Does Cancer Start in the Womb? Altered Mammary Gland Development and Predisposition to Breast Cancer due to in Utero Exposure to Endocrine Disruptors

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Abstract We are now witnessing a resurgence of theories of development and carcinogenesis in which the environment is again being accepted as a major player in phenotype determination. Perturbations in the fetal environment predispose an individual to disease that only becomes apparent in adulthood. For example, gestational exposure to diethylstilbestrol resulted in clear cell carcinoma of the vagina and breast cancer. In this review the effects of the endocrine disruptor bisphenol-A (BPA) on mammary development and tumorigenesis in rodents is used as a paradigmatic example of how altered prenatal mammary development may lead to breast cancer in humans who are also widely exposed to it through plastic goods, food and drink packaging, and thermal paper receipts. Changes in the stroma and its extracellular matrix led to altered ductal morphogenesis. Additionally, gestational and lactational exposure to BPA increased the sensitivity of rats and mice to mammaryotropic hormones during puberty and beyond, thus suggesting a plausible explanation for the increased incidence of breast cancer.

Keywords Xenoestrogen · Progesterone receptor · Beaded duct · Ecological developmental biology · Tissue organization field theory · Neoplasia

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Abbreviations

EDC Endocrine disrupting chemical
BPA Bisphenol-A
DES diethylstilbestrol
FDA United States Food and Drug Administration
DMBA dimethylbenzanthracene
DDT dichlorodiphenyltrichloroethane
PCB polychlorinated biphenyl
SMT somatic mutation theory
TOFT tissue organization field theory
bw bodyweight
CDC United States Centers for Disease Control and Prevention
GD Gestational day
E embryonic day
ERs Estrogen receptors
PR progesterone receptor
PND postnatal day
ECM extracellular matrix
TEB terminal end bud
SPARC secreted protein acidic and rich in cysteine
EPA United States Environmental Protection Agency

Introduction

Perturbations in the fetal environment predispose individuals to disease that will only manifest in adulthood [1, 2]. This fact has prompted scientists to hypothesize that fetal exposure to environmental estrogens is an underlying cause of the increased incidence of uterine leiomyomas, testicular cancer and breast cancer observed in European and US populations over the last 50 years [3–5].

Estrogen levels in the fetal environment have long-term consequences regarding the risk of developing breast cancer during adult life [6–8]. Given the long latency period between
exposure and effect, epidemiological studies designed to explore this hypothesis have used prenatal markers of in utero estrogen exposure because direct estrogen measurements are not available from birth records. Dizygotic twin pregnancy, which is associated with high estrogen levels, and pre-eclampsia, which is associated with low levels, were used as surrogates for high and low estrogen exposure, respectively. Dizygotic birth correlated with increased risk of breast cancer in the offspring while pre-eclampsia was associated with lower risk [7, 9, 10]. Direct evidence of a link between prenatal estrogen exposure and breast cancer risk has been gathered from the cohort of women born to mothers treated with the potent synthetic estrogen diethylstilbestrol (DES) during pregnancy. DES was administered to women to prevent miscarriages. In 1971 the Food and Drug Administration (FDA) issued a Drug Bulletin advising physicians to stop prescribing DES to pregnant women because a rare neoplasia, clear cell adenocarcinoma of the vagina, was reported to occur in young women exposed in utero to DES [11, 12].

Breast cancer risk at 40 years of age and older is 2.5 fold higher in DES-exposed women than in unexposed women of the same age [13]. In rats, prenatal exposure to DES also resulted in increased mammary cancer incidence during adulthood when these animals were challenged with the chemical carcinogen dimethylbenzanthracene (DMBA) at puberty [14]. DES was administered to rats at pharmacological doses to mimic its medical use.

In addition to exposures to natural and pharmacological estrogens, there is the inadvertent and continuous exposure of human fetuses to endocrine disrupting chemicals (EDCs) released in the environment. EDCs like the herbicide atrazine, dioxins resulting from incineration and fuel combustion and the surfactant perfluorooctanoic acid delay mammary gland development in rodents as a result of gestational and lactational exposures [15–17]. Exposure to estrogens such as BPA and DES resulted in the long-term increase in the number of epithelial structures and the development of pre-cancerous and cancerous lesions in the mammary glands of rodents that manifested in adulthood [18, 19]. To date, BPA is the best-studied EDC and is the only one for which effects of exposure have been described at multiple time points spanning fetal development and postnatal life.

The Complexity of Endocrine Disruptors

In the Statement of Principles from The Endocrine Society, an EDC is defined as “an exogenous chemical, or mixture of chemicals, that interferes with any aspect of hormone action” [20]. Several EDCs were found to have multiple hormonal activities. For example, the pesticide DDT is an estrogen agonist, while one of its metabolites is anti-androgenic [21]. BPA has estrogenic activity and is a thyroid hormone antagonist [22, 23]. Xenoestrogens are usually less potent than estradiol regarding their binding affinity to nuclear estrogen receptors (ER); effects manifesting at low doses are explained by the fact that they act additively with endogenous estrogens [24]. Xenoestrogens bind to plasma carrier proteins with significantly lower affinities than those of natural estrogens, and thus are more readily available to target organs [25]. Furthermore, some xenoestrogens like BPA produced low-dose effects by acting through ER located in the plasma membrane, including the ERα and β and the estrogen-binding protein GPR30 [26, 27].

Medical Hypotheses Derived from the Concept of Endocrine Disruption

The “EDC hypothesis” was first proposed at the Wingspread Conference held in Racine, Wisconsin, in 1991 to explain the detrimental effects observed in exposed wildlife and the likelihood of harm to humans [28]. The proceedings from this conference [31] prompted medical hypotheses proposing that the increased incidence of uterine leiomyoma, testicular and breast cancer, of malformations of the male genital tract and of the decreased sperm quality observed in European populations over the previous 50 years were due to fetal exposure to EDCs [3, 5, 29].

Fetal Origins of Cancer: Theoretical Foundations

The hypothesis that gestational exposure to EDCs predisposes an individual to cancer, challenged the entrenched beliefs that mammalian development is the execution of a genetic program, and that only mutagenic agents can cause cancer [30, 31]. The theoretical bases of this novel perspective are briefly presented below.

Is Development the Unfolding of a Program or an Open Ended Process?

The adoption of animal models such as Drosophila, Xenopus, and rodents that reproduce all year long in laboratory conditions promoted a concentration on genetics with the exclusion of evolution and ecology from embryology; this approach plus the success of molecular biology led to the notion of a “developmental program” whereby genes had a privileged role on the determination of the phenotype; F. Jacob expressed this concept succinctly: “For it is during embryonic development that the instructions contained in the genetic program of an organism are expressed, that the genotype is converted into phenotype”[32]. The idea of a
“program” is contested by i) gene splicing which challenges the one-to-one correlation between gene, mRNA and protein, a sine qua non of programmability, from gene to phenotype, and ii) stochastic gene expression, which is contrary to a Laplacian determination inherent in the notion of a program [33]. These arguments buttress the relevance of the environment in the expression of phenotypes, which is equally important to those of genes and other resources present in the embryo, and justifies why Medicine is now embracing the old tradition of ecological developmental biology [34].

Carcinogenesis: Development Gone Awry?

For about a century, it has been assumed that carcinogenesis is a cell-based process caused by DNA mutations in a single founder cell and that those mutations cause uncontrolled cell proliferation [35]. These are the tenets of the somatic mutation theory (SMT) [36]. When it became evident that non-mutagenic agents also cause cancer, a “course correction” was proposed whereby changes in the epigenome play a central role through dysregulated cell proliferation [30, 37]. An alternative theory of carcinogenesis, i.e., the tissue organization field theory (TOFT), instead, places carcinogenesis at the tissue level of organization, a concept spanning embryology and pathology [38]. The TOFT posits that the persistence of morphogenetic fields throughout adult life orchestrate histogenesis and organogenesis before birth and tissue remodeling and regeneration during postnatal life [39]. From this perspective, carcinogens cause neoplastic development by altering the reciprocal interactions between the mesenchyme/stroma and the parenchyma in a tissue or organ.

Experiments were conducted to resolve the controversy of whether the target of the carcinogen was a cell in the rat mammary gland epithelium or the stroma of the gland [40]. Recombination of stroma exposed to a carcinogen with normal, unexposed epithelial cells resulted in neoplasms in the epithelium [40]. The reverse combination did not. This observation suggests that the stroma, rather than individual cells in the epithelium, is the target of carcinogens. Moreover, the cancer phenotype was reversed when mammary cancer epithelial cells were placed into a normal, unexposed mammary stroma [41]. Equally significant, normal mammary epithelial cells can also contribute to the normalization of cancerous epithelial cells [42]. These results point to the contextuality of the neoplastic phenotype and the centrality of stroma-epithelium interaction in carcinogenesis.

A Model for Fetal Origin of Cancer: The Xenoestrogen BPA

BPA was described as a “synthetic estrogenic agent” in 1936 [43]; however, it was never used as an estrogen due to its low potency when compared to DES which synthesis and characterization was published in 1938 [44]. Twenty years later, polycarbonate plastics, which are made from BPA monomers, were introduced. Worldwide, approximately 3 million metric tons of BPA are produced per year [45]. Besides food and beverage polycarbonate containers, BPA is also used in some cash register receipts, medical devices, contact lenses and other consumer products. As a result, humans are routinely exposed, most likely throughout life [46].

BPA is present in most human tissues and fluids including blood, breast milk, and amniotic fluid, typically at levels of approximately 1 ng/ml [47]. In this review we focus on the effects observed at doses that result in plasma levels below or equal to those found in humans; these doses are referred to as “low dose” [48]. The pharmacokinetics of BPA appear to be similar across mice, rats, non-human primates and humans [48].

Exposure to BPA Alters Fetal Mammary Gland Morphogenesis

The mammary gland develops through complex reciprocal interactions between mesenchyma and epithelium [49, 50] (see also Propper et al. in this volume). Based on our published experimental data showing morphological alterations in both compartments [51, 52] and that of others showing that estrogen receptors are only expressed in the stroma during fetal development [53, 54], we hypothesized that BPA, acting as an estrogen, would interfere with those reciprocal interactions. BPA was administered to mouse dams from gestational day (GD) 8 to GD18 through subcutaneous osmotic pumps delivering 0.25 μg/kg maternal bw/day (from here on μg/kg bw/day); this BPA dose is 1000-fold lower than that needed to produce levels comparable to those found in human plasma. Mammary glands of embryos at day 18 (E18) were examined. In the primary periductal mesenchyme, where nuclear ER α and β [54] and trans-membrane ER GPR30 are expressed [52], BPA altered extracellular matrix (ECM) organization [55]. Collagen fiber density increased in the stroma abutting the epithelium and decreased in the loose connective tissue further away. There was a marked decrease of tenasin C (TnC) in the periductal stroma (Fig. 1) [52]. The number of cells containing lipid droplets in the periductal stroma increased, and accelerated adipocyte differentiation was also observed in the presumptive fat pad. Additionally, the cell number/area was significantly decreased in the
presumptive fat pad and the proportion of Bax-positive cells was increased, suggesting increased apoptotic activity. Transcriptome analysis of the primary periductal stroma confirmed an increased expression of adipogenesis regulatory genes [PPARγ, low density lipoprotein receptor (Ldlr), G protein-coupled receptor 81 (GPR81), and Fabp4] while a decreased expression of ECM components, such as TnC, was observed [52].

Within the epithelium, BPA increased the area subtended by the ductal tree, the ductal extension (distance from the nipple to the furthest point of growth) and delayed lumen formation [51]. At the transcriptome level, there was an increased expression of anti-apoptotic genes [52]. Because mammary gland development is dependent on reciprocal interactions between these compartments, the changes in the primary periductal stroma involving ECM and adipogenesis, as well as the advanced fat pad maturation, may be responsible for the altered growth and the delayed lumen formation recorded in the ducts (Fig. 2).

When the mammary glands were grouped by the fetal position in the uterus, significant differences became apparent between females positioned between two males (2 M) and females placed between 2 females (0 M), suggesting hormone mediation of this differential effect. In the control group, the epithelium of the 2 M glands had more branches and larger area than the 0 M. Treatment with BPA obliterated this difference; that is, the epithelium of the 0 M glands of BPA exposed females subtended larger areas and branched more than the non-exposed 0 M [55].

![Image](image_url)

**Fig. 1** Effects of exposure to 250 ng BPA/kg/day from E8 to E18 on mice. Trichrome stained 5 μm section of control (a, e) and BPA-exposed (b, f) E18 mammary glands (EP denotes epithelium). Collagen stains deep blue; note the reduction of collagen deposition in the mesenchyme distal to the epithelium. Immunohistochemical localization of TnC in controls (c) and BPA-exposed (d) mammary glands. Red alkaline phosphatase staining of TnC is observed in the stroma surrounding the epithelial ducts. The density of collagen in the entire stromal compartment is significantly decreased in BPA-exposed females compared with controls, $p=0.010$ (g). However, the density of collagen within 10 μm of the epithelial ducts is significantly increased in BPA females, $p=0.042$ (h). All scale bars represent 100 μm. i Quantification showed that the number of adipocytes was significantly increased within 1 mm from the developing epithelium in BPA-exposed females (squares), compared with controls (circles). **P <0.02; ***P <0.005. Copyright 2007, The Endocrine Society. (c) and (i) reprinted with permission [55]
Does BPA Act as an Estrogen in the Fetal Mammary Gland?

BPA displays estrogenic activity at low concentrations (1–100 nM) [26, 27] and interferes with thyroid hormone action at higher concentrations (100 nM–10 μM) [56]. In order to examine whether the effects of BPA were due to its estrogenic effect, the transcriptomes of the periductal stroma and the epithelium of mouse embryos exposed in utero from E8 to E19 to a reference estrogen, i.e., 10 ng ethinylestradiol/kg bw/day, were compared to those of embryos exposed to 0.25 μg BPA/kg bw/day [52]. The similarity of the E19 transcriptomes of these two compounds strongly suggests that BPA acts mainly as an estrogen [52]. Of note, the similarity of the transcription profiles of the two hormones is higher in the epithelium than the periductal stroma [52]. These observations suggest that the mesenchyme, which expresses both nuclear ERs [54] and GPR30 [52], responds to different estrogenic agents in distinct manners but “integrates” estrogenic effects of diverse substances into a common set of “instructions” for the epithelium.

Effects of Perinatal BPA Exposure Manifest in Adult Life

The post-natal mammary glands of CD1 female mice exposed pre-natally (0.025 or 0.25 μg BPA/kg bw/day through osmotic pumps) had a significantly enhanced response to estrogens when administered after ovariectomy [57]. At post-natal day (PND) 30 the mammary glands of intact animals exposed prenatally to BPA exhibited an increased number of terminal end buds (TEBs) relative to the ductal area [57], decreased apoptosis in the TEBs and an increased number of epithelial cells expressed progesterone receptor (PR) [57]. A study using C57BL/6 mice exposed through drinking water from conception to weaning found a dose dependent increase in TEBs and the mRNA expression levels of estrogen-regulated genes such as amphiregulin [58] and secretory leukocyte protease inhibitor (unpublished observation, Briskin). At 3 months of age, a significant increase of the mammary cell number was observed in both the epithelial and the stromal compartments. There was an increase in the number of PR-positive cells in mammary epithelium at 6 and 12 months of age (Fig. 3). The response to progesterone by mammary epithelial organoids obtained from 3 month-old animals exposed from conception to weaning to BPA was also increased when compared to unexposed controls [58].

Progesterone increases lateral branching [59]. The increase in the proportion of epithelial cells expressing PR and their organization in clusters at 1 month of age was followed by a significant increase of lateral branching observed at 4 months of age in mice exposed gestationally to BPA [57] (Fig. 4). At 6 months of age, there was an overall increase in epithelial structures including terminal ends and a premature appearance of alveolar buds which are normally associated with pregnancy in mice [60]. These findings are consistent with the persistence of an increased number of epithelial cells expressing PR at 6 and 12 months of age [58]. When gestational exposure was extended through the lactation period, mammary glands of the offspring of BPA-treated mothers had an increased volume fraction of alveolar buds at 3 and 9 months of age, and an increased volume fraction of ducts at 9 months of age; these changes occurred at doses of 0.25 μg BPA/kg bw/day.
Animals exposed to 2.5 μg/kg bw/day also showed an increase in alveolar buds at 9 months of age [61]. Perinatal exposure to estrogens also induced intraductal hyperplasias, which in whole mounts is recognized by the appearance of "beaded ducts". Animals exposed pre- and post-natally (up to day 16) to low doses of BPA ranging from 0.25 to 25 μg BPA/kg bw/day developed beaded ducts [61] (Fig. 5).

In rats, as in mice, pre-natal exposure (from E10 to birth) to 250 μg BPA/kg bw/day via gavage resulted in an increase in the number of TEBs at PND21, an increased number of terminal ducts at PND 21 and 100, and an increased number of lobules type 1 at PND 35. Although morphological changes were only observed with the 250 μg/kg bw/day exposure, changes in gene expression were observed after exposure to both doses [62], albeit in a dose-dependent and age-dependent manner. The proteomic profile of the mammary glands at PND 21 identified 11 proteins differentially expressed in the BPA group when compared to controls [63]. At PND 50, 10 proteins were differentially expressed in the BPA animals compared to control; they included vimentin, adiponectin, desmin and the matricellular protein SPARC (secreted protein acidic and rich in cysteine) which affects ECM composition and collagen assembly [64]. In sum, BPA induces alterations in the mammary gland that are manifested regardless of route of exposure (oral vs. subcutaneous), the type of exposure (bolus vs. continuous), the timing of exposure (in utero vs. postnatal) and the species (mouse vs. rat). The outcomes can be either obvious, such as altered tissue architecture, or more subtle, such as differential gene and/or protein expression.

**Exposure to BPA During Organogenesis Predisposes to Mammary Gland Neoplasia**

The main risk factor for breast cancer is lifetime exposure to ovarian hormones. Mammary glands of gestationally BPA-exposed mice showed an enhanced sensitivity to estradiol and to progesterone [57, 58]. BPA-exposed mice had an increased number of TEBs, TEB area, TEB density and ductal extension [65]. Intraductal hyperplasias, a pre-cancerous lesion that gives rise to mammary adenocarcinomas
after transplantation into syngeneic mice were observed in female mice exposed from E8 through weaning to 0.25, 2.5 and 25 µg BPA/kg BW/day. These pre-cancerous lesions appeared in 3 month-old mice exposed to the lowest dose [61] and were characterized by the presence of epithelial cells growing inside the ductal lumen or even spanning the entire luminal diameter. The proliferation index of these cells was 5 times higher than that of cells in normal ducts. Interestingly, epithelial cells from alveolar buds adjacent to the hyperplastic ducts also had a high proliferation index. The stroma associated with intraductal hyperplasias showed macrophages and mast cells and it was high in fibrous collagen [61].

To explore the hypothesis that developmental exposure to low doses of BPA induces mammary neoplasias long after the exposure ended, we used a rat model because their mammary glands respond to carcinogens by developing tumors that better mimic the human breast disease regarding estrogen dependence and histopathology [66, 67]. BPA was administered to fetuses at doses ranging from 2.5 to 1000 µg/kg bw/day; this exposure resulted in the development of carcinomas in situ in the mammary glands of 33% of the rats exposed to 250 and 1,000 µg BPA/kg bw/day 2 while none of the unexposed animals developed carcinomas in situ [68]. Neoplasias were observed in young adult rats (PND 50 and 95).

![Proposed causal links tying BPA exposure, mammary gland development and carcinogenesis. BPA binds to the ERs present in the primary mesenchyme which alters the peri-ductal stroma, increasing peri-ductal collagen deposition and thus tissue rigidity. Increased rigidity is known to block or delay lumen formation. BPA also induces adipocyte differentiation in the primary peri-ductal stroma and fat pad, which in turn causes increased duct elongation and branching. These changes lead to an increased sensitivity to mammotrophic hormones such as estrogens and progesterone and likely to prolactin. The solid arrows link observations at E18 with postulated causal links. Dashed arrows link the observed effects at E18 with effects observed during puberty and adulthood. Not represented here are the effects of BPA on the hypothalamus, where it alters the control of ovarian cyclicity and likely the control of prolactin production.](image-url)
Fetal exposure to BPA also increased by 3-4 fold the number of intraductal hyperplasias [69]. Lesions in BPA-exposed animals were highly proliferative and contained numerous ERα positive cells [68, 70], suggesting that the proliferative activity in these lesions may be estrogen-mediated. As mentioned previously, rat mammary carcinomas as well as those in humans are predominantly estrogen-dependent, a feature that strengthens the relevance of these findings.

Is BPA a Carcinogen?

According to the Environmental Protection Agency (EPA), a carcinogen is a chemical or physical agent capable of causing cancer [http://www.epa.gov/saitoxes/nata/gloss1.html], a definition that does not specify the mechanism(s) by which the cancer is induced or the time lapse between exposure and diagnosis; it only identifies the consequence of an insult. Thus, by this definition BPA is a carcinogen.

Linking Existing Data into an Explanation of BPA-Driven Mammary Carcinogenesis

In the embryo, BPA binds to ER present in the primary mesenchyme and induces changes in gene expression that lead to accelerated maturation of periductal adipocytes which in turn, accelerates ductal growth and branching [31, 51]. Simultaneous and profound changes in the primary periductal matrix composition and organization lead to alterations of its biomechanical properties, which in turn affect the growth and branching of the epithelium [71, 72]. Because ERs are not expressed in the embryonic mammary epithelium, the induction of anti-apoptotic genes (which mediate delayed lumen formation) suggests mediation by biochemical factors (morphogens) secreted by the carcinogen-damaged periductal stroma and/or by biomechanical factors [73] (Fig. 6). Altered tissue development and remodeling of the mammary gland during each ovarian cycle may also contribute to neoplastic development. This remodeling activity is likely to be augmented due to the increased sensitivity to estradiol and progesterone observed in adult animals who were exposed to BPA from E8 to PND16 [57, 58]. In addition, morphological changes observed during adulthood such as increased abundance of alveolar buds and lobulo-alveolar structures suggest the presence of increased prolactin plasma levels and/or increased sensitivity to prolactin. BPA alters the architecture of the hypothalamic nuclei [74], which may generate altered circulating hormone levels that would facilitate the development of intra-ductal hyperplasias and carcinomas in situ. The involvement of both the mammary gland and the hypothalamus is supported by the fact that the mammary gland phenotype is more severe in mice exposed pre- and postnatally to BPA (from E8 to PND 16) which spans a critical window for differentiation of cyclic vs. tonic gonadotropin release. Such exposed animals showed changes in a hypothalamic region essential for cyclic gonadotropin release [74] and altered estrous cycles [75, 76], which in turn, may abnormally affect the development and the remodeling of the mammary gland, thus exacerbating the observed neoplastic phenotypes of these animals.

Conclusions

The causal link between fetal exposure to estrogens and the development of breast cancer that was first suggested by epidemiologists has now been confirmed by the increased risk to develop breast cancer during adulthood of women exposed to DES during their fetal life. Fetal and neonatal exposures to EDCs cause persistent alterations in the mammary glands of rodents, including pre- and neoplastic lesions, long after the exposure ended. In the case of BPA, mammary neoplasias may have their origin in the altered mammary morphogenesis that occurs during fetal and neonatal exposure. The data obtained from laboratory animals support the extrapolation that exposure to BPA and other xenoestrogens during organogenesis in humans contributes to the increase in the incidence of breast cancer observed over recent decades.

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