# Bisphenol S instead of bisphenol A: a story of reproductive disruption by regretable substitution – a review

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ABSTRACT: A range of substances that are released into the environment, foodstuffs and drinking water as a result of human activity were originally considered relatively harmless, and it was only later that their adverse effects were discovered. In general the use of such substances is currently restricted, and they are often replaced by other substances. This applies also in the case of a range of endocrine disruptors. These substances have the capacity to disturb the balance of physiological functions of the organism on the level of hormonal regulation, and their pleiotropic spectrum of effects is very difficult to predict. Endocrine disruptors include the currently intensively studied bisphenol A (BPA), a prevalent environmental pollutant and contaminant of both water and foodstuffs. BPA has a significantly negative impact on human health, particularly on the regulation mechanisms of reproduction, and influences fertility. The ever increasingly stringent restriction of the industrial production of BPA is leading to its replacement with analogues, primarily with bisphenol S (BPS), which is not subject to these restrictions and whose impacts on the regulation of reproduction have not yet been exhaustively studied. However, the limited number of studies at disposal indicates that BPS may be at least as harmful as BPA. There is therefore a potential danger that the replacement of BPA with BPS will become one of the cases of regrettable substitution, in which the newly used substances manifest similar or even worse negative effects than the substances which they have replaced. The objective of this review is to draw attention to ill-advised replacements of endocrine disruptors with substances whose effects are not yet tested, and which may represent the same risks for the environment, for the reproduction of males and females, and for human health as have been demonstrated in the case of the originally used substances.

Keywords: human health; environment; endocrine disruptor; reproduction; oocyte; sperm

# INTRODUCTION

Many substances have been introduced into use with great hopes, only for it to be demonstrated

earlier or later that they are harmful to the environment and/or human health. Notorious cases include the mass use of DDT as an insecticide (http://apps. who.int/iris/handle/10665/40018), thalidomide

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as a drug for pregnant women (McBride 1961), or more recently neonicotinoid insecticides used for the protection of fields against seed-destroying insects (Blacquiere et al. 2012). Substances whose negative effects on the environment or human health were detected only after a long period of use also include endocrine disruptors (Damstra et al. 2002).

The detection of the negative effects of abundantly used substances leads to a dramatic restriction of their use and their substitution with other substances. In a range of cases this brings about a genuine improvement. For example, chromated copper arsenate (CCA) used for wood preservation was demonstrated to be a substance with carcinogenic effects, and as a result was replaced with alkaline copper quaternary (ACQ). ACQ does not contain arsenic or chrome, and although it is just as effective as CCA against wood destroying arthropods, its impacts on the environment and human health are fundamentally less serious (Landrigan et al. 2004).

On the other hand, we have been witnesses to substitutions of harmful substances which have later been shown to be highly problematic. For example, 2,3-butanedione, which occurs naturally in butter, has been produced synthetically and added to foods in order to impart a buttery flavour. When it was demonstrated that 2,3-butanedione damaged lung tissue, it was replaced by 2,3-pentanedione, which however was subsequently proven to have similar negative effects on lung tissue as 2,3-butanedione (Hubbs et al. 2012). There are far more similar examples of "regrettable substitutions" (Fahrenkamp-Uppenbrink 2015; Zimmerman and Anastas 2015). In these cases, negative impacts on reproduction are often subsequently detected. For example, in the case of pyrethroids, which replaced older insecticide agents such as organocholorines, organophosphates or carbamates, and which were considered harmless to mammals, negative impacts were demonstrated on the maturation of mammal oocytes (Petr et al. 2013).

From the perspective of reproductive risks, the substitution of bisphenol A (BPA), a widely used component of plastics and many other materials, with its analogue bisphenol S (BPS) appears to be potentially problematic. BPA has been proven to be a strong endocrine disruptor, and its use has been restricted. Many products are sold with a "BPA-free" guarantee. Because BPA is substituted in a range of cases by BPS, these products are not however "bisphenol-free" (Glausiusz 2014), and their use

may be linked to significant reproductive risks. The aim of this review is to point to the replacement of BPA by BPS as a "regrettable substitution".

# **Endocrine disruptors**

A less harmful substitute is currently searched for a number of substances that had previously been considered safe from a toxicological perspective and finally appeared to exert various negative effects on health. This category of compounds includes substances referred to summarily as endocrine disruptors (Clayton 2011). According to the US Environmental Protection Agency, endocrine disrupting chemicals (EDCs) are defined as "exogenous agent(s) that interfere(s) in synthesis, secretion, transport, metabolism, binding action, or elimination of natural blood-borne hormones that are present in the body and are responsible for homeostasis, reproduction, and developmental processes" (Diamanti-Kandarakis et al. 2009).

EDCs manifest a range of particular properties. Their hormone-like effects may be suppressed or may fade away entirely in the case that the concentration of EDCs is higher than the physiological level of their hormonal counterpart. This ability of agents to attain paradoxically stronger effects in low doses than in high ones (vom Saal and Welshons 2005) is termed the "low dose effect" (Grasselli et al. 2010; Vandenberg et al. 2012). The low dose hypothesis posits that exogenous chemicals that interact with hormone action can do so in a quite specific manner. In accordance with that, mentioned traditional toxicological endpoints are not capable to preclude adverse outcome, as EDCs act with dose responses, that are nonlinear and potentially non-monotonic (Vandenberg et al. 2012). In the case the relationship between dose and response is nonlinear, any prediction is even more complex. Therefore, the low dose definition was extended by the effects of non monotonic response curves. The mechanisms responsible for the non-linear effects are described in detail (Vandenberg et al. 2012), usually in connection with an interaction between a ligand (hormone or EDC) and a hormone receptor (Vandenberg 2014).

Non-linear dose-response patterns are commonly observed with endogenous and synthetic agonists (e.g. numerous drugs, hormones, peptides) that activate and inhibit receptor-mediated signal pathways that affect various biological functions



Figure 1. Chemical structure of bisphenol A (A), bisphenol S (**B**), bisphenol F (**C**)

(Calabrese and Baldwin 2001; Calabrese 2005). However, EDCs can also produce non monotonic dose responses in which the slope of the curve changes sign over the course of the dose-response (www. who.int/ceh/publications/endocrine/en/index.html) and low dose effects are described for the majority of EDCs (Birnbaum 2012; Vandenberg et al. 2012, 2013; Zoeller et al. 2012; Bergman et al. 2013).

The concept of endocrine-disrupting chemicals was proposed after these compounds had been observed to affect various reproductive functions in wildlife and humans (Colborn et al. 1993). The influence of several EDCs was demonstrated on the course of development of male gametes, sperm (Li et al. 2011; Knez et al. 2014) and female gametes, oocytes, as well as embryonic development of males and females (Mok-Lin et al. 2010; Xiao et al. 2011). Moreover, the effect of EDCs on the reproduction of adult individuals, including transgenerational inheritance, has been described (Susiarjo et al. 2015; Ziv-Gal et al. 2015). Therefore, reproductive functions represent crucial targets of the EDCs' negative effects. Recently intensively studied EDCs, interfering with the regulation of physiological reproductive processes, include bisphenols, a family of chemical compounds with two hydroxyphenyl functional groups (Figure 1).

# **Bisphenol** A

An example of a widely used substance, in which endocrine-disrupting properties were detected only later, is bisphenol A (BPA, 4,4'-(propane-2,2diyl)diphenol) (Vandenberg et al. 2009). BPA was first synthesized in 1891, and as early as in 1936 it was demonstrated that it imitates the activity of the hormone estradiol (Dodds and Lawson 1936). Despite a very strong estrogen activity, BPA has been commercially used since 1957, and despite the fact that its endocrine-disrupting activity was discovered (Krishnan et al. 1993), BPA has become a high production volume chemical (Wang et al. 2012). Worldwide annual production, which in the case of BPA reached 4.6 million t in 2012, is constantly increasing. Its production was estimated at 5.4 million t in 2015 (Merchant Research & Consulting, http://mcgroup.co.uk/researches/bisphenol-a-bpa).

BPA is present especially in polycarbonate plastics, epoxide resins, and several paper products (Ehrlich et al. 2014), and as a result it is used in a variety of commonly used consumer products such as thermal recipes, cosmetics, dental materials, medicinal tubes, utensils, toys, baby feeding bottles and dummies, etc. Heat, UV radiation, alkaline treatment or intensive washing causes a release of BPA monomer. It is estimated that the worldwide release of BPA into the environment is almost half million kg per year (Mileva et al. 2014).

BPA is released into the environment either directly from chemical, plastic coating, and staining manufacturers, from paper or material recycling companies, foundries which use BPA in casting sand, or indirectly leaching from plastic, paper, and waste in landfills (Yang et al. 2015). BPA passes into foodstuffs or water directly from the lining of food and beverage cans, where it is used as an ingredient in the plastic used to protect the food from direct contact with the can (Goodson et al. 2002; Vandenberg et al. 2009). The main path of human exposure is the consumption of such contaminated foodstuffs, drinking water or via dermal contact with thermal paper and cosmetics or inhalation (Miyamoto and Kotake 2005; Huang et al. 2012).

It is therefore not surprising that a range of studies have now demonstrated the presence of BPA in human tissue. Levels of BPA have been tested in various populations worldwide, and the presence of BPA was demonstrated in 92.6% of Americans (Wetherill et al. 2007) and 90% of Canadians (Bushnik et al. 2010). Levels of BPA have been demonstrated in various biological matrices, most frequently in urine (Casas et al. 2013; Salgueiro-Gonzalez et al. 2015), but also in blood serum. Within the human reproductive system, levels of BPA have been confirmed for example in testicle tissue, seminal plasma (Manfo et al. 2014), in ovarian follicular fluid (Ikezuki et al. 2002), mother's

Sample	Level of BPA	References
Blood (ng/ml)	12.4–14.4	Bushnik et al. (2010)
Maternal blood (ng/ml)	0.63-14.36	Yamada et al. (2002)
Fetal blood (ng/ml)	0.2–9.2	Schonfelder et al. (2002)
Urine (ng/ml)	0.02-21.0	Liao et al. (2012c)
Saliva (ng/ml)	0.3	Joskow et al. (2006)
Follicular fluid (ng/ml)	$2.4 \pm 0.8$	Ikezuki et al. (2002)
Amniotic fluid (ng/ml)	1.1-8.3	Ikezuki et al. (2002)
Placental tissue (ng/g)	1.0 - 104.9	Schonfelder et al. (2002)
Breast milk (ng/ml)	0.5–1.3	Mendonca et al. (2014)
Semen plasma (pg/ml)	66 (fertile men) 132–179 (infertile men)	Vitku et al. (2015)

Table 1. Bisphenol A (BPA) levels in human fluids

milk, fetal plasma (Shonfelder et al. 2002), amniotic fluid (Yamada et al. 2002; Edlow et al. 2012), and the placenta (Jimenez-Diaz et al. 2010; Cao et al. 2012) (Table 1). Several studies have demonstrated a direct correlation between exposure of the mother and the BPA level of the fetus (Ikezuki et al. 2002; Kuruto-Niwa et al. 2007). BPA may permeate the placenta and thus influence the development of the fetus (Edlow et al. 2012; Corbel et al. 2014). Newborns may then be further exposed to the effect of BPA during breastfeeding due to the presence of BPA in mother's milk (Mendonca et al. 2014).

The effects of BPA on humans are dependent not only on the dose, but also on the window of exposure.

Exposure to BPA in the prenatal and neonatal period probably affects the human organism in the most receptive period (Fernandez et al. 2014).

# **Mechanism of BPA action**

A typical feature of endocrine disruptors is their wide spectrum of outcomes (Figure 2). Combination of their action in various target systems in the organism is one of causes of their nonlinear effects. In this respect, BPA acts as a typical endocrine disruptor with multi-level impacts (Khan and Ahmed 2015). Nongenomic effects of BPA have been described, thus influencing cellular signalling



Figure 2. Possible mechanisms of bisphenol action and its potential impact on human health

(Nakagawa and Tayama 2000), as well as genomic, which affect transcription regulation (Trapphoff et al. 2013), and also epigenetic, responsible for the methylation and acetylation of DNA and core histones (Bromer et al. 2010). It is precisely pronounced estrogen activity of BPA *in vitro* (vom Saal et al. 2007; Wetherill et al. 2007) and *in vivo* that contributes to its immense potential to afflict the hormonal system and act as an endocrine disruptor.

BPA inhibits the activity of natural endogenous estrogens and thus disrupts estrogen nuclear hormone receptor action (Kitamura et al. 2005; Wetherill et al. 2007; Grignard et al. 2012). BPA affects hormonal homeostasis, for example through bonding to the classic nuclear estrogen receptors  $\alpha$ ,  $\beta$ ,  $\gamma$  (ER $\alpha$ , ER $\beta$ , ER $\gamma$ ), where it manifests a combination of agonistic and/or antagonistic actions in dependence on the target tissue, cell types, ER subtypes, and differential cofactors recruited by ER-ligand complexes (Kurosawa et al. 2002). BPA also bonds to non-classical membrane ERs and causes activation of the nuclear receptor gamma (Takayanagi et al. 2006; Matsushima et al. 2007).

BPA has been identified as an antagonist of androgen receptors (Kitamura et al. 2005; Wetherill et al. 2007; Vinggaard et al. 2008; Molina-Molina et al. 2013). Its anti-androgenic activity has been documented in several studies, but with changing values of the maximum inhibition concentration (Xu et al. 2005; Bonefeld-Jorgensen et al. 2007). In contrast with other known androgen receptor antagonists, BPA inhibits the effective nuclear translocation of the androgen receptors, and disrupts their function by means of a number of mechanisms (Teng et al. 2013). The endocrine-related BPA action mechanism also involves a reduction of aromatase expression (Zhang et al. 2011; Chen et al. 2014) and a decrease in aromatase activity in vitro (Bonefeld-Jorgensen et al. 2007). Within this context, it is of interest that a decline in the synthesis of testosterone and estradiol in vivo has been documented following exposure to BPA (Akingbemi et al. 2004).

The epigenetic mechanisms of the effect of BPA include the alteration of certain DNA methylation samples (Dolinoy et al. 2007; Susiarjo et al. 2013). Prenatal exposure to BPA alters the expression of genes coding individual subtypes of ERs in a sexand brain region-specific manner (Kundakovic et al. 2013) and disrupts the normal development of the placenta (Susiarjo et al. 2013). As a result, it is possible that BPA predetermines the response to

steroid hormones in the very early phase of development (Wilson and Sengoku 2013). It has been documented that BPA also disrupts the gene expression of the regulating factors that control the stability and flexibility of epigenetic regulation, and as a result has an adverse influence on the development of functions of the controlling organ of hormonal regulation, the hypothalamus (Warita et al. 2013). The impacts of these changes have transgenerational effects (Manikkam et al. 2013).

Further demonstrated actions of BPA in the organism include the bonding to the glucuronide receptor, suppression of the transcription receptor of the thyroid hormone, reduction of the transport of cholesterol via the mitochondrial membrane, increase of oxidation of fatty acids, stimulation of prolactin release (Machtinger and Orvieto 2014) or an agonistic effect on the human pregnane X receptor (Sui et al. 2012).

#### **BPA and human health**

With such a wide spectrum of effects, it is evident that BPA has a negative influence on human health. Frequently discussed themes include the possible association of BPA for example with obesity (Trasande et al. 2012), diabetes (Lang et al. 2008), neurobehavioural disorders (Jasarevic et al. 2011), cancer (Jenkins et al. 2011), hepatic (Peyre et al. 2014) and cardiovascular diseases, hypertension, and disorders of the thyroid gland function (Rochester 2013; Wang et al. 2013).

Especially in the area of reproduction in both animal models and in humans, a wide range of negative influences of BPA have been observed (Kwintkiewicz et al. 2010; Trapphoff et al. 2013; Zhang et al. 2014). BPA has varied and complex mechanisms of action that may interfere with normal reproductive development and functions. In both males and females, BPA interferes with hormonal regulation and influences the hypothalamic-pituitary-gonadal axis on all levels (Navarro et al. 2009; Patisaul et al. 2009; Xi et al. 2011).

#### Influences of BPA on reproduction of males

As a rule, endocrine-disrupting substances have pronounced impacts on the reproduction of both sexes. Several studies have shown detrimental effects of BPA on spermatogenesis and semen quality in fishes. The number of mature and im-

mature spermatozoa was decreased and increased, respectively (Sohoni et al. 2001) and also the sperm motility and concentration were reduced (Lahnsteiner et al. 2005). There is a large evidence that BPA can induce sex reversal from male to female in aquatic animals. Changes in sex ratio were observed at zebrafish during embryonic development (Drastichova et al. 2005) and *Xenopus* larvae through metamorphosis (Kloas et al. 1999).

Experimental studies on the effects of BPA on the reproduction of male rodents have revealed an adverse influence on the development of testes (Vrooman et al. 2015) and on the spermatogenesis of adult individuals following prenatal in utero or early postnatal exposure. Exposure to BPA during the period of development of the testes is frequently linked to a range of negative effects in adult testes, e.g. decreased levels of testicular testosterone, decreased weights of the epididymis and seminal vesicles, a decrease in daily sperm production per gram testis, and increased weights of the prostate and preputial (Richter et al. 2007). Vrooman et al. (2015), with the help of transplantation of spermatogonia from the testes of mice exposed to the action of BPA into mice which were not exposed, demonstrated permanent damage to spermatogenesis. The influence of the exposure of adult rodents to BPA on the quality of sperm was also studied (Peretz et al. 2014).

Despite the differences in the experimental designs used, certain findings appear repeatedly, especially reduction in the number of sperm, reduction in the motility of sperm, increased amount of apoptotic cells in the seminiferous tubules, changes in the levels of hormones and steroid enzymes, and damage to the DNA of sperm (Peretz et al. 2014).

Contemporary studies confirm that rodents are not relevant for predicting the effect of low BPA concentrations on the endocrine function of human fetal testis (N'Tumba-Byn et al. 2012). In a comparative study by Maamar et al. (2015), the influence of BPA was studied both on rats and on human fetal testes, and it was determined that in both cases BPA had dose-dependent anti-androgenic effects. Nevertheless, the authors urge caution in interpreting the results obtained on rodents and their application in human medicine (Maamar et al. 2015).

Unfortunately, there is only a limited number of studies that have observed the influence of exposure to BPA on the quality of sperm in adult humans. In men exposed to BPA in the workplace and patients in reproduction centres, a higher level of BPA in urine was linked to a lower number, concentration, and motility of sperm (Knez et al. 2014; Lassen et al. 2014). Nevertheless, in a study conducted by Mendiola et al. (2010) on fertile men, the concentration of BPA in urine did not correlate with changes in semen parameters, despite the fact that a significant correlation was observed between the level of BPA in urine and the volume of seminal plasma or markers of free testosterone (Mendiola et al. 2010).

The following cohort study examined the relationship between the concentration of BPA in urine and the level of reproductive hormones and semen in a group of 308 young healthy men. It was determined that the concentration of BPA strictly correlates with higher levels of selected circulating reproductive hormones and reduced motility of sperm. The results indicated that the exposure to BPA on the level of environment has an anti-androgenic and/or anti-estrogenic effect due to the effect of BPA on the level of receptors. The anti-estrogenic effect on the level of the epididymis also explains the determined low mobility of the sperm (Lassen et al. 2014).

#### Influences of BPA on reproduction of females

BPA markedly influences not only the reproduction of males, but also the reproduction of females. In both *in vitro* and *in vivo* studies, the influence of BPA has been demonstrated on fertility, function of the womb i.e. formation of benign and malignant lesions (Newbold et al. 2009), disruption apoptosis of the uterine epithelium during estrus (Mendoza-Rodriguez et al. 2011), function of ovaries and quality of oocytes (Peretz et al. 2014), and defective folliculogenesis (Santamaria et al. 2016). In females it is precisely the ovaries that are the key organ responsible for reproductive and endocrine functions, and BPA is frequently indicated as an ovarian toxicant. BPA afflicts not only the overall morphology and weight of the ovaries (Suzuki et al. 2002; Santamaria et al. 2016) but also demonstrably reduces the quality of oocytes in both animal and human models (Machtinger and Orvieto 2014).

During the course of the maturation of mouse oocytes *in vitro* following treatment with BPA, changes were documented in the configuration of the meiotic spindle resulting in errors in chromosome segregation and hyperploidy frequencies in mouse

oocytes (Hunt et al. 2003). Similary, it was reported that BPA exposure altered chromosome and spindle organization which resulted in hyperploidy of mouse oocytes during meiosis (Can et al. 2005) and it was also demonstrated that low BPA doses are related with aberration during meiotic prophase, including increased incidence of recombination (Susiarjo et al. 2007) and failure formation of primordial follicle by inhibiting meiotic progression of oocytes (Zhang et al. 2012). In contrast, Eichenlaub-Ritter and her colleagues found no evidence that low BPA doses increased hyperploidy at meiosis II. On the other hand they observed cell cycle delay and meiotic spindle abnormalities, changes in the distribution of pericentriolar material and chromosome alignment (Eichenlaub-Ritter et al. 2008). Exposure of mice, from mid-gestation to birth, causes synaptic abnormalities in oocytes and an increased amount of recombination between homologous chromosomes. It is also of interest that identical effects have been observed in homozygous mice with an intentionally disrupted gene coding the ER $\beta$ . In mouse oocytes, epigenetic changes have also been documented following cultivation of follicles in the presence of BPA, in which a disruption of the configuration of chromosomes took place, as well as disorders of meiosis caused by faulty genomic imprinting and altered posttranslational modification of histones (Trapphoff et al. 2013). Chronic exposure of oocytes was linked to an increased incidence of aberrant metaphases II and prematurely segregated chromatids (Pacchierotti et al. 2008).

Bovine oocytes cultivated in the presence of BPA have also manifested disorders of the meiotic spindle and the chromosomal configuration (Ferris et al. 2015). In Barbary Macaques, negative effects of BPA have been demonstrated in various stages of the oogenesis of developing ovaries. Oocytes in the prophase of meiosis and in fetal ovaries exhibited an increased number of recombination, and an increased number of abnormally formed follicles containing multiple oocytes was recorded in perinatal ovaries (Hunt et al. 2012).

Similary as in the aforementioned studies on rodents, cattle, and primates, an increased number of crossing over and degenerations in oocytes have been determined also in human oocytes cultivated *in vitro* in the presence of BPA (Brieno-Enriquez et al. 2011). In connected studies it has been demonstrated that the exposure of human oocytes to BPA is linked to up-regulation of genes involved in meiotic processes connected to double strand breaks repair progression (Brieno-Enriquez et al. 2012). A non-linear response to BPA doses on the incidence of MII oocytes with aligned chromosomes has also been determined (Machtinger et al. 2013). The changes which have been recorded in the development of oocytes exposed to bisphenol may lead to disorders in the development of embryos, fetal loss or genetic disorders (Rama Raju et al. 2007; Ye et al. 2007; Tomari et al. 2011). The result of maternal exposure to BPA may be the disruption of the entire oogenesis in the developing ovary (Susiarjo et al. 2007).

A number of cohort studies have been focused on groups of persons who undergo treatment for infertility through *in vitro* fertilization (IVF). The measured levels of BPA in these persons were examined in connection with the ovarian response, quality of embryos and implantation. A reduced ovarian response was linked to a reduced success rate of IVF (Mok-Lin et al. 2010). BPA also disrupted embryonal development of fish via delay hatching, yolk reabsorption, and larval growth of trouts (Aluru et al. 2010), moreover lethality in zebrafish larvae increased (Chan and Chan 2012).

There is only a limited number of studies which have observed the effects of BPA on the development and quality of mammalian blastocysts. Failure of embryonic development to mouse blastocyst stage has been demonstrated after exposure of females to BPA (Xiao et al. 2011). Disorder of implantation of mouse blastocysts was also demonstrated by Borman et al. (2015).

In human, Bloom et al. (2011) state a correlation between the concentration of BPA in the urine of men, though not in women, and a decline in the quality of embryos generated by IVF. By contrast, in a study performed by Knez et al. (2014), which confirms changes to the semen quality of men with a determined environmental level of BPA, undisrupted development of embryos into blastocysts is described. As against this finding, in women who have undergone IVF, a correlation has been demonstrated between the concentration of BPA in urine and a change to the formation of blastocysts, though a reduced quality of embryos was not recorded (Ehrlich et al. 2012).

#### The advent of BPS

The above-stated facts led to the necessity for stringent regulation of the use of BPA, and in a

range of cases its substitution with another chemical. On the basis of the effects on human health and reproduction demonstrated with the help of standardized toxicological testing procedures, government agencies in the United States (the US Environmental Protection Agency, USEPA), Canada (Health Canada), and Europe (the European Food Safety Authority, EFSA) have established tolerable daily intake levels, ranging from 25 to 50 µg BPA/kg of body weight (BW) per day (Rochester 2013). With regard to the fact that several studies have demonstrated BPA low dose effects (Vandenberg et al. 2012), and that this possibility is unfortunately not taken into account in the approach of "traditional" toxicological studies, in which low doses are not generally subjected to examination (Vandenberg et al. 2012; Rochester 2013), scientists have expressed concerns that the "safe" cut-off set for BPA is too high (vom Saal and Hughes 2005). In 2010 the Canadian government prohibited the import, sale, and advertisement of baby feeding bottles containing BPA. The European Union responded with a prohibition of the manufacture of baby feeding bottles with BPA, which was passed in 2011 (Commission Directive 2011). The Food and Drug Administration (FDA) has indicated BPA as a "chemical of concern", and in July 2012 a blanket prohibition of BPA in baby feeding bottles and sippy cups was recommended (FDA 2011). However, new data and refined methodologies have led EFSA experts to considerably reduce the safe level of BPA from 50 µg/kg of BW/day to 4 μg/kg of BW/day (EFSA 2014).

With regard to these restrictions and societal pressures, manufacturers of plastics are now forced to seek an alternative product which can replace BPA. It is in the interest of chemical concerns that the substitute which replaces BPA is inert or at least far less toxic than BPA. Nevertheless, new chemicals introduced onto the market are frequently untested, and may be equally or more harmful than the originals, which are ultimately termed "regrettable substitutions" (Rochester and Bolden 2015), as has been the case of a number of perfluorinated chemicals (Howard 2014), pesticides (Coggon 2002), and self-extinguishing compounds (Bergman et al. 2012). Manufacturers seeking BPA alternatives have turned primarily to bisphenol S (BPS, 4,4'-sulfonyldiphenol) (see Figure 1), a structural analogue of BPA, to produce "BPA-free" products (Grignard et al. 2012; Barrett 2013). BPS is chemically more stable, worse in terms of biodegradability than BPA, and shows better dermal penetration than BPA (Ike et al. 2006; Danzl et al. 2009; Liao et al. 2012a, b). It is disconcerting that these properties may lead to a longer or higher body burden or bioavailability of BPS versus BPA (Helies-Toussaint et al. 2014). For these reasons, too, at present the replacement of BPA with BPS is considered a "regrettable substitution" (Fahrenkamp-Uppenbrink 2015; Zimmerman and Anastas 2015). With regard to the increase in production of BPS and the indispensability of bisphenols in the production of plastics, it is unfortunately possible to expect the same widespread use of BPS as in the case of BPA (Liao et al. 2012c). Now the presence of BPS can be expected in almost all the consumer goods here in which BPA was initially used (Mathew et al. 2014), for example as a wash fastening agent in clearing products, an electroplanting solvent, and a constituent of phenolic resins (Rochester and Bolden 2015).

One of the major industries that have replaced BPA due its high occurrence (~3–22 g/kg) is that of thermal paper (Mathew et al. 2014). In the USA, Korea, Vietnam, Japan, and China (Liao et al. 2012c), BPS has been detected in several different "BPA free" paper products, including receipts and paper money (Liao et al. 2012a). The presence of BPS has been determined in tinned foodstuffs (Vinas et al. 2010). The occurrence of BPS has also been determined in indoor dust (Liao et al. 2012b), in fluvial water (Ike et al. 2006), surface water, and waste waters (Song et al. 2014) (Table 2).

The main pathway to the human body is dermal, dust ingestion, and dietary exposures (Liao et al. 2012b). Unfortunately, for example thermal paper carries BPS into all recycled paper products, making dermal exposure inevitable. Massive exposure of the population to the effects of environmental BPS has been demonstrated in a number of different countries. Within the range of 0.02–21 ng/ml (0.8–84nM) it has been detected in human urine samples originating from seven Asian countries and the USA (Liao et al. 2012a) in 81% of analyzed samples. In the following study the presence of BPS in urine was demonstrated in residents living near a manufacturing plant in south China in a concentration of 0.029 ng/ml (Yang et al. 2015).

#### **Biological effects of BPS**

Although nowhere near as much information is available about BPS as about the endocrine-

Table 2. Bisphenol S (BPS)	levels in the personal	care products and environment
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Sample	Level of BPS	References
Canned food (ng/g)	8.9–17	Vinas et al. (2010)
Thermal paper (mg/g)	0.0000138-22.0	Liao et al. (2012c)
Tickets (µg/g)	0.183-5.93	Liao et al. (2012c)
Currency bills (µg/g)	0.00-6.26	Liao et al. (2012c)
Other paper product types (µg/g)	0.00-8.38	Liao et al. (2012c)
Indoor dust (µg/g)	0.34	Liao et al. (2012b)
Municipal sawage sludge (ng/g dry weight)	0.17-110.00	Song et al. (2014)
River water (ng/l)	0.29-18.99	Yang et al. (2014)

disrupting effects of BPS, the substitution of BPA with BPS is raising concerns. The limited number of studies available at the present time, dealing with the biological interactions of BPS with the organism, indicate that BPS is also capable of imitating properties of hormones, interacting with ER (Delfosse et al. 2012; Rosenmai et al. 2014; Le Fol et al. 2015), and direct binding to nuclear ERs (Yamasaki et al. 2004) and serum albumins (Mathew et al. 2014) has been confirmed.

Some *in vitro* studies have demonstrated a weaker estrogen activity of BPS than the activity manifested by estradiol (Kuruto-Niwa et al. 2010; Grignard et al. 2012; Molina-Molina et al. 2013; Rochester and Bolden 2015). By contrast, a study conducted by Vinas and Watson (2013a, b) demonstrated the same or higher estrogen effectiveness than estradiol, BPS was capable of stimulating the membrane receptor pathways ordinarily up-regulated by estradiol. After exposure to BPS there are also changes in the expression of aromatase, the key enzyme in the synthesis of estradiol (Kinch et al. 2015).

Like in the case of BPA, the androgenic activity of BPS was confirmed (Kitamura et al. 2005), and subsequently its anti-androgenic activity as well (Molina-Molina et al. 2013). These observations in vitro have also been confirmed by in vivo studies. Chen et al. (2002) described acute toxicity of BPS in Daphnia magna and at the same time also demonstrated estrogen activity of BPS in vitro. Yamasaki et al. (2004) documented estrogen activity of BPS in vivo in rats with the assistance of postnatal exposure to BPS, which in both low and high doses induced the growth of the womb (Owens and Ashby 2002). An in vivo study on the effect of BPS in zebrafish documented not only changes in the mass of the gonads and plasmatic levels of estrogen and testosterone, but also a marked disruption of reproduction. The study of Qiu and colleagues evaluated the impact of BPA and BPS on the reproductive neuroendocrine system during zebrafish embryonic development, and explored potential mechanisms of action associated with ER, thyroid hormone receptor, and enzyme aromatase pathways. All of these pathways were necessary to observe the full effects of BPS on the changes in gene expression in the reproductive neuroendocrine axis (Qiu et al. 2016). These data were substantiated by a decrease in egg production and hatchability and an increasing number of embryo malformations (Ji et al. 2013). These observations were later extended upon by increased time to hatch, reduced number of sperm, increasing number of female to male ratio, and changes in the levels of testosterone, estradiol, and vitellogenin (Naderi et al. 2014). In further experiments provided in cell cultures it has been demonstrated that BPS acts cytotoxically, genotoxically (Lee et al. 2013), and mutagenically (Fic et al. 2013).

The reason for these negative effects may be for example binding to serum albumins or DNA damage and subsequent influencing of several signal cascades anywhere within the organism (Lee et al. 2013; Mathew et al. 2014). Exposure to BPS disrupts cellular signalling in the apoptotic and survival pathways (Salvesen and Walsh 2014). Evidently, it is possible to expect the interference of BPS in signal pro-apoptotic pathways and signal cascades described also in gametes, leading to an altered cell cycle and cell death (Nevoral et al. 2013; Sedmikova et al. 2013). Further studies focused on the mechanism of BPS action are needed for a full understanding its negative effect on reproduction on the gamete level and cell cycle regulation.

In respect to previous regrettable substitution, another bisphenols, such as bisphenol F (BPF, bis(4-hydroxyphenyl)methane; see Figure 1), do

not seem to be a suitable alternative. In addition to BPA and BPS, BPF has been described as endocrine disruptor as well (Perez et al. 1998). Surprisingly, natural presence of BPF has recently been observed in mustard and, therefore, it is a frequent compound of foodstuff (Zoller et al. 2016). Hence, BPF regulation is ambiguous for its chronical intake by a major part of human population (Dietrich and Hengstler 2016).

# CONCLUSION

At present we are witnessing the substitution of BPA with BPS in a whole range of materials, and BPS is becoming a standard component of several products. BPS is a substance which is structurally very similar to BPA, it shows analogous effectiveness and mechanism of in vitro action. Biological changes occurring in the range of typical human exposures were documented at doses below those used in traditional toxicology. On the basis of the described comparisons, it is possible to expect that BPS, like BPA, is an endocrine disruptor, and that it may have similar targets and manner of action in vivo and may influence physiological processes on several levels. With regard to its slower degradation, BPS may act for a longer time in the organism and thus interfere with the regulation of reproduction of mammals in a yet more dangerous manner than has been demonstrated by a range of studies in the case of BPA.

The alarming results of the first reproduction studies on BPS have generated an acute need for a wider and at the same time more detailed assessment of the impacts of BPS, with emphasis on the area of reproduction of mammals, which is entirely lacking at present. Should this not materialize, due to the increasing industrial production of BPS caused by the need to replace BPA, unfortunately BPS may within the foreseeable future become just as great an environmental health risk as BPA. There is a need for very intensive research and subsequently also legislative measures in order to ensure that BPS will not become another "regrettable substitution" with pronounced negative impacts on the environment and on human health, including negative impacts on reproduction.

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## REFERENCES

- Akingbemi B.T., Sottas C.M., Koulova A.I., Klinefelter G.R., Hardy M.P. (2004): Inhibition of testicular steroidogenesis by the xenoestrogen bisphenol A is associated with reduced pituitary luteinizing hormone secretion and decreased steroidogenic enzyme gene expression in rat Leydig cells. Endocrinology, 145, 592–603.
- Aluru N., Leatherland J.F., Vijayan M.M. (2010): Bisphenol A in oocytes leads to growth suppression and altered stress performance in juvenile rainbow trout. PLoS ONE, 5, e10741.
- Barrett J.R. (2013): Assessing the safety of a replacement chemical: nongenomic activity of bisphenol S. Environmental Health Perspectives, 121, 97.
- Bergman A., Ryden A., Law R.J., de Boer J., Covaci A., Alaee M. (2012): A novel abbreviation standard for organobromine, organochlorine and organophosphorus flame retardants and some characteristics of the chemicals. Environment International, 49, 57–82.
- Bergman A., Heindel J.J., Jobling S., Kidd K.A., Zoeller R.T. (eds) (2013): State of the Science of Endocrine Disrupting Chemicals – 2012. World Health Organization, Geneva, Switzerland/United Nations Environment Programme, Nairobi, Kenya.
- Birnbaum L.S. (2012): Environmental chemicals: evaluating low-dose effects. Environmental Health Perspectives, 120, 143–144.
- Blacquiere T., Smagghe G., van Gestel C.A., Mommaerts V. (2012): Neonicotinoids in bees: a review on concentrations, side-effects and risk assessment. Ecotoxicology, 21, 973–992.
- Bloom M.S., vom Saal F.S., Kim D., Taylor J.A., Lamb J.D., Fujimoto V.Y. (2011): Serum unconjugated bisphenol A concentrations in men may influence embryo quality indicators during in vitro fertilization. Environmental Toxicology and Pharmacology, 32, 319–323.
- Bonefeld-Jorgensen E.C., Long M., Hofmeister M.V., Vinggaard A.M. (2007): Endocrine-disrupting potential of bisphenol A, bisphenol A dimethacrylate, 4-n-nonylphenol, and 4-n-octylphenol in vitro: new data and a brief review. Environmental Health Perspectives, 1, 69–76.
- Borman E.D., Foster W.G., Greenacre M.K., Muir C.C., de Catanzaro D. (2015): Stress lowers the threshold dose at which bisphenol A disrupts blastocyst implantation, in conjunction with decreased uterine closure and e-cadherin. Chemico-Biological Interactions, 237, 87–95.
- Brieno-Enriquez M.A., Robles P., Camats-Tarruella N., Garcia-Cruz R., Roig I., Cabero L., Martinez F., Caldes M.G. (2011): Human meiotic progression and recombination are affected by Bisphenol A exposure during in vitro

human oocyte development. Human Reproduction, 26, 2807–2818.

Brieno-Enriquez M.A., Reig-Viader R., Cabero L., Toran N., Martinez F., Roig I., Garcia Caldes M. (2012): Gene expression is altered after bisphenol A exposure in human fetal oocytes in vitro. Molecular Human Reproduction, 18, 171–183.

- Bromer J.G., Zhou Y., Taylor M.B., Doherty L., Taylor H.S. (2010): Bisphenol-A exposure in utero leads to epigenetic alterations in the developmental programming of uterine estrogen response. The FASEB Journal, 24, 2273–2280.
- Bushnik T., Haines D., Levallois P., Levesque J., van Oostdam J., Viau C. (2010): Lead and bisphenol A concentrations in the Canadian population. Health Reports, 21, 7–18.
- Calabrese E.J. (2005): Toxicological awakenings: the rebirth of hormesis as a central pillar of toxicology. Toxicology and Applied Pharmacology, 204, 1–8.
- Calabrese E.J., Baldwin L.A. (2001): The frequency of U-shaped dose-responses in the toxicological literature. Toxicological Sciences, 62, 330–338.
- Can A., Semiz O., Cinar O. (2005): Bisphenol-A induces cell cycle delay and alters centrosome and spindle microtubular organization in oocytes during meiosis. Molecular Human Reproduction, 11, 389–396.
- Cao X., Zhang J., Goodyer C.G., Hayward S., Cooke G.M., Curran I.H.A. (2012): Bisphenol A in human placental and fetal liver tissues collected from Greater Montreal area (Quebec) during 1998–2008. Chemosphere, 89, 505–511.
- Casas M., Valvi D., Luque N., Ballesteros-Gomez A., Carsin A.E., Fernandez M.F., Koch H.M., Mendez M.A., Sunyer J., Rubio S., Vrijheid M. (2013): Dietary and sociodemographic determinants of bisphenol A urine concentrations in pregnant women and children. Environment International, 56, 10–18.
- Chan W.K., Chan K.M. (2012): Disruption of the hypothalamic-pituitary-thyroid axis in zebrafish embryo-larvae following waterborne exposure to BDE-47, TBBPA and BPA. Aquatic Toxicology, 108, 106–111.
- Chen M.Y., Ike M., Fujita M. (2002): Acute toxicity, mutagenicity, and estrogenicity of bisphenol A and other bisphenols. Environmental Toxicology, 17, 80–86.
- Chen S., Zhou D., Hsin L.Y., Kanaya N., Wong C., Yip R., Sakamuru S., Xia M., Yuan Y.C., Witt K., Teng C. (2014): AroER tri-screen is a biologically relevant assay for endocrine disrupting chemicals modulating the activity of aromatase and/or the estrogen receptor. Toxicological Sciences, 139, 198–209.
- Clayton R. (ed.) (2011): Endocrine Disrupters in the Environment. Foundation for Water Research, Marlow, UK, 3–22.
- Coggon D. (2002): Work with pesticides and organophosphate sheep dips. Occupational Medicine, 52, 467–470.

- Colborn T., vom Saal F.S., Soto A.M. (1993): Developmental effects of endocrine disrupting chemicals in wildlife and humans. Environmental Health Perspectives, 101, 378–384.
- Corbel T., Gayrard V., Puel S., Lacroix M.Z., Berrebi A., Gil S., Viguie C., Toutain P.L., Picard-Hagen N. (2014): Bidirectional placental transfer of Bisphenol A and its main metabolite, Bisphenol A-Glucuronide, in the isolated perfused human placenta. Reproductive Toxicology, 47, 51–58.
- Damstra T., Barlow S., Bergman A., Kavlock R., Van Der Kraak G. (eds) (2002): Global assessment of the state-of-thescience of endocrine disruptors. International Programme on Chemical Safety, World Health Organization. Available from http://www.who.int/ipcs/publications/new\_issues/ endocrine\_disruptors/en/ (accessed Aug 1, 2002).
- Danzl E., Sei K., Soda S., Ike M., Fujita M. (2009): Biodegradation of bisphenol A, bisphenol F and bisphenol S in seawater. International Journal of Environmental Research and Public Health, 6, 1472–1484.
- Delfosse V., Grimaldi M., Pons J.L., Boulahtouf A., le Maire A., Cavailles V., Labesse G., Bourguet W., Balaguer P. (2012): Structural and mechanistic insights into bisphenols action provide guidelines for risk assessment and discovery of bisphenol A substitutes. Proceedings of the National Academy of Sciences of the United States of America, 109, 14930–14935.
- Diamanti-Kandarakis E., Bourguignon J.P., Giudice L.C. (2009): Endocrine-disrupting chemicals: an Endocrine Society scientific statement. Endocrine Reviews, 30, 293–342.
- Dietrich D.R., Hengstler J.G. (2016): From bisphenol A to bisphenol F and a ban of mustard due to chronic low-dose exposures? Archives of Toxicology, 90, 489–491.
- Dodds E.C., Lawson W. (1936): Synthetic estrogenic agents without the phenanthrene nucleus. Nature, 137, 996.
- Dolinoy D.C., Huang D., Jirtle R.L. (2007): Maternal nutrient supplementation counteracts bisphenol A-induced DNA hypomethylation in early development. Proceedings of the National Academy of Sciences of the United States of America, 104, 13056–13061.
- Drastichova J., Svobodova Z., Groenland M., Dobsikova R., Zlabek V., Weissova D. (2005): Effect of exposure to bisphenol A and 17b-estradiol on the sex differentiation in zebrafish (Danio rerio). Acta Veterinaria Brno, 74, 287–291.
- Edlow A.G., Chen M., Smith N.A., Lu C., McElrath T.F. (2012): Fetal bisphenol A exposure: concentration of conjugated and unconjugated bisphenol A in amniotic fluid in the second and third trimesters. Reproductive Toxicology, 34, 1–7.
- Ehrlich S., Williams P.L., Missmer S.A., Flaws J.A., Ye X., Calafat A.M., Petrozza J.C., Wright D., Hauser R. (2012):

Urinary bisphenol A concentrations and early reproductive health outcomes among women undergoing IVF. Human Reproduction, 27, 3583–3592.

- Ehrlich S., Calafat A.M., Humblet O., Smith T., Hauser R. (2014): Handling of thermal receipts as a source of exposure to bisphenol A. Journal of the American Medical Association, 311, 859–860.
- Eichenlaub-Ritter U., Vogt E., Cukurcam S., Sun F., Pacchierotti F., Parry J. (2008): Exposure of mouse oocytes to bisphenol A causes meiotic arrest but not aneuploidy. Mutation Research, 651, 82–92.
- Commission Directive (2011): Commission Directive 2011/8/EU of 28 January 2011 amending Directive 2002/72/EC as regards the restriction of use of Bisphenol A in plastic infant feeding bottles. Official Journal of the European Union, L 26, 11–14.
- EFSA (2014): Bisphenol A: EFSA consults on assessment of risks to human health. European Food Safety Authority. Available from www.efsa.europa.eu (accessed Jan 17, 2014).
- Fahrenkamp-Uppenbrink J. (2015): Using chemical design to avoid regrets. Science, 347, 1213.
- FDA (2011): Bisphenol A (BPA). US Food and Drug Administration. Available from http://www.fda.gov/Food/ FoodborneIllnessContaminants/ChemicalContaminants/ ucm166145.htm (accessed March 24, 2011).
- Fernandez M.F., Roman M., Arrebola J.P., Olea N. (2014): Endocrine disruptors: time to act. Current Environmental Health Reports, 1, 325–332.
- Ferris J., Favetta L.A., King W.A. (2015): Bisphenol A exposure during oocyte maturation in vitro results in spindle abnormalities and chromosome misalignment in Bos taurus. Cytogenetic and Genome Research, 145, 50–58.
- Fic A., Zegura B., Sollner Dolenc M., Filipic M., Peterlin Masic L. (2013): Mutagenicity and DNA damage of bisphenol A and its structural analogues in HepG2 cells. Archives of Industrial Hygiene and Toxicology, 64, 189–200.
- Glausiusz J. (2014): Toxicology: the plastics puzzle. Nature, 508, 306–308.

Goodson A., Summerfield W., Cooper I. (2002): Survey of bisphenol A and bisphenol F in canned foods. Food Additives and Contaminants, 19, 796–802.

Grasselli F., Baratta L., Baioni L., Bussolati S., Ramoni R., Grolli S., Basini G. (2010): Bisphenol A disrupts granulosa cell function. Domestic Animal Endocrinology, 39, 34–39.

Grignard E., Lapenna S., Bremer S. (2012): Weak estrogenic transcriptional activities of Bisphenol A and Bisphenol S. Toxicology in Vitro, 26, 727–731.

Helies-Toussaint C., Peyre L., Costanzo C., Chagnon M.C., Rahmani R. (2014): Is bisphenol S a safe substitute for bisphenol A in terms of metabolic function? An in vitro study. Toxicology and Applied Pharmacology, 280, 224–235.

- Howard G.J. (2014): Chemical alternatives assessment: the case of flame retardants. Chemosphere, 116, 112–117.
- Huang Y.Q., Wong C.K.C., Zheng J.S., Bouwman H., Barra R., Wahlstrom B., Wong M.H. (2012): Bisphenol A (BPA) in China: a review of sources, environmental levels, and potential human health impacts. Environment International, 42, 91–99.
- Hubbs A.F., Cumpston A.M., Goldsmith W.T., Battelli L.A., Kashon M.L., Jackson M.C., Frazer D.G., Fedan J.S., Goravanahally M.P., Castranova V., Kreiss K., Willard P.A., Friend S., Schwegler-Berry D., Fluharty K.L., Sriram K. (2012): Respiratory and olfactory cytotoxicity of inhaled 2,3-pentanedione in Sprague-Dawley rats. The American Journal of Pathology, 181, 829–844.
- Hunt P.A., Koehler K.E., Susiarjo M., Hodges C.A., Ilagan A., Voigt R.C., Thomas S., Thomas B.F., Hassold T.J. (2003): Bisphenol A exposure causes meiotic aneuploidy in the female mouse. Current Biology, 13, 546–553.
- Hunt P.A., Lawson C., Gieske M., Murdoch B., Smith H., Marre A., VandeVoort C.A. (2012): Bisphenol A alters early oogenesis and follicle formation in the fetal ovary of the rhesus monkey. Proceedings of the National Academy of Sciences of the United States of America, 109, 17525–17530.
- Ike M., Chen M.Y., Danzl E., Sei K., Fujita M. (2006): Biodegradation of a variety of bisphenols under aerobic and anaerobic conditions. Water Science and Technology, 53, 153–160.
- Ikezuki Y., Tsutsumi O., Takai Y., Kamei Y., Taketani Y. (2002): Determination of bisphenol A concentrations in human biological fluids reveals significant early prenatal exposure. Human Reproduction, 17, 2839–2841.
- Jasarevic E., Sieli P.T., Twellman E.E., Welsh T.H., Schachtman T.R., Roberts R.M. (2011): Disruption of adult expression of sexually selected traits by developmental exposure to bisphenol A. Proceedings of the National Academy of Sciences of the United States of America, 108, 11715–11720.
- Jenkins S., Wang J., Eltoum I., Desmond R., Lamartiniere C.A. (2011): Chronic oral exposure to bisphenol A results in a nonmonotonic dose response in mammary carcinogenesis and metastasis in MMTV-erbB2 mice. Environmental Health Perspectives, 119, 1604–1609.
- Ji K., Hong S., Kho Y., Choi K. (2013): Effects of bisphenol S exposure on endocrine functions and reproduction of zebrafish. Environmental Science and Technology, 47, 8793–8800.
- Jimenez-Diaz I., Zafra-Gomez A., Ballesteros O., Navea N., Navalon A., Fernandez M.F., Olea N., Vilchez J.L. (2010): Determination of Bisphenol A and its chlorinated derivatives in placental tissue samples by liquid chromatography-tandem mass spectrometry. Journal of Chromatography B, 878, 3363–3369.

- Joskow R., Barr D.B., Barr J.R., Calafat A.M., Needham L.L., Rubin C. (2006): Exposure to bisphenol A from bis-glycidyl dimethacrylate-based dental sealants. The Journal of the American Dental Association, 137, 353–362.
- Khan D., Ahmed S.A. (2015): Epigenetic regulation of non-lymphoid cells by Bisphenol-A, a model endocrine disrupter: potential implications for immunoregulation. Frontiers in Endocrinology, 6, 91.
- Kinch C.D., Ibhazehiebo K., Jeong J.H., Habibi H.R., Kurrasch D.M. (2015): Low-dose exposure to bisphenol A and replacement bisphenol S induces precocious hypothalamic neurogenesis in embryonic zebrafish. Proceedings of the National Academy of Sciences of the United States of America, 112, 1475–1480.
- Kitamura S., Suzuki T., Sanoh S., Kohta R., Jinno N., Sugihara K., Yoshihara S., Fujimoto N., Watanabe H., Ohta S. (2005): Comparative study of the endocrine-disrupting activity of bisphenol A and 19 related compounds. The Journal of Toxicological Sciences, 84, 249–259.
- Kloas W., Lutz I., Einspanier R. (1999): Amphibians as a model to study endocrine disruptors: II. Estrogenic activity of environmental chemicals in vitro and in vivo. Science of the Total Environment, 225, 59–68.
- Knez J., Kranvogl R., Breznik B.P., Voncina E., Vlaisavljevic V. (2014): Are urinary bisphenol A levels in men related to semen quality and embryo development after medically assisted reproduction? Fertility and Sterility, 101, 215–221.
- Krishnan A., Stathis P., Permuth S., Tokes L., Feldman D. (1993): Bisphenol-A – an estrogenic substance is released from polycarbonate flasks during autoclaving. Endocrinology, 132, 2279–2286.
- Kundakovic M., Gudsnuk K., Franks B., Madrid J., Miller R.L., Perera F.P., Champagne F.A. (2013): Sex-specific epigenetic disruption and behavioral changes following low-dose in utero bisphenol A exposure. Proceedings of the National Academy of Sciences of the United States of America, 110, 9956–9961.
- Kurosawa T., Hiroi H., Tsutsumi O., Ishikawa T., Osuga Y., Fujiwara T., Inoue S., Muramatsu M., Momoeda M., Taketani Y. (2002): The activity of bisphenol A depends on both the estrogen receptor subtype and the cell type. Endocrine Journal, 49, 465–471.
- Kuruto-Niwa R., Tateoka Y., Usuki Y., Nozawa R. (2007): Measurement of bisphenol A concentrations in human colostrum. Chemosphere, 66, 1160–1164.
- Kuruto-Niwa R., Nozawa R., Miyakoshi T., Shiozawa T., Terao Y. (2010): Estrogenic activity of alkylphenols, bisphenol S, and their chlorinated derivatives using a GFP expression system. Environmental Toxicology and Pharmacology, 19, 121–130.

- Kwintkiewicz J., Nishi Y., Yanase T., Giudice L.C. (2010): Peroxisome proliferator-activated receptor-γ mediates bisphenol A inhibition of FSH-stimulated IGF-1, aromatase, and estradiol in human granulosa cells. Environmental Health Perspectives, 118, 400–406.
- Lahnsteiner F., Berger B., Kletz M., Weismann T. (2005): Effect of bisphenol A on maturation and quality of semen and eggs in the brown trout, Salmo trutta f. fario. Aquatic Toxicology, 75, 213–224.
- Landrigan P.J., Kimmel C.A., Eskenazi B. (2004): Children's health and the environment: public health issues and challenges for risk assessment. Environmental Health Perspectives, 112, 257–265.
- Lang I.A., Galloway T.S., Scarlett A., Henley W.E., Depledge M., Wallace R.B. (2008): Association of urinary bisphenol A concentration with medical disorders and laboratory abnormalities in adults. The Journal of the American Medical Association, 300, 1303–1310.
- Lassen T.H., Frederiksen H., Jensen T.K., Petersen J.H., Joensen U.N., Main K.M., Andersson A.M. (2014): Urinary bisphenol A levels in young men: association with reproductive hormones and semen quality. Environmental Health Perspectives, 122, 478–484.
- Lee S., Liu X., Takeda S., Choi K. (2013): Genotoxic potentials and related mechanisms of bisphenol A and other bisphenol compounds: a comparison study employing chicken DT40 cells. Chemosphere, 93, 434–440.
- Le Fol V., Ait-Aissa S., Cabaton N., Dolo L., Grimaldi M., Balaguer P., Perdu E., Debrauwer L., Brion F., Zalko D. (2015): Cell-specific biotransformation of benzophenone-2 and bisphenol-S in zebrafish and human in vitro models used for toxicity and estrogenicity screening. Environmental Science and Technology, 49, 3860–3868.
- Li D.K., Zhou Z., Miao M., He Y., Wang J., Ferber J., Herrinton L.J., Gao E., Yuan W. (2011): Urine bisphenol-A (BPA) level in relation to semen quality. Fertility and Sterility, 95, 625–630.
- Liao C., Liu F., Alomirah H., Loi V.D., Mohd M.A., Moon H.B. (2012a): Bisphenol S in urine from the United States and seven Asian countries: occurrence and human exposures. Environmental Science and Technology, 46, 6860–6866.
- Liao C., Liu F., Guo Y., Moon H.B., Nakata H., Wu Q. (2012b): Occurrence of eight bisphenol analogues in indoor dust from the United States and several Asian countries: implications for human exposure. Environmental Science and Technology, 46, 9138–9145.
- Liao C., Liu F., Kannan K. (2012c): Bisphenol S, a new bisphenol analogue, in paper products and currency bills and its association with bisphenol A residues. Environmental Science and Technology, 46, 6515–6522.

- Maamar M., Lesne L., Desdoits-Lethimonier C., Coiffec I., Lassurguere J., Lavoue V., Deceuninck Y., Antignac J.P., Le Bizec B., Perdu E., Zalko D., Pineau C., Chevrier C., Dejucq-Rainsford N., Mazaud-Guittot S., Jegou B. (2015): An investigation of the endocrine-disruptive effects of bisphenol A in human and rat fetal testes. PLoS ONE, 10, e0117226.
- Machtinger R., Orvieto R. (2014): Bisphenol A, oocyte maturation, implantation, and IVF outcome: review of animal and human data. Reproductive BioMedicine On-line, 29, 404–410.
- Machtinger R., Combelles C.M., Missmer S.A., Correia K.F., Williams P., Hauser R., Racowsky C. (2013): Bisphenol-A and human oocyte maturation in vitro. Human Reproduction, 28, 2735–2745.
- Manfo F.P., Jubendradass R., Nantia E.A., Moundipa P.F., Mathur P.P. (2014): Adverse effects of bisphenol A on male reproductive function. Reviews of Environmental Contamination and Toxicology, 228, 57–82.
- Manikkam M., Tracey R., Guerrero-Bosagna C., Skinner M.K. (2013): Plastics derived endocrine disruptors (BPA, DEHP and DBP) induce epigenetic transgenerational inheritance of obesity, reproductive disease and sperm epimutations. PLoS ONE, 8, e55387.
- Mathew M., Sreedhanya S., Manoj P., Aravindakumar C.T., Aravind U.K. (2014): Exploring the interaction of bisphenol-S with serum albumins: a better or worse alternative for bisphenol A? The Journal of Physical Chemistry B, 118, 3832–3843.
- Matsushima A., Kakuta Y., Teramoto T., Koshiba T., Liu X., Okada H., Tokunaga T., Kawabata S., Kimura M., Shimohigashi Y. (2007): Structural evidence for endocrine disruptor bisphenol A binding to human nuclear receptor ERR gamma. The Journal of Biochemistry, 142, 517–524.
- McBride W.G. (1961): Thalidomide and congenital abnormalities. The Lancet, 278, 1358.
- Mendiola J., Jorgensen N., Andersson A.M., Calafat A.M., Ye X., Redmon J.B. (2010): Are environmental levels of bisphenol A associated with reproductive function in fertile men? Environmental Health Perspectives, 118, 1286–1291.
- Mendonca K., Hauser R., Calafat A.M., Arbuckle T.E., Duty S.M. (2014): Bisphenol A concentrations in maternal breast milk and infant urine. International Archives of Occupational and Environmental Health, 87, 13–20.
- Mendoza-Rodriguez C.A., Garcia-Guzman M., Baranda-Avila N., Morimoto S., Perrot-Applanat M., Cerbon M. (2011): Administration of bisphenol A to dams during perinatal period modifies molecular and morphological reproductive parameters of the offspring. Reproductive Toxicology, 31, 177–183.

- Mileva G., Baker S.L., Konkle A.T., Bielajew C. (2014): Bisphenol-A: epigenetic reprogramming and effects on reproduction and behaviour. International Journal of Environmental Research and Public Health, 11, 7537–7561.
- Miyamoto K.I., Kotake M. (2005): Estimation of daily bisphenol A intake of Japanese individuals with emphasis on uncertainty and variability. Environmental Sciences: an International Journal of Environmental Physiology and Toxicology, 13, 15–29.
- Mok-Lin E., Ehrlich S., Williams P.L., Petrozza J., Wright D.L., Calafat A.M., Ye X., Hauser R. (2010): Urinary bisphenol A concentrations and ovarian response among women undergoing IVF. International Journal of Andrology, 33, 385–393.
- Molina-Molina J.M., Amaya E., Grimaldi M., Saenz J.M., Real M., Fernandez M.F., Balaguer P., Olea N. (2013): In vitro study on the agonistic and antagonistic activities of bisphenol-S and other bisphenol-A congeners and derivatives via nuclear receptors. Toxicology and Applied Pharmacology, 272, 127–136.
- Naderi M., Wong M.Y., Gholami F. (2014): Developmental exposure of zebrafish (Danio rerio) to bisphenol-S impairs subsequent reproduction potential and hormonal balance in adults. Aquatic Toxicology, 148, 195–203.
- Nakagawa Y., Tayama S. (2000): Metabolism and cytotoxicity of bisphenol A and other bisphenols in isolated rat hepatocytes. Archives of Toxicology, 74, 99–105.
- Navarro V.M., Sanchez-Garrido M.A., Castellano J.M., Roa J., Garcia-Galiano D., Pineda R., Tena-Sempere M. (2009): Persistent impairment of hypothalamic KiSS-1 system after exposures to estrogenic compounds at critical periods of brain sex differentiation. Endocrinology, 150, 2359–2367.
- Nevoral J., Krejcova T., Petr J., Melicharova P., Vyskocilova A., Dvorakova M., Sedmikova M. (2013): The role of nitric oxide synthase isoforms in aged porcine oocytes. Czech Journal of Animal Science, 58, 453–459.
- Newbold R.R., Jefferson W.N., Padilla-Banks E. (2009): Prenatal exposure to bisphenol A at environmentally relevant doses adversely affects the murine female reproductive tract later in life. Environmental Health Perspectives, 117, 879–885.
- N'Tumba-Byn T., Moison D., Lacroix M., Lecureuil C., Lesage L. (2012): Differential effects of bisphenol A and diethylstilbestrol on human, rat and mouse fetal Leydig cell function. PLoS ONE, 7, e51579.
- Owens J.W., Ashby J. (2002): Critical review and evaluation of the uterotrophic bioassay for the identification of possible estrogen agonists and antagonists: in support of the validation of the OECD uterotrophic protocols for the laboratory rodent. Critical Reviews in Toxicology, 32, 445–520.

- Pacchierotti F., Ranaldi R., Eichenlaub-Ritter U., Attia S., Adler I.D. (2008): Evaluation of aneugenic effects of bisphenol A in static and germ cells of the mouse. Mutation Research, 651, 64–70.
- Patisaul H.B., Todd K.L., Mickens J.A., Adewale H.B. (2009): Impact of neonatal exposure to the ERα agonist PPT, bisphenol-A or phytoestrogens on hypothalamic kisspeptin fiber density in male and female rats. Neurotoxicology, 30, 350–357.
- Peretz J., Vrooman L., Ricke W.A., Hunt P.A., Ehrlich S., Hauser R., Padmanabhan V., Taylor H.S., Swan S.H., VandeVoort C.A., Flaws J.A. (2014): Bisphenol A and reproductive health: update of experimental and human evidence. Environmental Health Perspectives, 122, 775–786.
- Perez P., Pulgar R., Olea-Serrano F., Villalobos M., Rivas A., Metzler M., Pedraza V., Olea N. (1998): The estrogenicity of bisphenol A-related diphenylalkanes with various substituents at the central carbon and the hydroxy groups. Environmental Health Perspectives, 106, 167–174.
- Petr J., Chmelikova E., Zalmanova T., Tumova L., Kheilova K., Kucerova-Chrpova V., Jilek F. (2013): Pyrethroids cypermethrin, deltamethrin and fenvalerate have different effects on in vitro maturation of pig oocytes at different stages of growth. Animal, 7, 134–142.
- Peyre L., Rouimi P., de Sousa G., Helies-Toussaint C., Carre B., Barcellini S., Chagnon M.C., Rahmani R. (2014): Comparative study of bisphenol A and its analogue bisphenol S on human hepatic cells: a focus on their potential involvement in nonalcoholic fatty liver disease. Food and Chemical Toxicology, 70, 9–18.
- Qiu W., Zhao Y., Yang M., Farajzadeh M., Pan C., Wayne N.L. (2016): Actions of bisphenol A and bisphenol S on the reproductive neuroendocrine system during early development in zebrafish. Endocrinology, 157, 636–647.
- Rama Raju G.A., Prakash G.J., Krishna K.M., Madan K. (2007): Meiotic spindle and zona pellucida characteristics as predictors of embryonic development: a preliminary study using PolScope imaging. Reproductive Biomedicine Online, 14, 166–174.
- Richter C., Birnbaum L.S., Farabollini F., Newbold R.R., Rubin B.S., Talsness C.E., Vandenbergh J.G., Walser-Kuntz D.R., vom Saal F.S. (2007): In vivo effects of bisphenol A in laboratory rodent studies. Reproductive Toxicology, 24, 199–224.
- Rochester J.R. (2013): Bisphenol A and human health: a review of the literature. Reproductive Toxicology, 42, 132–155.
- Rochester J.R., Bolden A.L. (2015): Bisphenol S and F: a systematic review and comparison of the hormonal activity of bisphenol A substitutes. Environmental Health Perspectives, 123, 643–650.

- Rosenmai A.K., Dybdahl M., Pedersen M., van Vugt-Lussenburg B.M.A., Wedebye E.B., Taxvig C., Vinggaard A.M. (2014): Are structural analogues to bisphenol a safe alternatives? Toxicological Sciences, 139, 3–47.
- Salgueiro-Gonzalez N., Turnes-Carou I., Vinas-Dieguez L., Muniategui-Lorenzo S., Lopez-Mahia P., Prada-Rodriguez D. (2015): Occurrence of endocrine disrupting compounds in five estuaries of the northwest coast of Spain: ecological and human health impact. Chemosphere, 131, 241–247.
- Salvesen G.S., Walsh C.M. (2014): Functions of caspase 8: the identified and the mysterious. Seminars in Immunology, 26, 246–252.
- Santamaria C., Durando M., Munoz de Toro M., Luque E.H., Rodriguez H.A. (2016): Ovarian dysfunction in adult female rat offspring born to mothers perinatally exposed to low doses of bisphenol A. The Journal of Steroid Biochemistry and Molecular Biology, 158, 220–230.
- Schonfelder G., Wittfoht W., Hopp H., Talsness C.E., Paul M., Chahoud I. (2002): Parent bisphenol A accumulation in the human maternal–fetal–placental unit. Environmental Health Perspectives, 110, 703–707.
- Sedmikova M., Petr J., Dorflerova A., Nevoral J., Novotna B. (2013): Inhibition of c-Jun N-terminal kinase (JNK) suppresses porcine oocyte aging in vitro. Czech Journal of Animal Science, 58, 535–545.
- Sohoni P.C.R.T., Hurd K., Caunter J., Hetheridge M., Williams T., Woods C., Evans M., Toy R., Gargas M., Sumpter J.P. (2001): Reproductive effects of long-term exposure to bisphenol A in the fathead minnow (Pimephales promelas). Environmental Science and Technology, 35, 2917–2925.
- Song S., Song M., Zeng L., Wang T., Liu R., Ruan T. (2014): Occurrence and profiles of bisphenol analogues in municipal sewage sludge in China. Environmental Pollution, 186, 14–19.
- Sui Y., Ai N., Park S., Rios-Pilier J., Perkins J.T., Welsh W.J., Zhou C. (2012): Bisphenol A and its analogues activate human pregnane X receptor. Environmental Health Perspectives, 120, 399–405.
- Susiarjo M., Hassold T.J., Freeman E., Hunt P.A. (2007): Bisphenol A exposure in utero disrupts early oogenesis in the mouse. PLoS Genetics, 3, e5.
- Susiarjo M., Sasson I., Mesaros C., Bartolomei M.S. (2013): Bisphenol A exposure disrupts genomic imprinting in the mouse. PLoS Genetics, 9, e1003401.
- Susiarjo M., Xin F., Bansal A., Stefaniak M., Li C., Simmons R.A., Bartolomei M.S. (2015): Bisphenol A exposure disrupts metabolic health across multiple generations in the mouse. Endocrinology, 156, 2049–2058.
- Suzuki A., Sugihara A., Uchida K., Sato T., Ohta Y., Katsu Y., Watanabe H., Iguchi T. (2002): Developmental effects of

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perinatal exposure to bisphenol-A and diethylstilbestrol on reproductive organs in female mice. Reproductive Toxicology, 16, 107–116.

- Takayanagi S., Tokunaga T., Liu X., Okada H., Matsushima A., Shimohigashi Y. (2006): Endocrine disruptor bisphenol A strongly binds to human estrogen-related receptor gamma (ERRgamma) with high constitutive activity. Toxicology Letters, 167, 95–105.
- Teng C., Goodwin B., Shockley K., Xia M., Huang R., Norris J., Merrick B.A., Jetten A.M., Austin C.P., Tice R.R. (2013): Bisphenol A affects androgen receptor function via multiple mechanisms. Chemico-Biological Interactions, 203, 556–564.
- Tomari H., Honjou K., Nagata Y., Horiuchi T. (2011): Relationship between meiotic spindle characteristics in human oocytes and the timing of the first zygotic cleavage after intracytoplasmic sperm injection. Journal of Assisted Reproduction and Genetics, 28, 1099–1104.
- Trapphoff T., Heiligentag M., Hajj N.E., Haaf T., Eichenlaub-Ritter U. (2013): Chronic exposure to a low concentration of bisphenol A during follicle culture affects the epigenetic status of germinal vesicles and metaphase II oocytes. Fertility and Sterility, 100, 1758–1767.
- Trasande L., Attina T.M., Blustein J. (2012): Association between urinary bisphenol A concentration and obesity prevalence in children and adolescents. The Journal of the American Medical Association, 308, 1113–1121.
- Vandenberg L.N. (2014): Non-monotonic dose responses in studies of endocrine disrupting chemicals: bisphenol A as a case study. Dose-Response, 12, 259–276.
- Vandenberg L.N., Maffini M.V., Sonnenschein C., Rubin B.S., Soto A.M. (2009): Bisphenol-A and the great divide: a review of controversies in the field of endocrine disruption. Endocrine Reviews, 30, 75–95.
- Vanderberg L.N., Colborn T., Hazes T.B., Heindel J.J., Jacobs D.R., Lee D.H., Shioda T., Soto A.M., vom Saal F.S., Welshons W.V., Zoeller R.T., Myers J.P. (2012): Hormones and endocrine-disrupting chemicals: low-dose effects and nonmonotonic dose response. Endocrine Reviews, 33, 378–455.
- Vandenberg L.N., Colborn T., Hayes T.B., Heindel J.J., Jacobs D.R., Lee D.H., Myers J.P., Shioda T., Soto A.M., Vom Saal F.S., Welshons W.V., Zoeller R.T. (2013). Regulatory decisions on endocrine disrupting chemicals should be based on the principles of endocrinology. Reproductive Toxicology, 38, 1–15.
- Vinas P., Campillo N., Martinez-Castillo N., Hernandez-Cordoba M. (2010): Comparison of two derivatizationbased methods for solid-phase microextraction-gas chromatography-mass spectrometric determination of bisphenol A, bisphenol S and biphenol migrated from

food cans. Analytical and Bioanalytical Chemistry, 397, 115–125.

- Vinas R., Watson C.S. (2013a): Bisphenol S disrupts estradiol-induced nongenomic signaling in a rat pituitary cell line: effects on cell functions. Environmental Health Perspectives, 121, 352–358.
- Vinas R., Watson C.S. (2013b): Mixtures of xenoestrogens disrupt estradiol-induced non-genomic signaling and downstream functions in pituitary cells. Environmental Health, 12, 26.
- Vinggaard A.M., Niemela J., Wedebye E.B., Jensen G.E. (2008): Screening of 397 chemicals and development of a quantitative structure–activity relationship model for androgen receptor antagonism. Chemical Research in Toxicology, 21, 813–823.
- Vitku J., Sosvorova L., Chlupacova T., Hampl R., Hill M., Sobotka V., Heracek J., Bicikova M., Starka L. (2015):
  Differences in bisphenol A and estrogen levels in the plasma and seminal plasma of men with different degrees of infertility. Physiological Research, 64, 303–311.
- vom Saal F.S., Hughes C. (2005): An extensive new literature concerning low-dose effects of bisphenol A shows the need for a new risk assessment. Environmental Health Perspectives, 113, 926–933.
- vom Saal F.S., Welshons W.V. (2005): Large effects from small exposures. II. The importance of positive controls in low-dose research on bisphenol A. Environmental Research, 100, 50–76.
- vom Saal F.S., Akingbemi T.B., Belcher S.M., Birnbaum L.S., Crain D.A., Eriksen M., Farabollini F., Guillette Jr. L.J., Hauser R., Heindel J.J., Ho S.-M., Hunt P.A., Iguchi T., Jobling S., Kanno J., Keri R.A., Knudsen K.E., Laufer H., LeBlanc G.A., Marcus M., McLachlan J.A., Myers J.P., Nadal A., Newbold R.R., Olea N., Prins G.S., Richter C.A., Rubin B.S., Sonnenschein C., Soto A.M., Talsness C.E., Vandenbergh J.G., Vandenberg L.N., Walser-Kuntz D.R., Watson C.S., Welshons W.V., Wetherill Y., Zoeller R.T. (2007): Chapel Hill bisphenol A expert panel consensus statement: integration of mechanisms, effects in animals and potential to impact human health at current levels of exposure. Reproductive Toxicology, 24, 131–138.
- Vrooman L.A., Oatley J.M., Griswold J.E., Hassold T.J., Hunt P.A. (2015): Estrogenic exposure alters the spermatogonial stem cells in the developing testis, permanently reducing crossover levels in the adult. PLoS Genetics, 11, e1004949.
- Wang F., Hua J., Chen M., Xia Y., Zhang Q., Zhao R., Zhou W., Zhang Z., Wang B. (2012): High urinary bisphenol A concentrations in workers and possible laboratory abnormalities. Occupational and Environmental Medicine, 69, 679–684.

- Wang T., Lu J., Xu M., Xu Y., Li M., Liu Y., Ning G. (2013): Urinary bisphenol A concentration and thyroid function in Chinese adults. Epidemiology, 24, 295–302.
- Warita K., Mitsuhashi T., Ohta K., Suzuki S., Hoshi N., Miki T., Takeuchi Y. (2013): Gene expression of epigenetic regulatory factors related to primary silencing mechanism is less susceptible to lower doses of bisphenol A in embryonic hypothalamic cells. The Journal of Toxicological Sciences, 38, 285–289.
- Wetherill Y.B., Akingbemi B.T., Kanno J., McLachlan J.A., Nadal A., Sonnenschein C., Watson C.S., Zoeller R.T., Belcher S.M. (2007): In vitro molecular mechanisms of bisphenol A action. Reproductive Toxicology, 24, 178–198.
- Wilson M.E., Sengoku T. (2013): Developmental regulation of neuronal genes by DNA methylation: environmental influences. International Journal of Developmental Neuroscience, 31, 448–451.
- Xi W., Lee C.K.F., Yeung W.S.B., Giesy J.P., Wong M.H., Zhang X., Wong C.K. (2011): Effect of perinatal and postnatal bisphenol A exposure to the regulatory circuits at the hypothalamus–pituitary–gonadal axis of CD-1 mice. Reproductive Toxicology, 31, 409–417.
- Xiao S., Diao H., Smith M.A., Song X., Ye X. (2011): Preimplantation exposure to bisphenol A (BPA) affects embryo transport, preimplantation embryo development, and uterine receptivity in mice. Reproductive Toxicology, 32, 434–441.
- Xu L.C., Sun H., Chen J.F., Bian Q., Qian J., Song L., Wang X.R. (2005): Evaluation of androgen receptor transcriptional activities of bisphenol A, octylphenol and nonylphenol in vitro. Toxicology, 216, 197–203.
- Yamada H., Furuta I., Kato E.H., Kataoka S., Usuki Y., Kobashi G., Fujimoto S. (2002): Maternal serum and amniotic fluid bisphenol A concentrations in the early second trimester. Reproductive Toxicology, 16, 735–739.
- Yamasaki K., Noda S., Imatanaka N., Yakabe Y. (2004): Comparative study of the uterotrophic potency of 14 chemicals in a uterotrophic assay and their receptor-binding affinity. Toxicology Letters, 146, 111–120.
- Yang O., Kim H.L., Weon J.I., Seo Y.R. (2015): Endocrinedisrupting chemicals: review of toxicological mechanisms using molecular pathway analysis. Journal of Cancer Prevention, 20, 12–24.

- Yang Y., Lu L., Zhang J., Yang Y., Wu Y., Shao B. (2014): Simultaneous determination of seven bisphenols in environmental water and solid samples by liquid chromatography–electrospray tandem mass spectrometry. Journal of Chromatography A, 1328, 26–34.
- Ye J., Coleman J., Hunter M.G., Craigon J., Campbell K.H.S., Luck M.R. (2007): Physiological temperature variants and culture media modify meiotic progression and developmental potential of pig oocytes in vitro. Reproduction, 133, 877–886.
- Zhang H.Q., Zhang X.F., Zhang L.J., Chao H.H., Pan B., Feng Y.M., Shen W., Li L., Sun X.F. (2012): Fetal exposure to bisphenol A affects the primordial follicle formation by inhibiting the meiotic progression of oocytes. Molecular Biology Reports, 39, 5651–5657.
- Zhang T., Qin X.S., Zhou Y., Zhang X.F., Wang L.Q., Felici M., Chen H., Qin G.Q., Shen W. (2014): Di-(2-ethylhexyl) phthalate and bisphenol A exposure impairs mouse primordial follicle assembly in vitro. Experimental and Molecular Mutagenesis, 55, 343–353.
- Zhang X., Chang H., Wiseman S., He Y., Higley E., Jones P., Wong C.K.C., Al-Khedhairy A., Giesy J.P., Hecker M. (2011): Bisphenol A disrupts steroidogenesis in human H295R cells. Toxicological Sciences, 121, 320–327.
- Zimmerman J.B., Anastas P.T. (2015): Toward designing safer chemicals. Science, 347, 215.
- Ziv-Gal A., Wang W., Zhou C., Flaws J.A. (2015): The effects of in utero bisphenol A exposure on reproductive capacity in several generations of mice. Toxicology and Applied Pharmacology, 284, 354–362.
- Zoeller R.T., Brown T.R., Doan L.L., Gore A.C., Skakkebaek N.E., Soto A.M., Woodruff T.J., vom Saal F.S. (2012): Endocrine-disrupting chemicals and public health protection: a statement of principles from the Endocrine Society. Endocrinology, 153, 4097–4110.
- Zoller O., Bruschweiler B.J., Magnin R., Reinhard H., Rhyn P., Rupp H., Zelter S., Felleisen R. (2016): Natural occurrence of bisphenol F in mustard. Food Additives and Contaminants: Part A, 33, 137–146.

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