stained with hematoxylin and eosin and examined and Masson’s trichrome histologically. Sections belonging to the control group showed the presence of oocytes in clusters separated by thin fibrous septa. Degeneration oocytes, empty follicles surrounded by follicular cells without gonocytes in the center were observed in ovarian stroma. Blood vessels in the ovarian stroma were prominent and congested. Around a bunch of follicles total architectural disarray was observed although on special staining fibrosis was not evident. As pregnant women are constantly exposed, effect of DIBP on ovary of a developing fetus would denote the long term consequence in future generations (Fig. 5, Ref. 39). Full Text in PDF www.elis.sk.

Key words: ovary, rat, diisobutyl phthalate, ovarian follicle, embryology, gonocytes.

Phthalates are a class of widely used industrial compounds known technically as dialkyl or aryl esters of 1, 2-benzenedicarboxylic acid. The common forms of phthalates are Diethylhexyl phthalate (DHEP), Diallyl phthalates (DAP), Di-n-propyl phthalates (DPP), Di-n-butyl phthalates (DBP), and Diisobutyl phthalates (DIBP).

Phthalates with a variety of toxicological properties are used in day to day life. These phthalates are nearly omnipresent in modern society and are found in the plastic items, vinyl flooring, wall coverings, lubricants, adhesives, detergents, nail polish, hair spray and car wash (1).

The adverse effects produced by phthalates are changes in morphology, physiology, growth, development or life span of an organism which results in an impairment of functional capacity and ability to compensate for additional stress (2).

Diisobutyl phthalate

DIBP (phthalic acid, di isobutyl ester 1,2-benzenedicarboxylic acid, bis-(2-methyl propyl) ester di-(isobuty1)-1,2-benzene dicarboxylate) is the branched isomer of DBP and an additive used to keep the plastics soft or more flexible (plasticizer) often in combination with other phthalates. DIBP is used in nitrocellulose plastic, nail polish, explosive materials, lacquer manufacturing, consumer products, blood bags, pharmaceuticals and automobile parts. They are ubiquitous in our environment and most people including pregnant women and their fetuses are exposed to multiple phthalates at a time. Saillenfait et al estimated that exposure of pregnant women to DIBP was 0.12 μg/kg/day with maximum of 2.9 μg/kg/day (3).

Several studies have shown that DIBP exposure in humans are generally low, near the limit of detection, though a small percentage of people working in adhesive industries and pharmaceutical industries are exposed to higher levels of DIBP.

The dermal absorption of DIBP has been assessed, along with other phthalates by applying on back and urinary and fecal excretion was assessed (4).

In humans, DIBP is metabolized to monoisobutyl phthalate (MIBP) which can be detected in the urine (5). A recent human study showed urinary MIBP concentrations in mothers were inversely related to anogenital index in male offspring. Total and free mono-n-butyl phthalate in human urine samples after medica-
tion of a di iso butyl phthalate containing capsule was determined by Seckin et al (6).

Singh et al (1972) administrated 0.375, 0.75 and 1.25 ml/kg body weight of DIBP by intraperitoneal route on gestation day 5, 10, 15 to the pregnant Sprague dawley rat. They found fetal mortality, growth retardation and skeletal abnormalities (7).

According to data produced by the Australian Government Department Of Health And Ageing, the toxic level of DIBP by Intraperitoneal route (LD50) is 4.5 mg/kg body weight of rat or greater than 1600mg/kg body weight of rat (8).

By the gavage route the rat showed toxicities at different quantities of DIBP. At and above 500 mg/kg body weight the mother rat showed transient decrease in body weight. At 750 mg/kg and 1000 mg/kg body weight the parent rat showed marked embryo lethality and teratogenicity. At and above 500mg/kg body weight, the fetal rat showed decrease in the body weight and male fetuses with undiscented testis were seen at 500 mg/kg body weight (3).

DIBP administrated by gavage is embryo toxic and teratogenic, and affects the developing male reproductive tract. The male fetuses with undiscented testes and visceral and skeletal malformations i.e., fused sternebrae occurred at a significantly higher frequency. Two skeletal variations were, retarded ossification and impairment of reproductive function in fetal rats and may have greater than 1600mg/kg body weight of rat (8).

In an experiment conducted in pregnant Wistar rats, gestation day 20/21 rather than 19 appeared to be the optimal time for investigating changes in anogenital distance together with reductions in testicular testosterone production and testicular expression of Ins1 and genes related to steroidogenesis. PPARα mRNA levels were reduced by DIBP at gestational day 19 in testis and liver. In females, DIBP increased anogenital distance and increased ovarian aromatase mRNA levels (10).

In a study by Seckin et al (6), DIBP on developing ovary of Wistar rat. Our working hypothesis is DIBP affects intraperitoneal ovarian development in rat. Objectives were to determine the effect of single dose of DIBP on developing ovary of Wistar rat. We investigated the effect of DIBP on the development of ovary in Wistar rat. Our working hypothesis is DIBP affects intraperitoneal ovarian development in rat. The aim of the study was to determine the effect of single dose of DIBP on developing ovary of Wistar rat. Our working hypothesis is DIBP affects intraperitoneal ovarian development in rat.
tive is to study the effect of DIBP as a teratogen on early ovarian development.

Materials and methods

108 Female Wistar rats of an average weight of 200 gm and an average age of 120 days were used in this study and they were housed individually in plastic cages in noise-free, air conditioned animal house with temperature maintained at 75 °F and on a light dark cycle of 12:12 hours. Humidity was maintained with a minimum of 50 %. Rats were fed on diet pellets (Hindustan Lever, Bombay, India) and tap water ad libitum and treated with utmost human care. The experiment was carried out with prior approval from the institutional animal ethical committee. The female rats in their pro-oestrous were caged overnight with males of the same stock (Female : Male = 3 : 1). Presence of sperms in the vaginal smear on the following morning confirmed start of gestation and the day was numbered as the day ‘zero’ of pregnancy.

Experimental Groups

One hundred and eight adult pregnant Wistar rats were used in the present work. They were divided into two main groups. Group I (54 rats) were treated with equal amount of distilled water and serve as control. Group II (54 rats) were given intraperitoneal injection of DIBP on day 10, 12 and 14 of gestation. The animals of treatment group (n = 6) were administrated DIBP intraperitoneally in a single dose with the help of a sterile tuberculin syringe. Along with this experimental group, a control group was also maintained and administered equal amount of distilled water alone or left uninjected. DIBP was obtained from Durga laboratories, Manglore, India.

The pregnant rats were sacrificed by cervical dislocation on day 20 of pregnancy. Pups were collected through laparotomy. Ovaries were dissected out approaching from anterior abdominal wall. They were fixed in formalin, embedded in paraffin and sectioned at 8μm thickness. Sections were stained with haematoxylin and eosin stain and Masson’s trichrome and examined under microscope. The photomicrographs for histological studies were taken with the help of a Photomicroscope.

Results

The microscopical examination of haematoxylin and eosin stained sections of the ovary belonging to the control group showed the presence of oocytes in clusters separated by thin fibrous septa. Degeneration of oocytes was observed in all experimental groups (Fig. 1). Pyknotic and irregular nuclei of oocytes indicated that they were in the process of degeneration. This process of degeneration ultimately resulted in death of the central oocytes. Empty follicles surrounded by follicular cells without gonocytes in the center were observed in ovarian stroma (Fig. 2). Blood vessels in the ovarian stroma were prominent and congested as compared to control (Fig. 3). Damaged follicles were noted and all these together gave an appearance of total architectural disarray with elongated cells giving a hint towards increased fibrosis (Fig. 4). However, fibrosis was not observed amongst damaged follicles as observed on Masson’s trichrome stained slides (Fig. 5). Severity of findings was also tabulated.

Discussion

In the present study, DIBP caused loss of germ cells leading to empty follicles and eventual loss and regression in size of follicles. As a repair mechanism to the insult, prominent and congested vasculature was noted. Also the process of restoration is ongoing; fibrosis is not yet evident replacing the lost follicles as seen in Masson’s trichrome stained slides. Widespread architectural disarray resulted from the above mentioned insult.

Lee et al reported, loss of germ cell development increasing with higher doses of DBP, leading to entire loss of spermatozoa from the tubule, a Sertoli-cells only appearance was observed. Giant cell formation was often a feature associated with loss of germ cells (31). DIBP in high dose group caused complete loss of germ cells in testis. Seminiferous tubular hypoplasia was observed in different doses. In few cases lower doses caused total tubular necrosis (3). Borch et al observed DIBP caused central location of gonocytes in seminiferous tubules and multinuclear gonocytes (11).

DBP and DIBP induced changes in gametes can be explained by their DNA mutating capacity. Kleinsasser et al. reported, using an in vitro comet assay, that DNA damage (single-strand breaks) was significantly induced by DIBP (35μumol/mL) in human oropharyngeal and nasal mucosa (32). In further work, Kleinsasser et al found that DIBP induced strand breaks in DNA, in both blood lymphocytes and normal mucosal cells from the oropharynx or larynx of patients of head and neck cancer (33).

DIBP is potentially dangerous because it has a similar structure to androgenic hormones in the humans. DIBP was negative for estrogenic activity in a yeast two-hybrid assay (34) and showed extremely weak estrogenic activity in recombinant yeast assay (35). DIBP (up to 10–5 M) had no binding affinity for the estrogen receptor α or β in vitro (36) but was also found to induce estrogen receptor α-mediated estrogenic activity and possess anti-androgenic activity in vitro but showed no activity towards ERβ in CHO-K1 cells (37).

According to Kemper and Peters, on day 10, germ cells migrate towards gonadal ridge, on day 12, they are localized either in the mesenchyme between gut and future genital ridges or in the future genital ridges and on day 13 almost all PGC had reached the well-developed genital ridges, therefore, day 14 is the period after germ cells have settled down in ovary (16). Degeneration of gonocytes was observed mostly when they were affected by the insult while in the process of migration and proliferation, rather than after settlement in gonadal ridge when their rate of proliferation reduces. However, the highest dose affected them considerably. Degenerated gonocytes ultimately gave rise to empty follicles. The germ cells were selectively affected as they remain in highly proliferating state at the period of insult. According to Merchant, PGCs do not contain reserve glycogen and lipid inclusions and move by amoeboid movement which require consumption of energy (13).
Gondos observed vacuoles in the cytoplasm immediately adjacent to granulosa cell membranes as well as cytoplasm lying adjacent to the plasma membrane of germ cells show some pinocytic vacuoles associated with phospholipid bodies (38). This arrangement suggests possibility of synthesis of material by granulosa cells, which transfer them directly across the narrow intercellular space for the use of germ cells having limited capacity for synthesis and secretion (39). These intercellular junctions may be involved not

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<th>Normal/Degenerating oocyte</th>
<th>prominent congested blood vessels</th>
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**Fig. 1.** Photomicrograph showing sections of fetal ovaries with normal oocytes in control and degenerating oocytes in different experimental groups (white arrow). H & E × 400.

**Fig. 3:** Photomicrograph showing sections of fetal ovaries with normal vasculature in control and prominent congested vessels in different experimental groups (white arrow). H & E × 400.

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<th>Normal/Empty follicle</th>
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**Fig. 2.** Photomicrograph showing sections of fetal ovaries with normal oocytes in control and empty follicles surrounded by follicular cells in different experimental groups (white arrow). H & E × 400.

**Fig. 4.** Photomicrograph showing sections of fetal ovaries with normal fibrous septa in control and architectural disarray in different experimental groups (white arrow). H & E × 400.
only in facilitating the migration of germ cells but also in germ cell-somatic cell interaction for providing exogenous substances (gases and nutrients). Therefore, the PGC, which are dependent on surrounding somatic cells for the transfer of nutrient and gases, might have got affected by the agent. Prominence and congestion of vasculature and ultimate architectural disarray was prominent with the highest dose of DIBP on 10th and 14th day and the lower 2 doses in 12th day exposure group. Probably 12th day was the most vulnerable period but effects with highest dose were less conspicuous because compensatory mechanisms took over to repair the damage. Although haematoxylline and eosine stained slides gave a hint towards increased fibrosis, it was negated on Masson’s trichrome stained slides. Possibly the time gap between the injection and observation was too short for fibrosis to develop.

Our results can be explained based on previous studies where it has been shown that DIBP is known inducer of DNA mutation and stimulates androgenicity and inhibits estrogenicity. On one hand, it has been shown that DIBP causes loss of germ cells, detaches the germ cells from basement membrane which ultimately would result in impending loss in experimental testis (3, 11, 31).

Conclusion

Our study indicates that DIBP causes germ cell loss and damage to healthy tissue and total architectural disarray in developing ovary when administered in early stages of gestation. Effect of DIBP on ovary of a developing fetus would denote the long term consequence in future generations.

References

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