PRENATAL EXPOSURE TO DI-(2-ETHYLHEXYL) PHthalate AND HIGH-FAr DIET
SYNERGISTICALLY DISRUPTS MOUSE FETAL OOGENESIS AND AFFECTS
FOLLICULOGENESIS

BY

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ABSTRACT
Di-(2-ethylhexyl) phthalate (DEHP) is a chemical that is widely used as a plasticizer. Plasticizers are chemicals that are added to materials to increase flexibility, durability, and strength. DEHP is abundant in products such as food storage containers, containers of personal care products, cosmetics, children’s toys, medical tubing, and blood storage bags. DEHP is known as an endocrine disrupting chemical and has the ability to leach out to the environment from products readily. Humans are exposed to DEHP through oral ingestion, inhalation, dermal contact, and during medical procedures. Exposure to DEHP alters ovarian function in humans. Additionally, foods high in fat content, regularly found in the western diet, are potential disruptors of fetal ovarian function. Due to DEHP’s high lipophilicity, high-fat foods can be easily contaminated with DEHP. Therefore, exposure to DEHP and a high-fat diet are both health concerns, especially in pregnant women, and the effects of these exposures on fetal oocyte quality and quantity should be elucidated. The goal of this study was to determine if there are synergistic effects of DEHP exposure at an environmentally relevant level (20 µg/kg body weight/day) and high-fat diet on oogenesis and folliculogenesis. To fulfill that goal, female CD-1 mice were fed with a high-fat diet (45 kcal% fat) or a control diet (10 kcal% fat) one week before mating and during pregnancy and lactation. The pregnant mice were orally dosed with DEHP (20 µg/kg body weight/day) or tocopherol-stripped corn oil (vehicle control) from E10.5 to E18.5 or until litter birth. Immunostaining was carried out to monitor meiotic gene silencing and synapsis in E18.5 oocytes, after making meiotic chromosomal spreads. At 8 days postpartum (dpp) and 21 dpp, F1 ovaries were harvested and subjected to histological evaluation. We found that treatment with an environmentally relevant dosage of DEHP and consumption of high-fat diet relevant to a western diet significantly increases synapsis defects in meiosis and affects the development of preantral follicles in the F1 generation.
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CHAPTER 1
INTRODUCTION¹

Mammalian females are born with a fixed number of primordial follicles within the ovaries. These primordial follicles either developed into antral follicles, ovulated and fertilized or eliminated from the follicle pool through atresia. The size of the oocyte pool at birth determines the female’s reproductive lifespan [1].

Oogenesis is the process where a primordial germ cell developed into a mature oocyte. During fetal development, diploid primordial germ cells entered meiosis. Meiosis is a special type of cell division that gives rise to haploid gametes from diploid primordial germ cells [2,3]. The main stages of meiosis include prophase I, metaphase I, telophase I, anaphase I, interkinesis, prophase II, metaphase II, anaphase II, and telophase II [4]. In the fetal ovaries, these oocytes progress through meiotic prophase I that includes several substages: leptotene, zygotene, pachytene, and diplotene stages. Around birth, all the oocytes are arrested at late diplotene stage, called dictyate [1]. Shortly before ovulation, oocytes resume meiosis I and then arrest at metaphase II until fertilization. Without fertilization, mammalian oocytes are unable to complete meiosis II [5].

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Exposure to environmental chemicals can cause adverse effects on reproductive systems [6], and this phenomenon is known as reproductive toxicity. Indeed, exposure to certain chemical substances causes adverse effects on oogenesis [7]. Meiosis-disrupting toxicants can cause a wide variety of effects on F1 females, including infertility and abnormal pregnancy outcomes. Further, exposure to some reproductive toxicants may cause effects on unexposed future generations which are known as transgenerational effects [8].

Plasticizers such as phthalates are major known reproductive toxicants [9]. Phthalates are a colorless, odorless, lipophilic family of synthetic chemicals that are used as plasticizers to increase flexibility, strength, and stability in plastics [7,10,11]. Phthalates are classified as endocrine disrupting chemicals in both male and female reproductive systems [12,13]. Billions of tons of phthalates are produced annually worldwide, of which di-(2-ethylhexyl) phthalate (DEHP) is one of the most widely produced phthalates [14,15]. DEHP is predominately used in various building materials, medical devices, common household items, pesticides, and lubricants. On average, the United States produces nearly 300 million pounds of phthalates including DEHP each year [16]. Humans are readily exposed to DEHP because of its ability to migrate through the materials and leach out into the environment. Thus, people are continuously exposed to DEHP via ingestion, inhalation, and dermal absorption [15]. Indeed, the continuous exposure of DEHP has been evidenced by the fact that 100% of tested human urine samples contain DEHP and its metabolites [17]. The estimated volume of daily human exposure to DEHP is 3-30 µg/kg/day [18]. Furthermore, DEHP and its metabolites have been identified as top contaminants in various tissues, such as ovarian follicular fluid, blood, umbilical cord blood, amniotic fluid, and breast milk [19–22].
DEHP exposure during pregnancy can adversely affect fetal oocyte meiosis. Meiotic defects can lead to infertility and birth defects. Previous studies have shown that DEHP exposure is linked to the decreased synthesis of estradiol and a delay in the meiosis I progression in fetal oocytes [23]. Further, fetal ovaries exposed to DEHP in-vitro inhibited meiotic progression from pachytene to diplotene stage of prophase I [24]. DEHP impairs the repair of DNA double-stranded breaks and causes a delay of oogenesis via a checkpoint at pachynema [24]. Although previous studies show that in-vitro exposure of DEHP inhibits meiotic progression in ovarian tissue culture, they have not assessed the effects of prenatal DEHP exposure at an environmentally relevant dose in-vivo. Thus, I tested the hypothesis that prenatal exposure to DEHP can disrupt female meiosis in-vivo.

In addition to being exposed to environmental chemicals such as DEHP, pregnant women often consume high-fat foods. Maternal consumption of high-fat diet resulted in fetal oocyte reduction and impaired follicle growth in rats [25]. Interestingly, the abnormal spindle formation induced by an obesogenic diet could still be observed in mouse oocytes even after reversing the obesogenic diet back to a regular diet [26]. However, the synergistic effects of DEHP + high-fat diet on fetal reproductive systems have not been studied. Thus, I investigated whether the combination of DEHP exposure and consumption of a high-fat diet can disrupt the female fetal meiosis and follicle development. We exposed pregnant mice to either vehicle or DEHP (20µg/kg body weight per day) starting from 10.5 days post coitum (dpc). The dams were treated with control diet or high-fat diet 7 days before they were paired for mating. We found that DEHP + high-fat diet synergistically disrupts the homologous chromosome synapsis in fetal oocytes and affects the development of follicles in the F1 offspring.
CHAPTER 2

Materials and Methods:

Reagents

A stock solution of DEHP (99% purity, Sigma-Aldrich) was prepared using tocopherol-stripped corn oil (MP Bio Medicals) as the vehicle. The dose of DEHP selected in this study is 20 μg/kg/day. We used this dose because it is within the range of human exposure (3-30 μg/kg/day) and because the dose adversely affects ovarian morphology and function as well as reduces female fertility [27,28].

Animals

CD-1 mice (Charles River Laboratories, Wilmington, MA) were used in this study and housed in the Animal Care Facility at the University of Illinois Urbana-Champaign (UIUC). Animal handling and procedures were approved by the UIUC Institutional Animal Care and Use Committee. Mice were housed under 12 h dark/12 h light cycles. Both high-fat food (D12451; 45 kcal% fat, 20 kcal% protein, 35 kcal% carbohydrate) and control food (D12450B; 10 kcal% fat, 20 kcal% protein, 70 kcal% carbohydrate) were purchased from Research Diets, Inc., New Brunswick, NJ, USA. In these experiments, dams were treated with vehicle control + control diet (n = 5), 20 μg/kg/day DEHP + control diet (n = 4), vehicle control + high-fat diet (n = 4), or 20 μg/kg/day DEHP + high-fat diet (n = 5). The body weights of the female mice were measured before starting the special diet. Female mice were paired with unexposed males to mate and checked regularly for the presence of a copulatory plug, and the day when the copulatory plug was detected was considered 0.5 dpc. The body weight of female mice was recorded when a
copulatory plug was observed (Fig. 9). All female mice were given access to either control diet or high-fat diet starting 7 days before the mating day. The food consumption of F0 females was measured from 0 dpc until 18.5 dpc (Fig. 10). Pregnant female mice were dosed with vehicle control or 20 µg/kg/day DEHP, starting at 10.5 dpc to the day of birth. The body weights of females were measured at 10.5 dpc as well (Fig. 9). Treatments were continued until dams were euthanized by CO₂ inhalation at 18.5 dpc or until the pups were born. Fetal ovaries were collected from the F1 fetus at 18.5 dpc and used to prepare oocyte chromosomal spreads as described below. The dams that give birth continued to have access to their special diet (control diet or high-fat diet) until the F1 pups were weaned at 21 days postpartum (dpp). Body weights of F1 pups were measured at 8 dpp (Fig. 11).

Chromosomal Spreads and Immunofluorescent Staining

To prepare chromosomal spreads, pregnant dams (n= 4-5 dams/ treatment) were euthanized at 18.5 dpc. The female pups were removed from the dams, and their ovaries were collected and placed in phosphate-buffered saline (PBS). Hypotonic extraction buffer (HEB) was used to incubate the ovaries for 20 minutes. Each ovary was placed in 10 µl of sucrose on a depression slide and then minced with scalpel blades. Then, an additional 20 µl of sucrose was added and mixed to make a cell suspension. The cell suspension (6 µl) was added into 30 µl of 1% paraformaldehyde (PFA) with 0.1% Triton X-100 on a microscope slide where the oocytes were spread out. At least 2 different slides were made using one ovary from a fetus. Each slide contained a considerable number of oocyte nuclei. The slides containing the spread oocyte chromosomes were first subjected to blocking by adding antibody dilution buffer (ADB) (0.3% BSA, 10% normal goat serum, and 0.005% Triton-X-100 in TBS) twice (15 minutes each). The
blocked slides were incubated in two primary antibodies, rabbit anti-SYCP3 (Santa Cruz Biotechnology, sc-33195) at a dilution of 1:300 and mouse anti γH2AX (Millipore-Sigma, Cat. # 05-636)) at a dilution of 1:500 overnight. The slides were further blocked twice with ADB. The slides were subsequently incubated 1 hour at 37°C in a pair of goat secondary antibodies: anti-rabbit 488 (Molecular Probes, A11070; 1:1,000 dilution) and anti-mouse 594 (Molecular Probes, A11020; 1:1,000 dilution). After the immunostaining of meiotic cells on the glass slides, images were taken with a Nikon A1R confocal microscope and processed using NIS-Elements software. The total number of oocytes varies in each treatment group because F0 dams often do not have the same number of female fetuses. Also from each fetus, we imaged a different number of oocyte nuclei. A total number of 232 oocytes were imaged from Corn Oil + Control Diet group. A total number of 178 oocytes were imaged from the Control Diet + DEHP group. A total number of 151 oocytes were imaged from Corn Oil + High-fat Diet group. A total number of 351 oocytes were imaged from High-fat Diet + DEHP group.

**Ovarian Follicle Counts in F1 Females**

The ovaries were collected from 8 dpp and 21 dpp F1 females and fixed in Dietrich’s fixative. The fixed ovaries were embedded in paraffin wax, the 8 dpp ovaries were sectioned at 5 µm, and the 21 dpp ovaries were sectioned at 8 µm. Sections were mounted on glass slides and stained with hematoxylin and eosin. EVOS™ XL Core Imaging System was used to image every 5th section of the 8 dpp ovaries, and the images were used to count primordial, primary, and preantral follicles in a blind fashion by two individuals. Every 10th section of the 21 dpp ovaries was directly used to count the number of primordial, primary, preantral, and antral follicles under the microscope. The proportion of each follicle type was calculated by dividing the number of
each type of follicles from the total number of follicles per ovary. The following criteria were used to classify each type of follicle: primordial follicles contained a single layer of squamous granulosa cells, and an oocyte can be observed within the granulosa cell circle; primary follicles contained an oocyte with a single surrounding layer of cuboidal granulosa cells; preantral follicles had at least two layers of cuboidal granulosa cells and theca cell layers with an oocyte inside the cell layers; and antral follicles contained multiple layers of cuboidal granulosa cells with fluid-filled antral space(s), theca cell layers and an oocyte inside the cell layers [29,30]. Preantral and antral follicles without a clear oocyte nucleus were not counted. The follicle counting was conducted without the knowledge of the treatment groups.

_Fertility Tests of F1 Females_

We assessed the mating index, pregnancy rate, and gestational index of female F1 pups from each treatment group at 90 dpp. F1 female mice were mated with unexposed CD-1 male mice and monitored for the presence of copulatory vaginal sperm plug. After confirming the presence of copulatory vaginal sperm plug, the females were separated from the male, weighed, and individually housed. These female mice were weighed twice a week to confirm a successful pregnancy. We calculated the mating and gestational indexes as well as the pregnancy rate by using the following equations [31]:

Mating index = number of females with copulatory plugs/number of total females × 100
Pregnancy rate = number of pregnant females/number of total females × 100
Gestational index = number of females who delivered/number of pregnant females × 100
Gene Expression Analysis

Total RNA was isolated from 21 dpp F1 ovaries (3 ovaries from 3 different animals per treatment) using the RNeasy Micro Kit (Qiagen, Inc., Valencia, CA). Isolated RNA was reverse transcribed to complementary DNA (cDNA) using the SuperScriptIII kit based on the manufacturer’s protocol. The mRNA expression levels of genes encoding for enzymes in estradiol biosynthetic pathway were analyzed for each treatment group via quantitative real-time polymerase chain reaction (qPCR) using SsoFast™EvaGreen® Supermixes according to manufacturer’s protocol (BioRad). A standard curve was generated to calculate the efficiency of each primer set using six serial dilutions of a sample representing the treatment groups. The gene expression data obtained from each sample were normalized to the corresponding values of housekeeping gene beta-actin (Actb). Beta-actin was chosen as the reference gene because this housekeeping gene is typically used for normalizing gene expression data in mouse studies [32,33] and the expression of beta-actin did not differ among treatments. The genes we tested, participate in the estradiol biosynthetic pathway or the synthesis of estradiol precursor in the ovary (Fig. 7) [34]. The genes tested include: steroidogenic acute regulatory protein (Star), 17α-hydroxylase-17,20-desmolase (Cyp17a1), aromatase (Cyp19a1), cytochrome P450 cholesterol side-chain cleavage enzyme (Cyp11a1), 17β-hydroxysteroid dehydrogenase 1 (Hsd17b1), and 3β-hydroxysteroid dehydrogenase 1 (Hsd3b1). The qPCR primer sequences of the tested genes are listed in Table 1. The Pfaffl method was used to calculate individual relative fold changes [35]. The relative mRNA expression level was expressed as the mean fold change ± SD from 3 independent samples.
**Statistical Analysis**

Differences among groups were statistically analyzed by unpaired two-sample independent t-test or one-way analysis of variance (ANOVA). Mating and gestational indexes and the pregnancy rate were analyzed by using 1-tailed Fisher’s exact test. Mann Whitney U test was used to analyze the differences among groups when normality cannot be assumed. Raw data were used to analyze follicle counts at 8 dpp and 21 dpp statistically. All tests were conducted using GraphPad Prism 7 analysis software. Comparisons were considered significant at *P<0.05.*
CHAPTER 3

Results:

*Exposure to DEHP + high-fat diet increases meiotic silencing of unsynapsed chromatin in fetal oocytes*

During meiotic prophase I, homologous chromosomes pair up along their length and are stabilized via synapsis [the assembly of a proteinaceous complex called synaptonemal complex (SC)]. The process of synapsis and desynapsis can divide prophase I into several substages: leptonema, zygonema, pachynema, and diplonema/diakinesis. The axial elements of the SC are composed of cohesin proteins and meiosis-specific proteins, such as synaptonemal complex protein 3 (SYCP3) [36]. Immunostaining of oocyte chromosomal spreads for SYCP3 can be used to stage meiotic prophase I [37]. To further assess the condition of meiotic chromosomes, immunostaining of gamma-histone 2AX (γH2AX) can be used. γH2AX represents DNA damage signaling that detects DNA double-strand breaks and asynapsis of homologous chromosomes [38,39]. Asynapsis of autosomal chromosomes can be identified as γH2AX-rich chromatin domains known as meiotic silencing of unsynapsed chromatin (MSUC) [40,41].

We found that some oocytes have large sections of γH2AX staining, resembling the γH2AX staining of MSUC in the control group and treatment groups (Fig. 1). Analysis of MSUC revealed that at least 20% of the fetal oocytes contained MSUC in the three treatment groups: A) the group dosed with corn oil and fed with control diet; B) the group dosed with DEHP and treated with control diet; C) the group dosed with corn oil and treated with high-fat diet (Fig. 2A). However, approximately 40% of fetal oocytes from dams treated with DEHP + high-fat diet contained MSUC. The data from the control group, DEHP only group, and the high-fat diet only
group were further combined to form a “combined control” group to compare with the DEHP + high-fat diet group (Fig. 2B). F-test for the equality of variances was used to verify that the variances of the three groups did not differ significantly. Counted cells from each group were treated as identical after verification of equality of means and variances, and the combined total cells were used as the “sample” for the combined control. Oocytes in the DEHP + high-fat group contained a significantly higher percentage of cells with MSUC compared to the combined control group. The increased percentage of cells with MSUC held true for DEHP + high-fat group oocytes at pachynema, early diplonema, and late diplonema. These results indicate that the exposure of pregnant dams to a high-fat diet coupled with DEHP results in significant disruption of synapsis.

**DEHP + high-fat diet treatment does not alter the follicle number of F1 pups at 8 dpp**

Asynapsis-induced gene silencing in MSUC regions could result in oocyte loss [42]. To examine whether the elevated MSUC found in DEHP + high-fat diet treated group led to a smaller oocyte pool compared to other three treatment groups, we counted the numbers of primordial, primary, and preantral follicles at 8 dpp. Ovaries in all four treatment groups contained similar numbers of primordial, primary, and preantral follicles at 8 dpp (Fig. 3C). Calculating the proportion of each follicle type in the ovaries of all four groups yielded no significant differences (Fig. 3D). These results indicate that the increased MSUC in DEHP + high-fat diet group did not result in significant oocyte loss.
**DEHP + high-fat diet treatment affected the preantral follicle development in F1 ovaries at 21 dpp**

Ovaries in all four treatment groups at 21 dpp contained primordial, primary, preantral, and antral follicles, with the majority of follicles being at the primordial stage (Fig. 5A). The proportion of each follicle type present in the ovaries did not vary among the four groups (Fig. 5B). However, ovaries of F1 pups treated with DEHP + high-fat diet had a significantly higher number of preantral follicles than the ovaries of F1 pups in the other three groups at 21 dpp (Fig. 4D). These results suggest that the exposure to DEHP + high-fat diet affects preantral follicle development in F1 ovaries at 21 dpp.

**DEHP + high-fat diet does not alter the expression levels of major steroidogenic enzymes in 21-dpp ovaries**

Preantral and antral follicles are steroidogenically active. The increase of preantral follicle number in DEHP + high-fat diet ovaries may elevate the expression levels of estradiol and its precursor steroid hormones in the ovaries with the combined treatment compared to the controls. Thus, we examined the gene expression of the steroidogenic enzymes in 21 dpp ovaries (Table 1). We focused on these genes because previous studies have shown that DEHP and its primary metabolite, MEHP, alters expression of these genes in the estradiol biosynthetic pathway [19,43–45]. However, we did not observe any significant differences in the expression of the tested genes among the four treatment groups (Fig. 6). Our results differ from the previous studies because the amount of MEHP reaching the ovaries may be different between our study and the previous studies.
Female F1 pups exposed to DEHP and high-fat diet do not experience a decline in the mating index, pregnancy rate, and gestational index at 90 dpp

The disrupted oogenesis and altered folliculogenesis caused by DEHP + high-fat diet led us to test several fertility-related indices. None of the treatments affected mating index compared to the control group. In fact, the mating index in groups ranged from 83.33% to 100% (Fig. 8A). Similarly, none of the treatments affected pregnancy rates and gestational indices of F1 females in each group. F1 dams from all four groups became pregnant and gave birth successfully (Figs. 8B and 8C). Litter sizes among the F1 pups also did not differ significantly (p>0.40), with each mother giving birth to 10-15 pups on average (Fig. 8D). These data suggest that the effects of DEHP + high-fat diet on F1 oocytes during meiosis ultimately do not impact the fertility of F1 females at an early age.
CHAPTER 4

Discussion:

The purpose of this study was to determine the synergistic effects of high-fat consumption and environmentally relevant DEHP exposure on female fetal meiosis. Although a previous in-vitro study has shown that prenatal DEHP exposure (10 µM and 100 µM) results in a delay of meiotic progression and disruption of DNA damage repair [24], studies have not examined the effect of DEHP on meiosis in-vivo, and they have not examined high-fat diet as a combined factor. In our in-vivo study, prenatal DEHP exposure alone did not cause a noticeable delay in meiotic progression of F1 oocytes from DEHP-exposed dams. Our results differ from this in-vitro study probably because we used a different dose of DEHP (20 µg/kg/day), and because the DEHP exposure is acute in their in-vitro study while our in-vivo study carried out chronic DEHP treatment. In our in-vivo study, DEHP exposure is indirect to the fetal ovaries while DEHP exposure is direct in the in-vitro study [24]. Since DEHP is metabolized to MEHP in animals’ GI tract, it is possible that more MEHP reached fetal ovaries than DEHP in the in-vivo study. The bioactivity differences between DEHP and its metabolite MEHP may explain why we cannot recapitulate the meiotic arrest found by Liu et al. (2017) [24].

In this study, we also observed that the exposure of dams to a high-fat diet (45% calories from fat, relevant to a western diet) [46] and environmentally-relevant levels of DEHP results in disruption of synapsis during prophase I of meiosis, as evidenced by a significantly high number of γH2AX stained MSUC in the fetal oocytes. Due to the failure of synapsis, the chromatin in the unsynapsed region may trigger MSUC, which is known to silence genes [41,47]. The gene silencing caused by MSUC can trigger meiotic checkpoints and induce germ cell death [39].
Thus, if the majority of the oocytes in the ovary experience asynapsis, increased levels of MSUC may potentially affect the size of the oocyte pool by eliminating the affected oocytes via the DNA damage checkpoint [48]. This may lead to decreased reproductive lifespan in affected individuals [49].

To investigate later stage oocyte development and to examine if the presence of MSUC is associated with increased oocyte death, we evaluated folliculogenesis by counting the total number of follicles in F1 ovaries at 8 dpp and 21 dpp. Previous studies in adult mice have reported that acute exposure to 200 mg/kg/day and 500 mg/kg/day DEHP for 10 days resulted in decreased total follicle numbers after 9 months [16]. Surprisingly, even though we observed increased levels of MSUC from the DEHP + high-fat diet group (Fig. 2A), we did not see a decrease in the size of oocyte pool at both 8 dpp and 21 dpp (Figs. 3C, 5A). A size reduction of the oocyte pool and infertility are rarely observed unless most oocytes in the ovaries are extensively affected [49]. If the damage induced by DEHP + high-fat diet treatment is not severe enough to kill the defective oocytes, the oocyte pool reduction compared to control mice may not be observed.

Interestingly, when we counted the number of each follicle type (primordial, primary, preantral, and antral follicles) in F1 ovaries, we observed an increase in the average number of preantral follicles in the high-fat diet + DEHP-treated group at 21 dpp. This result demonstrates the synergistic effect of DEHP + high-fat diet on folliculogenesis. This result correlates with a previous study where the 21 dpp preantral follicle counts increased in two in-utero DEHP exposed groups (200 µg/kg/day and 500 mg/kg/day) [50]. It is possible that the combination of DEHP and high-fat diet allows more DEHP and its metabolites to reach the F1 fetal ovaries, thus producing the same phenotype despite the lower concentrations. The interaction between DEHP
and a high-fat diet may be explained by the lipophilicity of DEHP [51]. Consumption of a high-fat diet during the administration of DEHP may facilitate the biotransformation of this chemical.

DEHP and its major metabolite, MEHP, are both endocrine disruptors [52]. Therefore, to investigate whether DEHP affects steroidogenesis and folliculogenesis through the disruption of the endocrine system, we conducted qPCR analysis to measure the relative expression of various genes in estradiol biosynthetic pathway. Our results did not show any significant differences in expression of the tested genes (Fig. 6), suggesting that the higher number of preantral follicles may not have been caused by disruption of the estradiol biosynthetic pathway, at least at the level of gene expression.

It is still unclear why the preantral follicle number significantly increased in F1 ovaries at 21 dpp. It is possible that the combination of DEHP and high-fat diet accelerates the maturation of primordial and primary follicles into preantral follicles. However, we did not observe a corresponding decrease in the number of primordial and primary follicles in DEHP + high-fat diet treated ovaries. It is possible that accelerated follicle maturation does not significantly decrease the number of primordial and primary follicles due to the higher number of these follicles as compared to preantral and antral follicles. Another potential explanation for our findings is that prenatal exposure to DEHP + high-fat diet inhibits the natural follicular atresia of preantral follicles in the ovaries.

To analyze the effects of DEHP + high-fat diet on the reproductive capacity of the F1 generation, we recorded the mating and gestational indices, litter size, and pregnancy rate of 90 dpp F1 female mice. In a previous study in F1 female mice, prenatal exposure to 20 µg/kg/day and 200 µg/kg/day of DEHP resulted in slightly lower pregnancy rate without significant
differences compared to the controls [27]. In our study, no significant differences were observed in any of the four treatment groups. This suggests that the synapsis disruption observed in defective oocytes were likely screened and removed by perinatal and/or later-stage checkpoints [30].

Future studies should investigate the effects of higher levels of occupational [53] and medical [54] DEHP exposure and/or consumption of fat during pregnancy that is above the levels we have tested, since high levels of DEHP and high-fat diet exposure may still have the potential to cause adverse effects on the fertility of F1 generation. Since phthalate exposure is unavoidable and present worldwide, this study stresses the importance and need for future studies to elucidate the effects of phthalates during prenatal, postnatal, and multigenerational development, especially at higher doses.

A plausible mechanism of the action of DEHP + high-fat diet on meiosis

Since DEHP is a highly lipophilic molecule [55], there is a possibility that more DEHP and its metabolites reached the fetal ovaries when DEHP is consumed with fat. The observed MSUC in DEHP + high-fat diet exposed mice indicates synaptic defects in homologous chromosomes. These synaptic defects may have been caused by DEHP and its metabolites. The localization studies of estrogen receptor β (ERβ/ERS2) suggest expression of ERβ/ESR2 in premeiotic germ cells as well as in prophase gonads [56–58]. Also, it has been reported that mutation of ERβ/Esr2 gene disrupts synapsis in mouse oocytes [59]. Therefore, it is possible that DEHP and/or its metabolites may exert the observed meiotic synaptic defects by interfering with ERβ/ESR2 by
acting as an estrogen antagonist or an estrogen agonist and affecting estrogen receptor β signaling pathways.
CHAPTER 5

Conclusions:

This study provides evidence that prenatal exposure to a combination of 20 µg/kg/day DEHP and high-fat diet relevant to a western diet affects synapsis during meiotic prophase I of F1 female mice. Also, this combination affects the development of preantral follicles of F1 females at 21 dpp. However, prenatal exposure to DEHP + high-fat diet combination does not affect mating index, pregnancy rate, gestational index, and litter sizes of F1 females. The resting oocyte pools in both treated and untreated ovaries are large, which might lead to the non-significant results from fertility studies. The mechanisms behind the effects of DEHP + high-fat diet are still unknown, and further studies are required to understand these mechanisms.
## TABLE 1

Sequences of primer sets used for gene expression analyses

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Figure 1. Representative images for fetal oocytes at the pachynema (A), early diplonema (B), and late diplonema (C) containing MSUC (arrows). Scale bars = 10 µm.
Figure 2. (A) Percentage of pachytene, early diplotene, and late diplotene fetal oocytes with MSUC in A group (Control Diet + Corn Oil) (n = 232 oocytes), B group (Control Diet + DEHP) (n = 178 oocytes), C group (High-fat Diet + Corn Oil) (n = 151 oocytes), and D group (High-fat Diet + DEHP) (n = 351 oocytes). Differences within treatment groups in % MSUC at each stage were evaluated using the A group as a reference (*p<0.05, ***p<0.001). (B) Comparison of MSUC percentage of D group with combined A, B, and C controls (*p<0.05). Graphs represent means ± SEM.
Figure 3: (A) 8 dpp F1 ovarian sections in A group (Control Diet + Corn Oil) and (B) D group (High-fat Diet + DEHP) stained with hematoxylin and eosin. Some primordial (arrows), primary (arrowheads), and preantral (asterisks) follicles have been demarcated. (C) Counts of primordial, primary, and preantral follicles for each treatment group at 8 dpp (n = 3 dams/treatment group) (D) Follicular progression of 8 dpp ovaries, as measured by the proportions of primordial, primary, and preantral follicles in each treatment group. Proportions were gathered separately for each ovary. Graphs represent means ± SD.
Figure 4: 21 dpp F1 ovarian sections stained with hematoxylin and eosin in four treatment groups: (A) Control Diet + Corn Oil, (B) Control Diet + DEHP, (C) High-fat Diet + Corn Oil, and (D) High-fat Diet + DEHP. Some antral follicles (arrows) and preantral follicles (asterisks) have been demarcated.
Figure 5: (A) Counts of primordial, primary, and preantral follicles for each treatment group at 21 dpp (*p<0.05) (n = 4 dams/treatment group). (B) Follicular progression of 21 dpp ovaries, as measured by the proportions of primordial, primary, preantral, and antral follicles in each treatment group. Proportions were gathered separately for each ovary. Graphs represent means ± SD.
Figure 6. DEHP and high-fat diet do not alter the mRNA expression levels of steroidogenic enzymes in 21 dpp F1 ovaries. After each treatment, ovaries were collected and subjected to qPCR analysis for (A) StAR, (B) Cyp17a1, (C) Cyp19a1, (D) Cyp11a1, (E) Hsd17b1, and (F) Hsd3b1 mRNA expression levels. All gene expression values were normalized to the expression of β-actin (n = 3). Graphs represent means ± SD.
Figure 7. Steroid-hormone biosynthetic pathways in the ovary. Theca cells and granulosa cells in antral follicles participate in steroidogenesis. Metabolism of cholesterol to steroid hormones is regulated by steroidogenic enzymes STAR, CYP11A1, CYP17A1, CYP19A1, HSD3B1, and HSD17B1.
Figure 8. (A) Mating indices of female F1 pups at 90 dpp in the Control Diet + Corn Oil, Control Diet + DEHP, High-fat Diet + Corn Oil, and High-fat Diet + DEHP groups. (B) Pregnancy rates of female F1 pups at 90 dpp in the Control Diet + Corn Oil, Control Diet + DEHP, High-fat Diet + Corn Oil, and High-fat Diet + DEHP groups. (C) Gestational indices of female F1 pups at 90 dpp in the Control Diet + Corn Oil, Control Diet + DEHP, High-fat Diet + Corn Oil, and High-fat Diet + DEHP groups. (D) Average litter size of F1 dams in the Control Diet + Corn Oil, Control Diet + DEHP, High-fat Diet + Corn Oil and High-fat Diet + DEHP groups. Control Diet + Corn Oil (n = 5 F1 females), Control Diet + DEHP (n = 7 F1 females), High-fat Diet + Corn Oil (n = 7 F1 females), and High-fat Diet + DEHP (n = 7 F1 females). Graphs represent means ± SD.
Figure 9. Body weight of F0 females at the initiation of the diet treatment, at 0 dpc and at 10.5 dpc. Control Diet + Corn Oil (n = 5 F0 females), Control Diet + DEHP (n = 4 F0 females), High-fat Diet + Corn Oil (n = 4 F0 females), and High-fat Diet + DEHP (n = 5 F0 females). Graph represents means ± SD.
Figure 10. Food consumption of F0 females from 0 dpc to 18.5 dpc. Control Diet + Corn Oil (n = 5 F0 females), Control Diet + DEHP (n = 4 F0 females), High-fat Diet + Corn Oil (n = 4 F0 females), and High-fat Diet + DEHP (n = 5 F0 females). Graph represents means ± SD.
Figure 11. Body weight of F1 pups at 8 dpp. Control Diet + Corn Oil (n = 10, F1 pups), Control Diet + DEHP (n = 15, F1 pups), High-fat Diet + Corn Oil (n = 8, F1 pups), and High-fat Diet + DEHP (n = 14, F1 pups). Graph represents means ± SD.
References:


