**Gestational bisphenol A exposure and testis development**

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**Abbreviations:** BPA, bisphenol A; DES, diethylstilbestrol; EDC, endocrine disrupting chemical; ERα, estrogen receptor alpha; ERβ, estrogen receptor beta; ERE, estrogen-response element

Virtually all humans are exposed to bisphenol A (BPA). Since BPA can act as a ligand for estrogen receptors, potential hazardous effects of BPA should be evaluated in the context of endogenous estrogenic hormones. Because estrogens are metabolized in the placenta, developing fetuses are normally exposed to very low endogenous estrogen levels. BPA, on the other hand, passes through the placenta and might have distinct adverse consequences during the sensitive stages of fetal development. Testicular gametogenesis and steroidogenesis begin early during fetal development. These processes are sensitive to estrogens and play a role in determining the number of germ stem cells, sperm count, and male hormone levels in adulthood. Although studies have shown a correlation between BPA exposure and perturbed reproduction, a clear consensus has yet to be established as to whether current human gestational BPA exposure results in direct adverse effects on male genital development and reproduction. However, studies in animals and in vitro have provided direct evidence for the ability of BPA exposure to influence male reproductive development. This review discusses the current knowledge of potential effects of BPA exposure on human health, in particular, on male reproductive health and whether gestational exposure adversely affects testis development.

**Introduction**

Testicular dysgenesis syndrome (TDS) includes hypospadias, combined with cryptorchidism, impaired spermatogenesis, and an increased risk of testicular germ cell cancer. TDS is often present in males with rare genetic abnormalities such as 45 X0 or 46 XY karyotypes. However, over the past 50 y the incidence of individuals with symptoms of TDS has been reported to be rapidly increasing in the general population (for a review, see ref. 1). One hypothesis that accounts for the increasing incidence of TDS is exposure to environmental compounds that disturb estrogen or androgen signaling during critical developmental periods.

It is known from work in experimental animals that exposure to high concentrations of estrogen during fetal development can result in phenotypes resembling TDS. In humans, gestational exposure to the estrogenic compound diethylstilbestrol (DES) results in abnormalities of the male and female reproductive tract along with infertility in women and possibly in men (for a review, see ref. 2). Estrogenic compounds are present in the environment and are likely to result in human exposure. In water and river sediments, biologically relevant concentrations of both natural and synthetic estrogens, such as bisphenol A (BPA), are readily detectable.3 This review discusses the current knowledge of the potential effects of BPA exposure on human health, in particular, on male reproductive health and whether gestational exposure adversely affects testis development.

**BPA Exposure**

BPA is one of the most highly produced chemicals worldwide. It was first synthesized in 1891 and its estrogenic capacity was defined by experiments in rats in 1938.4 It was proposed for clinical use as an estrogen, however, the more potent estrogenic compound, DES, was synthesized that same year and BPA was never used clinically. Instead, polymerized BPA was found to be useful for manufacturing polycarbonate plastics and epoxy resins. BPA has been in commercial use as such since 1937, and its production has been reported to be rapidly increasing in the general population (for a review, see ref. 1). One hypothesis that accounts for the increasing incidence of TDS is exposure to environmental compounds that disturb estrogen or androgen signaling during critical developmental periods.

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While polymeric BPA lacks estrogenic activity, the ester bond that links BPA monomers together is not stable and monomers are released with time. Monomeric BPA can then be absorbed from the plastic containers or cans into foods and drinks consumed by humans.6 Another major exposure route is dermal exposure to BPA from thermal papers used for receipts.7 Daily adult exposure of BPA is estimated to be between 0.05–10 μg/kg body weight per day, and BPA can be consistently detected in the blood, serum, urine, saliva, and other bodily fluids of virtually all humans.6,7 In addition, BPA is spread into the atmosphere.
through the burning of plastics, and is present at detectable concentrations throughout the world, including the polar regions.\textsuperscript{12}

Exposure to BPA starts before birth, as BPA can pass through the placenta.\textsuperscript{13} As BPA can be accumulated in fat tissues,\textsuperscript{14} the developing fetus may be subjected to elevated levels of BPA as maternal lipid stores are metabolized during pregnancy releasing deposited BPA. BPA is also secreted into breast milk, and formula-fed babies and young children are exposed through food, water, and plastic containers.\textsuperscript{15,16} Based on estimates obtained from food consumption and concentration data, infants have the highest BPA exposure in the general population.\textsuperscript{17} Thus, an urgent question is whether this early estrogenic exposure at sensitive developmental stages has adverse consequences.

### BPA Acts as an Estrogen

The structure of BPA makes it a ligand for estrogen receptor \(\alpha\) (ESR1/ER\(\alpha\)) and estrogen receptor \(\beta\) (ESR2/ER\(\beta\)).\textsuperscript{18,19} ERs regulate gene expression by forming hetero- or homodimers that interact with their cognate DNA estrogen-responsive elements (EREs), or by tethering to other DNA-binding proteins. In addition, BPA like other estrogens elicits “nongenomic” or “membrane-initiated” effects through ERs located in the cytoplasm or membrane, or through activation of the membrane-bound G protein-coupled estrogen receptor 1 (GPER1/GPR30) and subsequent activation of downstream cytoplasmic signaling cascades.\textsuperscript{20} BPA at higher concentrations can also modulate the activity of several other receptors, including the arylhydrocarbon receptor (AHR), all of the peroxisome proliferator-activated receptors (PPARs), the androgen receptor (AR), both thyroid hormone receptors (THRs/TRs), the pregnane X receptor (NR1I2/PXR), and the estrogen-related receptor gamma (ESRRG/ERR\(\gamma\)).\textsuperscript{21-23} Because of BPA’s capacity to interfere with hormone signaling, it is categorized as an endocrine disrupting chemical (EDC).

### Tissue- and Cell-Specific Effects of BPA

Tissue-specific effects of BPA primarily depend on its uptake and expression of the two ERs. The ERs have both cell-specific and receptor-specific actions, and their tissue distribution differs. For example, based on their respective mRNA levels in adult human tissue, ER\(\alpha\) is readily detected in the uterus, ovary, muscle, mammary gland, pituitary gland, spleen, thymus, prostate, adrenal gland, heart, and kidney; while ER\(\beta\) is primarily detected in the testis and lung.\textsuperscript{24} The effect of BPA on a specific tissue is also dependent on its concentration within that tissue. In the zebrafish model, general estrogen-induced activation during early development, based on transgenic ERE-luciferase reporter, is detected in the liver, pancreas, brain, and kidney.\textsuperscript{25} Subsequently, BPA exposure in zebrafish disrupted axonal growth of primary and secondary motoneurons, and activated expression of the liver-specific and estrogen-regulated vitellogenin 1 (\textit{vtg1}) gene.\textsuperscript{26,27} In rat uterus and vagina, both ER\(\alpha\)-expressing tissues, BPA produces physiologic effects identical to those elicited by estradiol.\textsuperscript{28} It is clear that BPA can activate the ERs at relatively low concentrations and that exposure can impact target organs that express either ER.

It has been suggested that BPA may induce BPA-specific target gene expression via ERs, and that its effect in this way would differ from that of other estrogens. In a recent study, we compared BPA-induced gene regulation with that of 17\(\beta\)-estradiol and phytoestrogens in human ER\(\alpha\)-positive MCF-7 breast cancer cells. We found that identical genes were regulated by BPA, phytoestrogens, and 17\(\beta\)-estradiol.\textsuperscript{29} Although MCF-7 cells also express the AR, PPAR\(\alpha\)/\(\gamma\), TR\(\alpha\)/\(\beta\), GPER1, and AHR,\textsuperscript{30-32} we found that the expression of all genes regulated at 24 h BPA treatment were inhibited by co-treatment with the ER\(\alpha\) antagonist ICI.\textsuperscript{29} This suggests that the transcriptional activity elicited by BPA in these cells was mediated through ER\(\alpha\), and that there were no unique BPA target genes that estrogen did not activate. We also demonstrated that BPA and phytoestrogens acted in an additive manner.\textsuperscript{29} The additive effect of BPA and phytoestrogens raises concerns that the feeding of phytoestrogen-rich soy-based formula to infants may enhance the effect of BPA exposure.

### Health Consequences of BPA Exposure

The levels of BPA that humans are exposed to are relatively low, and BPA binds to ERs 100 to 1000-fold less efficiently than 17\(\beta\)-estradiol.\textsuperscript{19,29} Thus, in theory, BPA should have negligible human health effects at the current exposure levels. Multiple studies have, however, shown that low level BPA exposure can alter many physiological processes, and epidemiologic evidence supports that BPA is related to altered neuronal and brain development, immune, ovarian, and metabolic diseases (for a recent review, see ref. 33).

One reason why BPA exposure appears to elicit more severe health consequences than predicted from its exposure levels may be due to gestational exposure, as illustrated in Figure 1. Both male and female fetuses have very low endogenous levels of estrogens: maternal 17\(\beta\)-estradiol is metabolized by the placenta, and 17\(\beta\)-estradiol in the fetus is sequestered by fetal \(\alpha\)-fetoprotein and not active. BPA, on the other hand, passes through the placenta and binds only weakly to serum proteins.\textsuperscript{33,34} As a result, even low-level BPA exposure can readily exceed the normal estrogen levels in utero, thereby exposing the developing fetus to greater estrogenic activity than expected. Fetal tissues express both ER\(\alpha\) and/or ER\(\beta\) and in utero exposure to DES, which has a three or four times higher affinity for both ERs than 17\(\beta\)-estradiol itself,\textsuperscript{35} results in adverse effects (reviewed in ref. 2). DES was given to many pregnant women during the 1940s to the 1970s in an effort to reduce the risk of habitual and spontaneous abortion, and to make babies stronger at birth. This caused a 40-fold increased risk of rare vaginal/cervical clear cell adenocarcinoma in girls born to these women, making DES one of the first transplacental carcinogens discovered in humans.\textsuperscript{36} Later studies showed that gestational DES exposure also increased the risk of cervical squamous cell dysplasia and breast cancer in adulthood, along with abnormalities of the reproductive tract and infertility.\textsuperscript{37,38}
Abnormal activation of ERs by BPA during pre-natal and early development may have similar adverse consequences that, even if small, are potentially problematic given the large population numbers that are exposed. In animals, maternal exposure to BPA (2.4–20 μg/kg) accelerates weight gain and puberty in females along with altered development of the mammary gland and uterus. Developmental BPA exposure has also been shown to result in stable epigenetic changes in offspring. The increased level of BPA exposure in industrial countries coincides with a lowering of semen quality and an increase in male reproductive disorders. Moreover, three reports describe a significant correlation between BPA levels in men and lower sperm quality measures (for a recent review, see ref. 33), and one study has linked parental BPA exposure to shorter anogenital distance in sons. These associations have led to concerns that BPA exposure may affect male reproductive health.

**Estrogen Signaling Impacts Testis Development**

Normal development of the testis is critical for male reproduction. The testis begins gametogenesis and steroidogenesis during early fetal development, and these processes play a role in determining the number of germ stem cells, sperm count, and male hormone levels in adulthood. Testis development is in fact very sensitive to estrogens: ERβ mRNA is detected in mouse gonocytes as early as 14 d post-conception and disappears by post-natal day 26, while ERα is expressed in the interstitial compartment of the testis, as illustrated in Figure 2. Inactivation of ERβ in the mouse leads to a 50% increase in the numbers of gonocytes at 2 and 6 d after birth, with no change in Sertoli cell number or Leydig cell number. Inactivation of the ERα increases testosterone production as early as 13.5 dpc.

Men exposed to DES before birth have an increased risk for non-cancerous epididymal cysts, fibrotic testis, and undescended testicles (cryptorchidism), along with an increased risk of hypogonadism and low testosterone levels. Early studies found decreased fertility and lower sperm count in men exposed to DES in utero, but a 40 y follow-up study found no increased risk of infertility. A later analysis, however, indicated that DES may have a negative effect on sperm count only if administered during the first trimester of pregnancy. While gestational exposure to potent estrogens has major disadvantages for male reproductive development, BPA is far less potent than DES, and exposure to BPA occurs at significantly lower doses. Although studies have shown a correlation, a consensus does not yet exist as to whether or not current human BPA exposure results in adverse effects on male genital development and reproduction. However, studies in animals have provided direct evidence for the ability of BPA exposure to influence male reproductive development.

**BPA Exposure and Testis Development**

BPA exposure appears to affect male development over a wide range of species and models. In the wild, EDC-induced feminization of male fish has been demonstrated for several species. In animal experiments, male goldfish exposed to BPA exhibit modulation of AR, ER-regulated genes, and testicular responses. In carp, BPA exposure (from 1 μg/L) caused testicular changes, including loss of typical lobular structure, reduction of spermatogenic cysts, and lobule diameter. In mice, high-dose BPA exposure (480 and 960 mg/kg/day at postnatal days 31–44) resulted in underdeveloped testes, apoptotic Leydig and germ cells, and disruption of spermatogenesis. Meanwhile, postnatal exposure to lower BPA doses relevant to human exposure (20 and 40 μg/kg/day from day 3) resulted in slowed meiotic progression of germ cells after 5 wk treatment along with decreased quality and quantity of spermatozoa in 7-wk-old mice. Gestational BPA exposure at higher concentrations have also been shown to affect testis development in rodents: Exposed mice (5 or 50 mg/kg, days 7 and 14 of pregnancy) exhibited lower epididymal sperm counts, sperm with abnormal morphology and reduced mobility, disruption of sperm differentiation, and decreased numbers of Sertoli cells. Lower exposure levels (2 and 20 μg/kg at gestational days 11–17) were found to result in longer anogenital distances. In rats, the testis effects of a mixture of plastic-derived compounds (including 25 mg/kg BPA per day, embryonic days 8 to 14) exhibited transgenerational epigenetic inheritance.
In our previous study, we explored the effects of gestational BPA exposure on testis. The dose, 10 µg/ml in 1% ethanol in the mothers’ drinking water, results in BPA accumulation in mouse tissues comparable with concentrations found in humans. Dams from two mouse strains, C57BL/6 (B6) and SJL/JCrHsd (SJL), were treated with BPA or control from preconception through weaning. We found no significant changes in weight-corrected anogenital distance or in testis wet weight. However, degeneration of the testes, determined by greater levels of germ cell apoptosis compared with baseline, along with a notable loss of germ cells and general disruption of normal maturation, was observed more often in BPA-exposed mice, and was most common in the later VIII–XII stages of spermatogenesis. Our data suggest potential developmentally induced alterations in germ cell maturation, and support the findings from multiple groups that gestational BPA exposure can induce changes in the testis.

Other studies, however, have found no effects of BPA exposure in rodents over a wide range of concentrations and endpoints. The outcome appears to depend on both the experimental design and potential genetic effects. With regard to the latter, neonatal exposure to potent and environmental estrogens, including tamoxifen, has been shown to lead to strain-specific differences in male reproductive organ abnormalities.

Direct BPA Exposure Affects the Function of Testicular Cells

In testicular cells, high-dose BPA exposure (100 µM) can inhibit the activity of 11β-hydroxysteroid dehydrogenases (HSD11b1/11β-HSD) in both humans and rats. In cultured Leydig cells from the mouse fetal testis, BPA exposure (10 µM) had a negative effect on testosterone production and this effect was maintained in the absence of ERα. While BPA concentrations as low as 10 nM affected the function of cultured human fetal Leydig cells, concentrations of 10 µM BPA were required in rodent cells, suggesting that the human testis may be more sensitive to BPA exposure than the testis of mice and rats. In the murine spermatogonial GC-1 cell line (ERβ-negative), indications that low-dose BPA exposure (0.1–10 nM) induce proliferation through ERα and GPER1-signaling have been presented, while another study reported that BPA did not activate the GPER1 pathway in spermatozoa. BPA has also been found to inhibit the activity of ATP-binding cassette, sub-family G (WHITE), member 2, also known as breast cancer-resistance protein (ABCG2/BCRP), a transporter expressed in the blood-testis barrier that transports e.g., steroids.
**Direct and Gestational BPA Exposure Affects Gene Expression in the Testis**

BPA exposure can lead to transcriptional changes in rodent male reproductive organs. Direct BPA exposure of adult rat testes (50 mg/kg, 6 d per week for 8 wk) led to decreased expression of testicular steroidogenic enzymes, along with altered n-6 fatty acid composition and antioxidant enzyme levels.90 Similar results were observed in another study of BPA-exposed adult rats (5 mg/kg/day for 8 wk) where Star and Cyp450ssc (Cyp11a) expression was increased, and Ar, 3β-HSD (Hsd3b), 17β-HSD (Hsd17b), and aromatase (Cyp19a1) expression was decreased.68 In mice, upregulation of Fas, Fasl, and Casp3 expression was noted in the testis after BPA exposure (480 and 960 mg/kg/day).51 Young male mice exposed to BPA in drinking water (50 μg/ml) exhibited changes of both ERα and ERβ expression in testes 8 wk after exposure, and in pooled testis samples of mice treated with higher doses of BPA (50 mg/kg, administered twice) downregulation of Msi1h, Ncoa1, Nid1, Hspb2, and Gata6 were detected using a testis-focused small microarray.55,69 However, the latter changes were not confirmed by qPCR, and variability between individuals was not assessed. Newborn mice exposed to a low concentration of BPA (20 and 40 μg/kg/day from postnatal day 3) showed increased testicular expression of ERα after 5 wk treatment,60 and mice exposed to a mixture containing BPA and phthalates during gestation (1–10 mg BPA/kg/day) exhibited decreased expression levels of anti-Mullerian hormone (Amh), Ar, Cyclin A1 (CcnA1), and Star in the testis.70 Gestational BPA-exposure of rats (0.02, 0.5, 400 mg/kg/day, from gestation day 11) resulted in dose-related decrease of Star in the testis at gestation day 20.21

We investigated the result of gestational and early BPA exposure on gene expression in the adult testis of B6 mice (as described above) using microarray analysis. We found subtle changes in gene expression suggesting dysregulation of cell proliferation, spermatogenesis, and apoptosis, consistent with histopathological analyses and increased levels of germ cell apoptosis.54 We further observed downregulation of platelet-derived growth factor α (Pdgfa), a gene well documented to be critical for proper testes development and function.22 This downregulation was confirmed by qPCR in individual mice, and it aligns with previous reports of BPA and other estrogens stimulating testicular gonocyte proliferation in a PDGF-dependent manner in vitro.73 Previous finding that BPA exposure can alter the expression of receptors for PDGF in neonatal testes indicates the dysregulation of the PDGF-signaling pathway as a general theme in BPA-mediated endocrine disruption of the testes.74

**Discussion**

Combined, current data and literature suggest that BPA exposure can induce histological changes in the testis along with dysregulated proliferation and apoptosis, repression of steroidogenesis, and reduce the number of germ cells, at least in rodents. These effects may involve different molecular pathways including ERα-, ERβ-, and GPER1-mediated pathways, and affect Pdgfa expression and steroidogenesis enzymes.

It’s possible that systemic effects of BPA through the neuro-endocrine system also contribute to testicular degeneration and function. In adult rare minnow *Gobiocypris rarus*, a freshwater teleost, BPA disrupted the gonadotropin-releasing hormone system in the brain.75 Mice exposed to BPA in early life (postnatal day) exhibited altered ERα and ERβ expression in the hypothalamus-pituitary-ovary axis, and BPA exposure during perinatal and postnatal periods resulted in the upregulation of Kiss1, Gnrb1, and follicle stimulating hormone (FSHβ) mRNA in pups.76 It is not clear whether these changes are induced through interaction with ERs in the testes or is a systemic effect originating from BPA affecting the hypothalamus/pituitary axis. Experiments using tissue-specific ERα- and ERβ-knockout animals are required to address these possibilities.

Multiple reports have implicated BPA exposure in a wide variety of physiological abnormalities, and many countries have banned the use of BPA in food packaging.27 Yet, compared with other xenoestrogens, such as DES, there is often a lack of consensus regarding the effects of BPA. This may be in part due to the fact that (1) many different exposure protocols and animal models are used, (2) the exposure concentrations of BPA and time points vary widely from study to study, (3) that some physiological responses to BPA do not always show classical dose dependence, and (4) that responsiveness to both potent and environmental estrogens is genetically controlled.78-83 Further, monomeric form of BPA can react with other molecules and form a number of derivatives. For example, BPA in drinking water can react with chlorine resulting in chlorinated aqueous BPA which has other molecular properties.84 Chlorinated BPA has been detected in human adipose and placental tissues, dichlorinated BPA being the most abundant form.85,86 Also, halogenated analogs of BPA, such as brominated and chlorinated BPAs, are produced today as flame retardants. The highly produced tetrabromobisphenol A (TBBPA) can be dehalogenated by microorganisms in contaminated sediments from streambeds to form BPA.87 The relevance for human exposure of modified BPA molecules is currently unknown. Toxicology testing today is largely performed on one compound at a time. However, the effect of a compound needs to be examined in its environmental context, e.g., in combination with other man-made compounds or natural compounds, and genetic context. There is a concern that different EDCs act in synergy and that the effects seen in humans are a result of the so-called cocktail effect. There is a lack of studies analyzing the specific effects and combined exposure to environmental estrogenic ligands and the genetic control of responsiveness to individual and combined exposures. We recently investigated the impact of combined BPA and phytosterogens exposure and detected additive activation of ER signaling.29 Other studies have shown that mixtures of EDCs produce large effects, even when each chemical is present at doses below when they individually have any measurable effects (reviewed in ref. 88). Furthermore, comparatively little is known about mixtures composed of chemicals from different classes of EDCs acting through different molecular pathways. A recent
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Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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